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Local and regional specialization in plant-pollinator networks

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Abstract

 Specialization of species is often studied in ecology but its quantification and meaning is disputed. More recently, ecological network analysis has been widely used as a tool to quantify specialization, but here its true meaning is also debated. However, irrespective of the tool used, the geographic scale at which specialization is measured remains central. Consequently, we use data sets of plant-pollinator networks from Brazil and the Canary Islands to explore specialization at local and regional scales. We ask how local specialization of a species is related to its regional specialization, and whether or not species tend to interact with a non-random set of partners in local communities. Local and regional specialization were strongly correlated around the 1:1 line, indicating that species conserve their specialization levels across spatial scales. Furthermore, most plants and pollinators also showed link conservatism repeatedly across local communities, and thus seem to be constrained in their fundamental niche. However, some species are more constrained than others, indicating true specialists. We argue that several geographically separated populations should be evaluated in order to provide a robust evaluation of species specialization.

Keywords: community, scale, spatial variation, partner fidelity, link conservatism

Introduction

 The Earth's biodiversity is shaped by a plethora of interactions between species that may range from being relatively specialized to relatively generalized, and the exact nature of these interactions depend upon a variety of ecological and evolutionary factors. Many of these types of interactions are common and ecologically vital; pollination by animals for example is the norm for an estimated 87.5% of the c. 352,000 species of flowering plants (Ollerton et al. 2011). Extensive work based on visitation data between plants and pollinators indicate that such interactions tend to be more generalized than previously thought, even for tropical systems (Ollerton and Cranmer 2002, Schleuning et al. 2012, Waser et al. 1996). However, recent studies, considering visitation frequency and service effectiveness, suggest stronger fidelity between interaction partners and that binary visitation data might overestimate generalization levels (King et al. 2013, Rosas-Guerrero et al. 2014 – though see Ollerton et al. 2015). Furthermore, several studies indicate that species can appear as generalists while actually being composed of specialist populations, or even generalist populations composed of specialist individuals (Araujo et al. 2008, Bolnick et al. 2002, Devictor et al. 2010, Dupont et al. 2011, Fox and Morrow 1981, Tur et al. 2014). Such studies suggest that phylogenetic and geographic scale are highly relevant when studying specialization, and sampling of multiple populations is necessary to reliably determine the niche breath of a species to account for cross-community variation (Carstensen et al. 2014, Fox and Morrow 1981, Ollerton et al. 2007, Ollerton et al. 2009, Poisot et al. 2015, Trøjelsgaard et al. 2015).

 The fundamental taxonomic niche of a species describes the set of potential interaction partners as determined by functional traits (Junker et al. 2013). Conversely, the realized taxonomic niche is the subset of species within a given community with which the focal species interacts. Thus, while functional traits do not necessarily enable us to predict the occurrence of a given interaction, they do partly define the set of possible interactions (Morales-Castilla et al. 2015) and

 trait complementarity does seem to consistently govern general interaction patterns within communities (Carstensen et al. 2016). The fundamental and realized niche concepts are related to specialization at the level of species and populations respectively, and the fundamental niche is seldom fully realized in any single local community because of differences in relative abundances, species composition and other biotic and abiotic factors (Burkle et al. 2016, Carstensen et al. 2014, Trøjelsgaard et al. 2015).

 Here, we explore species specialization across interacting plants and pollinators within two contrasting regions: Brazilian *campo rupestre*, or rupestrian grasslands, and the Canary Islands. Both datasets consist of seven spatially separated community-level plant-pollinator networks, sampled in a standardized way within their regions. Using these data we explore the relationship between local and regional specialization, and investigate whether this relationship differ between these two contrasting study regions, assuming the ecological specialization framework (Armbruster 2017). Specifically we ask: 1) Are local and regional measures of specialization correlated? 2) Do species interact locally with a non-random subset of the available species in the community?

 We expect that the specialization level of a species is conserved across populations, but that the identity of interaction partners might change. If so, species would appear more generalized at the regional level compared to the local level.

Methods

Study sites and field observations

 Data were collected in October-December 2012 from seven sites of rupestrian grasslands, or *campo rupestre*, in the National Park of Serra do Cipó, SE Brazil (Carstensen et al. 2014, 2016). *Campo rupestre* is a species-rich vegetation of mostly shrubs and herbs associated with rocky outcrops surrounded by sandy and stony grasslands (Silveira et al. 2016). Sites were 1.4-8.5 km

 apart within an altitudinal range of 1073-1260 m a.s.l. One site was sampled per day with a weekly rotation among sites. Attempting to observe all flowering plant species in each site, plant individuals were observed in 15 min intervals (totalling 252 hours of observation), recording all visitors touching the reproductive floral parts (for more details see Carstensen et al. 2014). For each 15-min interval randomly selected plant individuals were observed, and if individuals of the same species were in close proximity of each other they were often surveyed simultaneously. Flower visitors were collected for taxonomic identifications by experts. For each site, the data were summarized in a quantitative interaction matrix expressing the frequency of interactions between pairs of plants and pollinators. This resulted in seven spatially separated networks with a total of 101 plant species and 201 pollinator species.

 Data were also collected from five islands within the Canarian archipelago (El Hierro, La Gomera, Tenerife (two sites), Gran Canaria and Fuerteventura) and a single site in Western Sahara located at the West African coast close to the Canary Islands (Trøjelsgaard et al. 2013). In total this gave us seven spatially separated pollination networks from habitats characterized as semi-arid and dominated by shrubs. The Canary Island networks were 53-455 km from each other and separated by ocean, except the two sampling sites at Tenerife, which, conversely, were separated by El Pico del Teide (3718 m a.s.l.). All flowering perennial plant species were surveyed for flower visitors in intervals of 15 min (totalling 296 hours of observation) in January-March 2010 through flower- based focal observations in a similar way as for the Brazilian data set (for more details see Trøjelsgaard et al. 2013). Most plant species were observed in 15-min intervals approximately 8 99 times (mean $= 7.0$, Std. dev. $= 1.8$). Randomly selected individuals were preferably chosen for each of the individual 15-min surveys, although low abundance of some species precluded this practice. Sometimes multiple individuals of the same species were surveyed simultaneously depending on their spatial aggregations, and also depending on the total number of flowers per individual.

 Approximately 1300 flower visitors were collected for taxonomic verifications by entomological experts, allowing species identification of many pollinators, while some were grouped as morphospecies. Overall we scored interactions between 39 plant species and 249 pollinator species and, similar to the Brazilian dataset, all seven networks from the Canary Islands were quantitative interaction matrices expressing the frequency of interactions between plants and pollinators.

Specialization and interaction richness

 For each plant and pollinator species observed in more than one site, we defined a metaweb of species *s* (Dunne 2006) as the network consisting of all species occurring in the sites in which *s* occurs, as well as all the interactions between them. In other words, it is the regional network of interactions for species *s* and its co-occurring species. We quantified specialization using the index d'_{s} , which describes the deviation of interactions from what would be expected under neutral 115 conditions. Instead of counting the number of interaction partners, d'_{s} measures the exclusiveness of a species' interactions and is insensitive to variation in observation frequencies across species 117 (Blüthgen et al. 2006). Specialization, d'_{s} , was calculated using the *dfun* function in the package 'Bipartite' in R (Dormann 2011); and for all species we calculated both a regional specialization 119 level using the metaweb of species s ($d'_{s,meta}$) as well as an average local specialization level using 120 the local networks in which species *s* occurred $(d'_{s,local})$.

 We tested the correlation between local and regional specialization using the Pearson correlation coefficient. Through simple linear regression, we further tested whether the regional specialization was affected by the number of sites in which a species occurred, or rather, the number of potential interaction partners with which it co-occurred across the region.

 To quantify if regional interaction richness of a species differed from what could be expected if species interacted randomly with the available set of partners, we first calculated the

 regional degree for each species, γ*s*, which is the total number of different interaction partners across all sites in which the species was observed (i.e. the degree in the metaweb for species *s*). We 129 then compared the empirical γ_s with a null model where the identity of partners in the local networks was randomized while conserving the local number of interactions for each species. New interaction partners were drawn with a probability proportional to their abundance (measured as total number of interactions, i.e. the marginal sums in the quantitative interaction matrix) among all potential partner species in a local community, and random partners were drawn with replacement until the empirical number of interactions was reached. In this way, the total number of local interactions (i.e. the marginal sum) was retained, but the regional degree was allowed to change as species were allocated new interaction partners among all potential partners at a local site. This procedure was repeated 1000 times for each species, which allowed us to calculate a mean (γ*s,random*) and standard deviation (*SDrandom*) for each species. Subsequently we compared empirical regional species degrees (γ*s*) with the randomized regional degrees (γ*s,random*) by using z-scores [(γ*^s* - γ*s,random*)/*SDrandom*]. If the z-score was less than -1.96, or larger than 1.96, the difference between the empirical and random regional degree was deemed statistically significant. Moreover, negative values suggest that a species is more constrained in its choice of interaction partners than if partner identities were determined solely by availability (abundance) (we call this 'link conservatism'), while positive values suggest that the species is more opportunistic than expected (we call this 'link opportunism'). By incorporating species abundance as a way of selecting partners in the null-model, the procedure leans towards the neutral perspective which assumes that interactions between species is largely determined by species abundance (see e.g. Canard et al. 2014, Poisot et al. 2015). It is important to note that the null-model ignores phenological or morphological constraints (e.g. Olesen et al. 2011), and therefore may overestimate the availability of potential partners.

Results

160 Regional ($d'_{s,meta}$) and local ($d'_{s,local}$) specialization were significantly and strongly correlated in 161 the Brazilian networks for both plants and pollinators (Pearson correlation, $r = 0.76$ and 0.83 162 respectively, $p < 0.001$, Fig. 1A) as well as in the Canary Islands networks (Pearson correlation, $r =$ 0.76 and 0.89 respectively, p < 0.001, Fig. 1B). In Brazil, the number of co-occurring species (i.e. the number of potential interaction partners in the metaweb) had no effect on regional specialization 165 values for plants ($p > 0.7$, $R^2 < 0.01$), but correlated significantly with regional specialization of 166 pollinators, albeit with a very poor fit ($p < 0.05$, $R^2 = 0.05$, indicated by size of data points in Fig. 1A). In the Canary Islands the number of co-occurring species had no effect on regional 168 specialization for neither pollinators nor plants ($p > 0.2$, $R^2 < 0.01$). The null model analysis showed similar results for the Brazilian and Canary Islands networks (Fig. 2). In both systems, the majority of the plant species showed link conservatism because 81% (Brazil) and 57% (Canary Islands) of the plants had significantly fewer regional interaction partners than expected if partner identity were determined solely by availability (abundance). That is, these

species tend to conserve their interaction partners across sites. The majority of the pollinators also

 showed link conservatism in Brazil but not in the Canary Islands as 58% and 46% of the pollinators, respectively, had significantly fewer regional interaction partners. Finally, the pollinators deviation from randomness (i.e. the z-scores, with negative and positive values being suggestive of link conservatism and opportunism, respectively) correlated significantly and 178 negatively with regional specialization in both Brazil (Pearson correlation, $r = -0.34$, $p < 0.001$) and 179 the Canary Islands (Pearson correlation, $r = -0.30$, $p = 0.009$), while the relationship where non-180 significant for plants in both Brazil (Pearson correlation, $r = -0.25$, $p = 0.14$) and the Canary Islands 181 (Pearson correlation, $r = 0.16$, $p = 0.50$) (Fig. 3).

Discussion

 The results from the two regions were surprisingly similar. Local and regional specialization were strongly correlated in both regions. Plants were generally more specialized than pollinators (Fig. 1) despite a general trend of having more interaction partners (Fig. 2). A correlation between local and regional specialization was expected, however, a strong correlation close to 1:1 indicates that local 188 and regional specialization, in terms of d'_{s} , is largely interchangeable, and that species in plant- pollinator networks are consistent in their specialization level across scales. By aggregating the specialization level across geographically separated sampling sites, the current procedure attempted to capture the differences in interactions that can be found between populations and individuals (see e.g. Araujo et al. 2008, Tur et al. 2014). In the Canary Islands the geographical distance between sites varied from 52 to 456 km, while the distance in Brazil varied between 1.4 and 8.5 km. Yet, the rocky outcrops found in the *campo rupestre*, within which the surveys were done in Brazil, can to some extent be seen as isolated habitat patches within a matrix of grassland, and the actual isolation between the populations is likely larger than the geographical distances dictates. However, we cannot preclude that increasing the geographical extent so that we covered even more distant

 populations, especially in Brazil, would lead to different results. Nevertheless, it seems encouraging that the relationship between local and regional specialization were almost identical when comparing the Brazilian and Canarian communities, which suggest that this pattern might actually be a general trend. More studies from different communities are needed, however, to explore the generality of this finding.

 Larger distances and increased isolation between sites could result in local population- specific adaptations within species, and thus translate into a larger inter-island variability in interaction partners, which ultimately would result in higher opportunism at the regional scale (Thompson 2005, Trøjelsgaard et al. 2015). Indeed, this could potentially partly explain why a lower percentage of plants and animals showed link conservatism in the Canary Island compared with the networks in Brazil. In fact, it is important to note that the Brazilian and Canarian communities also differ in a number of other aspects. For example, across the sampling sites the 210 ratio of pollinators: plants were on average 4.4 ± 1.0 (mean \pm SD) in the Canary Islands, and 2.6 ± 0.4 in Brazil. A lower diversity of plants in the Canarian communities may hamper the pollinators potential to show a strong fidelity across sites when compared with a null model. However, if partner diversity was the sole explanation, we would expect the Canarian plants to show a stronger fidelity as they have more partners to choose from and, therefore, a larger potential to deviate from the null model. Nonetheless, as mentioned, both Canarian plants and pollinators had a lower percentage of link conservative species compared with the species found in Brazil. Thus, partner diversity (i.e. the pollinators:plants ratio) is likely not the sole explanation. Another important aspect could be species turnover (see also Carstensen et al 2014, Trøjelsgaard et al. 2015). That is, if there is a strong turnover in partners from site to site (or island to island) the potential for link conservatism diminishes. The plant communities were on average more similar across the Canary 221 Islands (average Sorensen similarity across sites \pm SD, Canary Islands = 0.44 \pm 0.13; Brazil = 0.33

 222 ± 0.14), while the pollinator communities on average were more similar in Brazil (average Sorensen 223 similarity across sites \pm SD, Canary Islands = 0.24 \pm 0.12; Brazil = 0.43 \pm 0.06). Thus, while the high turnover of pollinators at the Canary Island might explain why a lower percentage of the plants show significant link conservatism, this does not explain the patterns observed for the pollinators. Finally, the studied communities also differ in e.g. species composition, climate conditions, and other biotic and abiotic aspects, which makes it difficult to isolate a single common mechanism as responsible for the difference in link conservatism between the Canarian and Brazilian communities.

 Overall, for specialist species the foraging and interaction choices are most likely a consequence of trait complementarities, while generalists could be expected to forage or interact mainly according to relative abundances of potential partners. Link conservatism should therefore be higher for specialist species, as we have confirmed (Fig. 3). Indeed, as we argue below, link 234 conservatism and d'_{s} should be evaluated together to reliably indicate true specialists.

 Our results show that most species have significantly fewer regional interactions than expected based on our null model. Thus, at the local scale species are constrained in their fundamental niche, indicating innate restrictions in their interaction partners, likely because of trait complementarity (Olesen et al. 2011, Santamaria and Rodríguez-Girones 2007). Species with the largest negative z-values tend to interact repeatedly with the same species across sites, and they therefore show high link conservatism (see also Trøjelsgaard et al. 2015). This indicates stronger constraints on the fundamental niche. Even within the species that show significant link conservatism (z-score < -1.96) some species show much stronger constraints, indicating true specialists. Interestingly, the level of conservatism seems unrelated to regional interaction richness (γ*s*) (Fig. 2). We argue that labeling a species as specialist is not necessarily dependent only upon it having few interaction partners, but that interactions are repeatedly formed with the same subset of

 species, and thus showing a strong signal of link conservatism. This might add another dimension to the discussion about species specialization vs. generalization (Ollerton et al. 2007, Rosas-Guerrero et al. 2014, Schleuning et al. 2012, Waser 2006), as the categorization as specialist or generalist might be strongly dependent upon the spatial scale at which species are investigated. For pollinators there was a significant correlation between z-scores and the measurement of regional specialization (1) $(d'_{s,meta})$ in both Brazil and the Canary Island, which further supports the designation of true specialists. It is here important to re-emphasize that our null model was framed around the neutral theory suggesting that biotic interaction to a large extent is governed by species abundance (Canard et al. 2014, Poisot et al. 2015). By ignoring any phenological and morphological constraints we may overestimate the availability of potential partners, and some species may appear more conservative than if such constraints were also taken into account. Importantly, however, large deviation form randomness also occurred among species having many regional interaction partners, where there is little evidence of phenological and morphological constraints, as they interact with a large proportion of the available partners. Therefore, incorporating such constraints would probably have minimal impact for at least some of the species. Still, with good phenological resolution and detailed morphological trait data the impact of these constraints could be explored further. Also, our approach could be further applied to investigate the role of floral traits in defining visitation fidelity, revisiting the discussion on the generalization in pollination system towards the tropics (Ollerton et al. 2009, Rosas-Guerrero et al. 2014, Schleuning et al. 2012), with consequences for plant reproductive success and adaptation to environmental changes.

 The interaction richness of a species is ultimately limited by the availability of potential partners at a given site, which, in turn, is determined by functional traits (Ibanez et al. 2016). However, according to our results, it would be sensible to distinguish between niche breath and specialization, because link conservatism (z-scores) and niche breath (regional interaction

290 Figure 1. Mean local specialization $(d'_{s,local})$ and regional specialization $(d'_{s,meta})$ are strongly positively correlated in both Brazil and the Canary islands. Plants and pollinators are represented by blue and red points, respectively. The size of the data points indicate the number of co-occurring species (i.e. the number of potential interaction partners in the metaweb). Note that plants are generally more specialized than pollinators.

 Figure 2. Null model analysis on expected regional interaction richness vs. observed regional 297 interaction richness (y_s) . A z-score below -1.96 means that the species has significantly fewer regional interaction partners than expected from random (link conservatism), and colored data points fall below this threshold. Density curves indicate the distribution of points across the spectra. Note that plants generally have more interaction partners (higher interaction richness) than pollinators. Plants and pollinators are represented by blue and red points, respectively.

303 Figure 3. Z-scores and regional specialization $(d'_{s,meta})$ show significantly negative correlations for 304 pollinators in both Brazil ($r = -0.34$, $p < 0.001$) and the Canary Islands ($r = -0.30$, $p = 0.009$). These are highlighted with trend lines. For the plant species, however, the relationships were non- significant in both Brazil and the Canary Islands and trend lines have been omitted. Plants and pollinators are represented by blue and red points, respectively.

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Fig 1

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