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Species mobility and landscape context determine the importance of local and landscape-level attributes: Insights from a large-scale natural experiment

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1 **ABSTRACT:**

2 Conservation strategies to tackle habitat loss and fragmentation require actions at local (e.g.
3 improving/expanding existing habitat patches) and landscape level (e.g. creating new habitat
4 in the matrix). However, the relative importance of these actions for biodiversity is still
5 poorly understood, leading to debate on how to prioritise conservation activities. Here, we
6 assess the relative importance of local *vs.* landscape-level attributes in determining the use of
7 woodlands by bats in fragmented landscapes; we also compare the role of habitat amount in
8 the surrounding landscape *per se vs.* a combination of both habitat amount and configuration
9 and explore whether the relative importance of these attributes varies with species mobility
10 and landscape context. We conducted acoustic surveys in 102 woodland patches in the UK
11 which form part of the Wren project (www.wren-project.com), a large-scale natural
12 experiment designed to study the effects of 160 years of woodland creation on biodiversity
13 and inform landscape-scale conservation. We used multivariate analysis and a model-
14 selection approach to assess the relative importance of local (e.g. vegetation structure) and
15 landscape-level (e.g. amount/configuration of surrounding land types) attributes on bat
16 occurrence and activity levels. Species mobility was an important trait determining the
17 relative importance of local *vs.* landscape-level attributes for different bat species. Lower
18 mobility species were most strongly influenced by local habitat quality; the landscape became
19 increasingly important for higher mobility species. At the landscape-scale, a combination of
20 habitat amount and configuration appeared more important than habitat amount alone for
21 lower mobility species, whilst the opposite was observed for higher mobility species.
22 Regardless of species mobility, landscape-level attributes appeared more important for bats in
23 a more homogeneous and intensively farmed landscape. Conservation strategies involving
24 habitat creation and restoration should take into account the mobility of target species and

25 prioritise landscape-level actions in more homogeneous and intensively farmed landscapes
26 where habitat loss and fragmentation have been more severe.

27

28 Keywords: Bats, Chiroptera, connectivity, ecological networks, fragmentation, landscape-
29 scale conservation, natural experiment, woodland creation, WrEN project.

30

31 **1. INTRODUCTION:**

32 Habitat loss and fragmentation are amongst the main threats to biological diversity (Haddad
33 *et al.* 2015). Reversing these trends is therefore a high conservation priority, and becoming
34 increasingly important with the need to make species populations more resilient to climate
35 change (e.g. by facilitating inter-population movements and providing more resources to
36 support population recovery; Newson *et al.* 2014). Conservation strategies to tackle habitat
37 fragmentation often include creating new habitat patches, expanding, connecting and
38 restoring existing patches, and increasing the permeability of the surrounding matrix to
39 increase the persistence and movement of species across the landscape (Lawton *et al.* 2010).
40 This approach requires actions at both the local and landscape level. However, the relative
41 importance of these actions is still poorly understood and much debated (e.g. Hodgson *et al.*
42 2011; Fahrig 2013; Hanski 2015). This hampers the translation of ecological theory into
43 practice and highlights the urgent need for scientific evidence to inform conservation and
44 ensure that habitat creation and restoration activities are implemented in the most effective
45 manner to benefit biodiversity.

46 Studies on habitat fragmentation can provide insights into what factors are likely to be
47 important for habitat creation and restoration (e.g. Humphrey *et al.* 2015). However, their
48 focus is mainly on the capacity of species to persist in habitat fragments, rather than on their
49 ability to colonise and capitalise on new patches. Therefore, there is a need for ecological
50 studies informing landscape-scale conservation to investigate the effects of habitat restoration
51 and creation. Additionally, such studies should focus on landscapes at sufficiently large
52 spatial scales to ensure ecological realism and incorporate appropriately long time scales to
53 account for the considerable lag in ecosystem development and colonisation associated with
54 habitat restoration and creation (Watts *et al.* 2016).

55 The WrEN project (Watts *et al.* 2016) is a large-scale natural experiment designed to study
56 the effects of 160 years of woodland creation on biodiversity in UK landscapes; WrEN aims
57 to inform conservation activities by quantifying the relative effects of different local and
58 landscape-level attributes on woodland biodiversity. Historically, woodland has been
59 severely affected by habitat loss and fragmentation, with worldwide deforestation resulting in
60 a 50% decrease in woodland cover over the last three centuries (Ramankutty & Foley 1999;
61 ‘woodland’ is the term commonly used in the United Kingdom to describe any forested area,
62 for convenience we use this term hereafter in the paper). More recently, deforestation rates
63 have slowed down; net woodland loss has halved over the last 25 years and woodland extent
64 has increased in many countries, particularly in temperate regions, largely through deliberate
65 planting often encouraged through the provision of financial incentives to landowners
66 (Keenan *et al.* 2015). It is often assumed that creating new woodlands will benefit
67 biodiversity. However, despite an overall increase in woodland area over recent decades,
68 many species (particularly those associated with native semi-natural woodland) continue to
69 undergo severe population declines in many places (e.g. Burns *et al.* 2013). A sound
70 ecological knowledge of the factors likely to enhance biodiversity in newly planted
71 woodlands is of key importance to inform future conservation strategies if these are to benefit
72 biodiversity. However, slow rates of vegetation succession, together with significant time
73 lags in species colonisation and capitalisation of new habitat patches, have resulted in a lack
74 of empirical studies assessing the long-term value of woodland creation and the relative
75 importance of local (e.g. patch size and quality) vs. landscape-level (e.g. degree of
76 connectivity) characteristics for biodiversity.

77 In the UK, long-term deforestation has been so severe that woodland cover dropped from a
78 post-glacial high of 70% to a low of 5% in 1900 (Watts 2006). Since then it has increased to
79 13%, mainly as a result of woodland planting and restoration programmes that started in the

80 mid-19th century and increased in the 20th century (Quine *et al.* 2013). These historical
81 changes in woodland cover have resulted in landscapes containing new woodland patches of
82 varying ages, sizes and degrees of isolation. Many of these new woodlands were established
83 on agricultural land without remnant woodland biodiversity. Therefore, the occurrence and
84 relative abundance of species within these sites represents successful colonisation,
85 presumably mediated by attributes of the sites themselves and the landscapes around them.
86 This array of new woodland sites forms the basis of the WrEN project (Watts *et al.* 2016).

87 We have selected bats as one of the WrEN taxonomic study groups because many bat species
88 are woodland specialists (i.e. are strongly dependent on woodland for roosting and/or
89 foraging) whilst others frequently use woodland for roosting, foraging or as commuting
90 habitat alongside other land types (Lacki, Hayes & Kurta 2007). Therefore, bats have been
91 badly affected by worldwide deforestation (Mickleburgh, Hutson & Racey 2002) and
92 fragmentation (e.g. Meyer & Kalko 2008). Many bat species are highly mobile and so are
93 influenced by both woodland extent and configuration at relatively large spatial scales (e.g.
94 up to 5 km; Klingbeil & Willig 2009; Boughey *et al.* 2011a; Fuentes-Montemayor *et al.*
95 2013). Bats' responses to habitat fragmentation often depend on species life-history traits,
96 such as roosting and foraging ecology. In general, slow-flying species adapted to forage in
97 cluttered environments, are particularly sensitive to habitat fragmentation (e.g. Meyer &
98 Kalko 2008), whilst fast-flying species adapted to forage in uncluttered environments are
99 relatively tolerant of fragmentation processes (e.g. Klingbeil & Willig 2009). Local woodland
100 attributes (e.g. vegetation structure) are also important in determining the availability and
101 quality of habitat for bats. In general, mature broadleaved woodlands with high availability of
102 large diameter snags and dense canopy cover are associated with high bat activity levels
103 (Lacki, Hayes & Kurta 2007).

104 We studied temperate bat assemblages in historic woodland creation sites as part of the
105 WrEN project. Our objectives were to:

- 106 i) Assess the relative importance of local *vs.* landscape-level habitat characteristics
107 on species occurrence and activity levels.
- 108 ii) Compare the role of habitat amount in the surrounding landscape *per se vs.* a
109 combination of both habitat amount and configuration in determining species
110 occurrence and activity levels.
- 111 iii) Examine how variables identified in i) and ii) vary between bat species with
112 differing mobility.

113 We expect the relative importance of local *vs.* landscape-level attributes to depend on species
114 mobility (hypothesis 1), with higher mobility species being more strongly influenced by their
115 surrounding landscape than lower mobility species for which local woodland character might
116 be more important (Meyer & Kalko 2008; Fuentes-Montemayor *et al.* 2013). At the
117 landscape scale, we expect amount of surrounding woodland to be more important for higher
118 mobility species, whereas a combination of surrounding woodland amount and configuration
119 will be more important for lower mobility species (hypothesis 2), as the latter are more
120 limited by dispersal and often reluctant to fly across open spaces (Entwistle, Racey &
121 Speakman 1996; Frey-Ehrenbold *et al.* 2013).

122 Additionally, we identify specific local (e.g. woodland patch size and vegetation structure)
123 and landscape-level (e.g. proportion of different habitat types in the surrounding matrix)
124 attributes influencing the occurrence and/or activity levels of temperate bats. Finally, we use
125 our findings to provide recommendations to inform landscape-scale conservation strategies
126 which aim to combat habitat loss and fragmentation.

127

128 **2. MATERIALS AND METHODS:**

129

130 **2.1. Study area and site selection protocol:**

131 The WrEN woodland sites are located in two study areas in Scotland (*ca.* 7,335 km²) and
132 England (*ca.* 8,570 km²) (Figure 1). Both areas are dominated (>70%) by agricultural land
133 and represent fairly typical lowland landscapes in these countries. We used digital maps
134 within a GIS environment (ArcGIS 10.2; ESRI) and a systematic site-selection process to
135 identify 102 broadleaved woodland patches (>80% broadleaved canopy cover according to
136 the National Forest Inventory; Forestry Commission 2012) created over the past 160 years on
137 what was previously agricultural land (see Watts *et al.* 2016 for a detailed description of the
138 site-selection protocol). These sites were selected to range in area (0.5 to 30 ha), age (10 to
139 160 years old), amount of woodland within the surrounding landscape (0.4 to 17%
140 broadleaved woodland within a 3 km buffer) and isolation from the next nearest broadleaved
141 woodland (10 to 1570 m). We surveyed woodlands of different character evenly throughout
142 the duration of the field seasons and across the study areas, avoiding any seasonal or spatial
143 bias. The majority of study sites were at least 3 km (a minimum of 1 km) from each other.

144

145 ### Figure 1 approximately here ###

146

147 **2.2. Characterisation of local-level attributes:**

148 We conducted field surveys to characterise the vegetation structure of all woodland patches.
149 Vegetation surveys were conducted along edge-to-interior transects. We established points
150 every 15 m along each transect and used the point-centred quarter method to collect the
151 following data: tree species richness, tree density and tree diameter at breast height (DBH;
152 only trees ≥ 7 cm DBH were measured). Each point also served as the corner of a 10 m \times 10

153 m quadrat which was used to visually assess understory cover (%). Canopy cover (%) was
154 assessed using a sighting tube with an internal crosshair; if the crosshair intersected canopy
155 vegetation, presence of canopy was recorded (Fuentes-Montemayor *et al.* 2013). This was
156 repeated 10 times at 1 m intervals perpendicular to each of the transect-points. We used the
157 OS historic maps collection (EDINA 2013) to determine the age of each woodland patch (i.e.
158 the time period when each woodland patch ‘appeared’ in maps - see Watts *et al.* 2016 for
159 further details). We used digital maps (National Forest Inventory) and GIS software to
160 measure the area and shape (patch perimeter divided by the minimum perimeter possible for a
161 maximally compact patch of the same area; McGarigal & Cushman 2002) of each woodland.
162 We also used Land Cover Map 2007 data (Morton *et al.* 2011) to measure the proportion of
163 woodland edge adjacent to each of the following land cover types: 1) agricultural areas (i.e.
164 arable land or agriculturally improved grassland), 2) semi-natural vegetation (e.g. rough-
165 grassland and scrub), 3) urban areas (e.g. buildings and roads) and 4) water. Improved
166 grassland is distinguished from semi-natural grasslands based on its higher productivity and
167 lack of winter senescence. The proportion of woodland edge adjacent to water approximated
168 zero and was excluded from further analyses.

169

170 **2.3. Characterisation of landscape-level attributes:**

171 We used GIS software to create buffers of different sizes (100, 250, 500, 1000, 1500, 2000,
172 2500 and 3000 m) around each of the 102 woodland patches. These spatial scales were
173 selected to represent distances travelled during foraging trips (i.e. from a roost to a feeding
174 site) by low (e.g. *Plecotus auritus*), intermediate (e.g. *Pipistrellus pygmaeus*) and higher
175 mobility bat species (e.g. *Pipistrellus pipistrellus*; Entwistle, Racey & Speakman 1996;
176 Nicholls & Racey 2006) in agricultural landscapes. Within each buffer category we measured
177 the proportion of land covered by woodlands of the following categories: 1) any woodland

178 (including broadleaved, conifer and mixed woodland), 2) broadleaved woodland (>80%
179 canopy covered by broadleaved trees) and 3) ancient woodland (i.e. native woodland areas
180 which have remained woodland since at least 1600 in England and 1750 in Scotland). We
181 also calculated a measure of inter-patch connectivity between our study sites and surrounding
182 woodland patches of each of the woodland categories outlined above. This measure is based
183 on the Incidence Function Model (Hanski 1994; Moilanen & Hanski 2001; Moilanen &
184 Nieminen 2002). The sum contribution from all surrounding woodland patches of each
185 category was calculated based on their size and distance from the target patch (i.e. each of our
186 study sites), assuming that 5% of dispersers would potentially reach each of the previously
187 defined buffer sizes (i.e. 100, 250, 500, 1000, 1500, 2000, 2500 and 3000 m). Thus the
188 contribution from each surrounding patch declines along a negative exponential dispersal
189 function to the target patch. We also measured the Euclidean distance to the nearest
190 woodland of each category. For woodland categories 1) and 2) we used data from the
191 National Forest Inventory and for 3) we used the Ancient Woodland Inventory (Forestry
192 Commission 2011). In addition, we used Land Cover Map 2007 (Morton *et al.* 2011) data to
193 quantify the proportion of land covered by agricultural areas, semi-natural vegetation, urban
194 areas and water bodies (as described in Section 2.2) in the matrix surrounding each woodland
195 patch within each buffer size.

196

197 **2.4. Bat surveys and sound analyses:**

198 We conducted bat surveys between June and August in 2013 (in Scotland n=31) and 2014
199 (Scotland n=35; England n=36) using ultrasonic detectors which recorded simultaneously at
200 the edge and ‘interior’ (i.e. 50m away from the edge) of each woodland patch. Each site was
201 surveyed once, starting 45 minutes after sunset (to avoid recording bats commuting to feeding
202 sites) and finishing 4 hours later. We recognise that a single visit to each woodland provides

203 only a coarse description of local bat assemblages and that species with low detectability (e.g.
204 *P. auritus*) might have been missed at some sites; we adopted this approach in order to
205 maximise the number/range of sites surveyed and statistical power, and because previous
206 work has indicated that it can successfully identify influential habitat characteristics (e.g.
207 Fuentes-Montemayor *et al.* 2013; Lintott *et al.* 2015). SongMeter SM2BAT+ and SongMeter
208 SM3 detectors (Wildlife Acoustics Inc.) were used in 2013 and 2014 respectively. Whilst the
209 performance of the two detector types might differ, woodlands of different character (e.g.
210 sizes, shapes and degree of isolation) were evenly surveyed throughout the two field seasons
211 and across the study areas to avoid any seasonal or spatial bias. Sound files were recorded in
212 WAC format and later converted to zero crossing for sound analysis using Analoow 4.1
213 (Titley Electronics). Files which required further inspection for species identification were
214 converted to WAV format and analysed using BatSound 4.2 (Pettersson Elektronik AB). We
215 analysed all bat recordings manually to identify bat species and quantify activity (numbers of
216 bat passes, ≥ 2 echolocation calls within 1 s of each other; Walsh & Harris 1996). Of the six
217 bat genera present in the study area *Pipistrellus*, *Plecotus* and *Barbastella* were identified to
218 species; *Myotis*, *Nyctalus* and *Eptesicus* were identified to genus due to similarities in call
219 structure between species. *Eptesicus serotinus* calls can sometimes be difficult to distinguish
220 from *N. leisleri*; when this was the case these two species were grouped into a single category
221 but due to small sample size (<50 bat passes in total) were later excluded from analyses.
222 *Barbastella* and *Eptesicus* only occur in England (Richardson 2000).

223

224 **2.5. Statistical analyses:**

225 All statistical analyses were conducted in R v3.2.2 within Rstudio v0.99.484 using the MASS
226 v7.3-43 and MuMIn v1.15.1 packages (Barton 2015; R Core Team 2015; RStudio Team
227 2015; Venables & Ripley 2002). We ran separate analyses for the two study areas given that:

228 1) the relative abundance of different bat species differs in England and Scotland (Richardson
229 2000); and 2) land-use intensity is higher in England than in Scotland (e.g. higher proportion
230 of farmland tilled annually and lower proportion of land covered by woodland in England
231 than in Scotland; Robinson & Sutherland 2002; Watts 2006).

232

233 **2.5.1. Selection of landscape-level variables:**

234 Given the high degree of collinearity between proportions of land covered by a particular
235 land type across different spatial scales, we conducted Generalised Linear Models (GLMs)
236 using bat activity of each species per site as a response variable (species presence in the case
237 of *P. auritus* due to low activity levels) and the proportion of land covered by each land type
238 category at each spatial scale as the only explanatory variable in each model. Models were
239 fitted using Negative Binomial error distributions (except for *P. auritus* where a Binomial
240 error distribution was used; Crawley 2013). For each land type category (i.e. any woodland,
241 broadleaved woodland, ancient woodland, agricultural, semi-natural, urban and water; for
242 descriptions see Section 2.2) we selected the spatial scale with the largest coefficient to be
243 included in further analyses.

244

245 **2.5.2. Principal components analysis:**

246 We used Principal Components Analysis (PCA) to reduce the number of explanatory
247 variables to be included in statistical models. We adopted this approach because we were
248 primarily interested in the relative influence of local *vs.* landscape-level attributes on different
249 bat species, depending on their mobility, rather than on the effects of specific variables which
250 have already been identified as predictors of bat occurrence/activity in fragmented landscapes
251 (e.g. Fuentes-Montemayor *et al.* 2013). To test hypothesis 1, we conducted a PCA for all
252 local-level attributes (described in Section 2.2 and Table 1) and separate PCAs for landscape-

253 level attributes (described in Sections 2.3, 2.5.1 and Table 1). To test hypothesis 2, we
254 conducted two sets of landscape-level PCAs, one incorporating metrics relating to the amount
255 of surrounding woodland, and the other to woodland inter-patch connectivity (as described in
256 Section 2.3) to account for both amount and configuration of surrounding woodland.
257 Variables in PCAs were scaled to standardise the weights of components. Details of PCA
258 analyses are presented in Appendix S1.

259

260 ### Table 1 approximately here ###

261

262 **2.5.3. Model specification and selection approach:**

263 We conducted GLMs to quantify the relative importance of local vs. landscape-level
264 attributes on activity levels or presence of each species. Data recorded at woodland ‘edge’
265 and ‘interior’ were pooled. Full models incorporated Principal Components (PCs; see Section
266 2.5.2) explaining $\geq 10\%$ variation in the data for either ‘local’, ‘landscape’ or both ‘local and
267 landscape’ level attributes. In addition, date (days since first bat survey of the season) and
268 minimum temperature per night (data obtained from Met Office) were included as covariates
269 in the full models. For Scotland, year (2013 or 2014) was included as a factor; however,
270 preliminary analyses indicated that temperature in Scotland was significantly lower in 2014
271 than in 2013 ($t = -3.90$; coefficient \pm SE = -2.83 ± 0.73 ; $P = <0.001$); therefore only one of
272 these two predictors was included in each model at a time during the model selection
273 procedure. We then followed an information-theoretic model-selection approach based on
274 AICc (Akaike Information Criterion corrected for small sample sizes) to compare all
275 candidate models (i.e. all possible combinations of predictors included in a particular model)
276 to identify the best model (i.e. lowest AICc value) for each response variable (Burnham &
277 Anderson 2002). In the Results section we report Δ AICc (relative to the model with the

278 lowest AICc within a particular set of models) and R^2 values (Likelihood-ratio based pseudo-
279 R^2 ; not adjusted). For illustrative purposes, we also present parameter estimates (\pm SE) and P
280 values obtained from the best models for each response variable; these were not used during
281 the model-selection process.

282

283 3. RESULTS:

284 We surveyed a total of 66 sites in Scotland and 36 sites in England; we detected six
285 species/genera and counted 56,543 bat passes during a total of 816 recording hours. These
286 were soprano pipistrelles (*Pipistrellus pygmaeus*; 48.1% of all bat passes), common
287 pipistrelles (*Pipistrellus pipistrellus*; 41.0%), *Myotis* sp. (2.1%), *Nyctalus* sp. (1.0%), brown
288 long-eared bats (*Plecotus auritus*; 0.2%), Nathusius pipistrelles (*Pipistrellus nathusii*; 0.1%)
289 and barbastelle bats (*Barbastella barbastellus*; <0.1%) (Table 2). *Myotis nattereri* and *M.*
290 *daubentonii* were identified to species level in some cases, but these data were insufficient for
291 species-specific analyses and were pooled to genus level for further analyses. Due to small
292 sample size *P. nathusii* and *B. barbastellus* were excluded from analyses. Models with the
293 lowest AICc for each response variable usually explained between 35% and 66% of data
294 variation, except for *P. auritus* in Scotland ($R^2 = 17\%$) and *Myotis* sp. (R^2 Scotland = 16%; R^2
295 England = 25%) which had lower explanatory power (Figure 2).

296

297 ### Table 2 approximately here ###

298 ### Figure 2 approximately here ###

299

300 3.1. Relative importance of local vs. landscape-level habitat characteristics:

301 In Scotland, models which included only local-level variables were better (i.e. had lower
302 AICc and higher R^2 values) than those including only landscape-level variables for *P. auritus*,

303 *P. pygmaeus* and *P. pipistrellus* (Figure 2a,c,e); the opposite was observed for *Myotis* sp.
304 (Figure 2g). In England, local-only models were better than landscape-only models for *P.*
305 *auritus* and *P. pipistrellus* (Figure 2b,f), but the opposite was observed for *P. pygmaeus*,
306 *Myotis* sp. and *Nyctalus* sp. (Figure 2d,h,i). However, in both study areas the best models (i.e.
307 lowest AICc and highest R^2 values) for most bat species included both local and landscape-
308 level predictors, except for *P. auritus* (Scotland) where the best model included local
309 variables only and *Myotis* sp. (England) where the best model included landscape variables
310 only (Figure 2). Where landscape-level variables were retained in the best models, these
311 usually incorporated woodland inter-patch connectivity metrics, except for *P. pipistrellus*
312 (Scotland) and *Myotis* sp. (Scotland and England), where the proportion of woodland in the
313 landscape provided a better model fit (Table 3).

314

315 #### Table 3 approximately here ####

316

317 **3.2.Effects of specific local and landscape-level habitat characteristics on bats:**

318 In the majority of cases, PCs 1-4 explained $\geq 10\%$ of variation in the data (for both local and
319 landscape-level PCAs) and were therefore included in further analyses; in two cases (for *P.*
320 *pipistrellus* Scotland and *Nyctalus* sp. England) Landscape-PC5 was also included. Total
321 variance explained by these ranged from 66% to 79% in Scotland and 74% to 86% in
322 England (see Appendix S1 for species-specific figures illustrating the most influential PC
323 axes). Since we conducted two sets of landscape-level PCAs (see Section 2.5.2), in the
324 following paragraphs we differentiate between ‘Landscape-PCs (amount)’ (i.e. PCs
325 incorporating metrics relating to the amount of surrounding woodland) and ‘Landscape-PCs
326 (configuration)’ (i.e. PCs incorporating woodland inter-patch connectivity metrics).

327 After accounting for the effects of year, date and temperature, the occurrence of *P. auritus*
328 was negatively related to Local-PC2 in Scotland (although model R^2 was low) and to Local-
329 PC3 in England (Table 3; Appendix S1), mainly indicating a higher probability of occurrence
330 in woodlands bordering agricultural land rather than semi-natural habitat (in Scotland) and a
331 relatively open canopy, low tree densities and low proportion of urban edge (in England).
332 Local-PC4 and Landscape-PC3 (configuration) were also marginally related (positively and
333 negatively, respectively) to the occurrence of *P. auritus* in England, indicating that this
334 species is more likely to occur in woodlands with large amounts of understorey, relatively
335 small trees (i.e. small DBH mean), that primarily border agricultural land and with larger
336 amounts of water within 250m.

337 Activity levels of *P. pygmaeus* in Scotland were related to Local-PC1 (negatively) and
338 Landscape-PC1 and PC2 (configuration) (positively and negatively, respectively), indicating
339 higher activity in older woodlands with lower tree densities, larger trees (large DBH mean)
340 and high degree of variation in tree size (large tree DBH SD) and in woodlands with larger
341 amounts of semi-natural vegetation within 1 km and larger amounts of well-connected
342 broadleaved and ancient woodlands within 1 and 3 km respectively. In England, *P. pygmaeus*
343 activity was negatively related to Local-PC3 and Landscape-PC4 (configuration) indicating
344 higher activity levels in woodlands with a relatively open canopy, lower tree densities,
345 relatively low proportions of urban edge and large amounts of water within 2 km.

346 The activity of *P. pipistrellus* in Scotland was negatively related to Local-PC1 and PC4 and
347 to Landscape-PC5 (amount), indicating higher activity levels in older woodlands with larger
348 trees (large DBH mean), high degree of variation in tree size (large tree DBH SD), lower tree
349 densities, relatively little understorey, a large proportion of urban edge and low amounts of
350 ancient woodland within 500 m. In England, *P. pipistrellus* activity levels were negatively
351 related to Local-PC1 and Landscape-PC3 (configuration), indicating similar patterns to those

352 observed in Scotland at the local scale (i.e. higher activity in older woodlands with larger
353 trees, high degree of variation in tree size and lower tree densities) and also higher activity
354 levels in woodlands farther away from ancient woodland and with a larger proportion of
355 water within 100 m.

356 *Myotis* bats activity levels in Scotland were only significantly related (negatively) to
357 Landscape-PC4 (amount) indicating higher activity in woodlands with a larger proportion of
358 water within 100 m and closer to other broadleaved woodland. In England, a positive
359 relationship with Landscape-PC3 (amount) indicated mainly higher activity in woodlands
360 with a smaller proportion of urban areas within 1500 m, a larger proportion of agricultural
361 land within 1500 m and closer to other woodland, particularly broadleaved.

362 The activity of *Nyctalus* bats in England was negatively related to Local-PC3 and PC1 and
363 Landscape-PC3 and PC1 (configuration), indicating higher activity in older woodlands with
364 larger trees and higher variation in tree size, with a relatively open canopy, lower tree
365 densities and smaller proportion of urban edge, and in woodlands with larger amounts of
366 water within 2500 m and poorly connected woodland in the surrounding landscape
367 (particularly ancient woodland within 250 m and broadleaved woodland within 1000 m).

368

369 **4. DISCUSSION:**

370 In this study we assessed bat occurrence and activity levels in secondary woodland patches
371 created over the last 160 years as part of a large-scale natural experiment in two distinct study
372 landscapes. Our primary interest was to evaluate the relative influence of local (i.e. woodland
373 patch size, age and characteristics/quality) vs. landscape-level attributes (i.e. woodland
374 amount, degree of isolation and nature of the surrounding matrix) on bats (objective i). Very
375 few studies have simultaneously incorporated this full range of factors likely to influence

376 biodiversity in fragmented woodland landscapes (<30% of studies have assessed five or more
377 of these variables in tandem; Humphrey *et al.* 2015). We also compared the role of habitat
378 amount in the surrounding landscape *per se* vs. a combination of both habitat amount and
379 configuration in determining species occurrence and activity levels (objective ii). Finally, we
380 explored the influence of species mobility on the relative importance of these factors for bats
381 (objective iii). Bat species detected during our study ranged from relatively low (e.g. *P.*
382 *auritus*) to high (e.g. *Myotis* sp.) mobility. In accordance with our hypotheses, this trait was
383 important in determining the relative importance of local vs. landscape-level attributes for
384 different bat species (hypothesis 1) and, at the landscape-scale, of woodland amount vs.
385 configuration (hypothesis 2). Although we did not formally assess landscape-moderated
386 effects (e.g. Tschardt *et al.* 2012), our findings support the idea that the effects of habitat
387 fragmentation/restoration processes depend on the landscape context (e.g. vary in landscapes
388 with different proportions of suitable habitat; Andr n 1994). In general, our study sites in
389 England were embedded in more intensively farmed landscapes with a lower proportion of
390 semi-natural vegetation and lower degree of woodland connectivity than our study sites in
391 Scotland (Watts *et al.* 2016), where landscapes tend to be more heterogeneous. These
392 differences were consistently reflected in a higher importance of landscape-level attributes
393 for bats in England than in Scotland.

394

395 **4.1 Bat mobility and the relative importance of local vs. landscape-level attributes:**

396 The occurrence of *P. auritus* was influenced largely by local-level attributes. These are
397 relatively low mobility bats spending most of their time foraging within 500 m from their
398 roosts (Entwistle, Racey & Speakman 1996), mainly in woodlands with large amounts of
399 well-developed understorey (Murphy, Greenaway & Hill 2012). Their higher habitat

400 specificity and lower mobility in comparison to other bat species might explain the relatively
401 low importance of the surrounding landscape; as these bats are often reluctant to fly across
402 open spaces and are restricted to woodland remnants, local habitat characteristics become
403 more important. The best model for *P. auritus* in Scotland included only local variables,
404 whilst the best model for England included local and landscape-level variables (although only
405 local variables were significant predictors), which suggests the relative importance of the
406 surrounding landscape is higher in more intensive agricultural landscapes such as central
407 England.

408 Activity levels of pipistrelle bats (*P. pygmaeus* and *P. pipistrellus*) in both study areas were
409 best explained by models including both local and landscape-level components. These
410 species have intermediate mobility (e.g. the maximum straight-line distance a bat has been
411 recorded from a roost during foraging trips in agricultural landscapes is 2300 m for *P.*
412 *pygmaeus* and 3700 m for *P. pipistrellus*; Nicholls & Racey 2006). Our findings are in
413 accordance with previous studies showing that these species are influenced by both local
414 habitat attributes (e.g. woodland vegetation structure and patch configuration) and the
415 surrounding landscape (e.g. amount of surrounding woodland; Fuentes-Montemayor *et al.*
416 2013). For both pipistrelle species the amount of data variation explained by landscape-level
417 metrics relative to that explained by local-level attributes was higher in England than in
418 Scotland, suggesting that the relative importance of the surrounding landscape is higher
419 where fragmentation is more severe and land-use more intensive. Therefore, both species
420 require a conservation approach involving habitat creation and management at local and
421 landscape scales, but managing the landscape seems particularly important in more
422 homogenous landscapes of intensive agriculture.

423 The activity of *Myotis* bats was mainly influenced by landscape-level attributes. Previous
424 work carried out in woodlands within our study area in Scotland (Fuentes-Montemayor *et al.*

425 2013) suggests that most of the *Myotis* bats recorded during our surveys are likely to be *M.*
426 *nattererii*. Whilst the specific foraging habitat preferences of different *Myotis* species present
427 in the area might differ, these are all relatively high mobility species which can commute
428 long distances from their roosts to their foraging sites (e.g. *M. nattereri* up to 6000 m; Smith
429 & Racey 2008). Although the best models for these species did not retain any significant
430 local-level woodland attributes, previous studies have shown that the activity of *Myotis* bats
431 is influenced by some local habitat characteristics (e.g. higher activity levels in woodlands
432 with higher tree densities), but that the importance of these is lower than that of landscape-
433 level attributes (Fuentes-Montemayor *et al.* 2013). Similar to *Pipistrellus* species, for *Myotis*
434 bats the amount of data variation explained by the landscape relative to that explained by
435 local-level attributes was much higher in England than in Scotland.

436 Activity levels of *Nyctalus* bats were influenced by both local and landscape-level
437 components. These bats display highest activity levels over lakes, rivers and intensive cattle
438 grazing land (Vaughan, Jones & Harris 1997), while radio-tracking studies suggest a
439 preference for pasture and broadleaved woodland areas (Waters, Jones & Furlong 1999;
440 Mackie & Racey 2007). *Nyctalus* bats roost predominantly in tree cavities (Ruczyński &
441 Bogdanowicz 2008) and are highly mobile, often foraging further than 6000 m from their
442 roosts (Mackie & Racey 2007). This high mobility might explain why they were influenced
443 by their surrounding landscape. However, information on these species' habitat associations
444 at the local scale is scarce (although they are selective when choosing tree roosts; Ruczyński
445 & Bogdanowicz 2008) which makes it harder to interpret the observed influence of local-
446 level attributes on these species.

447

448 **4.2 Woodland amount vs. amount and configuration:**

449 For most bat species, a combination of woodland amount and configuration in the
450 surrounding landscape (measured as inter-patch connectivity) appeared to be more important
451 than a measure of only woodland amount in the landscape, except for *P. pipistrellus*
452 (Scotland) and *Myotis* sp. (Scotland and England) for which the opposite was observed. Our
453 findings suggest that lower mobility species respond strongly to habitat configuration (Hanski
454 2015), whereas for higher mobility species the amount of habitat in a landscape outweighs
455 the importance of habitat configuration (Fahrig 2013). Higher mobility species, which are
456 able to fly longer distances across a variety of land cover types (such as *P. pipistrellus*), are
457 likely to find suitable foraging sites as long as they have a sufficient amount of woodland in
458 the landscape, even if it is poorly connected. Despite the amount of woodland being more
459 important than configuration for *P. pipistrellus* in Scotland, these bats often make use of
460 linear landscape elements (e.g. tree lines and hedgerows) which might enhance woodland
461 connectivity in the landscape and allow access to feeding sites over wider areas (Boughey *et al.*
462 *al.* 2011b; Frey-Ehrenbold *et al.* 2013). Interestingly, however, woodland configuration did
463 appear to be more important for *P. pipistrellus* in England than Scotland; this difference
464 might be a result of the landscape in England being more dominated by intensive agricultural
465 land with a lower proportion of woodland cover, making the effects of woodland
466 fragmentation stronger (Andr n 1994).

467 As expected, woodland configuration was more important than woodland amount *per se* for
468 lower mobility species such as *P. auritus* and *P. pygmaeus*. Short-range echolocating bats
469 such as *P. auritus* have been shown to focus foraging activity in well-connected landscapes
470 (Frey-Ehrenbold *et al.* 2013). *Pipistrellus pygmaeus* have also been shown to be influenced
471 by habitat connectivity; for instance, they make use of hedgerows, particularly if these
472 contain trees and are relatively close to woodland areas (Boughey *et al.* 2011b). Woodland
473 connectivity at large spatial scales (e.g. within 3 km) has also been shown to increase the

474 probability of occurrence of *P. pygmaeus* relative to that of other more mobile species such as
475 *P. pipistrellus* (Lintott *et al.* 2016).

476 For *Nyctalus* bats, woodland configuration also appeared more important than just the
477 amount of woodland in the surrounding landscape; this association was negative, however,
478 indicating higher activity levels in poorly connected landscapes. This contrasts with previous
479 studies showing higher activity levels of these species in well-connected landscapes (Frey-
480 Ehrenbold *et al.* 2013), although other studies suggest that these species' preference for
481 specific foraging sites is unlikely to be affected by connectivity (Mackie & Racey 2007).

482

483 **4.3 Bat species-specific responses to local woodland character and surrounding** 484 **landscape:**

485 Most bat species were influenced by local woodland characteristics in a similar way. For
486 instance, *P. pygmaeus*, *P. pipistrellus* and *Nyctalus sp.* all displayed higher activity levels or
487 were more likely to occur in older woodlands with larger trees, higher structural
488 heterogeneity (i.e. larger variability in tree diameter), lower tree densities and a relatively
489 open canopy. Activity of both *Pipistrellus* species was higher in woodlands with relatively
490 little understorey, whilst *P. auritus* was marginally more likely to be present in woodlands
491 with larger amounts of understorey. Our findings are consistent with these species' habitat
492 preferences, which include foraging in woodlands with low tree densities and open
493 understorey (for *P. pygmaeus* and *P. pipistrellus*; Fuentes-Montemayor *et al.* 2013), old
494 forests with native tree species and a dense and diverse understorey (for *P. auritus*; Murphy,
495 Greenaway & Hill 2012) and roosting in woodlands with old, tall and thick trees (for
496 *Nyctalus* bats; Ruczyński & Bogdanowicz 2008). Our findings also suggest that there might
497 be long time lags before newly created woodlands (which are slowly-developing ecosystems)

498 reach successional stages with a habitat structure likely to provide suitable resources for bats.
499 In addition, the activity or probability of occurrence of most species (i.e. *P. auritus*, *P.*
500 *pygmaeus* and *Nyctalus sp.*) was higher in woodlands with a lower proportion of urban edge,
501 although the opposite was observed for *P. pipistrellus*. Previous studies have reported the
502 avoidance of urban areas by *Nyctalus* bats (i.e. *N. leisleri*; Waters, Jones & Furlong 1999),
503 whilst high amounts of urban development around woodlands have been shown to decrease
504 the probability of occurrence of *P. pygmaeus* when compared to *P. pipistrellus* (which seem
505 better adapted to urban environments; Lintott *et al.* 2015, 2016).

506 Bats' responses to landscape-level attributes were species-specific and more variable. In
507 general, bat activity was higher in woodlands with larger amounts of water in the surrounding
508 landscape. These findings are consistent with many bat species' preference for water/riparian
509 areas (e.g. *P. pygmaeus*, Nicholls & Racey 2006; *Myotis* bats, Russ & Montgomery 2002;
510 *Nyctalus* bats, Vaughan, Jones & Harris 1997). Woodland, particularly broadleaved
511 woodland, has been described as one of the most important habitats for many bat species (e.g.
512 Walsh & Harris 1996; Vaughan, Jones & Harris 1997). However, previous studies have
513 reported negative associations between activity levels of some species (e.g. *P. pipistrellus*)
514 and the amount of woodland in the landscape, suggesting that bats might use woodland more
515 intensively in areas where it is scarce (Fuentes-Montemayor *et al.* 2013).

516

517 **4.4. Conservation and management implications:**

518 Bats' responses to local and landscape-level attributes were dependent on species mobility;
519 lower mobility species were most strongly influenced by local habitat quality, whilst the
520 surrounding landscape became increasingly important for higher mobility species (sections
521 3.1 & 4.1). Therefore, actions to enhance bat populations in fragmented landscapes should

522 involve both local and landscape-scale habitat creation and management. We identified
523 specific local and landscape-level habitat characteristics influencing the occurrence and
524 activity levels of different bat species (section 3.2 & 4.3). Our findings provide evidence to
525 suggest that:

- 526 • Local habitat quality should be enhanced, for instance by protecting mature
527 woodlands, managing younger woodlands to promote the development of large trees,
528 relatively low tree densities and open canopies, and maintaining low levels of
529 urbanisation in the immediate surroundings of woodland patches. Increasing
530 understorey cover would benefit gleaning bats but negatively affect aerial hawkers;
531 this highlights the importance of ensuring structural heterogeneity (either within or
532 between patches) to benefit a wider range of species.
- 533 • The surrounding landscape should be improved by increasing the amount of
534 broadleaved woodland in the landscape. However, our findings indicate that whilst
535 this might be an adequate conservation strategy for higher mobility species,
536 improving woodland configuration (for example by spatially targeting woodland
537 planting or creating wooded linear corridors between habitat patches to increase
538 connectivity) seems particularly important for lower mobility species (section 4.2).
539 Maintaining and enhancing the amount and quality of ponds and rivers in the
540 landscape would also benefit most bat species.
- 541 • Landscape-scale management will be particularly important in more homogeneous
542 and intensively farmed landscapes where woodland loss and fragmentation have been
543 more severe.

544 Our findings provide evidence on the local and landscape-level factors likely to enhance
545 biodiversity in newly planted woodlands and are highly relevant to conservation actions
546 focused on the creation and restoration of habitats to reconnect fragmented landscapes. Many

547 of the patterns we observed in bats are broadly similar to those displayed by other taxa (e.g.
548 birds; Dolman *et al.* 2007) and our recommendations are likely to create woodland patches
549 valuable not only for bats, but also for other species. Current work by the authors (e.g. Watts
550 *et al.* 2016) focuses on surveying WrEN sites for a wide range of taxa with different life-
551 history traits whose populations are likely to respond differently to changes in the structure,
552 management and spatial configuration of woodlands and their surrounding landscape, and at
553 different spatial and temporal scales. Using this approach we hope to identify potential
554 differences in the requirements of different taxonomic or functional groups, as well as
555 attempt to draw out general recommendations for conserving woodland biodiversity.

556

557 **4.5. Conclusion:**

558 Conservation strategies to tackle habitat fragmentation require actions at both local and
559 landscape level. However, the relative importance of these actions is still poorly understood
560 and much debated, hampering the translation of ecological theory into practice. Using a
561 large-scale natural experiment (WrEN), we assessed the relative influence of local *vs.*
562 landscape-level attributes on the occurrence and activity levels of a range of bat species of
563 different mobility, and within two distinct study landscapes of varying land-use intensity. Our
564 results indicate that species mobility is an important trait determining the relative importance
565 of local *vs.* landscape-level attributes for different bat species; lower mobility species are
566 most strongly influenced by local habitat quality, whilst the surrounding landscape becomes
567 increasingly important for higher mobility species. In addition, for lower mobility species a
568 combination of amount and configuration of habitat in the surrounding landscape appears
569 more important than habitat amount only, whilst the opposite was observed for higher
570 mobility species. Our results also show that, regardless of species mobility, landscape-level

571 attributes appear more important for bats in more homogeneous and intensively farmed
572 landscapes. Our findings are highly relevant to inform landscape-scale conservation, and
573 suggest that conservation strategies involving habitat creation and restoration activities
574 should take into account the mobility of target species and prioritise landscape-level actions
575 in more homogeneous landscapes where woodland loss and fragmentation have been more
576 severe.

577

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585 For further information and updates on the WrEN project visit <http://wren-project.com/>.

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Table 1. Local and landscape-level attributes measured for all woodland sites and included in PCAs.

ID	Variable type	Variable	Description	Obtained from
1	Local	Tree species richness	Number of tree species.	Field surveys
2	Local	Tree density	Number of trees per hectare.	Field surveys
3	Local	Tree DBH (mean)	Tree diameter at breast height (average value; used as indicator of tree size).	Field surveys
4	Local	Tree DBH (SD)	Tree diameter at breast height (standard deviation; used as indicator of structural heterogeneity).	Field surveys
5	Local	Understorey cover	Proportion of understorey cover in 10x10 m quadrats (average value). Uses Domin scale.	Field surveys
6	Local	Canopy cover	Proportion of vegetation cover in canopy (average value).	Field surveys
7	Local	Patch age	Years since woodland patch appeared on historic maps.	Historic maps
8	Local	Patch area	Area of woodland patch (ha).	Digital maps / GIS
9	Local	Patch shape	Patch perimeter divided by the minimum perimeter possible for a maximally compact patch of the same area.	Digital maps / GIS
10	Local	Agricultural edge %	Proportion of woodland edge adjacent to agricultural land.	Digital maps / GIS
11	Local	Semi-natural edge %	Proportion of woodland edge adjacent to semi-natural habitats.	Digital maps / GIS
12	Local	Urban edge %	Proportion of woodland edge adjacent to urban areas.	Digital maps / GIS
13	Landscape	Woodland (any type) % ^a	Proportion of landscape covered by woodland of any type.	Digital maps / GIS
14	Landscape	Woodland (broadleaved) % ^a	Proportion of landscape covered by broadleaved woodland.	Digital maps / GIS
15	Landscape	Woodland (ancient) % ^{a, b}	Proportion of landscape covered by ancient woodland.	Digital maps / GIS
16	Landscape	Woodland (any type) - Inter-patch connectivity ^a	Connectivity index based on the distance between the target woodland patch and all surrounding woodland patches (of any type) weighted by their area.	Digital maps / GIS
17	Landscape	Woodland (broadleaved) - Inter-patch connectivity ^a	Connectivity index based on the distance between the target woodland patch and all surrounding broadleaved woodland patches weighted by their area.	Digital maps / GIS
18	Landscape	Woodland (ancient) - Inter-patch connectivity ^{a, b}	Connectivity index based on the distance between the target woodland patch and all surrounding ancient woodland patches weighted by their area.	Digital maps / GIS
19	Landscape	Distance to nearest woodland (any type)	Distance (m) to nearest woodland of any type.	Digital maps / GIS
20	Landscape	Distance to nearest woodland (broadleaved)	Distance (m) to nearest broadleaved woodland.	Digital maps / GIS
21	Landscape	Distance to nearest woodland (ancient)	Distance (m) to nearest ancient woodland.	Digital maps / GIS
22	Landscape	Agricultural matrix % ^a	Proportion of landscape covered by agricultural land.	Digital maps / GIS
23	Landscape	Semi-natural matrix % ^a	Proportion of landscape covered by semi-natural habitats.	Digital maps / GIS
24	Landscape	Urban matrix % ^a	Proportion of landscape covered by urban areas.	Digital maps / GIS
25	Landscape	Water matrix % ^a	Proportion of landscape covered by water bodies.	Digital maps / GIS

^a Calculated within buffers of 100, 250, 500, 1000, 1500, 2000, 2500 and 3000 m.

^b Native woodland areas which have remained woodland since at least 1600 in England and 1750 in Scotland; excludes plantations on ancient woodland sites.

Table 2. Bat species/genera detected in woodland sites through acoustic surveys.

	Number of sites detected ^b	Total bat passes	Bat passes (% of total)
<i>Pipistrellus pygmaeus</i>	95	27,183	48.07
<i>Pipistrellus pipistrellus</i>	92	23,014	40.70
<i>Pipistrellus</i> sp. ^a	85	4,363	7.72
<i>Myotis</i> sp.	70	1,171	2.07
<i>Nyctalus</i> sp.	31	582	1.03
<i>Plecotus auritus</i>	26	95	0.17
<i>Pipistrellus nathusii</i>	5	52	0.09
<i>Barbastella barbastellus</i>	3	9	0.02
Other / unidentified	32	74	0.13
Total	102	56,543	100

^a Includes *Pipistrellus* bats (i.e. *P. pygmaeus*, *P. pipistrellus* and *P. nathusii*) which could not be confidently identified to species level.

^b A total of 102 sites surveyed; at least one bat pass was detected in all sites.

Table 3. Parameter estimates (\pm SE) obtained from GLMs using local and landscape-level Principal Components as predictors and bat activity or species presence as response variables. Only top models (i.e. lowest AICc) are shown. All models were fitted using Negative Binomial error distributions (except for *P. auritus* where a Binomial error distribution was used).

	Local	Landscape
<i>P. auritus</i> occurrence		
Scotland	PC2 (-0.59 \pm 0.29) *	-
England	PC3 (-2.42 \pm 1.22) * PC4 (1.60 \pm 0.95) [†]	PC1-configuration (-2.20 \pm 1.30) [†] PC3-configuration (-1.90 \pm 1.19) ^{ns}
<i>P. pygmaeus</i> activity		
Scotland	PC1 (-0.23 \pm 0.09) **	PC1-configuration (+0.20 \pm 0.08) * PC2-configuration (-0.33 \pm 0.12) **
England	PC3 (-0.47 \pm 0.20) *	PC4-configuration (-0.75 \pm 0.22) ***
<i>P. pipistrellus</i> activity		
Scotland	PC1 (-0.66 \pm 0.12) *** PC4 (-0.47 \pm 0.18) **	PC5-amount (-1.05 \pm 0.29) ***
England	PC1 (-0.69 \pm 0.12) ***	PC3-configuration (-0.70 \pm 0.17) ***
<i>Myotis</i> sp. activity		
Scotland	PC1 (-0.19 \pm 0.13) ^{ns}	PC4-amount (-0.58 \pm 0.21) **
England	-	PC3-amount (+0.94 \pm 0.24) ***
<i>Nyctalus</i> sp. activity		
Scotland	NA	NA
England	PC1 (-0.27 \pm 0.11) * PC3 (-0.57 \pm 0.16) ***	PC1-configuration (-0.25 \pm 0.12) * PC3-configuration (-0.64 \pm 0.18) ***

Significance values: *** $p \leq 0.001$, ** $p \leq 0.01$, * $p \leq 0.05$ and [†] $p \leq 0.1$, ^{ns} $p \geq 0.1$. NA: insufficient data for analyses.

Figure 1. Map showing the location of the two study landscapes and woodland sites within them.

Figure 2. Summary of GLMs using ‘local’, ‘landscape’ and ‘local and landscape’ level Principal Components as predictors and bat activity or species presence as response variables. Bars represent model R^2 and dots are $\Delta AICc$ values (relative to model with the lowest AICc for that response variable). Only top models are shown for each category.

Figure 1.



Figure 2.

