

Spatiotemporal changes in flying insect abundance and their functional diversity as a function of distance to natural habitats in a mass flowering crop

Benoît Geslin, Melissa Oddie, Morgane Folschweiller, Gaëlle Legras, Colleen

L. Seymour, F.J.Frank van Veen, Elisa Thébault

▶ To cite this version:

Benoît Geslin, Melissa Oddie, Morgane Folschweiller, Gaëlle Legras, Colleen L. Seymour, et al.. Spatiotemporal changes in flying insect abundance and their functional diversity as a function of distance to natural habitats in a mass flowering crop. Agriculture, Ecosystems & Environment, 2016, 229, pp.21-29. 10.1016/j.agee.2016.05.010 . hal-01327223

HAL Id: hal-01327223 https://hal.sorbonne-universite.fr/hal-01327223v1

Submitted on 6 Jun 2016

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

1 ORIGINAL PAPER

2	Spatiotemporal changes in flying insect abundance and their functional diversity as a
3	function of distance to natural habitats in a mass flowering crop
4	
5	Benoît Geslin ^{a,b} , Melissa Oddie ^c , Morgane Folschweiller ^{b,d} , Gaëlle Legras ^{b,e} , Colleen L.
6	Seymour ^{f,g} , F.J. Frank van Veen ^c and Elisa Thébault ^b
7	
8	^a Institut Méditerranéen de Biodiversité et d'Ecologie marine et continentale (IMBE), Aix
9	Marseille Université, CNRS, IRD, Avignon Université, Faculté des Sciences et Techniques
10	St-Jérôme, Avenue Escadrille Normandie Niémen, Case 421, F-13 397 Marseille Cedex 20,
11	France
12	^b Institute of Ecology and Environmental Sciences - IEES Paris (UMR 7618, UPMC, CNRS,
13	IRD, INRA, UPEC, Paris Diderot), Université Pierre et Marie Curie, 7 quai St Bernard, 75005
14	Paris, France
15	^c Centre of Ecology and Conservation, College of Life and Environmental Sciences,
16	University of Exeter, Penryn Campus, Penryn, Cornwall, TR10 9FE, UK
17	^d Laboratoire de Zoologie, University of Mons, Mons, Belgium
18	^e University of French Polynesia - UMR-241 EIO (UPF, IRD, Ifremer, ILM) Papeete, French
19	Polynesia
20	^f South African National Biodiversity Institute, Kirstenbosch Gardens, PVT Bag X7,
21	Claremont, 7735, South Africa
22	^g DST/NRF Centre of Excellence at the Percy FitzPatrick Institute of African Ornithology,
23	Department of Biological Sciences, University of Cape Town, Rondebosch, 7701
24	Corresponding author: Benoît Geslin
25	E-mail: benoit.geslin@imbe.fr

26

27 Abstract

28 To meet the dietary requirements of a burgeoning human population, the demand for animal-29 dependent crops continues to grow. To meet the demand, intensive farming practices are used. 30 The gains in food production associated with agricultural intensification may be offset by its 31 detrimental effects on pollinator populations through natural habitat fragmentation and 32 pesticide use. Abundance and species richness of pollinators have been found to decrease with 33 increasing distance to natural habitat in agroecosystems, reducing crop yields. A key aspect of 34 crop pollination lies in the diversity of functional traits (functional diversity, FD) of flower-35 visitor communities within crop fields. Higher FD allows improved pollination success 36 through complementarity between flower-visitors' morphology, phenology and behaviour. 37 Many studies reported negative effects of increasing distance to natural habitats on the 38 abundance and richness of flower-visitor communities, but the link between FD and natural 39 habitat isolation is less well understood. Also, a more complete understanding of the 40 functional traits of flower-visitor communities within crops should consider potential 41 variations through time. Differences in resources availability between seasons are important 42 in tropical areas and could modify ecological responses of flower-visitor communities to 43 isolation. In this study, we surveyed the Hymenoptera and Diptera communities within mango 44 orchards of South Africa using pan traps at 100m, 200m and at the maximal distance possible 45 from any natural habitat. We measured the response of insect abundance, wing span and body 46 size as well as functional diversity to habitat isolation during mango flowering (dry season), 47 and during the wet season (after mango fruit harvest). Flying insect abundance decreased with 48 increasing distance to natural habitat during mango flowering, but no effect was detected 49 during the wet season. FD of flying insects declined with increasing distance to natural habitat 50 in both sampling periods. Insects captured during mango flowering were smaller but had

51	higher wing length/body length ratios than those caught during the wet season. This study
52	highlights that mango orchards are more inhospitable for flying insects during mango
53	flowering. This effect might be due to low palatability of mango flowers, or pesticide use in
54	mango fields. In order to maintain a high FD of flower-visiting species, and reduce the
55	detrimental effects of habitat isolation to ultimately ensure better crop pollination, we propose
56	establishment of patches of resource-rich habitats combined with judicious use of pesticides
57	within orchards.
58	
59	Keywords
60	Body size, Flower-visitors, Functional diversity, Isolation from natural habitat, Mangoes,
61	Morphological traits, Wing span

63 **1. Introduction**

64 To ensure nutritional security to a burgeoning human population, the demand for animalpollinated crops is continually increasing (Eilers et al. 2011; Ehrlich and Harte, 2015). To 65 66 meet this demand, agricultural production has intensified in recent decades through 67 conversion of large areas to monocultures, with concomitant loss of natural and semi-natural 68 areas, and increasing use of agrochemicals (Tscharntke et al. 2005; Kennedy et al. 2013). 69 However, given the reported negative effects of agricultural intensification on pollinator 70 populations (Kremen et al. 2002; Vanbergen et al. 2013), the benefits of intensification for 71 animal-pollinated crop yield might be negated by ensuing pollinator loss (Garibaldi et al. 72 2011a; Leonhart et al. 2013; Deguines et al. 2014). By pollinating crops, insects provide a 73 critical ecosystem service estimated to be worth more than €153 billion worldwide (Klein et 74 al. 2003; Gallai et al. 2008; Winfree 2008). The decline of pollinators owing to agricultural 75 intensification therefore raises concerns for food security (Aizen et al. 2008; Garibaldi et al. 76 2011a) and highlights the need for sustainable agriculture that ensures agricultural production 77 whilst conserving biodiversity (Garibaldi et al. 2015, 2016). 78 In agricultural landscapes, natural areas provide habitat for wild insects and constitute sources 79 of flower-visitors for crops (Kennedy et al. 2013; Morandin and Kremen, 2013). Many 80 flower-visitors, such as bees, are central place foragers and usually visit plants close to their 81 nests (Cresswell et al. 2000). As distant resources are more energetically costly to visit than 82 proximate resources, it is predicted that flowers isolated from natural areas will be less 83 frequently visited (Schmid-Hempel et al. 1985, 1986; Steffan-Dewenter and Tscharntke 84 1999). In croplands, species richness of flower-visitors, visitation rates, and pollination 85 services all tend to decline with distance to natural areas (e.g. Ricketts et al. 2008; Carvalheiro 86 et al. 2010; Garibaldi et al. 2011b). Whether or not flower-visitors disperse from natural 87 habitats into the adjacent crop depends on many factors such as floral resource abundance,

floral reward level or type of management (e.g., Kennedy et al. 2013). In this respect, 88 89 pollinator functional traits can be key. Most studies on the effect of distance to natural patches 90 of vegetation have not considered flower-visitor traits or their functional diversity (e.g. 91 Ricketts et al. 2008; Farwig et al. 2009; Carvalheiro et al. 2010, 2012; Garibaldi et al. 2011b; 92 but see Jauker et al. 2009; Williams et al. 2010; Benjamin et al. 2014). Yet traits such as 93 flower-visitor body size are likely to affect pollination success as these traits can influence 94 insect behaviour, foraging distances and pollen deposition (Hoehn et al. 2008). For example, 95 foraging distance is primarily dictated by pollinator body size, and small flower-visitors will 96 forage closer to their nests (Araújo et al. 2004; Greenleaf et al. 2007; Benjamin et al. 2014). 97 Moreover, several studies have shown that functional diversity (FD) of flower-visitor 98 communities enhances pollination by providing complementary pollination services, for 99 example through niche partitioning of resource use in time and space (Fontaine et al. 2006; 100 Hoehn et al. 2008; Albrecht et al. 2012; Fründ et al. 2013). Studying the effects of isolation 101 from natural areas on traits and FD of flower-visitors can thus shed light on the factors which 102 affect both crop pollination and the persistence of diverse flower-visitor communities in agro-103 ecosystems, informing effective land management strategies (Williams et al. 2010; Benjamin 104 et al. 2014).

105 Temporal dynamics of agricultural landscapes are also important. Cultivated areas exhibit 106 particularly large temporal variations in floral resources which could affect flower-visitor 107 communities (Westphal et al. 2003). Mass flowering crops such as mangoes (Mangifera 108 indica L., Anacardiaceae) constitute a super-abundant floral resource during a short period of 109 time, representing a resource pulse for flower-visitors (Orford et al. 2015). The relationship 110 between increasing distance to natural habitat and flower-visitor communities has been mostly 111 explored during crop mass flowering (e.g. Holzschuh et al. 2011) but negative effects of 112 distance on flower-visitors are likely to be stronger when the crop is not flowering, because

113 only flowering weeds then provide resources in cultivated fields. Seasonal variation of wild 114 floral resources and flower-visitor communities also contribute to temporal variation of 115 agroecosystems. Spatiotemporal turnover in flower-visitor assemblages varies between and 116 within years in temperate and tropical ecosystems (Oertli et al. 2005; Rollin et al. 2015; 117 Samnegård et al. 2015), with temporal changes in abundance or even functional traits of 118 flower-visitors varying because of food and nesting requirements (Tylianakis et al. 2005; 119 Rollin et al. 2015). Flower-visitor assemblages might respond differently to agricultural 120 perturbations in different seasons (Samnegård et al. 2015). For example, in tropical 121 environments, resources are more scarce and patchily-distributed during the dry season 122 relative to the wet season. Greater mobility may therefore be advantageous to crop flower-123 visitors during the dry season (see Samnegård et al. 2015). 124 To explore those questions, we studied the response of potential flower-visitors (hereafter 125 referred to as "flying insects") to increasing distance from natural vegetation in mango 126 orchards in north-eastern South-Africa in two different seasons. Mango is one of the most 127 important tropical fruits produced in the world, and is economically important for income and 128 employment-creation in the region (FAO, 2010). Research conducted in our study area in the 129 past has found clear effects of distance on pollination (Carvalheiro et al. 2010, 2012), pest 130 control (Henri et al. 2015) and bird assemblages (Ehlers Smith et al. 2015). 131 We used pan-traps to survey flying insects during mango flowering in the dry season (winter), 132 and during the wet season (summer, when mango is not flowering), along transects of 133 increasing distance to natural habitat. We hypothesised that the number of flying insects 134 caught in traps would decrease with distance to natural vegetation. If isolation from natural 135 areas is the only cause of the decrease in flying insect abundance, the effect should be 136 consistent between seasons. We also measured wing span and body length of each flying 137 insect and investigated distance effects on flower-visitor functional-trait diversity. We

- 138 hypothesised a decline of the FD with increasing distance from natural areas in both seasons,
- 139 since species loss caused by isolation is likely to result in loss of trait diversity. Finally, given
- 140 links between size and flight abilities, we predicted that average insect body size would be
- 141 larger with increasing distance to natural area.

142 **2. Materials and Methods**

143 2.1 Study site

144 Our study site was situated in the Kruger to Canyons Biosphere Region, Limpopo Province,

- 145 South Africa. This region includes agricultural areas and also two large protected areas
- 146 (Kruger National Park and Blyde River Canyon reserve; 24° 24'S 30°50'E). The entire area
- 147 retains more than 50% of intact vegetation unaffected by anthropogenic perturbations
- 148 (Coetzer et al. 2013).

149 Other than tourism, agriculture is an important source of employment in the area. Although in 150 some areas, there are small subsistence farms, in our study area, the farms are composed of 151 sets of large blocks of monoculture stands of mango, citrus or avocado. Mango farms 152 consisted of multiple plots of 0.66 to 3 ha (mean \pm sd= 1.39 \pm 0.87) separated by windbreaks 153 of tall *Casuarina sp.* trees. All the study farms provided similar water and nutrient supply and 154 made intensive use of herbicides (e.g., glycophosphates) and insecticides (neonicotinoids or 155 organophosphates). The sets of blocks of mango plantation are bordered by patches of natural 156 vegetation that vary in size, but are usually at least 250 m wide. These patches of natural 157 vegetation have not been ploughed, and differ from natural vegetation in neighbouring Kruger 158 National Park in that livestock have replaced wild herbivores. The vegetation in the area is 159 Granite Lowveld (Mucina and Rutherford, 2006), a savannah in which the woody component 160 is dominated by Acacia (Senegalia) nigrescens and Sclerocarya birrea, with a herbaceous 161 layer that includes Digitaria eriantha, Panicum maximum and Aristida congesta. 162 The primary flower-visitors to mango in this area include Diptera (mostly Calliphoridae and 163 Empididae families), and Hymenoptera (Apis mellifera, Ceratina spp. bees and species of 164 Campenotus, Monomorium and Tetramorium ants; Carvalheiro et al. 2010, 2012). Mango is 165 self-compatible, but only to a small extent: a bagging experiment found that there was 21% 166 fruit set for inflorescences from which pollinators had been excluded (100% of fruit set in

open inflorescences see Carvalheiro et al. 2010). Mango flowers do produce nectar, but only asmall amount of pollen (Eardley et al. 2007).

169 The first surveys were performed during mango flowering (July-August 2013, dry season 170 hereafter), with maximum temperatures averaging around 26°C and minima of 9°C. The lack 171 of rain during this time means that few flowering species are in bloom for flower-visiting 172 insects. We selected five mango farms separated from each other by between three to thirteen 173 kilometres. Within each farm, we established two transects perpendicular to the edge between 174 mango and natural vegetation. Along each transect, we sampled insects at 5 sampling points, 175 at 50m from the edge with mango in the natural vegetation, at the field edge (0m) and at 100, 176 200m and the maximal distance possible from any natural habitat within mango orchards. 177 Distance to natural vegetation was measured as the distance from each set of traps within the 178 blocks of mango to the border of the nearest patch of natural vegetation. During April-May 179 2014, we re-sampled insects in two transects in each of the three most accessible farms. April-180 May coincides with the end of the wet season, when maximum and minimum temperatures 181 average 29°C and 12°C, respectively, and floral resources are fairly abundant, having 182 flourished during the rains.

At each sampling point, three coloured pan traps (radius=7.25 cm, depth=5 cm) painted with UV-reflecting paints (yellow, blue and white) were placed and filled with 400 ml of water and a drop of detergent (surfactant). Although pan trap sampling is known to under-sample some insect groups like large bees (*Bombus, Colletes*), this method is nevertheless considered the most efficient method to sample flying flower-visitors (Roulston et al. 2007; Westphal et al. 2008, Nielsen et al. 2011). It allows sampling of a large number of points simultaneously and uniformly, obtaining standardized estimates of flower visitor abundance and diversity.

190

191 2.2 Data collection

192 In dry and wet seasons, we performed three sampling sessions in each transect (one every two 193 weeks) where traps were set out in the field for 24 hours before being collected (N = 720 pan 194 traps). Traps were set on warm sunny days (minimum of 15°C, no wind and clear sky). Once 195 collected, insects were stored in 70% ethanol before being rinsed, identified at the order level 196 and measured. We focused on potential pollinators including Hymenoptera, Lepidoptera and 197 Diptera, which are groups that are considered as mango pollinators (Sung et al. 2006). 198 Morphometric measures of insects were also taken, *i.e.* wing length (distance between the 199 wing tips) and body length using digital callipers (Digit-Cal MK IV 599-571, Brown and 200 Sharpe, USA, 0.01 mm). 201 Finally, we assessed the number of mango flowers and wild flowers in a perimeter of 15 202 meters around each sampling point. Within mango orchards, the herbaceous cover was 203 dominated by non-native weeds (e.g. Tridax procumbens (L.) and Bidens pilosa (L.), both 204 Asteraceae). Since mangoes exhibit "big bang" flowering with a massive number of flowers 205 per tree, we estimated the number of mango flowers by counting the number of flowers on 206 three different panicles and multiplying the average by the number of panicles for each tree (a 207 mature tree has 600-1000 panicles; Manning, 1995; Sung et al. 2006).

208

209 2.3 Functional Diversity

Our dataset was comprised of individual insects, and not species, so we calculated functional diversity of the individuals found in our samples. We used order, body length, wing span and the ratio of body length to wing length as traits. Order was represented as a binary variable of one or zero for each of three categories (Diptera, Hymenoptera or Lepidoptera), and therefore order was represented by three "traits". We therefore down-weighted order to one third the value of the other traits to avoid biasing the analysis. We calculated the FDis (Functional Dispersion, Laliberté and Legendre 2010) measure of functional diversity, using the FD package in R (Laliberté et al. 2010). In this case, FDis is the mean distance of individuals to the centroid calculated for all individuals in multidimensional trait space. FDis is not affected by species richness (in our case, abundance), and is able to handle missing values and qualitative as well as quantitative traits (Laliberté and Legendre 2010).

222

223 2.4 Data analysis

We assessed how functional diversity and the value of each of the traits used to calculate functional diversity (i.e. body length, wing length, and ratio of wing length over body length) changed with season and habitat. Although functional diversity is a measure of the variation in these traits, changes in the average values of traits yields insights into how the traits themselves are affected by habitat and season. These two analyses therefore produce complementary information.

230 We assessed the effects of sampling season (dry or wet season), habitat (natural vegetation or 231 mango fields) and distance to natural habitat, as well as the interactions between season and 232 distance or habitat effects, on flower abundance, abundance and traits of flying insects and on 233 flying insect functional diversity. To do so, for each response variable, we performed a model 234 averaging approach which allows comparison of all possible submodels from a global model 235 containing all the predictors of interest, in order to identify the best set of models describing 236 the observed variation (Grueber et al. 2011). With this approach, it is possible to account for 237 model uncertainty and to assess the relative importance of various predictor variables. 238 To analyse patterns in flower abundance, traits of flying insects and functional diversity FDis, 239 we used linear mixed-effect models on log-transformed response variables (except for FDis) 240 using distance to natural vegetation, sampling season, type of habitat, interactions between

distance and sampling season, and interactions between type of habitat and sampling season
as fixed variables. Farm, transect nested within farm and date of sampling were included as
random effects to avoid pseudo-replication (Crawley, 2007). Additionally, in models with
insect traits as a response variable, we included insect order as a random effect to account for
differences in trait values between orders.

To assess variations in abundance of flying insects, we carried out generalized linear mixedeffect models (GLMER) with a Poisson distribution and a log-link using the same fixed and random variables with the addition of flower abundance as a fixed variable. We included flower abundance as an explanatory variable because pan-traps are known to be less attractive to flower-visitors in resource-rich sites relative to resource-poor sites (Wilson et al. 2008). The GLMERs were corrected for overdispersion by including observation-level random factors (Harrison 2014).

253 For each response variable, we selected the set of models according to the Akaike information 254 criterion (AIC). More precisely, we kept the models that were within the range of four highest 255 AIC units and we used these "top models" to estimate the mean effects and confidence 256 intervals of each predictor variable using model averaging (natural average method, Grueber 257 et al. 2011). We also measured the relative importance of each predictor variable which is 258 expressed as the relative sum of the Akaike weights across all top models in which the 259 variable appears (relative importance is equal to 1 for variables present in all top models). Additionally, we calculated the marginal R^2 values (fixed effects) and conditional R^2 values 260 (R²) both fixed and random effects) to assess the amount of variance explained by the best 261 262 model (i.e. with highest AIC; Johnson 2014). All analyses were performed using the R 263 statistical interface (v 3.1.0, R Development Core Team, 2014); mixed models were 264 conducted using the package lme4 (Bates et al. 2014) and model comparison and averaging were performed with the functions dredge, get.models and model.avg (package MuMIn; 265

Barton 2016). Model predictor variables were also centered and standardized with the

267 function standardize (package arm; Gelman and Su, 2015) to facilitate comparison between

268 the relative strengths of parameter estimates (Grueber et al. 2011). R^2 values were calculated

with the function r.squared GLMM (package MuMIn; Barton 2016).

270

271 **3. Results**

272 3.1. Abundance of flowers

273 Distance to natural vegetation, as well as habitat type, sampling season and their interactions 274 (habitat×season and distance×season), were included in all top models describing the response 275 of flower abundance (Table 1). As expected, the abundance of flowers was greater in mango 276 fields in the dry than in the wet season, whereas we observed more flowers in the nearby 277 natural vegetation during the wet season (see parameter estimate and confidence interval of 278 the interaction between sampling season and habitat type in Table 1, Figure 1A). Flower 279 abundance was far higher in the mango fields than in the natural vegetation during the dry 280 season (Figure 1A), given that we did not observe any flowers in the neighbouring natural 281 vegetation in any of the sampling sessions in the five farms over the dry season. The 282 abundance of flowers also increased with increasing distance from natural vegetation (Figure 283 1A). The interaction between distance and season had only 35% relative importance to other predictor variables and the confidence interval for this parameter estimate included 0. Thus 284 285 there was little evidence that the effect of distance from natural vegetation on flower 286 abundance differed between seasons.

287

288 3.2 Abundance of flying insects

Distance to natural vegetation, sampling season, and the interaction between these twovariables, were included in all top models describing the response of abundance of flying

291 insects (Table 1). The abundance of flying insects was far higher during the wet season than 292 the dry season, which was expected since we captured 798 insects in the five farms during the 293 dry season but captured 1719 insects in only three of these farms during the wet season. 294 Abundance of flying insects declined with increasing distance from natural vegetation, but 295 only during the dry season (Table 1, Figure 1B). The difference in abundance of flying insects 296 between the two sampling periods was also more pronounced in mango fields (see parameter 297 estimate and confidence interval of the interaction between sampling season and habitat type 298 in Table 1), but there was weaker evidence for this effect (55% relative importance). Lastly, 299 there was very little evidence of an effect of flower abundance on the abundance of flying 300 insects (39% relative importance to distance and season, and the confidence interval for this parameter estimate included 0). 301

302

303 3.3 Functional diversity

304 Distance to natural vegetation and sampling season were included in all top models describing 305 the response of functional diversity (Table 1). Functional diversity of flying insects was 306 higher in the wet season than during the dry season and FD declined with increasing distance 307 from natural vegetation in both seasons (Figure 1C). Functional diversity was also higher in 308 nearby natural vegetation than in mango fields (79% relative importance to distance and 309 season, Table 1). The interaction between distance and season had only 23% relative 310 importance and the confidence interval for this parameter estimate included 0. Thus, there was 311 little evidence that the negative effect of distance from natural vegetation on functional 312 diversity differed between seasons.

313

314 3. 4 Insect traits

315 Only sampling season was included in all top models describing the responses of body length, 316 wing span and ratio of wing span to body length of flying insects (Table 2). Both body length 317 and wing span of flying insects were on average smaller during the dry than during the wet 318 season, whereas the ratio of wing span to body length was larger during the dry season 319 (Figure 2). Body length and wing span of flying insects also differed between habitat types, 320 insects being on average larger in natural vegetation than in mango fields (Figure 2, Table 2). 321 Although body length and wing span of flying insects tended to decrease with increasing 322 distance to natural vegetation during the dry season (Figure 2), this effect was not significant 323 (the confidence interval for the corresponding estimate included 0). There was very little 324 evidence of any effect of habitat type and distance to natural vegetation on the ratio of wing span to body length of flying insects (Table 2). In addition, the marginal R^2 of the models 325 326 were very low (about 0.02, see Table 2), indicating that on average variance in insect traits 327 explained by effects of distance, habitat type and sampling season was only 2% whereas 328 random factors (which represented insect order, as well as sample position and date) 329 explained between 40 and 50% of the variation. 330

332 **4. Discussion**

333 This study highlights that mango orchards are more inhospitable for flying insects during the 334 dry (mango flowering) than during the wet season, despite the abundance of mango flowers 335 available in the orchards. We found declines in insect abundance with increasing distance to 336 natural areas during the dry season whereas no pattern was found during the wet season. The 337 functional diversity (FD) of flying insects declined with increasing distance to natural habitat 338 in both sampling periods, however, and insects were on average smaller in mango orchards 339 than in natural vegetation. Taken together, our results suggest impacts on crop yield, given 340 that as a rule, rate of flower visitation (Vázquez et al. 2005, Garibaldi et al. 2013) and 341 functional diversity (Fontaine et al. 2006, Hoehn et al. 2008, Albrecht et al. 2012, Fründ et al. 342 2013) increase crop fruit set. Creating patches of habitat rich in native floral resources that 343 flower before and during mango flowering within orchards could mitigate the detrimental 344 effects of isolation from natural habitats and increase mango production (Carvalheiro et al. 345 2012).

346

347 **4.1 Seasonal variation in flying insect abundance**

348 Agricultural landscapes show strong temporal variation in the floral resource offer. Mass 349 flowering crops, during their flowering season, are expected to exert a strong magnet effect on 350 pollinators from adjacent areas (Blitzer et al. 2012). We did not find this effect, however, and 351 this was somewhat surprising, given that there are few other floral resources in the landscape 352 at this time. It is possible that this pulse of floral resources is too short-lived to enable 353 invertebrate populations to respond to this sudden availability in floral resources. This finding 354 highlights the importance of considering seasonal variation in the value of agricultural patches 355 for pollinators.

356 The decline in insect abundance with increasing distance to natural vegetation that we 357 observed during the dry season is a pattern that has been observed in numerous studies in 358 agricultural systems (e.g. Ricketts et al. 2008; Garibaldi et al. 2011b). This pattern appears 359 stronger in tropical than temperate regions (Ricketts et al. 2008) and has been particularly 360 well documented by Carvalheiro et al. (2010) in our study system. This latter study notably 361 showed that even in a biodiversity rich area, with a high proportion of intact natural habitat, 362 the abundance of flying insects was impacted by isolation from natural habitats, which in turn 363 impacted crop fruit set. Because several flying insects such as bees are central place foragers 364 (*i.e.* individuals that return to their nests after foraging (for nectar and/or pollen; Williams and 365 Kremen, 2007), this decline in insect abundance with increasing distance to natural area is 366 commonly attributed to the distance from nesting sites and foraging resources available in 367 natural habitats (e.g. Ricketts et al. 2008; Farwig et al. 2009; Williams et al. 2010; Garibaldi 368 et al. 2011b; Samnegård et al. 2015).

369 But intriguingly, in complete contrast to the dry season when mango is flowering, we did not 370 observe any effect of distance to natural area on insect abundance during the wet season. 371 These contrasting results might be explained by the seasonal dynamics of wild floral 372 resources and pollinator communities. Flying insects were more abundant and had higher 373 functional diversity during the wet than dry season, which is expected since the wet season 374 coincides with spring/summer, when more floral resources are available and temperatures are 375 higher. The link between plant richness and abundance and flower-visitor richness and 376 abundance has been well documented in the scientific literature for croplands (e.g., Holzschuh 377 et al., 2007; Kennedy et al., 2013). The higher diversity of native flowering plant during the 378 wet season, and the increasing availability of weed flowers with increasing distance from 379 natural vegetation within mango orchards might explain the absence of a relationship between 380 isolation from natural habitat and insect abundance. It is also worth noting that mango flowers yield only small amounts of pollen, although they do offer some nectar (Eardley et al. 2007), and so are not very attractive to many insects (Free and Willams 1976). Thus, the lack of attractiveness of mango mass flowering might explain the strong negative effect of isolation from natural habitat we observed during the dry season. This idea is supported by an earlier study in which small planted patches of native flowering plants within mango orchards were associated with an increased abundance of flower-visitors to mango flowers, which mitigated to some extent the effects of isolation (Carvalheiro et al. 2012).

388 In addition to the limitations in floral resources, alternative explanations exist that might 389 explain seasonal variation in distance effects on flying insects in the crop. Pesticides are used 390 in mango fields, even during mango flowering. Many studies have reported on the negative 391 effect of pesticides on flower-visitors (Henry et al. 2012; Goulson et al. 2015; Stanley et al. 392 2015), and Carvalheiro et al. (2012) also previously highlighted the detrimental effect of 393 pesticide use on flying insect along gradients of increasing distance to natural area in our 394 study system. Varying pesticide applications might thus also explain the different effects of 395 distance to natural area on insect abundance between the two sampling seasons. 396 Pan-traps are widely considered to provide accurate surveys of bee assemblages (Roulston et 397 al. 2007, Westphal et al. 2008, Geslin et al. 2016), but are also prone to some bias. Pan-traps 398 under-sample large insects and their attractiveness can vary with the amount of flowers 399 locally present (Wilson et al. 2008, Popic et al. 2013). The relative attractiveness of pan-traps 400 decreases as floral resources increase because pan traps compete with flowers for flying insect 401 attention. However, it is unlikely that our results are an artefact of pan-trapping. We included 402 floral abundance as an explanatory variable in our statistical models, and this variable did not 403 have a significant effect on insect abundance. Secondly, in both seasons, more flowers were 404 observed in mango orchards compared to natural habitats and the decline in insect abundance

with increasing distance to natural habitat was observed uniquely during the dry season, withthe same capture method.

407

408 4.2 Functional diversity

409 Although flying insect abundance decreased with increasing distance from natural habitat 410 during the dry season only, declines in functional diversity with increasing isolation were 411 consistent in the two sampling seasons. A recent study highlighted that farmed landscapes 412 were detrimental to the maintenance of functional-trait diversity of flying insects (Forrest et 413 al. 2015), and our findings support this view.

414 FD is increasingly recognized as being more important than species diversity to ecosystem

415 functioning (Gagic et al. 2015) and has been suggested to be the major component of diversity

416 (Tilman et al. 1997; Hulot et al. 2000; Hoehn et al. 2008). The main reason for the importance

417 of FD lies in the complementarity of pollination services through niche partitioning of

418 resource use in time and space (Fontaine et al. 2006; Hoehn et al. 2008; Albrecht et al. 2012;

419 Fründ et al. 2013). The value of greater FD to pollination services has been demonstrated both

420 experimentally (Fontaine et al. 2006) and empirically (Hoehn et al. 2008), and a recent

421 example in apple orchards highlighted that fruit and seed set was mainly mediated by bee FD

422 (Martins et al. 2015, but see Garibaldi et al. 2015).

423 The loss of FD with increasing isolation can reduce the efficiency of pollination (Forrest et al.

424 2015; Wood et al. 2015) reducing yield for trees at the centre of mango orchards. Carvalheiro

425 et al. (2010, 2012), demonstrated that agricultural production and proximity to natural habitat

- 426 are positively correlated in the mango orchards we studied. Yet, easily implementable
- 427 management practices might ameliorate, to some extent, the negative effects of isolation.
- 428 Creating patches of flowering rich habitat has been shown to increase crop yield in mango
- 429 orchards (Carvalheiro et al. 2012). FD of flying insects has been shown to improve with only

modest enhancements of floral diversity (Orford et al. 2016). Given that isolation from natural
habitat together with loss of flower-visitor abundance and diversity are now well established
as critical drivers of limitation in crop yield (Garibaldi et al. 2016), we encourage the
establishment of those simple management practices to enhance food production.

434

435 **4.3 Traits of flying insects**

436 In contrast to functional diversity, there was weak evidence of distance effects on average 437 insect trait values, and the traits were only weakly affected by habitat type. Although a large 438 part of the variance in insect traits was related to size differences between orders, sampling 439 season had also a significant effect on flying insect average body length and wing span. 440 During the dry season in tropical environments, resources are scarcer and more patchily-441 distributed compared to the wet season (Samnegård et al. 2015). Because the foraging range 442 of flying insects is positively correlated with their body size (Araújo et al. 2004; Greenleaf et 443 al. 2007), we initially hypothesized that we would trap larger flying insect during the dry 444 season. However, our results show that insects were smaller during the dry season. This 445 finding is more consistent with the findings of Wray et al. (2014), who found large-bodied 446 species to be favoured by increasing availability of floral resources (although this was in a 447 landscape context of urbanisation). Indeed, large-bodied species may have larger resource 448 needs and smaller population sizes (Kremen and McGonigle, 2015), which could 449 disadvantage them during the dry season. We nevertheless did detect a greater ratio of wing to 450 body length during the dry season, which might indicate greater mobility of insects relative to 451 their size than during the wet season. Insect size differences between seasons might also arise 452 from differences in climatic conditions such as temperature. Some studies reported that higher 453 wing to thorax size ratios could be selectively advantageous at lower temperatures (e.g., 454 Azevedo et al. 1998), which is in agreement with what we found during the dry/winter season. 455 Species turnover between seasons likely explains the difference in insect sizes between 456 seasons. The distribution of insect body size/tongue size has been found to closely match the 457 distribution of flower corolla tube depth (Agosta and Janzen 2005, Stang et al. 2006; 2009) 458 and to determine the abundance of interactions in pollinator webs (Stang et al. 2006, Geslin et 459 al. 2013). The distribution of the size of insects might be due to the predominance of flowers 460 with short corollas during the dry season because mango flowers, as well as the dominant 461 non-native weeds T. procumbens and B. pilosa have short corollas. This is supported by the 462 fact that the few large individuals of flying insect we found during the dry season were 463 trapped in natural habitat where plants such as Grewia spp., have flowers that present a more 464 tubular and deep corolla structure. Finally, we observed larger insects, in both seasons, in the 465 natural vegetation compared to mango orchards. Some studies have found large-bodied flying 466 insects to be more prone to extinction in response to land use intensity and habitat loss 467 (Larsen et al. 2005, Bartomeus et al. 2013; Rader et al 2014), whereas others have found non-468 significant effects (Williams et al. 2010), and several authors found small insects to be more 469 negatively affected by isolation and habitat loss than larger ones (Bommarco et al. 2010; 470 Öckinger et al. 2010; Jauker et al. 2013). Here, large-bodied species seemed more sensitive to 471 agricultural perturbations than small bodied ones and were thus less abundant in mango 472 orchards. Moreover, flowers within mango orchards may be less attractive to large bodied 473 insects because of the nectar and pollen offerings of mango flowers relative to wild flower 474 species.

475

476 **5. Conclusion**

477 Our study provides new evidence of the detrimental effects of isolation from natural habitats,
478 even in a biodiversity rich area, on abundance and functional diversity of flying insects in
479 different seasons. Although these deleterious effects were season-dependent for insect

480 abundance (i.e. negative effects of distance from natural vegetation only during crop 481 flowering), functional diversity declined consistently with increasing distance from natural 482 habitat in both dry and wet seasons. This study thus sheds a light on the importance of 483 providing patches of resource-rich habitat within orchards over the year to mitigate the 484 detrimental effects of isolation and maintain functionally-diverse insect communities. Such 485 management practices are easily achievable in mango orchards and should be encouraged. 486 Finally, this study also underlines the temporal variability within a year of patch use by flying 487 insect communities within an agricultural landscape. Agricultural landscapes and mechanisms 488 that structure flying insect communities should thus be studied at various spatial and temporal 489 scales. We encourage new studies regarding links between morphological traits of insects and 490 agricultural intensification, contributing to a synthesis that can untangle the relative 491 importance of temporal and spatial contexts, crop and management practices. 492

493 Acknowledgments

- 494 We are grateful to Floriane Flacher, Lavhelesani Simba, Lyndre Nel, Anne-Sophie Bonnet-
- 495 Lebrun, Pierre Quévreux, Thomas Aubier and Kejun Zou for their help with data collection.
- 496 BG thanks Lucas A. Garibaldi and Boris Leroy for useful discussions. This work was
- 497 supported by the European Commission Marie Curie International Research Staff Exchange
- 498 Scheme (IRSES) ('NETWORK' project, Grant agreement: PIRSES-GA-2012-318929). .

499 **References**

- 500 Agosta, S.J., Janzen, D.H., 2005. Body size distributions of large Costa Rican dry forest
- 501 moths and the underlying relationship between plant and pollinator morphology. Oikos

502 108, 183–193. doi:10.1111/j.0030-1299.2005.13504.x

- 503 Aizen, M. A., Garibaldi, L.A., Cunningham, S. A., Klein, A.M., 2008. Long-term global
- 504 trends in crop yield and production reveal no current pollination shortage but increasing
- 505 pollinator dependency. Curr. Biol. 18, 1572–5. doi:10.1016/j.cub.2008.08.066
- 506 Albrecht, M., Schmid, B., Hautier, Y., Müller, C.B., 2012. Diverse pollinator communities
- 507 enhance plant reproductive success. Proc. Biol. Sci. 279, 4845–52.
- 508 doi:10.1098/rspb.2012.1621
- 509 Araújo, E., Costa, M., 2004. Body size and flight distance in stingless bees (Hymenoptera:
- 510 Meliponini): inference of flight range and possible ecological implications. Brazilian J.
 511 Biol. 64, 563–568.
- 512 Azevedo, Ricardo BR, Avis C. James, Jennie McCabe, and L. Partridge. 1998. Latitudinal
- 513 variation of wing: thorax size ratio and wing-aspect ratio in Drosophila melanogaster.
- 514 Evolution 5: 1353-1362.
- 515 Bartomeus, I., Park, M.G., Gibbs, J., Danforth, B.N., Lakso, A.N., Winfree, R., 2013.
- 516 Biodiversity ensures plant-pollinator phenological synchrony against climate change.
- 517 Ecol. Lett. 16, 1331–8. doi:10.1111/ele.12170
- 518 Barton K., 2016. MuMIn: Multi-Model Inference. R package version 1.15.6. https://CRAN.R-
- 519 project.org/package=MuMIn
- 520 Bates, D., M. Maechler, B. Bolker, and S. Walker., 2014. lme4:Linear mixed-effects models
- 521 using Eigen and S4. R package version 1.1-5.

Benjamin, F., R. Reilly, J., Winfree, R., 2014. Pollinator body size mediates the scale at
which land use drives crop pollination services. J. Appl. Ecol. 51, 440–449.

524 doi:10.1111/1365-2664.12198

- 525 Blitzer, E.J., Dormann, C.F., Holzschuh, A., Klein, A.M., Rand, T.A., Tscharntke, T., 2012.
- 526 Spillover of functionally important organisms between managed and natural habitats.
- 527 Agric. Ecosyst. Environ. 146, 34–43. doi:10.1016/j.agee.2011.09.005
- 528 Bommarco, R., Biesmeijer, J.C., Meyer, B., Potts, S.G., Pöyry, J., Roberts, S.P.M., Steffan-
- 529 Dewenter, I., Ockinger, E., 2010. Dispersal capacity and diet breadth modify the
- response of wild bees to habitat loss. Proc. Biol. Sci. 277, 2075–2082.
- 531 doi:10.1098/rspb.2009.2221
- 532 Carvalheiro, L.G., Seymour, C.L., Veldtman, R., Nicolson, S.W., 2010. Pollination services
- decline with distance from natural habitat even in biodiversity-rich areas. J. Appl. Ecol.
 47, 810–820. doi:10.1111/j.1365-2664.2010.01829.x
- 535 Carvalheiro, L.G., Seymour, C.L., Nicolson, S.W., Veldtman, R., 2012. Creating patches of
- 536 native flowers facilitates crop pollination in large agricultural fields: mango as a case
- 537 study. J. Appl. Ecol. 49, 1373–1383. doi:10.1111/j.1365-2664.2012.02217.x
- 538 Coetzer, K.L., Erasmus, B.F.N., Witkowski, E.T.F., Reyers, B., 2013. The race for space:
- 539 Tracking land-cover transformation in a socio-ecological landscape, South Africa.
- 540 Environ. Manage. 52, 595–611. doi:10.1007/s00267-013-0094-9
- 541 Crawley, M. (2007). The R book. John Wiley and Sons Ltd., Chichester, UK, 949 pp.
- 542 Cresswell, J.E., 2000. A comparison of bumblebees' movements in uniform and aggregated
- 543 distributions of their forage plant. Ecol. Entomol. 25, 19–25. doi:10.1046/j.1365-
- 544 2311.2000.00230.x

- 545 Deguines, N., Jono, C., Baude, M., Henry, M., Julliard, R., Fontaine, C., 2014. Large-scale
 546 trade-off between agricultural intensification and crop pollination services. Front. Ecol.
 547 Environ. 12, 212–217. doi:10.1890/130054
- 548 Eardley, C., B. Gemmill-Herren, M. Gikungu, R. Kagioya, W. Kinuthia, P. K. Kwapong, D.
- Mar-tins, L. Njoroge, and G. Tribe. 2007. Crops, browse and pollinators in Africa: an
 initial stock-taking. FAO, Rome.
- Eilers, E.J., Kremen, C., Smith Greenleaf, S., Garber, A.K., Klein, A.-M., 2011. Contribution
 of pollinator-mediated crops to nutrients in the human food supply. PLoS One 6, e21363.
- 553 doi:10.1371/journal.pone.0021363
- Ehlers Smith, Y.C., Ehlers Smith, D.A., Seymour, C.L., Thébault, E., van Veen, F.J.F., 2015.
- 555 Response of avian diversity to habitat modification can be predicted from life-history
- traits and ecological attributes. Landsc. Ecol. 30, 1225–1239. doi:10.1007/s10980-015-
- 557 0172-x
- Ehrlich, P.R., Harte, J., 2015. Opinion: To feed the world in 2050 will require a global
 revolution. PNAS 112, 14743–14744. doi:10.1073/pnas.1519841112
- 560 Farwig, N., Bailey, D., Bochud, E., Herrmann, J.D., Kindler, E., Reusser, N., Schüepp, C.,
- 561 Schmidt-Entling, M.H., 2009. Isolation from forest reduces pollination, seed predation
- and insect scavenging in Swiss farmland. Landsc. Ecol. 24, 919–927.
- 563 doi:10.1007/s10980-009-9376-2
- 564 Forrest, J.R.K., Thorp, R.W., Kremen, C., Williams, N.M., 2015. Contrasting patterns in
- species and functional-trait diversity of bees in an agricultural landscape. J. Appl. Ecol.
 52, 706–715. doi:10.1111/1365-2664.12433
- 567 Free, J.B., Williams, I.H. 1976. Insect pollination of Anacardium occidentale L., Mangifera
- 568 *indica* L., *Blighia sapida* Koenig and *Persea americana* Mill. Tropical Agriculture, 53,
- 569 125–139.

- 570 Fründ, J., Zieger, S.L., Tscharntke, T., 2013. Response diversity of wild bees to overwintering
 571 temperatures. Oecologia 173, 1639–1648. doi:10.1007/s00442-013-2729-1
- 572 Fontaine, C., Dajoz, I., Meriguet, J., Loreau, M., 2006. Functional diversity of plant-pollinator
- 573 interaction webs enhances the persistence of plant communities. PLoS Biol. 4, e1.
- 574 doi:10.1371/journal.pbio.0040001
- 575 Gagic, V., Bartomeus, I., Jonsson, T., Taylor, A., Winqvist, C., Fischer, C., Slade, E.M.,
- 576 Steffan-dewenter, I., Emmerson, M., Potts, S.G., Tscharntke, T., Weisser, W.,
- 577 Bommarco, R., 2015. Functional identity and diversity of animals predict ecosystem
- 578 functioning better than species-based indices.
- 579 doi:http://dx.doi.org/10.1098/rspb.2014.2620
- 580 Gallai, N., Salles, J.-M., Settele, J., Vaissière, B.E., 2008. Economic valuation of the
- 581 vulnerability of world agriculture confronted with pollinator decline. Ecol. Econ. 68,

582 810–821. doi:10.1016/j.ecolecon.2008.06.014

- 583 Garibaldi, L. A, Aizen, M. A, Klein, A.M., Cunningham, S. a, Harder, L.D., 2011a. Global
- 584 growth and stability of agricultural yield decrease with pollinator dependence. Proc.
- 585 Natl. Acad. Sci. U. S. A. 108, 5909–5914. doi:10.1073/pnas.1012431108
- 586 Garibaldi, L. A, Steffan-Dewenter, I., Kremen, C., Morales, J.M., Bommarco, R.,
- 587 Cunningham, S. A, Carvalheiro, L.G., Chacoff, N.P., Dudenhöffer, J.H., Greenleaf, S.S.,
- 588 Holzschuh, A., Isaacs, R., Krewenka, K., Mandelik, Y., Mayfield, M.M., Morandin, L.
- 589 A, Potts, S.G., Ricketts, T.H., Szentgyörgyi, H., Viana, B.F., Westphal, C., Winfree, R.,
- 590 Klein, A.M., 2011b. Stability of pollination services decreases with isolation from
- natural areas despite honey bee visits. Ecol. Lett. 14, 1062–72. doi:10.1111/j.1461-
- 592 0248.2011.01669.x
- 593 Garibaldi, L. a, Steffan-Dewenter, I., Winfree, R., Aizen, M. a, Bommarco, R., Cunningham,
- 594 S. A, Kremen, C., Carvalheiro, L.G., Harder, L.D., Afik, O., Bartomeus, I., Benjamin, F.,

595	Boreux, V., Cariveau, D., Chacoff, N.P., Dudenhöffer, J.H., Freitas, B.M., Ghazoul, J.,
596	Greenleaf, S., Hipólito, J., Holzschuh, A., Howlett, B., Isaacs, R., Javorek, S.K.,
597	Kennedy, C.M., Krewenka, K.M., Krishnan, S., Mandelik, Y., Mayfield, M.M., Motzke,
598	I., Munyuli, T., Nault, B. a, Otieno, M., Petersen, J., Pisanty, G., Potts, S.G., Rader, R.,
599	Ricketts, T.H., Rundlöf, M., Seymour, C.L., Schüepp, C., Szentgyörgyi, H., Taki, H.,
600	Tscharntke, T., Vergara, C.H., Viana, B.F., Wanger, T.C., Westphal, C., Williams, N.,
601	Klein, A.M., 2013. Wild pollinators enhance fruit set of crops regardless of honey bee
602	abundance. Science 339, 1608–11. doi:10.1126/science.1230200
603	Garibaldi, L. A., Bartomeus, I., Bommarco, R., Klein, A.M., Cunningham, S. A., Aizen, M.
604	A., Boreux, V., Garratt, M.P.D., Carvalheiro, L.G., Kremen, C., Morales, C.L., Schüepp,
605	C., Chacoff, N.P., Freitas, B.M., Gagic, V., Holzschuh, A., Klatt, B.K., Krewenka, K.M.,
606	Krishnan, S., Mayfield, M.M., Motzke, I., Otieno, M., Petersen, J., Potts, S.G., Ricketts,
607	T.H., Rundlöf, M., Sciligo, A., Sinu, P.A., Steffan-Dewenter, I., Taki, H., Tscharntke, T.,
608	Vergara, C.H., Viana, B.F., Woyciechowski, M., 2015. Trait matching of flower visitors
609	and crops predicts fruit set better than trait diversity. J. Appl. Ecol. 52, 1436–1444.
610	doi:10.1111/1365-2664.12530
611	Garibaldi, L.A., Carvalheiro, L.G., Vaissiere, B.E., Gemmill-Herren, B., Hipolito, J., Freitas,
612	B.M., Ngo, H.T., Azzu, N., Saez, A., Astrom, J., An, J., Blochtein, B., Buchori, D.,
613	Garcia, F.J.C., Oliveira da Silva, F., Devkota, K., Ribeiro, M. d. F., Freitas, L.,
614	Gaglianone, M.C., Goss, M., Irshad, M., Kasina, M., Filho, A.J.S.P., Kiill, L.H.P.,
615	Kwapong, P., Parra, G.N., Pires, C., Pires, V., Rawal, R.S., Rizali, A., Saraiva, A.M.,
616	Veldtman, R., Viana, B.F., Witter, S., Zhang, H., 2016. Mutually beneficial pollinator
617	diversity and crop yield outcomes in small and large farms. Science 351, 388–391.
618	doi:10.1126/science.aac7287

- 619 Gelman A., Su Y-S., 2015. arm: Data Analysis Using Regression and Multilevel/ Hierarchical
 620 Models. R package version 1.8-6. https://CRAN.R-project.org/package=arm
- 621 Geslin, B., Gauzens, B., Thébault, E., Dajoz, I., 2013. Plant Pollinator Networks along a
- 622 Gradient of Urbanisation. PLoS One 8, e63421. doi:10.1371/journal.pone.0063421
- 623 Geslin, B., Le Féon, V., Kuhlmann, M., Vaissière, B.E., Dajoz, I., 2016. The bee fauna of
- 624 large parks in downtown Paris , France. Ann. la Société Entomol. Fr.
- 625 doi:10.1080/00379271.2016.1146632Greenleaf, S.S., Williams, N.M., Winfree, R.,
- 626 Kremen, C., 2007. Bee foraging ranges and their relationship to body size. Oecologia
- 627 153, 589–596. doi:10.1007/s00442-007-0752-9
- 628 Goulson, D., Nicholls, E., Botias, C., Rotheray, E.L., 2015. Bee declines driven by combined
- 629 stress from parasites, pesticides, and lack of flowers. Science (80-.). 347, 1255957–
- 630 1255957. doi:10.1126/science.1255957
- Grueber, C.E., Nakagawa, S., Laws, R.J. & Jamieson, I.G., 2011 Multi- model inference in
- ecology and evolution: challenges and solutions. J. of Evo. Biol., 24, 699–711
- Henri, D.C., Jones, O., Tsiattalos, A., Thébault, E., Seymour, C.L., van Veen, F.J.F., 2015.
- 634 Natural vegetation benefits synergistic control of the three main insect and pathogen
- pests of a fruit crop in southern Africa. J. Appl. Ecol. 52, 1092–1101. doi:10.1111/1365-
- 636 2664.12465
- 637 Henry, M., Béguin, M., Requier, F., Rollin, O., Odoux, J.-F., Aupinel, P., Aptel, J.,
- 638 Tchamitchian, S., Decourtye, A., 2012. A common pesticide decreases foraging success
- and survival in honey bees. Science 336, 348–50. doi:10.1126/science.1215039
- Hoehn, P., Tscharntke, T., Tylianakis, J.M., Steffan-Dewenter, I., 2008. Functional group
- 641 diversity of bee pollinators increases crop yield. Proc. Biol. Sci. 275, 2283–91.
- 642 doi:10.1098/rspb.2008.0405

643	Holzschuh, A., Steffan-dewenter, I., Kleijn, D., Tscharntke, T., 2007. Diversity of flower-
644	visiting bees in cereal fields : effects of farming system, landscape composition and
645	regional context 41-49. doi:10.1111/j.1365-2664.2006.01259.x
646	Holzschuh, A., Dormann, C.F., Tscharntke, T., Steffan-Dewenter, I., 2011. Expansion of
647	mass-flowering crops leads to transient pollinator dilution and reduced wild plant
648	pollination. Proc. Biol. Sci. 278, 3444-51. doi:10.1098/rspb.2011.0268
649	Hulot, F. D., Lacroix, G., Lescher-Moutoué, F. & Loreau, M., 2000 Functional diversity
650	governs response to nutrient enrichment. Nature, 405, 340-344.
651	Jauker, F., Diekötter, T., Schwarzbach, F., Wolters, V., 2009. Pollinator dispersal in an
652	agricultural matrix: opposing responses of wild bees and hoverflies to landscape
653	structure and distance from main habitat. Landsc. Ecol. 24, 547–555.
654	doi:10.1007/s10980-009-9331-2
655	Johnson, P.C.D., 2014. Extension of Nakagawa & Schielzeth's R 2 GLMM to random slopes
656	models. Methods Ecol. Evol. 5, 944–946. doi:10.1111/2041-210X.12225
657	Kennedy, C.M., Lonsdorf, E., Neel, M.C., Williams, N.M., Ricketts, T.H., Winfree, R.,
658	Bommarco, R., Brittain, C., Burley, A.L., Cariveau, D., Carvalheiro, L.G., Chacoff, N.P.,
659	Cunningham, S.A., Danforth, B.N., Dudenhöffer, JH., Elle, E., Gaines, H.R., Garibaldi,
660	L.A., Gratton, C., Holzschuh, A., Isaacs, R., Javorek, S.K., Jha, S., Klein, A.M.,
661	Krewenka, K., Mandelik, Y., Mayfield, M.M., Morandin, L., Neame, L.A., Otieno, M.,
662	Park, M., Potts, S.G., Rundlöf, M., Saez, A., Steffan-Dewenter, I., Taki, H., Viana, B.F.,
663	Westphal, C., Wilson, J.K., Greenleaf, S.S., Kremen, C., 2013. A global quantitative
664	synthesis of local and landscape effects on wild bee pollinators in agroecosystems. Ecol.
665	Lett. 16, 584–599. doi:10.1111/ele.12082

- 666 Klein, A.-M., Steffan-Dewenter, I., Tscharntke, T., 2003. Fruit set of highland coffee
- 667 increases with the diversity of pollinating bees. Proc. Biol. Sci. 270, 955–61.

668 doi:10.1098/rspb.2002.2306

- 669 Klein, A.-M., Vaissière, B.E., Cane, J.H., Steffan-Dewenter, I., Cunningham, S. A, Kremen,
- C., Tscharntke, T., 2007. Importance of pollinators in changing landscapes for world
 crops. Proc. Biol. Sci. 274, 303–13. doi:10.1098/rspb.2006.3721
- 672 Kremen, C., M'Gonigle, L.K., 2015. Small-scale restoration in intensive agricultural
- 673 landscapes supports more specialized and less mobile pollinator species. J. Appl. Ecol.
- 674 n/a-n/a. doi:10.1111/1365-2664.12418
- Kremen, C., Williams, N.M., Thorp, R.W., 2002. Crop pollination from native bees at risk
- 676 from agricultural intensification. Proc. Natl. Acad. Sci. U. S. A. 99, 16812–16816.
- 677 doi:10.1073/pnas.262413599
- Laliberté, E., Legendre. P. 2010. A distance-based framework for measuring functional
 diversity from multiple traits. Ecology 91: 299-305
- 680 Laliberté, E., Legendre, P., Bill Shipley, 2010. Measuring functional diversity (FD) from
- 681 multiple traits, and other tools for functional ecology. http://cran.r-
- 682 project.org/web/packages/FD/FD.pdf
- Larsen, T.H., Williams, N.M., Kremen, C., 2005. Extinction order and altered community
- 684 structure rapidly disrupt ecosystem functioning. Ecol. Lett. 8, 538–547.
- 685 doi:10.1111/j.1461-0248.2005.00749.x
- 686 Leonhardt, S.D.S., Gallai, N., Garibaldi, L.A., Kuhlmann, M., Klein, A.-M., 2013. Economic
- 687 gain, stability of pollination and bee diversity decrease from southern to northern
- 688 Europe. Basic Appl. Ecol. 14, 461–471. doi:10.1016/j.baae.2013.06.003
- 689 Manning, R. 1995. Honeybee pollination technical data. Western Australia Department of
- 690 Agriculture, Bulletin No. 4298

Martins, K.T., Gonzalez, A., Lechowicz, M.J., 2015. Pollination services are mediated by bee
functional diversity and landscape context. Agric. Ecosyst. Environ. 200, 12–20.

693 doi:10.1016/j.agee.2014.10.018

- Morandin, L. A., Kremen, C., 2013. Hedgerow restoration promotes pollinator populations
- and exports native bees to adjacent fields. Ecol. Appl. 23, 829–839. doi:10.1890/12-

696 1051.1

- Mucina L., Rutherford M. C., 2006, The vegetation of South Africa, Lesotho and Swaziland
 Strelitzia 19. South African National Biodiversity Institute, Pretoria. 807pp.
- Nielsen, A., Steffan-Dewenter, I., Westphal, C., Messinger, O., Potts, S.G., Roberts, S.P.M.,
- 700 Settele, J., Szentgyörgyi, H., Vaissière, B.E., Vaitis, M., Woyciechowski, M., Bazos, I.,
- 701 Biesmeijer, J.C., Bommarco, R., Kunin, W.E., Tscheulin, T., Lamborn, E., Petanidou, T.,
- 702 2011. Assessing bee species richness in two Mediterranean communities: Importance of
- habitat type and sampling techniques. Ecol. Res. 26, 969–983. doi:10.1007/s11284-011-
- 704 0852-1
- 705 Öckinger, E., Schweiger, O., Crist, T.O., Debinski, D.M., Krauss, J., Kuussaari, M., Petersen,
- J.D., Pöyry, J., Settele, J., Summerville, K.S., Bommarco, R., 2010. Life-history traits
- 707 predict species responses to habitat area and isolation: a cross-continental synthesis.

708 Ecol. Lett. 13, no-no. doi:10.1111/j.1461-0248.2010.01487.x

709 Ollerton, J., Winfree, R., Tarrant, S., 2011. How many flowering plants are pollinated by

710 animals? Oikos 120, 321–326. doi:10.1111/j.1600-0706.2010.18644.x

- 711 Oertli, S., Müller, A., Dorn, S., 2005. Ecological and seasonal patterns in the diversity of a
- species-rich bee assemblage (Hymenoptera: Apoidea: Apiformes. Eur. J. Entomol. 102,
- 713 53–63. doi:10.1016/j.biocon.2005.05.014

714 Orford, K.A., Vaughan, I.P., Memmott, J., 2015. The forgotten flies: the importance of non-

715 syrphid Diptera as pollinators. Proc. Biol. Sci. 282, 20142934–.

716 doi:10.1098/rspb.2014.2934

- 717 Orford, K.A., Murray, P.J., Vaughan, I.P., Memmott, J., 2016. Modest enhancements to
- conventional grassland diversity improve the provision of pollination services. J. Appl.

719 Ecol. Online Only doi:10.1111/1365-2664.12608

Popic, T.J., Davila, Y.C., Wardle, G.M., 2013. Evaluation of common methods for sampling

invertebrate pollinator assemblages: net sampling out-perform pan traps. PLoS One 8,

722 e66665. doi:10.1371/journal.pone.0066665

723 R Development Core Team, 2014. R: A language and environment for statistical computing.

R Foundation for Statistical Computing, Vienna, Austria. URL: http://www.r-project.org.

Rader, R., Bartomeus, I., Tylianakis, J.M., Laliberté, E., 2014. The winners and losers of land

vse intensification: pollinator community disassembly is non-random and alters

functional diversity. Divers. Distrib. 20, 908–917. doi:10.1111/ddi.12221

- 728 Ricketts, T.H., Regetz, J., Steffan-Dewenter, I., Cunningham, S. A, Kremen, C., Bogdanski,
- A., Gemmill-Herren, B., Greenleaf, S.S., Klein, A.M., Mayfield, M.M., Morandin, L. A,
- 730 Ochieng', A., Potts, S.G., Viana, B.F., 2008. Landscape effects on crop pollination
- services: are there general patterns? Ecol. Lett. 11, 499–515. doi:10.1111/j.1461-

732 0248.2008.01157.x

733 Rollin, O., Bretagnolle, V., Fortel, L., Guilbaud, L., Henry, M., 2015. Habitat, spatial and

temporal drivers of diversity patterns in a wild bee assemblage. Biodivers. Conserv.
doi:10.1007/s10531-014-0852-x

736 Roulston, T.H., Smith, S.A., Brewster, A.L., Farm, B.E., Lane, B.F., 2007. A Comparison of

737 Pan Trap and Intensive Net Sampling Techniques for Documenting a Bee

738 (Hymenoptera : Apiformes) J. Kansas Entomol. Soc. 80, 179–181.

739	Samnegård, U., Hambäck, P. A, Eardley, C., Nemomissa, S., Hylander, K., 2015. Turnover in							
740	bee species composition and functional trait distributions between seasons in a tropical							
741	agricultural landscape. Agric. Ecosyst. Environ. 211, 185–194.							
742	doi:10.1016/j.agee.2015.06.010							
743	Schmid-Hempel, P., Kacelnik, A., Houston, A.I., 1985. Honeybees maximize efficiency by							
744	not filling their crop. Behav. Ecol. Sociobiol. 17, 61–66. doi:10.1007/BF00299430							
745	Schmid-Hempel, P., & Schmid-Hempel, R., 1986. Nectar-collecting bees use distance-							
746	sensitive movement rules. Animal behaviour, 34, 605-607. doi:10.1016/S0003-							
747	3472(86)80131-2							
748	Stang, M., Klinkhamer, P.G.L., Waser, N.M., Stang, I., van der Meijden, E., 2009. Size-							
749	specific interaction patterns and size matching in a plant-pollinator interaction web. Ann.							
750	Bot. 103, 1459–1469. doi:10.1093/aob/mcp027							
751	Stang, M., Klinkhamer, P.G.L., Van Der Meijden, E., 2006. Size constraints and flower							
752	abundance determine the number of interactions in a plant-flower visitor web. Oikos							
753	112, 111–121. doi:10.1111/j.0030-1299.2006.14199.x							
754	Stanley, D.A., Garratt, M.P.D., Wickens, J.B., Wickens, V.J., Potts, S.G., Raine, N.E., 2015.							
755	Neonicotinoid pesticide exposure impairs crop pollination services provided by							
756	bumblebees. Nature 528, 548–550. doi:10.1038/nature16167							
757	Steffan-Dewenter, I., Tscharntke, T., 1999. Effects of habitat isolation on pollinator							
758	communities and seed set. Oecologia 121, 432-440.							
759	Sung, I., Lin, M., Chang, C., Cheng, A., Chen, W., 2006. Pollinators and Their Behaviors on							
760	Mango Flowers in Southern Taiwan. Formos. Entomol. 170, 161–170.							
761	Tilman, D., Knops, J., Wedin, D., Reich, P., Ritchie, M., Siemann, E., 1997. The influence of							
762	functional diversity and composition on ecosystem processes. Science. 277, 1300–1302.							
763	doi:10.1126/science.277.5330.1300							

764	Tscharntke, T., Klein, A.M., Kruess, A., Steffan-Dewenter, I., Thies, C., 2005. Landscape
765	perspectives on agricultural intensification and biodiversity - Ecosystem service
766	management. Ecol. Lett. doi:10.1111/j.1461-0248.2005.00782.x

- 767 Tylianakis, J.M., Klein, A.M., Tscharntke, T., 2005. Spatiotemporal variation in the diversity
- of hymenoptera across a tropical habitat gradient. Ecology 86, 3296–3302.
- 769 doi:10.1890/05-0371
- 770 Vanbergen, A.J., and the Pollinator Initiative, 2013. Threats to an ecosystem service:
- pressures on pollinators. Front. Ecol. Environ. 11, 251–259. doi:10.1890/120126
- Vázquez, D.P., Morris, W.F., Jordano, P., 2005. Interaction frequency as a surrogate for the
- total effect of animal mutualists on plants. Ecol. Lett. 8, 1088–1094. doi:10.1111/j.1461-
- 774 0248.2005.00810.x
- Westphal, C., Steffan-Dewenter, I., Tscharntke, T., 2003. Mass flowering crops enhance
 pollinator densities at a landscape scale. Ecol. Lett. 6, 961–965. doi:10.1046/j.14610248.2003.00523.x
- 778 Westphal, C., Bommarco, R., Carré, G., Lamborn, E., Morison, N., Petanidou, T., Potts, S.G.,
- Roberts, S.P.M., Szentgyörgyi, H., Tscheulin, T., Vaissière, B.E., Woyciechowski, M.,
- 780 Biesmeuer, J.C., Kunin, W.E., Settele, J., Steffan-Dewenter, I., 2008. Measuring bee
- diversity in different European habitats and biogeographical regions. Ecol. Monogr. 78,
 653–671. doi:10.1890/07-1292.1
- Winfree, R., 2008. Pollinator-dependent crops: an increasingly risky business. Curr. Biol. 18,
 R968–9. doi:10.1016/j.cub.2008.09.010
- 785 Williams, N.M., Crone, E.E., Roulston, T.H., Minckley, R.L., Packer, L., Potts, S.G., 2010.
- Ecological and life-history traits predict bee species responses to environmental
- 787 disturbances. Biol. Conserv. 143, 2280–2291. doi:10.1016/j.biocon.2010.03.024

- Williams, N.M., Kremen, C., 2007. Resource distributions among habitats determine solitary
 bee offspring production in a mosaic landscape. Ecol. Appl. 17, 910–921.
 doi:10.1890/06-0269
- 791 Wilson, J.S., Griswold, T., Messinger, O.J., 2008. Sampling Bee Communities
- (Hymenoptera : Apiformes) in a Desert Landscape : Are Pan Traps Sufficient ? J. Kansas
 Entomol. Soc. 81, 288–300.
- Wood, S. a., Karp, D.S., DeClerck, F., Kremen, C., Naeem, S., Palm, C. A., 2015. Functional
- traits in agriculture: agrobiodiversity and ecosystem services. Trends Ecol. Evol. 30,
- 796 531–539. doi:10.1016/j.tree.2015.06.013

797	Table 1: Model selection for flower abundance, insect abundance and functional diversity. N
798	corresponds to the number of models selected in the top $4AIC_c$, R^2_m and R^2_c correspond
799	respectively to the marginal and conditional R^2 values of the best model. Est. corresponds to
800	the parameter estimate (for centralized and standardized predictor variables), CI to its 95%
801	confidence interval and Imp. to the parameter relative importance. * Mango field was the
802	reference habitat and [§] the dry season was the sampling season of reference.

	Flower abundance			Insect abundance			Functional diversity		
	(N = 2 models)			(N = 6 models)			(N = 6 models)		
	$R_{m}^{2} = 0.40$; $R_{c}^{2} = 0.68$			$R_{m}^{2} = 0.54$; $R_{c}^{2} = 0.67$			$R_{m}^{2} = 0.32$; $R_{c}^{2} = 0.41$		
Parameter	Est.	CI	Imp.	Est.	CI	Imp.	Est.	CI	Imp.
Intercept	2.867	(2.056, 3.679)		1.402	(1.120, 1.683)		0.069	(0.059, 0.082)	
Habitat [*]	-2.563	(-3.033, -2.093)	1.00	0.255	(-0.082, 0.592)	0.68	0.017	(0.0002, 0.033)	0.79
Season [§]	-1.427	(-1.756, -1.098)	1.00	1.393	(1.033, 1.751)	1.00	0.048	(0.033, 0.063)	1.00
Distance	0.538	(0.187, 0.889)	1.00	-0.632	(-0.940, -0.325)	1.00	-0.026	(-0.042, -0.011)	1.00
Habitat X Season	2.363	(1.632, 3.093)	1.00	-0.517	(-0.962, -0.072)	0.55	0.015	(-0.007, 0.038)	0.45
Distance X Season	0.326	(-0.346, 0.999)	0.35	0.687	(0.227, 1.148)	1.00	-0.006	(-0.036, 0.023)	0.23
Flower abundance				-0.119	(-0.328, 0.089)	0.39			

Table 2: Model selection for body length, wing span and ratio of wing span over body length.
Same legend as Table 1. * Mango field was the reference habitat and [§] the dry season was the
sampling season of reference.

	Body length (N = 7 models)			Wing span (N = 7 models)			Ratio of wing span over body length (N = 5 models)		
	$R_m^2 = 0$	0.037; R ² _c	= 0.45	$R_{m}^{2} = 0.015$; $R_{c}^{2} = 0.54$			$R_{m}^{2} = 0.02$; $R_{c}^{2} = 0.40$		
Parameter	Est.	CI	Imp.	Est.	CI	Imp.	Est.	CI	Imp.
Intercept	0.556	(0.271, 0.840)		0.805	(0.433, 1.177)		0.248	(0.155, 0.342)	
Habitat [*]	0.049	(0.0007, 0.097)	0.89	0.052	(0.006, 0.099)	0.93	-0.0004	(-0.013, 0.013)	0.24
Season [§]	0.168	(0.112, 0.225)	1.00	0.126	(0.067, 0.186)	1.00	-0.042	(-0.061, -0.022)	1.00
Distance	-0.047	(-0.109, 0.014)	0.68	-0.018	(-0.095, 0.028)	0.52	0.002	(-0.011, 0.015)	0.28
Habitat X Season	-0.023	(-0.10, 0.053)	0.32	-0.008	(-0.097, 0.048)	0.32	0.003	(-0.019, 0.025)	0.07
Distance X Season	0.060	(0.004, 0.117)	0.53	0.016	(-0.011, 0.109)	0.32	-0.009	(-0.028, 0.009)	0.10

810 Figures

811 **Figure 1**: Effect of distance to the natural habitat (m), habitat type and sampling period on: A.

812 abundance of flowers (log+1), B. abundance of insects, and C. functional diversity of insects.

813 Lines represent model predicted values. In all panels, mango flowering period (dry season) is

814 represented in dark grey and mango non-flowering period (wet season) is represented in light

815 grey.

816

817 Figure 2: Relationship between body length (mm) and wing span (mm) of flying insects (A),

818 and effect of distance to the natural habitat (m) and habitat type on insect body length (B).

819 Mango flowering period (dry season) is represented in dark grey and mango non-flowering

820 period (wet season) is represented in light grey.

Figure 1





Figure 2



825