Impact of land-use change on flower-visiting insect communities on an oceanic island

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Abstract. 1. Land-use change has profoundly impacted pollinator communities throughout the world. However, the processes through which it acts on pollinator diversity and composition are still poorly understood, especially in highly vulnerable island ecosystems.

2. In this study, we investigated the distribution, abundance, richness and composition of flower-visiting insects to assess their response to land-use change in Terceira Island (Azores).

3. Flower-visiting insects were sampled over 2 years using a standardised protocol along 50 transects across five different habitats corresponding to a land-use gradient. Insect species were classified as indigenous or exotics. We assessed changes across habitats using multiple diversity indices, species abundance distribution models (SAD) and species composition metrics (β-diversity), along with plant species composition.

4. We observed that indigenous flower-visiting insects were dominant, both in abundance and species richness, across the entire land-use gradient. Species diversity varied only slightly across the gradient. SADs were lognormal in all habitats, with very few truly common and rare flower-visiting insects and a prevalence of species of intermediate abundance. Species replacement was significantly higher mainly between the two most contrasting habitats (i.e. natural forests and intensive pastures) but was significantly correlated with species replacement of host plant species across the gradient.

5. Our results revealed that the Azorean flower-visiting insect communities were highly simplified across the entire gradient with little difference between habitats. In the absence of strong exotic competitors, indigenous flower-visiting insects expand their range and occupy new anthropogenic habitats, also facilitating the expansion of a large number of exotic plant species.

Key words. Beta diversity, community assembly, exotic species, flower-visiting insects, land-use change, native species, oceanic islands, pollinator networks, species abundance distribution.

Introduction

Land-use change is leading to the loss and degradation of natural habitats, resulting in the severe disruption of biodiversity processes and patterns throughout the world.
In particular, land-use change has profoundly impacted species ranges and abundances and is now recognised as a major driver of the current extinction crisis (Fahrig, 1997; Brooks et al., 2002). As a consequence, key ecosystem processes such as pollination interaction networks have been severely affected, with dramatic consequences for ecosystem functioning and the provision of goods and services for humans (Cane, 2001; Kearn, 2001; Warren et al., 2001; Williams et al., 2001; Kremen et al., 2002; Biesmeijer et al., 2006; Klein et al., 2007; Steffan-Dewenter & Westphal, 2008; Winfree et al., 2009; Potts et al., 2010; Groom & Schwarz, 2011; Rader et al., 2014). The implementation of appropriate management actions to mitigate the impact of human disturbance on pollination interaction networks requires, therefore, a better understanding of how species diversity, distribution and abundance patterns of pollinators are altered in response to land-use change.

Over the last century, island ecosystems have been disproportionately affected by anthropogenic alterations and a large proportion of recorded extinctions have taken place on islands (e.g. Cardoso et al., 2010b; Rando et al., 2013; Alcover et al., 2015; Régnier et al., 2015; Terzopoulou et al., 2015). Land conversion of native forest to agricultural and exotic forest is now recognised as one of the major causes of island biodiversity decline, with many extant species predicted to be committed to future extinction as a result of land-use change (‘the extinction debt’; Triantis et al., 2010). These profound changes are known to have impacted several components of island ecosystems (see Connor et al., 2012), but very little is known about the impact of land-use change on island ecological networks, and in particular, on pollinating insects.

Insects are responsible for 78–94% of pollination across all flowering plants, and 75% of global food crops (Klein et al., 2007; Ollerton et al., 2011; Winfree et al., 2011). Guaranteeing a diversity of pollinators, particularly the species with a high degree of specialisation (Steffan-Dewenter et al., 2006; Albrecht et al., 2012), is therefore crucial for maintaining gene flow and community stability in plant communities (Ricketts, 2004; Klein et al., 2007; Steffan-Dewenter & Westphal, 2008; Cranmer et al., 2012). Insular ecosystems usually support less complex networks with lower numbers of pollinator species, are mostly comprised of generalist species (Olesen et al., 2002; Whittaker & Fernández-Palacios, 2007) and have less redundancy between species in comparison with continental areas (Olesen et al., 2002). Thus, pollinator networks on oceanic islands are potentially highly vulnerable to any kind of disturbance (Traveset, 2002), and can be considered ideal model systems to evaluate the impact of land-use change on the diversity, distribution and abundance of pollinator species (Alarcón et al., 2014; Castro-Urgal & Traveset, 2014; Kaiser-Bunbury & Blüthgen, 2015; Traveset et al., 2015).

Over the last decade, a large range of negative impacts that can be attributed to land-use change have already been documented for pollinator communities. For instance, previous studies have identified a negative correlation between land-use intensity and the provision of functions sustained by pollinator species (Garibaldi et al., 2011; Winfree et al., 2011; Rader et al., 2014). With increasing land-use intensity, a clear increase in the dominance of common species has also been identified (Tylia-nakis et al., 2005), especially in small island populations that are more susceptible to the disruption of interaction networks (Kaiser-Bunbury et al., 2010). In response to intermediate disturbances, studies have also underlined the presence of an initial increase in local pollinator richness, but with some degree of regional homogenisation, as the few specialists are replaced by abundant, often invasive, generalists (Kremen, 2005; Rader et al., 2014).

In this study, we investigate the flower-visiting insect species communities of the Azores archipelago. Located in the North Atlantic Ocean, the archipelago is composed of nine main islands, all volcanic and of recent origin (the oldest island being 8.12 Myr BP). The Azorean climate is temperate oceanic, characterised by high levels of relative humidity and small temperature fluctuations. Since the 15th century and the arrival of humans to the Azores, the native semi-tropical evergreen laurel forest (Laurisilva), originally covering most of the surface area across the islands, has been gradually replaced by agricultural land uses (i.e. intensively managed pastures for cattle and semi-natural pastures) and exotic forest (plantations of introduced wood species). Most of the native forest is nowadays confined to Juniperus-Ilex montane forests, characterised by reduced tree stature (usually up to 5 m, rarely reaching 10 m) on shallow soil and rugged terrain at high altitude, mostly between 800 and 1000 m.a.s.l (Martins, 1993; Borges et al., 2005; Cardoso et al., 2009, 2010a; Elias et al., 2016). Recent investigation of the impact of land-use changes in the Azores has shown that native forests and intensively managed pastures are the most important habitats influencing arthropods species composition and diversity, playing a fundamental role as source habitats for endemic and exotic species respectively (Borges et al., 2008; Cardoso et al., 2009, 2010a). Intermediate-disturbed habitats, such as semi-natural pastures and exotic forests, also perform important functional roles, acting as corridors connecting native forest fragments for many indigenous arthropod species (Borges et al., 2008; Cardoso et al., 2009). However, despite the persistence of some Azorean native species in anthropogenic habitats (Fattorini et al., 2012), the large spread of exotic species throughout the landscape matrix tends to promote biotic homogenisation of arthropod species at both local and island scales (Florence et al., 2013).

In this contribution, we examine the impact of land-use change on flower-visiting insect species community structure in an Azorean island. Based on previous work on Azorean arthropod communities (Borges et al., 2008; Cardoso et al., 2009, 2010a; Florence et al., 2013, 2015), we predict that: (i) native habitats such as natural forest will support a higher abundance and richness of indigenous
flower-visiting insects in comparison to non-native land uses; (ii) species composition of flower-visiting insect communities will change from native habitats to non-native land-uses and (iii) the dominance of a few common and many rare flower-visiting insect species will increase as disturbance increases.

Methods

Study area, sampling and species identification

Our study was conducted on Terceira Island. Terceira is an island from the central group of the Azores archipelago, located in the North Atlantic Ocean (38°37'N–38°48'N, 27°02'W–27°23'W) with an Area 402 km² and maximum elevation of 1023 m. Field work was conducted from June to September 2013 and from July to October 2014, due to favourable weather conditions and relatively high number of plant species in the flowering period. For this study, we selected five distinct habitat types covering a large percentage of the total island area with, from the least to the most disturbed, natural forests (NatFor), naturalised vegetation areas (NatVeg), exotic forests (ExoFor), semi-natural pastures (SemiPast) and intensively managed pastures (IntPast) (Cardoso et al., 2013). Compared with previous ecological studies undertaken in the Azores (Borges et al., 2008; Cardoso et al., 2009, 2010a; Florencio et al., 2013, 2015), we added naturalised vegetation areas, dominated by Erica azorica, Pittosporum undulatum and Rubus hochstetterorum, as an important habitat for flower-visiting insects, because of its recent growing extent due to pasture abandonment and combination of native and exotic flora. Detailed features regarding each habitat type are outlined in Table S1.

In each habitat type, we chose 10 sites in which 10 m long line-transects (1 m width) were set up (Pollard & Yates, 1993), making a total of 50 transects located across the entire island (Fig. 1, see Table S2 for details). To select the 10 sites per habitat type, we tried to maximise the covered environmental diversity following Jiménez-Valverde and Lobo (2004) and Aranda et al. (2011). First, an environmental matrix for Terceira Island (see Borges et al., 2006) was compiled using climatic, topographic and geological variables with a resolution of 100 x 100 m. Using the k-means non-hierarchical clustering algorithm, we grouped all cells of each habitat type in 10 clusters, making a total of 50 clusters (5 habitats x 10 groups). For each cluster, we ordered the cells according to their distance to the group’s multidimensional centroid using Euclidean distance. The first cell in this ranking, deemed to be the most representative of the cluster, was chosen for sampling. If it was impossible to reach the selected cell in the field due to inaccessibility or lack of authorisation from land owners, the second cell was chosen and so forth.

Transect surveys were carried out once per year and repeated in the following year, in a randomised order, under sunlight (from 9 a.m. to 6 p.m) and only in sunny weather, with a duration of 180 minutes per transect. Transect location was selected to encompass spots of dense flowering. Each flower along every 10 m transect was surveyed for 4 minutes to guarantee effective contact of the insect; therefore, only insects probing for nectar or eating/collecting pollen (foraging) were recorded. Flower-visiting insects were observed and collected with a poouter when it was not possible to identify them in the field. The specimens collected were sorted first into morphospecies and later identified to species-level under the supervision of PAVB, following the taxonomic nomenclature in Borges et al. (2010). When species-level identification could not be resolved, individuals were identified to the lowest taxonomic unit possible and classified as morphospecies (see Acknowledgements). Voucher specimens and a reference collection were deposited in EDTP – Entomoteca Dalberto Teixeira Pombo, University of Azores, Angra do Heroísmo, Portugal. All species were classified as indigenous or exotic species. Indigenous species may be endemic (i.e. found only in the Azores) or native non-endemic (i.e. species that colonised the Azores by natural long-distance dispersal mechanisms). Exotic species are those whose original distribution range did not include the Azores and are believed to have arrived as a consequence of human activities; these species often have a cosmopolitan distribution (see Borges et al., 2010).

Data analysis

Initially, we performed several statistical comparison tests (Chi-square, paired t-test, analysis of variance (ANOVA) followed by Tukey HSD post hoc tests) to identify potential differences between the 2 years of sampling (2013–2014), regarding habitat types, taxonomic orders, colonisation status, abundance and richness. A statistically significant increase in species richness was observed between the 2 years (t = −4.4; P = 0.006; Table S5), which was primarily a result of the addition of rare species between years, although in absolute terms the increase was small. The difference in total abundance of individuals per species between years was found to be non-significant (t = 1.43; P = 0.22; Table S5). Therefore, in the following analyses, we combined data from the same transects of both years to obtain a better sampling completeness.

Using equivalent sampling effort in combination with the same standardised method in different habitat types may still result in differences in inventory completeness due to differences in the abundance of plant species in different transects. To analyse the variation in flower-visiting insect species accumulation between habitats and rule out possible biases in the sampling effort, we constructed species accumulation curves for the observed number of species, species richness estimates, singletons, and doubletons using the non-parametric estimators Chao1 and Jackknife 1 (Chao1 and Jack1, both abundance based). Species
accumulation curves were constructed by randomly selecting the order of transect addition at each iteration. We repeated this process 1000 times, and used the mean of the 1000 random runs. To analyse the estimators’ performance across all habitats, slopes were calculated along the entire curve. Sampling completeness was calculated in two ways: first, we calculated the ratio of observed richness to estimated richness ratio with Chao1, due to its higher precision (Hortal et al., 2006) and second, we recorded the final slope of species richness accumulation curves built with both observed and estimated richness as the inverse of the number of individuals needed to add the final single species to the accumulation curve (see Cardoso et al., 2009 for more details).

To investigate differences in flower-visiting insects’ diversity between habitats, we calculated the mean number of individuals, species richness and two commonly used diversity indices namely Shannon–Wiener (H’) and Pielou’s evenness (J’). In addition, we calculated the Berger–Parker dominance index (D), which expresses the proportional abundance of the most abundant species, presented in the inverse format (1/D), so that an increase in the value of the index accompanies an increase in diversity and a decrease in dominance (Magurran, 2004). To test for statistically significant differences in diversity between habitats, we applied one-way ANOVAs followed by Tukey HSD post hoc tests. ANOVAs were performed using generalised least square models (GLS; Pinheiro & Bates, 2000) to account for potential heteroscedasticity. Additionally, we also tested the ability of the GLS models to account for potential spatial structures by estimating the Moran’s I spatial autocorrelation index for GLS residuals using the latitude and longitude of each transect site. When the overall GLS was statistically significant, the Tukey’s post hoc test was used to identify statistically significant pairwise differences between habitats.

We studied the dissimilarity in flower-visiting species composition between sites of all habitat types using Jaccard’s index as an overall beta diversity measure (β_{total}), and decomposing it into its replacement (β_{rep}) and richness difference (β_{rich}) components (Carvalho et al., 2012; Cardoso et al., 2014). β diversity indices were computed using presence/absence data. We also computed β diversity with log-transformed abundance data (results not

Fig. 1. Land-use distribution map of Terceira Island with the selected sampling sites as black dots: NatFor (natural forests), SemiPast (semi-natural pastures), NatVeg (naturalised vegetation areas), ExoFor (exotic forests), IntPast (intensively managed pastures) [cartographic information from DROTRH (2008) and Gaspar (2007), see also Gaspar et al. (2011)]. [Colour figure can be viewed at wileyonlinelibrary.com]
shown), but the results were similar (Cardoso et al., 2015). Dissimilarity distances were visualised using non-metric multidimensional scaling ordinations (NMDS). To examine between-habitat differences in species composition, we used analysis of similarities (ANOSIM) using the three beta diversity components as dissimilarity measures, followed by post hoc tests with P-values adjusted using the Benjamini and Hochberg (1995) correction for multiple testing. We also computed $\beta_{\text{tot}}$, $\beta_{\text{rep}}$ and $\beta_{\text{rich}}$ for plant species composition and correlated each $\beta$ component of flower-visiting insects with its respective component for plants communities using Mantel tests with Spearman correlation.

In addition to examining patterns in flower-visiting species diversity and composition, we also explored variations in the species abundance distributions (SADs) of flower-visiting species (Matthews & Whittaker, 2015) across the five habitat types. To determine the shape of the SAD in each sample, we fitted logseries, lognormal and gambin SAD models to the observed abundance data, using both binned and unbinned data with the logseries and lognormal models, and only binned data with the gambin model (Matthews et al., 2014). The theoretical description of these SAD models and the complete methodological approach, including how the models were fitted and compared, and classification types of rare species is provided in Appendix S1 (see also Matthews & Whittaker, 2014).

All analyses were performed with Microsoft Excel, IBM SPSS 20.0 (Nie et al., 2011) and the R statistical environment (R Development Core Team, 2016) using the R packages BAT (Cardoso et al., 2015, 2016), vegan (Oksanen et al., 2013), polog (Grotn & Engen, 2009) and gambin (Matthews et al., 2014).

Results

Species composition

Insects visited 2134 flowers (49% of the 4354 sampled flowers) belonging to 48 plant species from 21 families. The number of plant species surveyed per habitat type was distributed as follows: 17 plant species (1134 flowers) were identified in NatFor, 27 plant species (815 flowers) in NatVeg, 26 plant species (820 flowers) in ExoFor, 15 plant species (828 flowers) in SemiPast and 14 plant species (757 flowers) in IntPast (see Table S4).

The sampled flower-visiting insects belonged to 54 species and morphospecies from four orders, namely, Coleoptera, Diptera, Hymenoptera and Lepidoptera (Table S3). The most representative group was Diptera, with 51% of the individuals, followed by Hymenoptera with 25%, Coleoptera with 18% and finally, the Lepidoptera with 6%. The most common species were Sepsis neocynipsea (Diptera) (17% of the individuals) and Anaspis proteus (Coleoptera) (16.5%), followed by Bombus ruderatus (6.3%), Apis mellifera (5%), Lasioglossum villosulum (all Hymenoptera) (4.5%) and Stomorhina lunata (Diptera) (4.6%) (Table S3). Flies (Diptera) were the most represented group in all habitats, invariably followed by bees (Hymenoptera) ($\chi^2 = 4.81$, d.f. = 12, $P = 0.96$). Sepsis neocynipsea (Diptera) had the highest number of individuals in three habitat types: NatVeg, SemiPast and IntPast, whereas Anaspis proteus (Coleoptera) was dominant in NatFor and Bombus ruderatus (Hymenoptera) in ExoFor.

At the island scale, we observed that the majority of flower-visiting insects were native non-endemic species (82.1%), while only a small percentage was endemic (5.4%) or exotic (12.5%). These proportions were similar throughout all habitats ($\chi^2 = 0.89$, d.f. = 8, $P = 1$), showing that indigenous species dominated flower-visiting insect’s communities across the entire gradient (Table 1). On the other hand, at the island scale, the majority of host plants were exotic species (75%), and a small percentage was native non-endemic (14.6%) or endemic (10.4%). These proportions slightly differed between habitats ($\chi^2 = 17.5$, d.f. = 8, $P = 0.025$), although the introduced plant species were dominant in all habitats with the exception of NatFor (Table 1).

Sampling completeness

The average numbers of flower-visiting insect species per habitat estimated by the Chao1 and Jack1 estimators were found to be close to the observed richness values (Table S6). Considering the estimates obtained with Chao 1, the sampling completeness values for each habitat

Table 1.

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varied between 98% for NatFor and 63% for SemiPast, with 90% for ExoFor, 87% for IntPast and 86% for NatVeg, all representing a good level of sampling completeness (Cardoso et al., 2009). The species accumulation curves (Fig. S1) approached an asymptote (with slope values between 0.002 and 0.08 by the end of the accumulation process) and the final slope values of the estimators’ curves were close to 0 for all habitats, which shows that the inventory was relatively complete in all habitats (Fig. S2).

**Insect diversity in the different habitats**

Mean number of individuals, species evenness ($J'$) and dominance (1/D) for flower-visiting insects did not show any significant differences between habitats ($F_{1,4} = 1.185, P = 0.330$; $F_{1,4} = 1.682, P = 0.171$ and $F_{1,4} = 2.513, P = 0.055$, respectively, Fig. 2a, d and e). However, species richness differed significantly between habitats ($F_{1,4} = 4.231, P = 0.005$) with NatFor being the richest habitat and NatVeg and SemiPast being the poorest (Fig. 2b). Shannon-Wiener $H'$ index differed marginally between habitats ($F_{1,4} = 2.711, P = 0.042$) with ExoFor being significantly more diverse than SemiPast (Fig. 2c). No spatial autocorrelation was detected in the residuals of the binned data (ANOSIM: $r = 0.019, P = 0.233$). Significant correlations were found between the flower-visiting insects and plant species of the three $eta$ measurements (Fig. 4a–c) with the pattern of $\beta_{total}$ being mostly driven by the $\beta_{rich}$ component.

**Habitat similarity**

Overall, the analysis of flower-visiting insects’ $eta$-diversity using Jaccard’s index ($\beta_{total}$) showed significant differences in composition between habitat types (ANOSIM: $r = 0.179, P = 0.001$, Fig. 3a) with values ranging from 0.835 between NatFor and IntPast to 0.794 between NatFor and ExoFor (Table S7). NatFor was significantly more dissimilar to all anthropogenic habitats (Post hoc ANOSIM $P < 0.05$, Table S8), while no significant differences were detected between anthropogenic habitats, except between ExoFor and SemiPast (Post hoc ANOSIM $P = 0.02$). $\beta_{total}$ had the dominant component of $\beta_{rich}$ with values ranging from 0.602 between ExoFor and both NatFor and NatVeg, to 0.494 between SemiPast and NatFor. $\beta_{repl}$ had lower but still significant importance (ANOSIM: $r = 0.061, P = 0.023$, Fig. 3b) in explaining $\beta$ diversity patterns. Significant differences in $\beta_{rich}$ values ranged from 0.316 between NatFor and SemiPast, to 0.192 between NatFor and ExoFor, but no significant difference between habitat types was found (ANOSIM: $r = 0.019, P = 0.233$). Significant correlations were found between the flower-visiting insects and plant species of the three $\beta$ measurements (Fig. 4a–c) with the pattern of $\beta_{total}$ being mostly driven by the $\beta_{rich}$ component.

**Species abundance distributions (SADs) and rarity patterns**

Considering the binned data, the gambin model provided the best fit to all five habitat types ($\Delta AIC_c = 0$, Table S9), although for the NatFor, the PLN had a $\Delta AIC_c < 2$. The PLN always provided a better fit to the binned data than the logseries. However, when the logseries and PLN were fitted to the unbinned data, the logseries provided a better fit to all five habitat types, indicating a greater number of rare species than predicted.
by the PLN (Table S9). The gambin model provided a good fit to the data in all habitat types according to the Pearson’s chi-square ($\chi^2$) goodness-of-fit test for NatFor: $\chi^2 = 6.376, \ P = 0.605$; NatVeg: $\chi^2 = 5.963, \ P = 0.31$; ExoFor: $\chi^2 = 1.568, \ P = 0.905$; SemiPast: $\chi^2 = 11.303, \ P = 0.079$ and IntPast: $\chi^2 = 2.656, \ P = 0.753$. The $\alpha$ parameter of the gambin model did not show substantial variations between habitats with values of 2.364 for

Fig. 3. Two-dimensional ordination solution using non-metric multidimensional scaling (NMDS) with the $\beta$ diversity measures $\beta_{\text{total}}$ (a) and $\beta_{\text{repl}}$ (b) for flower-visiting insects. Dots indicate transects, while lines delimit the smallest polygon that encloses all transects for a given habitat. The stress value of NMDS was 0.17 and 0.18 for $\beta_{\text{total}}$ and $\beta_{\text{repl}}$ respectively. NatFor (natural forest), NatVeg (naturalised vegetation areas), ExoFor (exotic forest), SemiPast (semi-natural pasture), and IntPast (intensively managed pasture). [Colour figure can be viewed at wileyonlinelibrary.com]

Fig. 4. Correlations between species composition ($\beta$ diversity) of flower-visiting insects and plants across the 50 transects. Correlations were performed with $\beta_{\text{total}}$ (a), $\beta_{\text{repl}}$ (b) and $\beta_{\text{rich}}$ (c). Spearman correlation coefficient and its associated $P$-values of the Mantel test are given on the top of each panel. NatFor (natural forest), NatVeg (naturalised vegetation areas), ExoFor (exotic forest), SemiPast (semi-natural pasture), and IntPast (intensively managed pasture).
NatFor, 2.348 for SemiPast, 3.244 for NatVeg, 4.502 for ExoFor and 3.965 IntPast. Alpha values in this range indicate positively skewed lognormal-like (i.e. more rare species than predicted by a standard lognormal model) to standard lognormal SAD shapes (Fig. 5), with the lower values of $\alpha$ in NatFor and SemiPast denoting a relatively higher proportion of rarer species in these two habitat types.

With regard to the species classified as common species (i.e. the 25% most abundant), there is only one habitat type with one species having more than 128 specimens: NatFor with *Anaspis proteus* (264 specimens). However, when considering regional abundance in the island, there are three true common species (*Sepsis neocynipsea* with 362 specimens, *Anaspis proteus* with 352 specimens and *Bombus ruderatus* with 134 specimens) (Fig. 5). The proportions of rare flower-visiting insect species represented in the first two bins of the SADs histograms in Fig. 5 were decomposed into pseudo-rare and regionally rare species. The pseudo-rare species are relatively high in numbers when data from all habitats are aggregated, but are rare in some particular habitats and are the species primarily responsible for the differences in proportions of rare species between habitat types. The regionally rare species, that is, the number of species with less than four individuals (Fig. 5 Island; i.e. the first quartile of available bins) only comprise five species. These are the truly rare species. All habitats revealed a high number of intermediate abundance species (Fig. 5), as is to be expected in lognormal-shaped SADs (Table 2).

**Discussion**

In this study, we documented the influence of different levels of disturbance on the distribution, composition, richness and abundance of flower-visiting insect species on an Azorean island. First, we revealed that the island flower-visiting insect community is dominated by widespread generalist native species of intermediate abundance, despite the high representation of exotic plant species. Second, we showed that the species diversity, species abundance distribution (SAD) and species composition of flower-visiting insect species vary only slightly across the land-use gradients. Species replacement was significantly higher mainly between the two most contrasting habitats (i.e. natural forests and intensive pastures). Finally, species composition of flower-visiting insects was

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**Fig. 5.** Species abundance distribution (SADs) histograms for flower-visiting insects, with predicted values of the gambin model (black dots), for all habitats, (a) natural forest (b), naturalised vegetation (c), exotic forest (d) semi-natural pasture (e), and intensively managed pasture (f). The following binning system was used: bin 1 corresponds to the number of species with 1 individual per species, bin 2 corresponds to the number of species with 2–3 individuals per species, bin 3 corresponds to the number of species with 4–7 individuals per species, etc. (see Gray et al., 2006 and Matthews et al., 2014). [Colour figure can be viewed at wileyonlinelibrary.com]
influenced by the distribution of host plant species regardless of the landscape matrix.

With the exception of the study of Olesen et al. (2002), to the best of our knowledge, there is no other study investigating flower-visiting insect communities in the Azores. In fact, although there are many studies investigating the impacts of land-use change on the community structure of pollinator insects on continental regions, such studies are scarce on oceanic islands. In one of the few examples, Sahari et al. (2010), in contrast to our results, showed that landscape change in Java Island (Indonesia) strongly affects insect pollinating species composition and richness with increasing rainforest isolation and land-use intensity, indicating significant changes in species composition between habitat types in the tropics, with emphasis on case studies of wild and crop plants from Indonesia.

**Insect diversity in the different habitats**

Our results demonstrated a surprising uniformity of several community metrics across the different habitats, suggesting that similar mechanisms may control flower-visiting species diversity across our land-use gradient. In most of the habitats, native non-endemic flies were the group with the largest number of species, a pattern already documented for island pollination networks (Castro-Urgeal & Traveset, 2014). Concerning our original aims and hypotheses, as expected, natural forest was found to be a favourable habitat for indigenous flower-visiting insects, although we did not observe statistical differences between habitat types in terms of abundance (Fig. 2a, S1 and Table S5). This could be explained by adaptation or cross-scale resilience and response diversity of the native flower-visiting insect species to non-native habitats (see also Winfree & Kremen, 2009; Cardoso et al., 2010a), a possible consequence of the island’s small area relative to the flower-visiting species’ foraging area (Miller et al., 2015) and loss of native habitats. Hence, these differences in insect flower-visiting community could have been also influenced by the variation in altitude through the different habitat types; native forest being always at higher altitude than intensively managed pastures (Table S1). In conclusion, and even considering that exotic plants dominate all habitats with the exception of native forest, indigenous flower-visiting insects’ diversity did not greatly vary, both in terms of abundance and species diversity, across the entire gradient.

**Habitat similarity**

As in previous studies focusing on the impact of land-use change in Azorean arthropod communities (e.g. Borges et al., 2008; Cardoso et al., 2009, 2010a; Meijer et al., 2011), native forest and intensively managed pasture showed the most contrasting flower-visiting species composition. This difference was mostly a consequence of replacement differences (species substitution), with only a minor contribution of species richness variation (Fig. 3). This result differs from previous work conducted with epigean arthropods in Terceira (see Cardoso et al., 2009), where strong differences in species composition were reported between all types of habitats. Our finding illustrates the need for further investigation concerning the role of landscape dynamics on Azorean insect pollinator species. The few differences in community composition reported across habitats could also be explained by the ‘habitat heterogeneity hypothesis’ (e.g. MacArthur & Wilson, 1967), where the flower-visiting species, due to low interspecific competition, predation and parasitic pressures (Olesen et al., 2002; Ribeiro et al., 2005) subdivide the landscape into suitable habitats (i.e. niche partitioning), based on plant communities (Tews et al., 2004; Cramer & Willig, 2005). In fact, flower-visiting species composition was found to be mostly influenced by host plant species composition across all habitats (Fig. 4). The fact that differences in flower-visiting insects’ composition correlated with differences in host plant species composition across habitat types (Fig. 4) implies that any changes in vegetation composition (i.e. replacement of native by exotic or invasive plants) might have a profound impact on pollinating insect community structure in the Azores. Interestingly, the high correlation between similarity values of plant and arthropod community structure was also observed by Borges (1999) for phytophagous insects and predatory arthropods from sown and semi-natural pastures in the Azores. In an additional study, Fründ et al. (2010) reported positive diversity correlation between 1764 individuals of 131 pollinator species with 77 plant species (n = 27 networks) across sites at a regional scale, even though only parts of the variation in bees and hoverfly diversity was explained by the diversity of flower species.

**Species abundance distributions (SADs) and rarity patterns**

The structure of flower-visiting insect species relative abundances did not differ substantially between habitats (Figs 2a and 5), in spite of the clear land-use gradient

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**Table 2. Number of regionally rare, habitat rare, pseudo-rare intermediate and common species of flower-visiting insects for NatFor (natural forests), SemiPast (semi-natural pastures), Nat-Veg (naturalised vegetation areas), ExoFor (exotic forests), and IntPast (intensively managed pastures) and island (region).**

<table>
<thead>
<tr>
<th>Habitats</th>
<th>Regionally rare</th>
<th>Pseudo-rare</th>
<th>Intermediate</th>
<th>Common</th>
</tr>
</thead>
<tbody>
<tr>
<td>Island</td>
<td>5</td>
<td>–</td>
<td>46</td>
<td>3</td>
</tr>
<tr>
<td>NatFor</td>
<td>1</td>
<td>13</td>
<td>26</td>
<td>1</td>
</tr>
<tr>
<td>NatVeg</td>
<td>2</td>
<td>14</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>ExoFor</td>
<td>2</td>
<td>12</td>
<td>26</td>
<td>0</td>
</tr>
<tr>
<td>SemiPast</td>
<td>0</td>
<td>11</td>
<td>19</td>
<td>0</td>
</tr>
<tr>
<td>IntPast</td>
<td>1</td>
<td>11</td>
<td>22</td>
<td>0</td>
</tr>
</tbody>
</table>

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present in Terceira, and the fact that previous studies have reported a clear effect of land-use change on SAD form for epigean arthropods on the same island (see Matthews et al., 2014). In fact, we found only slight variation in the form of the SAD between habitat types as highlighted by the small differences in gambin’s $\alpha$ values calculated using binned data, and the fact that the logseries model provided the best fit to the non-binned data from all five habitat types. The SAD form in the different habitat types was accurately assessed by the gambin model for which the range of $\alpha$ values were characteristic of lognormal-like SADs (Ugland et al., 2007; Matthews et al., 2014), albeit with a relatively higher than expected proportion of rare species in native forest and semi-natural pasture (Table 2; Fig. 5). These results reveal therefore that most flower-visiting species, across all habitats, were of intermediate abundance. This could be explained by the fact that Azorean communities are largely unsaturated with ample resources, both features associated with low competition for food (Preston, 1948; Borges et al., 2008; Miller et al., 2015) (Table S9: Fig. 5). We also documented the presence of rare species although these were mostly considered to be pseudo-rare species (i.e. these species are rare in a given habitat but more frequent in others) that were likely present due to source-sink dynamics across habitat types. Our work supports the view that Azorean arthropod communities are highly simplified, characterised by a dominance of generalist species (see also Olesen et al., 2002; Ribeiro et al., 2005; Whittaker & Fernández-Palacios, 2007; Travest et al., 2015), the presence of multiple local habitat pseudo-rarities, and few regionally rare species (see also Borges et al., 2008). Borges et al. (2008) also reported another example of a functional group in the Azorean arthropod community with a high proportion of pseudo-rare species, in that case spiders which, similarly to pollinator insects, have many species able to use diverse resources, not limited to one specific habitat.

Conclusions and future directions

Our finding supports the observations of Olesen et al. (2002) reported for a different Azorean island (Flores), where indigenous super-generalist species tend to include exotic plants in their set of pollinated plants without any clear evidence for a facilitation between exotic plant and pollinator species. Therefore, our findings emphasise the need for further studies on pollination networks on islands to investigate the spread of exotic and invasive plants by indigenous pollinating insects that could in turn threaten the endemic flora. Finally, also further work is needed to clarify whether the Azorean indigenous pollinating insect species are behaving as ‘jacks of all trades, masters of none’, that is, what is the efficacy of these species in pollen transport and plant reproduction in the Azores? Given that we documented only a slight variation in pollinator community according to a land-use gradient, we suggest this is a starting point for assessing the insects’ pollinators’ behaviour along a disturbance gradient in the other islands of the Azores archipelago, and compare it at island-level with Terceira flower-visiting insect communities. In conclusion, our study offers one of the first exhaustive assessments of the impact of land-use change on an Azorean island flower-visiting insect community, revealing (i) the influence of plant species composition on flower-visiting insect species composition, and (ii) providing evidence for potential occupation of native flower-visiting insects in new anthropogenic habitats.

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Supporting Information

Additional Supporting Information may be found in the online version of this article under the DOI reference: doi: 10.1111/icad.12216:

Appendix S1. Description of the species abundance distribution models and methodological approach.

Fig. S1. Species accumulation curves for several non-parametric estimators and for singletons and doubletons of the different habitat types.

Fig. S2. Slopes of species accumulation curves for non-parametric abundance-based estimators Chao1 and Jackknife 1 of the different habitat types.

Table S1. Habitat descriptions with information on the altitude, main plant species and disturbance index.

Table S2. Geographical coordinates for all sites in Terceira Island for each habitat type.

Table S3. Abundance and number of individuals per insect species/morphospecies in each habitat type.

Table S4. Number of flowers per plant species in each habitat type.

Table S5. Number of species/morphospecies studied in 2013 and 2014 years per each habitat type.

Table S6. Number of individuals, species/morphospecies, singletons and doubletons studied in each habitat type.

Table S7. Comparison of total beta diversity ($\beta_{\text{total}}$), replacement beta diversity ($\beta_{\text{rep}}$) and richness beta diversity ($\beta_{\text{rich}}$) values along a gradient of increasing dissimilarity for all transects considered together of each habitat.

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Table S8. P-values of the post hoc pairwise Analysis of similarities (ANOSIM) corrected for multiple tests.

Table S9. AICc values for the SAD model selection. The three SAD models (logseries, PLN and gambin) were fitted to flower-visiting insect data from five land-use types.

References


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