

Geographical Patterns in the Architecture of Neotropical Flower-visitor Networks of Hummingbirds and Insects

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Received 19 May 2020 / Accepted 16 August 2020 / Published xx September 2020

Communicated by Chih-Ming Hung

Geographical variations in environmental factors can affect species diversity and consequently influence the structure of interspecific ecological interactions. Relationships between flowering plants and animal flower visitors are among the most important ecological interactions and can structure and maintain ecological diversity in different environments. Additionally, many animal and plant species participate in these interactions, which shape the specific characteristics of these communities, both in terms of the responses of the interacting species involved and environmental differences. Therefore, in the present study we investigated geographical and environmental effects on the architecture of Neotropical flower-visitor networks of vertebrate and invertebrate animals. To this end, we used data regarding interaction networks available in the literature and constructed binary interaction networks of plants and plant-visitors (hummingbirds and insects) and tested the effects of altitude, latitude, vegetation type and number of plant families on the structure of these networks. In total, we analyzed 55 networks of flower-visitor interactions with 746 species of flower-visiting animals and 1,185 species of plants, totaling 5,463 distinct plant-animal interactions. In general, the architecture of flower-visitor networks varies along latitudinal and altitudinal gradients, with more pronounced effects for flower-insect networks where latitude influenced network size, modularity and nestedness and altitude influenced network size and connectance. Flower-hummingbird networks of open vegetation (grassland) were more modular than networks of other environments. The number of plant families positively influenced network size for insects and hummingbirds, and affected connectance and nestedness positively and modularity negatively in the

flower-insect networks. So, the patterns we found indicate that plant-visitor interactions in flower-insect and flower-hummingbird networks are differently affected by geographical and plant-related factors, possibly due to the differences in taxonomic and functional groups involved in these interactions.

Key words: Bees, Ecological services, Plant-animal interactions, Pollination, Tropical ecology.

Citation: Moreira LT, Falcão LAD, de Araújo WS. 2020. Geographical patterns in the architecture of Neotropical flower-visitor networks of hummingbirds and insects. *Zool Stud* **59**:0ii. doi:-.

BACKGROUND

Interactions between flowering plants and animal flower visitors represent one of the most important biological interactions and have fundamental consequences for the evolution and maintenance of ecological diversity (Senapathi et al. 2015). Studies indicate that about 300,000 species of angiosperm plants are pollinated by animals (Ollerton et al. 2011) and in tropical regions all plant species have some type of dependence on their flower visitors in some ecological communities (Rech et al. 2016). Therefore, ecological networks formed by flowering plants and their visitors in the Tropics have proven to be very diverse in species and very complex in interactions (review in Vizentin-Bugoni et al. 2018). The consensus in the literature is that plant-flower visitor interactions are very specialized in tropical communities, as discussed in recent thematic reviews (Ollerton 2017; Vizentin-Bugoni et al. 2018). In plant-flower visitor networks, species can be defined as specialists when they have a low number of interactions, while those with a high number of interactions are defined as generalists (Carstensen et al. 2018). Nevertheless, the hypothesis that at low latitudes plant-visitor interactions are more specialized has rarely been tested (Ollerton 2017), and the results were generally contradictory, due to the idiosyncrasy of the limited taxonomic groups that have been tested (*e.g.*, Olesen and Jordano 2002; Ollerton and Cranmer 2002; Biesmeijer et al. 2005).

The hypothesis that latitude and altitude can influence interactions is derived from the latitudinal and altitudinal gradients observed for species diversity (review in Hillebrand 2004; McCain and Grytnes 2010). A recent meta-analytical review corroborates the well-known pattern that the number of species diminishes from the equator towards the poles (Kinlock et al. 2017). There is also relative consensus regarding decreasing richness with increasing elevation (McCain

and Grytnes 2010). Latitude is expected to influence plant-visitor interactions because in very diverse communities, species tend to have more narrow ecological niches (Hillebrand 2004; Brown 2014). In this context, in tropical latitudes there is a general expectation that species of flower visiting animals frequent a low number of flower species and that each flowering plant receives few flower visitors (Vizentin-Bugoni et al. 2018). This high average species specialization means that plant-visitor networks at low latitudes have a loosely connected and very modular topology (Trøjelsgaard and Olesen 2013). A similar pattern is found for the altitudinal gradient, since considerable evidence indicates that at low altitude communities tend to be more diverse and with more specialized species (Cuartas-Hernández and Medel 2015). Despite the apparently clear patterns for latitudinal and altitudinal gradients in species diversity, studies on the effects of latitude and altitude on the structure of plant-visitor networks present contrasting results, both confirming (Trøjelsgaard and Olesen 2013; Cuartas-Hernández and Medel 2015) and contradicting expectations (Biesmeijer et al. 2005; Schleuning et al. 2012).

Plant-animal interactions also can be influenced by plant-related factors such as vegetation type and plant taxonomic diversity. For other ecological interactions, such as plant-herbivore networks, evidences point out that forest vegetation and open vegetation can differ significantly in the network structure (Araújo et al. 2020), which is related to the negative effect that the sclerophylly of open vegetation has on the palatability of plants for herbivores (Neves et al. 2010). For interactions between flowering plants and their visitors the effects of the vegetation type are expected because the higher level of environmental severity in the open vegetation can generate a greater environmental filter for the plant diversity (Kissling et al. 2008; Fründ et al. 2010; Laliberté et al. 2014), such as types of flowers. These environmental filters can also act on animal characteristics and restrict many interactions within plant-animal networks (Araújo et al. 2020). The taxonomic diversity of plants can also affect the structure of plant-flower visitor networks because each plant taxon (e.g., plant family) tends to have plant species with functionally and morphologically similar flowers (Albor et al. 2019). Thus, the greater the diversity of plant families in the network, the greater the diversity and specialization of the interactions of floral visitors must also be (Albor et al. 2019). Both types of vegetation and the taxonomic diversity of plants are expected to vary geographically, given that at low latitudes in the Neotropical region landscapes tend to be dominated by tropical rain forests that are extremely rich in plant species and families (Iwasa et al. 1995).

Various groups of invertebrate and vertebrate animals act as flower visitors and pollinating agents in Neotropical environments (Vizentin-Bugoni et al. 2018). Insects, especially Lepidoptera (butterflies and moths) and Hymenoptera (bees), are the most diverse invertebrates that visit flowers in the Neotropical region, and indeed worldwide (Ollerton 2017). Estimates show that in Brazil

alone are listed more than 26,000 species of lepidopterans and 3,000 species of bees (Lewinsohn et al. 2005). Among vertebrates, the most important group in this context is the birds, mainly the hummingbird family (Trochilidae), hosting the largest number of Neotropical flower visitors (Ollerton 2017), with 86 species recorded in Brazilian territory (Ficher et al. 2014). In the literature regarding flower-visitor interaction networks, studies comparing different taxonomic groups of visiting animals at the macroecological scale are scarce (*e.g.*, Zanata et al. 2017) and no study has focused on vertebrate and invertebrate animals simultaneously.

In the present study we investigated the geographical and environmental effects on the architecture of Neotropical flower-visitor networks of vertebrate and invertebrate animals. Thus, we compiled interactions between flowers and their visitors in communities composed of hummingbirds and insects. We characterized the plant–visitor networks using the topological descriptors network size, connectance, modularity, and nestedness (Dormann et al. 2009). The network size is measure of the number of species interacting in the network, and the connectance is a descriptor of the level of connectivity (*i.e.*, specialization) between these species (Antoniazzi et al. 2018). In turn, modularity and nestedness are measures of the modular (*i.e.*, occurrence of specialized subsets of interacting animals and plants) or nested (*i.e.*, species forming a single dense nest of interactions) arrangement of interactions between species within the network (Lewinsohn et al. 2006). Thus, we tested the following hypotheses: 1) plant-flower visitor networks are larger, less connected, more modular, and less nested at low latitudes; 2) altitude has a negative effect on network size and modularity and a positive effect on connectance and nestedness of plant-visitor networks; 3) plant-flower visitor networks of open vegetation are more specialized (*i.e.*, more diverse, less connected, less nested and more modular) than networks of forest vegetation; 4) plant taxonomic diversity have a positive effect on network size and modularity and a negative effect on connectance and nestedness of the networks; 5) plant-visitor networks composed of hummingbirds and insects have different response patterns because insects are more diverse in species and functional groups, and are expected to better reflect geographical and plant-related factors.

MATERIALS AND METHODS

Data collection

We compiled a comprehensive set of interaction data between flowering plants and their visitors of the Neotropical region (Table S1). We used data from interaction networks available on the Interaction Web Database of the National Center for Ecological Analysis and Synthesis

(www.nceas.ucsb.edu/interactionweb). Additional data were retrieved from the Google Scholar and Scopus databases using the following keywords: (plant*) AND (pollinator*) AND (floral visitors) AND (network* OR interaction*) AND (search OR list). The search for the data was carried out in December 2018 and all data available to date have been included. Additionally, we also carried out a search in the literature cited in macroecological studies on the flower-visitor interactions of insects (Biesmeijer et al. 2005) and birds (Zanata et al. 2017).

The following criteria were adopted for inclusion of studies: (1) provision of at least a basic description of the study area, containing a geographical coordinate; (2) indication of the species (or morphospecies) of flower visitors in each species (or morphospecies) of plant; (3) the network had at least five species of plants and five species of flower visitors, totaling at least 10 species; and (4) at least 80% of the visitors were identified to species level. The following data were extracted from the selected studies: geographical coordinates, altitude, country, type of vegetation and number of plant families.

Network measures

The compiled interaction data were used to build binary bipartite networks between flowering plant species their visitor species (Figs. 1, 2). We did not use quantitative data regarding interactions because for many networks this information was missing. In order to describe the structure of flower-visitor networks, we used the following network descriptors: network size, connectance, modularity and nestedness. These network descriptors were adopted because they are commonly indicated to describe the architecture of binary bipartite networks (review in Dormann et al. 2009) and they have been used in several recent studies investigating flower-visitor interaction networks (Cuartas-Hernández and Medel 2015; Zanata et al. 2017; Traveset et al. 2018; Zhao et al. 2019).

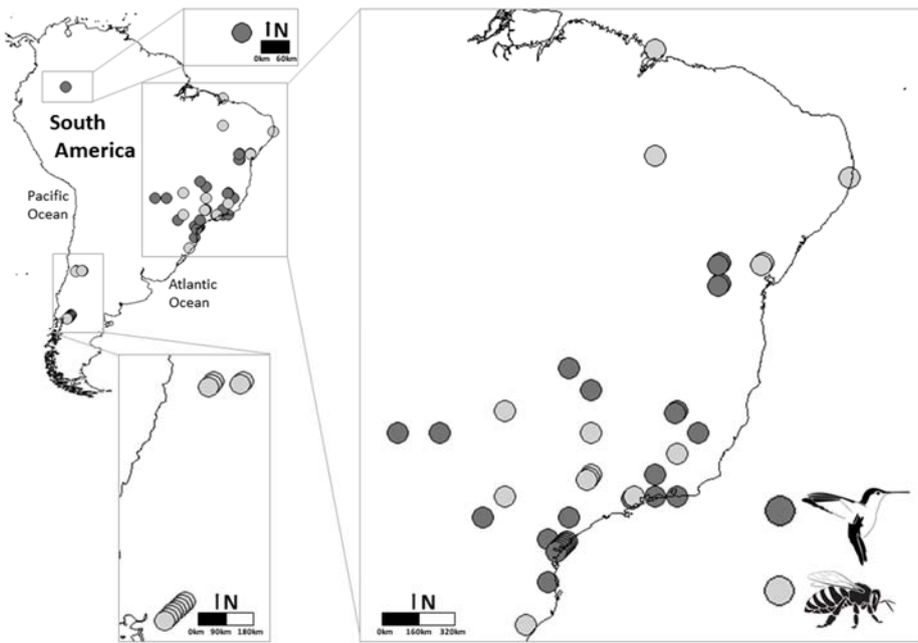


Fig. 1. Distribution of the 55 flower-visitor networks analyzed in this study. Dark gray circles represent flower-hummingbird networks and light gray circles the flower-insect networks. At this map scale some of flower-visitor networks are located so close to another that they are indistinguishable.

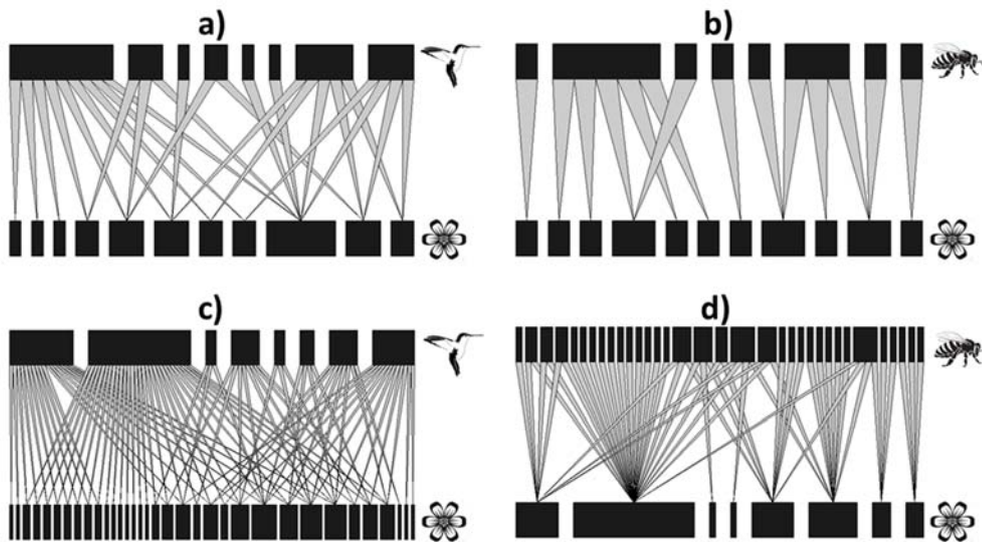


Fig. 2. Bipartite graphs showing the topological structure of examples of flower-visitor networks analyzed in this study. For each network, upper bars represent visitor species and lower bars represent flowering plant species. Bar thickness is proportional to the number of interactions of each species (drawn at different scales). a) flower-hummingbird network of Machado (2014); b) flower-insect network of Clemente (2017); c) flower-hummingbird network of Lasprilla (2003); d) flower-insect network of Vázquez and Simberloff (2002) (network 5). Networks a and b have the same total number of species (network size) and the same number of species at each trophic level

(plants and animals). Networks c and d have the same network size, although they have different numbers of species of plants and animals.

We calculated the network size by counting the total number of plants and animal species in each network (*i.e.*, the species richness). Network connectance was calculated as the ratio between the number of observed interactions and the number of possible interactions within the network (Dunne et al. 2002; Dormann et al. 2008). Connectance is an inverse measure of network specialization, and therefore greater connectance values result in lower network specialization (Araújo et al. 2015). To compute the network modularity, we used the bipartite modularity index Q (Barber 2007) through the DIRTLPAwb+ algorithm in order to detect network modules (Beckett 2016). Network nestedness was calculated using the Nestedness metric based on Overlap and Decreasing Fill (NODF) (Almeida-Neto et al. 2008). NODF accounts for the paired overlap and the decreasing fill of the matrix representing an interaction network, and takes values between 0 (perfectly non-nested) and 100 (perfectly nested). All networks were built and analyzed using the bipartite package in R (Dormann et al. 2008).

Data analyses

In addition to latitude and altitude, we also used the number of plant families and type of vegetation as explanatory variables for the flower-visitor network descriptors. We used the number of plant families as a measure of taxonomic diversity. The number of plant families also is an indirect measure of sampling effort (because to obtain networks with many plant families, more time and more sampling effort are required) and a proxy for the phylogenetic diversity in the network (since a positive correlation between the number of families and the phylogenetic diversity of plants is expected). The type of vegetation was determined for each network according to the description given by the authors of the original studies. We categorized vegetation into three types, based on the structure of the vegetation, which were grassland (vegetation predominantly composed of grasses and herbs, without canopy), savanna (vegetation with few trees, with open canopy) and forest (tree vegetation, with closed canopy).

We used generalized linear models followed by ANOVA to test the effects of latitude, altitude, type of vegetation and number of plant families on the descriptors of network structure (network size, connectance, modularity and nestedness). In order to control possible effects of network size on the network topology, we used network size as an explanatory variable in the models for connectance, modularity and nestedness (Dormann et al. 2017). Additionally, we have performed post-hoc contrast tests to highlight the differences in the network descriptors between

types of vegetation. We built different models for network descriptors of flower-hummingbird and flower-insect networks. The error distribution was assumed to be normal (Gaussian distribution) for all of the models. All statistical analyses were performed in R software (R Development Core Team 2020).

RESULTS

In total, we analyzed 55 networks of flower-visitor interactions (Fig. 1; Table S2) with 746 species of floral visiting animals and 1,185 species of plants, totaling 5,463 distinct plant-animal interactions. Of this total, 28 networks were based on hummingbirds with 429 plant species, 57 animal species and 1,411 distinct interactions. There were 27 insect-based networks in total, comprising 787 plant species, 689 visitor species and 4,052 distinct interactions. Flower-hummingbird networks ranged from 12 to 774 interactions (128.5 mean \pm 157.2 SD), while flower-insect networks ranged from 12 to 328 interactions (68.3 mean \pm 77.6 SD). Among hummingbirds, the species that interacted with the largest number of plant species were *Chlorostilbon lucidus* Shaw, 1812 ($n = 112$); *Thalurania glaucopis* Gmelin, 1788 ($n = 86$); and *Colibri serrirostris* Vieillot, 1816 ($n = 83$). Regarding insects, flower visitors were recorded in nine orders (Coleoptera, Diptera, Hemiptera, Hymenoptera, Lepidoptera, Odonata, Orthoptera, and Thysanoptera). Insect species that interacted with a greater number of plants were bees (Hymenoptera: Apidae), with an emphasis on *Apis mellifera* Linnaeus, 1758 ($n = 343$); *Trigona spinipes* Fabricius, 1793 ($n = 262$); and *Paratrigona subnuda* Schwarz, 1938 ($n = 114$).

The ranges of both latitude (0.04°S to 41.00°S) and altitude (5 to 3400 m) varied greatly between the networks analyzed. Likewise, different types of vegetation (grasslands, savannas, and forests) and a wide range of host plant families (1 to 56) were sampled in the studies compiled. The size of flower-hummingbird networks was positively affected by altitude (Fig. 3a), contrary to expected, and number of plant families (Fig. 3b), confirming our expectations (Table 1). As expected, the size of flower-insect networks was negatively affected by latitude (Fig. 3c, Table 2). On the other hand, network size of flower-insect networks was positively influenced by altitude (Fig. 3d) and number of plant families (Fig. 3e) (Table 2). The connectance of flower-hummingbird networks was not affected by any of the explanatory variables (Table 1), but the connectance of flower-insect networks was negatively influenced by altitude (Fig. 4a) and positively influenced by number of plant families (Fig. 4b) (Table 2), partially corroborating our hypothesis. Modularity in flower-hummingbird networks was affected only by type of vegetation (Table 1). As expected, the networks of open habitats (grassland) were more modular than networks of other environments

(Fig. 5a, Table 1). For insects, the network modularity had a positive relationship with latitude (Fig. 5b), and a negative relationship with number of plant families (Fig. 5c) (Table 2), contrary to our expectations. Concerning network nestedness, flower-hummingbird networks were not affected by any of the explanatory variables (Table 1), but flower-insect networks were negatively influenced by latitude (Fig. 6a) and positively influenced by number of plant families (Fig. 6b) (Table 2).

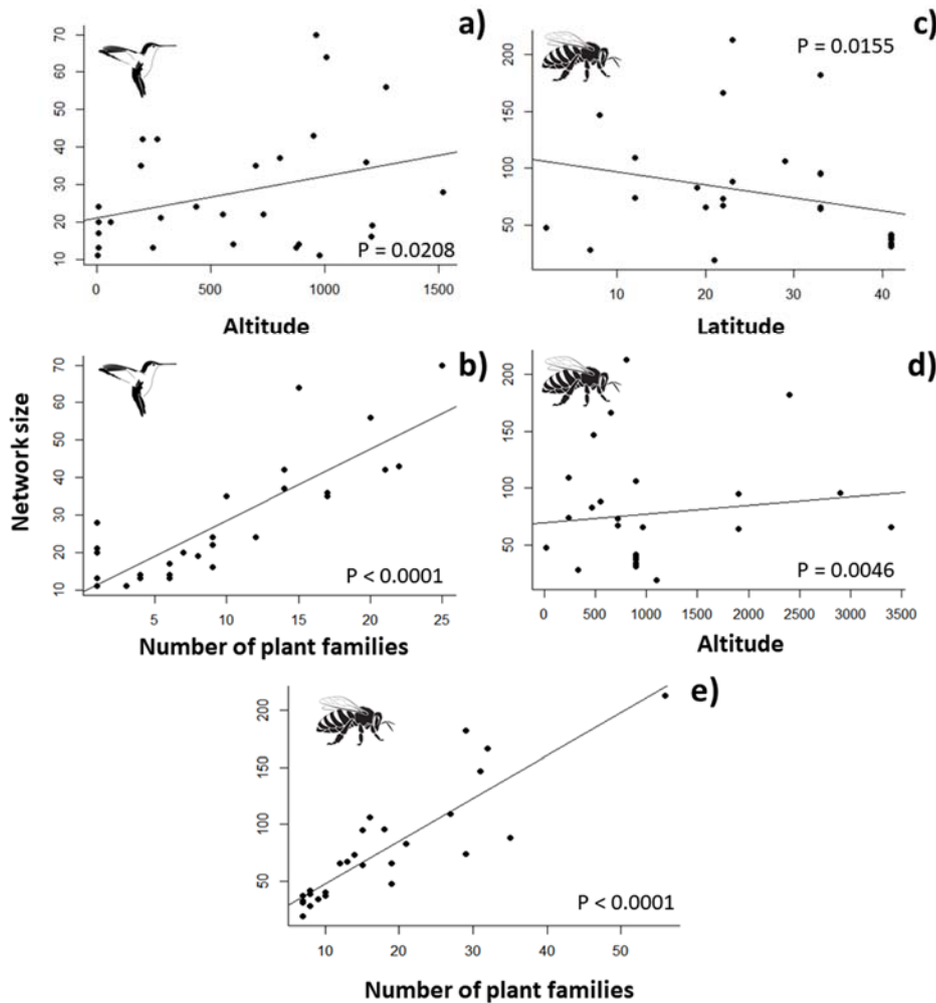


Fig. 3. Factors influencing the network size of flower-hummingbird and flower-insect networks.

Table 1. Generalized linear models of the effects of latitude, altitude, type of vegetation and number of plant families on the size, connectance, modularity and nestedness of Neotropical flower-hummingbird networks

Response variable	Explanatory variable	Df	Resid. Dev.	F-value	P-value
Network size	Latitude	23	6310.8	1.319	0.2650
	Altitude	22	5851.9	6.360	0.0208
	Vegetation type	20	5656.2	1.357	0.2814
	Number of plant families	19	1370.9	59.393	< 0.0001
Network connectance	Latitude	23	0.0842	0.2664	0.6121
	Altitude	22	0.0807	0.9742	0.3367

	Vegetation type	20	0.0763	0.5993	0.5598
	Number of plant families	19	0.0729	0.9139	0.3517
	Network size	18	0.0659	1.9336	0.1813
Network modularity	Latitude	23	0.1796	1.437	0.2462
	Altitude	22	0.1578	3.9513	0.0623
	Vegetation type	20	0.1049	4.7872	0.0215
	Number of plant families	19	0.1010	0.7118	0.4099
	Network size	18	0.0993	0.2992	0.5911
Network nestedness	Latitude	23	5366.3	2.9975	0.1005
	Altitude	22	5047.5	1.5176	0.2338
	Vegetation type	20	4273.8	1.8414	0.1873
	Number of plant families	19	4040.1	1.1128	0.3054
	Network size	18	3781.2	1.2321	0.2816

Table 2. Generalized linear models of the effects of latitude, altitude, type of vegetation and number of plant families on the size, connectance, modularity and nestedness of Neotropical flower-insect networks

Response variable	Explanatory variable	Df	Resid. Dev.	F-value	P-value
Network size	Latitude	24	53100.0	7.009	0.0155
	Altitude	23	47237.0	10.196	0.0046
	Vegetation type	21	43330.0	3.398	0.0537
	Number of plant families	20	11501.0	55.348	< 0.0001
Network connectance	Latitude	24	0.1164	0.2857	0.5992
	Altitude	23	0.0936	8.2197	0.0099
	Vegetation type	21	0.0923	0.2329	0.7944
	Number of plant families	20	0.0723	7.2036	0.0147
	Network size	19	0.0528	7.0064	0.0159
Network modularity	Latitude	24	0.2992	7.3854	0.0137
	Altitude	23	0.2938	0.6468	0.4312
	Vegetation type	21	0.2646	1.7285	0.2043
	Number of plant families	20	0.1610	12.2859	0.0024
	Network size	19	0.1602	0.0863	0.7721
Network nestedness	Latitude	24	4757.5	10.0281	0.0051
	Altitude	23	4294.9	4.0142	0.0596
	Vegetation type	21	3939.8	1.5406	0.2398
	Number of plant families	20	3180.9	6.5863	0.0189
	Network size	19	2189.4	8.6051	0.0085

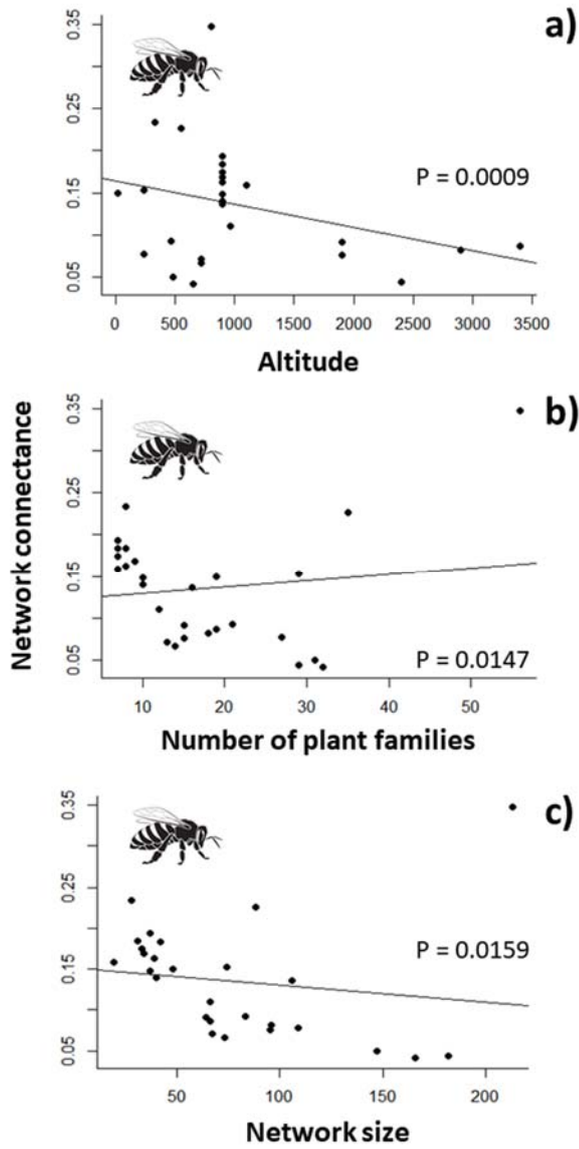


Fig. 4. Factors influencing the network connectance of flower-insect networks.

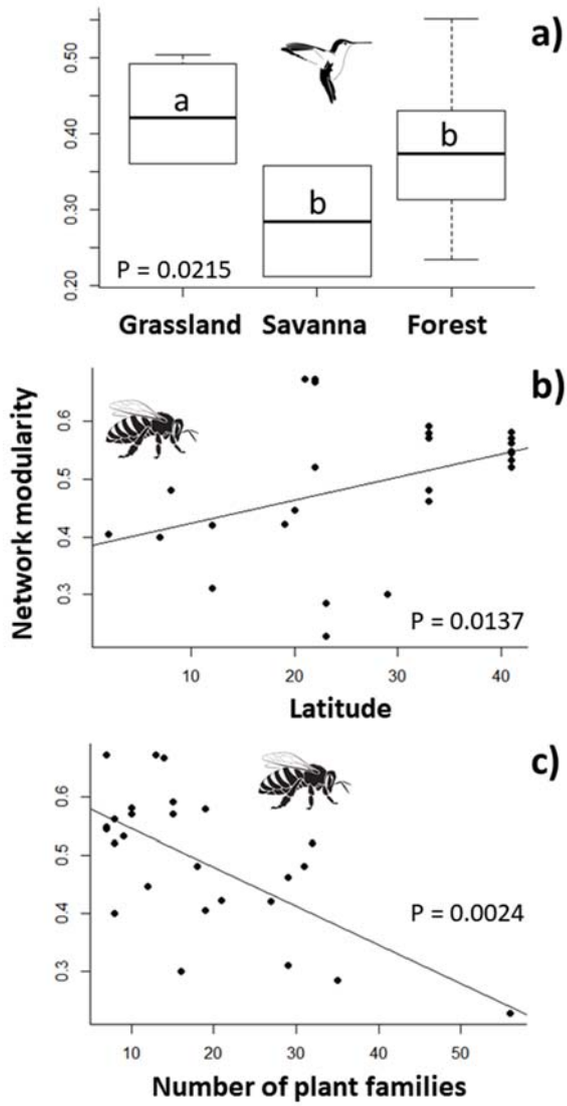


Fig. 5. Factors influencing the network modularity of flower-hummingbird and flower-insect networks.

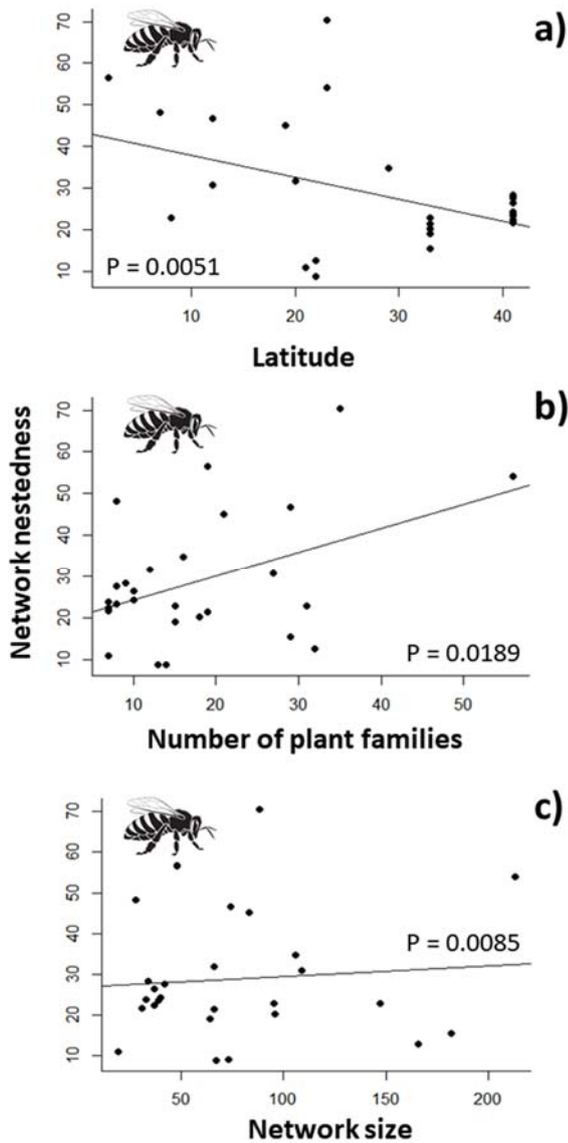


Fig. 6. Factors influencing the network nestedness of flower-insect networks.

DISCUSSION

Our results revealed that the effects of latitude and altitude were more pronounced for flower-insect networks. The latitude negatively influenced the network size, positively the modularity and negatively the nestedness, while the altitude positively affected the network size and negatively the connectance for insects. These results corroborate recent studies indicating geographical effects in the structure of plant-insect visitor networks (Cuartas-Hernández and Medel 2015; Kelly and Elle 2020). For flower-hummingbird networks the only effect that we observed was the unexpected positive relationship between network size and altitude. In this context, the results observed in our study point in the opposite direction to that which was expected, with lower network specialization at low latitudes and high altitudes. However, the patterns found indicate that

plant-visitor interactions in flower-insect and flower-hummingbird networks are differently organized along latitudinal and altitudinal ranges. We also found that the taxonomic diversity of plants positively affected size, connectance and nestedness and negatively affected the modularity of flower-insect networks. On the other hand, flower-hummingbird networks had size positively related to the taxonomic diversity of plants and were more modular in open habitats (grassland) than in the other vegetation.

Topological descriptors of flower-insect networks changed considerably with increasing latitude. The size of the networks (i.e., the richness of animals and plants in the community) was negatively affected by latitude in the flower-insect networks. This finding is in line with the well-documented pattern that species diversity decreases from the equator towards the poles (review in Kinlock et al. 2017). However, latitude influenced modularity positively and nestedness negatively, contrary to our expectations and the pattern found in many previous studies (Olesen and Jordano 2002; Trøjelsgaard and Olesen 2013; Zanata et al. 2017). It is important to note that these results are not due to the dependence of modularity and nestedness on the size or connectance of the networks, as observed in other studies (Dunne et al. 2002; González et al. 2015), since in our study these parameters displayed the opposite pattern or did not vary, respectively. On the other hand, our results corroborate the patterns found by Schleuning et al. (2012) for pollination networks and Dalsgaard et al. (2017) for dispersal networks, showing that specialization decreases toward tropical latitudes. These findings suggest that high tropical diversity can often generate less specialized topological patterns in ecological networks.

Altitude positively affected network size (species richness) in both flower-insect and flower-hummingbird networks, contrary to our expectations. In addition, we observed a negative effect of altitude on the connectance of flower-insect networks, also contradicting our hypothesis. We observed that the largest networks are located in a range that extends from 1000 to 2500 m in altitude (see Fig. 3). Considering that altitudes in our study ranged up to 3400 meters, our results indicate a peak of species richness (animals and plants) in intermediate to high altitudes. Our observations agree with Rahbek (1995), who, in a thorough review of the literature, showed that in tropical communities higher species diversity is very frequently reported at intermediate altitudes. This pattern is probably due to the intermediate altitudes having mixed environmental conditions that allow the occurrence of both species of higher altitudes and species typical of low altitude environments (Rahbek 1995). Although these are rare, some studies showed a positive correlation between altitude and species richness, as documented by Rohde (1992) for tropical birds. Similarly, Hortal et al. (2013) showed a positive hump-shaped relationship between elevation and bird species richness in Spain, which can be attributed to the greater diversity of habitats at intermediate elevations. In this context, another factor that may explain the larger species richness of the

networks at intermediate and high altitudes is the conservation status of these habitats, which in general tend to be much better preserved against human interventions (Paudel and Šipoš 2014). For connectance of flower-insect networks, which decreases with altitude, we believe that the observed pattern is a reflection of the size of the networks. Corroborating this, we found a negative correlation between the size and connectance of flower-insect networks, as expected. This finding corroborates previous studies with ecological networks (*e.g.*, Dunne et al. 2002; Dormann et al. 2009; Araújo et al. 2015), and can be explained by the number of possible interactions increasing much faster (*i.e.*, geometric progression) with the size of the networks, than the number of observed interactions (*i.e.*, arithmetic progression) (Dunne et al. 2002).

The architecture of Neotropical flower-visitor networks featuring both hummingbirds and insects showed some very interesting differences in responses to geographical and environmental variations. Flower-insect networks were much more variable along the latitudinal and altitudinal gradients (network size, connectance, modularity and nestedness) whereas flower-hummingbird networks varied only in network size. We believe that these differences can be attributed to the intrinsic characteristics of these networks. For example, networks compiled in our dataset of flower visiting insects were characterized by different groups of animals (*e.g.*, bees, butterflies and others) (Rech et al. 2014). This great diversity of taxonomic and functional groups within flower-insect networks can generate more variable responses along the geographic gradient (Adedoja et al. 2018). On the other hand, flower-hummingbird networks are made up of a single animal group, which results in strongly phylogenetically-structured interactions (González et al. 2015). This pattern suggests that, due to phylogenetic restrictions, flower-hummingbird networks tend to be structured independently of latitude and altitude, because we did not find any variation in the connectivity or in the arrangement of their interactions. Another factor may be the breadth of geographic distribution of the studies considered, because flower-insect networks were distributed over a wider latitudinal range (see Fig. 1), which allows for greater plasticity in responses. This observation indicates the need for further studies on interactions between flowering plants and hummingbirds (and other floral visiting vertebrates, such as other birds and bats) in Neotropical areas of higher altitudes and latitudes.

By using vegetation type and the number of plant families as explanatory variables in our analyses, we tested the possible plant-related effects related to the characteristics of the habitats and the taxonomic diversity (a proxy for sampling effort) of the studies, respectively. Type of vegetation influenced flower-hummingbird network modularity with networks of grassland habitats being more modular than that of other environments. As expected, this pattern likely is due to the hummingbird species that occur in open vegetation have more diverse and specialized diets, as a consequence of the severe environmental filters acting on plant-animal interactions (Araújo et al.

2020). Our finding is in concordance with study of Rodrigues and Rodrigues (2015) that, although performed on a local scale, points out differences in hummingbird diversity between forest patches and the open rocky field in Brazil. The number of plant families positively influenced network size for both flower-insect and flower-hummingbird networks. Exclusively for flower-insect networks, number of plant families affected connectance and nestedness positively and modularity negatively. These results indicate that the higher the taxonomic diversity (a proxy of the level of taxonomic inclusion), the greater the diversity of species and the generality of interactions recorded in the network. Contrary to expectations, our results point to a less specialized (that is, more connected and less modular) topology in networks with more plant families. One possible explanation for this pattern is that there is a high overlap in the floral visiting insects between different plant families, so that when the number of botanical families increases, the connectance and also the nestedness of the interactions increases. Anyway, our study corroborates previous studies indicating that taxonomic and sampling amplitudes are important factors influencing the topology of ecological networks (Nielsen and Bascompte 2007; Trøjelsgaard and Olesen 2013; Araújo et al. 2015).

CONCLUSIONS

Our findings revealed interesting and contrasting geographical patterns of Neotropical flower-visitor networks composed of insects and hummingbirds. Our results indicated that flower-insect networks vary geographically in both species richness and network architecture, but contrary to expectations, topological descriptors showed lower network specialization near the equator. Although unexpected, these results corroborate some studies showing that network specialization decreases toward tropical latitudes (Schleuning et al. 2012; Dalsgaard et al. 2017). Flower-hummingbird networks, on the other hand, varied in species richness, but the network structure remained unchanged along the latitudinal and altitudinal gradients. In conclusion, our results suggest that relations between geography and interaction specialization in Neotropical flower-visitor communities are very complex and can vary between taxonomic and functional zoological groups.

Acknowledgments: The authors are thankful to F. Costa and L. Leite for their valuable suggestions to manuscript; and the Programa de Pós-Graduação em Biodiversidade e Uso dos Recursos Naturais and State University of Montes Claros for providing logistical support.

Authors' contributions: LTM and WSA planned and conceived the study, LTM compiled the database, LTM and WSA performed data analyses LTM, LADF, and WSA wrote the manuscript.

Competing interests: LTM, LADF, and WSA declare that they have no conflict of interest.

Availability of data and materials: All of the authors agreed with the publication of the data (supplementary material).

Consent for publication: All of the authors agreed to publish the paper.

Ethics approval consent to participate: Not applicable.

REFERENCES

- Adedoja OA, Kehinde T, Samways MJ. 2018. Insect-flower interaction networks vary among endemic pollinator taxa over an elevation gradient. *PLoS ONE* **13**:e0207453. doi:10.1371/journal.pone.0207453.
- Albor C, García-Franco JG, Parra-Tabla V, Díaz-Castelazo C, Arceo-Gómez G. 2019. Taxonomic and functional diversity of the co-flowering community differentially affect *Cakile edentula* pollination at different spatial scales. *J Ecol* **107**:2167–2181. doi:10.1111/1365-2745.13183.
- Almeida-Neto M, Guimaraes P, Guimaraes Jr PR, Loyola RD, Ulrich W. 2008. A consistent metric for nestedness analysis in ecological systems: reconciling concept and measurement. *Oikos* **117**:1227–1239. doi:10.1111/j.0030-1299.2008.16644.x.
- Antoniazzi R, Dáttilo W, Rico-Gray V. 2018. A useful guide of main indices and software used for ecological networks studies. *In*: Dáttilo W, Rico-Gray V (eds) *Ecological networks in the tropics*, vol 1. Cham, Springer, pp. 185–196. doi:10.1007/978-3-319-68228-0_13
- Araújo WS, Vieira MC, Lewinsohn TM, Almeida-Neto M. 2015. Contrasting effects of land use intensity and exotic host plants on the specialization of interactions in plant-herbivore networks. *PLoS ONE* **10**:e0115606. doi:10.1371/journal.pone.0115606.
- Araújo WS, Freitas ÉVD, Silveira LT, Daud RD. 2020. Network structure of interactions between phytophagous mites and their host-plants in natural ecosystems in Brazil. *Syst Appl Acarol* **25**:821–832. doi:10.11158/saa.25.5.4.
- Barber MJ. 2007. Modularity and community detection in bipartite networks. *Phys Rev E* **76**:066102. doi:10.1103/PhysRevE.76.066102.

- Beckett SJ. 2016. Improved community detection in weighted bipartite networks. *R Soc Open Sci* **3**:140536. doi:10.1098/rsos.140536.
- Biesmeijer JC, Slaa EJ, Castro MSD, Viana BF, Kleinert ADM, Imperatriz-Fonseca VL. 2005. Connectance of Brazilian social bee: food plant networks is influenced by habitat, but not by latitude, altitude or network size. *Biota Neotrop* **5**:85–93. doi:10.1590/S1676-06032005000100010.
- Brown JH. 2014. Why are there so many species in the tropics? *J Biogeogr* **41**:8–22. doi:10.1111/jbi.12228.
- Carstensen DW, Trøjelsgaard K, Ollerton J, Morellato LPC. 2018. Local and regional specialization in plant–pollinator networks. *Oikos* **127**:531–537. doi:10.1111/oik.04436.
- Clemente MA, Campos NR, Viera KM, Del-Claro K, Prezoto F. 2017. Social wasp guild (Hymenoptera: Vespidae) visiting flowers in two of the phytophysiognomic formations: Riparian Forest and campos rupestres. *Sociobiology* **64**:217–224. doi:10.13102/sociobiology.v64i2.1364.
- Cuartas-Hernández S, Medel R. 2015. Topology of Plant - Flower-Visitor Networks in a Tropical Mountain Forest: Insights on the Role of Altitudinal and Temporal Variation. *Plos One* **10**:e0141804. doi:10.1371/journal.pone.0141804.
- Dalsgaard B, Schleuning M, Maruyama PK, Dehling DM, Sonne J, Vizentin-Bugoni J, Zanata TB, Fjeldså J, Böhning-Gaese K, Rahbek, C. 2017. Opposed latitudinal patterns of network-derived and dietary specialization in avian plant-frugivore interaction systems. *Ecography* **40**:1395–1401. doi:10.1111/ecog.02604.
- Dormann CF, Gruber B, Frund J. 2008. Introducing the bipartite Package: Analysing ecological networks. *R News* **8**:8–11.
- Dormann CF, Fründ J, Blüthgen N, Gruber B. 2009. Indices, graphs and null models: analyzing bipartite ecological networks. *Open J Ecol* **2**:7–24. doi:10.2174/1874213000902010007.
- Dormann CF, Fründ J, Schaefer HM. 2017. Identifying causes of patterns in ecological networks: opportunities and limitations. *Ann Rev Ecol Evol Syst* **48**:559–584. doi:10.1146/annurev-eolsys-110316-022928.
- Dunne JA, Williams RJ, Martinez ND. 2002. Food-web structure and network theory: the role of connectance and size. *Proc Natl Acad Sci USA* **99**:12917–12922. doi:10.1073/pnas.192407699.
- Ficher E, Araujo AC, Gonçalves F. 2014. Polinização por vertebrados. In: Rech AR, Agostini K, Machado ICS, Oliveira PEAM (eds) *Biologia da polinização*, vol. 1, Projeto Cultural, Rio de Janeiro, pp. 311–326.

- Fründ J, Linsenmair KE, Blüthgen N. 2010. Pollinator diversity and specialization in relation to flower diversity. *Oikos* **119**:1581–1590. doi:10.1111/j.1600-0706.2010.18450.x.
- González AMM, Dalsgaard B, Nogués-Bravo D, Graham CH, Schleuning M, Maruyama PK, Abrahamczyk S, Alarcón R, Araujo AC, Araújo FP, Azevedo SM, Baquero AC, Cotton PA, Ingversen TT, Kohler G, Lara C, Las-Casas FMG, Machado AO, Machado CG, Maglianesi MA, McGuire JA, Moura AC, Oliveira GM, Oliveira PE, Ornelas JF, Rodrigues da Cruz L, Rosero-Lasprilla L, Rui AM, Sazima M, Timmermann A, Varassin IG, Vizentin-Bugoni J, Wang Z, Watts S, Rahbek C, Martinez ND. 2015. The macroecology of phylogenetically structured hummingbird-plant networks. *Global Ecol Biogeogr* **24**:1212–1224. doi:10.1111/geb.12355.
- Hillebrand H. 2004. On the generality of the latitudinal diversity gradient. *Am Nat* **163**:192–211. doi:10.1086/381004.
- Hortal J, Carrascal LM, Triantis KA, Thébault E, Meiri S, Sfenthourakis S. 2013. Species richness can decrease with altitude but not with habitat diversity. *Proc Natl Acad Sci USA* **110**:E2149–E2150. doi:10.1073/pnas.1301663110.
- Iwasa Y, Kubo T, Sato K. 1995. Maintenance of forest species diversity and latitudinal gradient. *Vegetatio* **121**:127–134. doi:10.1007/BF00044678.
- Kelly T, Elle E. 2020. Effects of community composition on plant–pollinator interaction networks across a spatial gradient of oak-savanna habitats. *Oecologia* **193**:211–223. doi:10.1007/s00442-020-04661-5.
- Kinlock NL, Prowant L, Herstoff EM, Foley CM, Akin-Fajiyeh M, Bender N, Umarani M, Ryu HY, Şen B, Gurevitch J. 2018. Explaining global variation in the latitudinal diversity gradient: Meta-analysis confirms known patterns and uncovers new ones. *Global Ecol Biogeogr* **27**:125–141. doi:10.1111/geb.12665.
- Kissling WD, Field R, Böhning-Gaese K. 2008. Spatial patterns of woody plant and bird diversity: functional relationships or environmental effects? *Glob Ecol Biogeogr* **17**:327–339. doi:10.1111/j.1466-8238.2007.00379.x.
- Laliberté E, Zemunik G, Turner BL. 2014. Environmental filtering explains variation in plant diversity along resource gradients. *Science* **345**:1602–1605. doi:10.1126/science.1256330.
- Lasprilla LR. 2003. Interações planta/beija-flor em três comunidades vegetais da parte sul do Parque Nacional Natural Chiribiquete, Amazonas (Colômbia). PhD Dissertation, Universidade Estadual de Campinas.
- Lewinsohn TM, Freitas AVL, Prado PI. 2005. Conservation of terrestrial invertebrates and their habitats in Brazil. *Conserv Biol* **19**:640–645. doi:10.1111/j.1523-1739.2005.00682.x.

- Lewinsohn TM, Inácio Prado P, Jordano P, Bascompte J, Olesen J. 2006. Structure in plant–animal interaction assemblages. *Oikos* **113**:174–184. doi:10.1111/j.0030-1299.2006.14583.x.
- Olesen JM, Jordano P. 2002. Geographic patterns in plant–pollinator mutualistic networks. *Ecology* **83**:2416–2424. doi:10.1890/0012-9658%282002%29083%5B2416%3AGPIPPM%5D2.0.CO%3B2.
- Ollerton J, Winfree R, Tarrant S. 2011. How many flowering plants are pollinated by animals? *Oikos* **120**:321–326. doi:10.1111/j.1600-0706.2010.18644.x.
- Ollerton J. 2017. Pollinator diversity: distribution, ecological function, and conservation. *Annu Rev Ecol Evol Syst* **48**:353–376. doi:10.1146/annurev-ecolsys-110316-022919.
- Ollerton J, Cranmer L. 2002. Latitudinal trends in plant-pollinator interactions: are tropical plants more specialized? *Oikos* **98**:340–350. doi:10.1034/j.1600-0706.2002.980215.x.
- Machado CG. 2014. A comunidade de beija-flores e as plantas que visitam em uma área de cerrado ralo da Chapada Dimantina, Bahia, Brasil. *Bioscience J* **30**:1578–1587.
- McCain CM, Grytnes JA. 2010. Elevational Gradients in Species Richness. In: *Encyclopedia of Life Sciences*. John Wiley and Sons, Chichester, pp. 1–10.
- Neves FS, Araújo LS, Espírito-Santo MM, Fagundes M, Fernandes GW, Sanchez-Azofeifa GA, Quesada M. 2010. Canopy herbivory and insect herbivore diversity in a dry forest-Savanna transition in Brazil. *Biotropica* **42**:112–118. doi:10.1111/j.1744-7429.2009.00541.x.
- Nielsen A, Bascompte J. 2007. Ecological networks, nestedness and sampling effort. *J Ecol* **95**:1134–1141. doi:10.1111/j.1365-2745.2007.01271.x.
- Paudel PK, Šipoš J. 2014. Conservation status affects elevational gradient in bird diversity in the Himalaya: A new perspective. *Global Ecol Biogeogr* **2**:338–348. doi:10.1016/j.gecco.2014.10.012.
- Pinheiro M, Gaglianone MC, Nunes CEP, Sigrist MR. 2014. Polinização por abelhas. In: Rech AR, Agostini K, Machado ICS, Oliveira PEAM (eds) *Biologia da polinização*, vol 1. Projeto Cultural, Rio de Janeiro, pp. 207–233.
- R Development Core Team. 2020. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. <https://www.R-project.org>. Accessed dd mm year.
- Rahbek C. 1995. The elevational gradient of species richness: a uniform pattern? *Ecography* **18**:200–205. doi:10.1111/j.1600-0587.1995.tb00341.x.
- Rech AR, Agostini K, Machado ICS, Oliveira PEAM (eds). 2014. *Biologia da polinização*. Projeto Cultural, Rio de Janeiro.
- Rech AR, Dalsgaard B, Sandel B, Sonne J, Svenning JC, Holmes N, Ollerton J. 2016. The macroecology of animal versus wind pollination: ecological factors are more important than

- historical climate stability. *Plant Ecol Divers* **9**:253–262.
doi:10.1080/17550874.2016.1207722.
- Rodrigues LC, Rodrigues M. 2015. Floral resources and habitat affect the composition of hummingbirds at the local scale in tropical mountaintops. *Braz J Biol* **75**:39–48.
doi:10.1590/1519-6984.06913.
- Rohde K. 1992. Latitudinal gradients in species diversity: the search for the primary cause. *Oikos* **65**:514–527. doi:10.2307/3545569.
- Senapathi D, Biesmeijer JC, Breeze TD, Kleijn D, Potts SG, Carnevalheiro LG. 2015. Pollinator conservation—the difference between managing for pollination services and preserving pollinator diversity. *Curr Opin Insect Sci* **12**:93–101. doi:10.1016/j.cois.2015.11.002.
- Schleuning M, Fründ J, Klein AM, Abrahamczyk S, Alarcón R, Albrecht M, Andersson GK, Bazarian S, Böhning-Gaese K, Bommarco R. 2012. Specialization of mutualistic interaction networks decreases toward tropical latitudes. *Curr Biol* **22**:1925–1931.
doi:10.1016/j.cub.2012.08.015.
- Traveset A, Castro-Urgal R, Rotllàn-Puig X, Lázaro A. 2018. Effects of habitat loss on the plant–flower visitor network structure of a dune community. *Oikos* **127**:45–55.
doi:10.1111/oik.04154.
- Trøjelsgaard K, Olesen JM. 2013. Macroecology of pollination networks. *Global Ecol Biogeogr* **22**:149–162. doi:10.1111/j.1466-8238.2012.00777.x.
- Vázquez DP, Simberloff D. 2002. Ecological specialization and susceptibility to disturbance: conjectures and refutations. *Am Nat* **159**:606–623. doi:10.1086/339991.
- Vizentin-Bugoni J, Maruyama PK, de Souza CS, Ollerton J, Rech AR, Sazima M. 2018. Plant–pollinator networks in the tropics: a review. *In*: Dáttilo W, Rico-Gray V (eds) *Ecological networks in the tropics*, vol 1. Cham, Springer, pp. 73–91. doi:10.1007/978-3-319-68228-0_6.
- Zanata TB, Dalsgaard B, Passos FC, Cotton PA, Roper JJ, Maruyama PK, Fisher E, Schleuning M, Martín González AM, Vizentin-Bugoni J, Franklin D, Abrahamczyk S, Alarcon R, Araújo A, Araujo F, Azevedo-Júnior S, Baquero A, Böhning-Gaese K, Carstensen D, Chupil H, Coelho A, Faria R, Horak D, Ingwersen T, Janecek S, Kohler G, Las-Casas FM, Lopes A, Machado A, Machado CG, Machado IC, Maglianesi AM, Malucelli T, Mohd-Azlan A, Moura AC, Oliveira G, Oliveria PE, Ornelas JF, Riegert J, Rodrigues L, Lasprilla L, Rui AM, Sazima M, Schmid B, Sedlacek O, Timmermann A, Vollstädt M, Wang Z, Watts S, Rahbek C, Varassin IG. 2017. Global patterns of interaction specialization in bird–flower networks. *J Biogeogr* **44**:1891–1910.
- Zhao YH, Lázaro A, Ren ZX, Zhou W, Li HD, Tao ZB, Xu K, Wu ZK, Wolfe LM, Li D, Z, Wang H. 2019. The topological differences between visitation and pollen transport networks: a

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comparison in species rich communities of the Himalaya–Hengduan Mountains. *Oikos* **128**:551–562.

Supplementary materials

Table S1. References of the 55 flower-visitor networks used in the present study. (download)

Table S2. Characteristics of the 55 flower-visitor networks used in the present study. Reference of original studies are listed in the table S1. (download)