Evaluating factors that predict the structure of a commensalistic epiphyte–phorophyte network

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A central issue in ecology is the understanding of the establishment of biotic interactions. We studied the factors that affect the assembly of the commensalistic interactions between vascular epiphytes and their host plants. We used an analytical approach that considers all individuals and species of epiphytic bromeliads and woody hosts and non-hosts at study plots. We built models of interaction probabilities among species to assess if host traits and abundance and spatial overlap of species predict the quantitative epiphyte–host network. Species abundance, species spatial overlap and host size largely predicted pairwise interactions and several network metrics. Wood density and bark texture of hosts also contributed to explain network structure. Epiphytes were more common on large hosts, on abundant woody species, with denser wood and/or rougher bark. The network had a low level of specialization, although several interactions were more frequent than expected by the models. We did not detect a phylogenetic signal on the network structure. The effect of host size on the establishment of epiphytes indicates that mature forests are necessary to preserve diverse bromeliad communities.

1. Introduction

An important aim in community ecology is to identify the factors that drive the establishment of species interactions. Mutualisms, antagonisms and more recently commensalisms have been depicted as complex networks [1–3], and have revealed patterns of community organization common in ecological systems, such as the tendency of species to interact with subsets of the interaction partners of more generalized species (nestedness, [4]). Quantitative differences in network properties among these interaction types have been identified [5–7], indicating differences in the underlying structuring factors. If interactions are neutral, the structure of the network is explained by the random interaction among individuals in the community and species abundance determines network patterns [8]. However, deterministic factors can also affect the structure of a network. There has been debate on whether neutral or biological factors such as complementarity in species phenotypes determine structural patterns of networks [9–12]. Phylogenetic analyses can help understand network topology because evolutionary history can influence ecological interactions [13,14]. Recent advances have been made in identifying the factors that influence network structure in mutualisms [12,15–17] and antagonisms [18–20], though there is a lag in the study of commensalistic interactions. Network structures may be the result of multiple, hierarchical, non-exclusive interactions among factors [21], but few studies have assessed several factors simultaneously [12,15,18]. Identifying the factors that structure interaction...
networks provides novel insights into the ecological and evolutionary processes at the community level that shape interactions, with possible implications for the management and conservation of species, particularly for groups with high vulnerability to extinction.

Vascular epiphytes (henceforth referred to as epiphytes) are plants that establish obligate interactions with other plants (phorophytes) by using them as a substrate without parasitizing them [22]. This makes them a vulnerable group to anthropogenic disturbance, together with their long life cycles, slow growth and the stressing environment in which they live [22]. Epiphytes represent 10 per cent of the diversity of living vascular plants [23] attaining their highest diversity in the Neotropics [24]. Studies of networks have found high values of nestedness [3,5,25]; no phylogenetic signal in species interaction patterns [26]; and the generalization of species partly or fully explained by species abundance [5,25]. However, no attempt has been made to incorporate other explanatory variables that might determine the structure of epiphyte–phorophyte networks. Certain host traits are associated with the presence of epiphytes. For example, bark ornamentation affects seed establishment [27,28]; bark porosity affects the humidity of the substrate [28,29]; and the production of secondary compounds can inhibit the germination of epiphytes [29,30]. Host tree size is related to epiphyte diversity and abundance [27,31,32]. Larger trees have more complex structures, providing more microhabitats and more substrate area for seeds to land. Tree size is also related to age, and older trees tend to have more epiphytes than younger trees because they have been exposed for a longer period of time to the seed rain of epiphytes. Additionally, bigger branches provide more stability, as these branches are less likely to fail [33,34]. Wood density is expected to affect epiphyte–phorophyte interactions because it influences branch stability, and because species with dense wood have slower growth rates [20] that affects the time of exposure to seed rain when tree size is considered. Thus, the combination of traits in a host species will influence the colonization, establishment and success of the interaction.

One of the main focuses in species interactions is their specialization. The observed specificity of epiphytes on host species may be owing to their specialization or to sampling effects posed by community attributes of the flora, such as species richness and abundance [21,22]. Low diversity forests have only few potential phorophyte species, and epiphytes may appear as highly specialized. On the other hand, species in forests with high alpha diversity have low densities, and rare species are seldom recorded, appearing more specialized than common species. Community studies have only considered common host, and sometimes epiphyte species even in network studies [3,25,31,35,36], and have used null models incorporating species abundance to overcome this problem when assessing specialization [3,25,31,36]. A further limitation to assess specialization (or neutrality) has been the exclusion of absence data, i.e. trees that do not bear epiphytes, because the real abundance of host species or species that lack epiphytes is not considered [22,37] (but see [31]).

Here, we assess which factors predict the structure of a quantitative epiphyte–phorophyte network in a tropical dry forest in western Mexico. We hypothesize that bark texture, size of woody individuals and wood density of woody species, together with the abundance and spatial overlap of species, determine network level patterns of species interactions and specialization. We evaluate the relative contribution of these factors to network structure, and whether patterns of interactions are conserved in the phylogeny of epiphytes and phorophytes.

2. Material and methods

Epiphyte–phorophyte networks in the tropical dry forest offer a great opportunity to investigate which factors contribute to emergent patterns of network organization because epiphyte and woody individuals can be easily recorded during the leafless season of this low canopy forest. This is an important advantage because excluding species and individuals that do not interact precludes the appropriate assessment of factors that may limit the establishment of interactions. We focus on the epiphytic bromeliads of a tropical dry forest in western Mexico. Rather than trying to document all possible interactions in the system, we focused on assessing if the factors considered can explain the network structure given the pool of individuals and species at the study sites.

(a) Study area and field data

The work was conducted in conserved tropical dry forest at the Chamehuixmala Biosphere Reserve and its vicinity, in the central western coast of Mexico in the state of Jalisco (19°22’–19°35’S, 105°56’–105°03’W). The climate is very seasonal; the wet season occurs between July and October [38]. In this forest there are 421 woody species (potential phorophytes) reported and 29 vascular epiphyte species [39] distributed among the families Cactaceae (1 species), Orchidaceae (10 species) and Bromeliaceae (18 species). We focused on the Bromeliaceae, a monophyletic family of the monocots, which is the family with the highest epiphyte species richness and abundance in the region, and contributes importantly to the diversity of epiphytes in Neotropical forests [37].

We registered all interactions between bromeliad epiphytes and woody species at three 20 × 20 m plots (19°30.065’N, 105°02.584’ W; 19°30.532’N, 105°02.410’ W; 19°24.297’N, 104°58.968’ W). We registered all epiphytes larger than approximately 3 cm in height, which can be confidently seen and identified from the ground using binoculars. Data on epiphytic bromeliads was collected during the dry season of 2007/2008 (from November to May) when the canopy is leafless and bromeliads are easy to observe. Two Tillandsia species have the potential of secondary dispersal by asexual means (Tillandsia intermedia, R. Sayago 2007, personal observation; Tillandsia usneoides, [37]) when ramets (clones) detach. We defined an individual as an epiphyte physically separated from other epiphytes.

All the woody plants at the plots with a diameter at 1.3 m above ground (trees and shrubs) or at the base (lianas) greater than 2.5 cm (DBH henceforth) were marked with an aluminium tag. For each marked individual we collected the following information: (i) DBH, (ii) species identity, and (iii) the abundance of each epiphytic bromeliad species. Leaf and reproductive samples were collected for each woody species. Plant identification was conducted by the authors and corroborated with herbarium specimens preserved at the Chamehuixmala Biological Station (Universidad Nacional Autonoma de Mexico). We refer to phorophytes only when referring to woody individuals bearing epiphytes, and to woody species to refer to all potential phorophytes present at the plots.

Bark texture for all woody species (see the electronic supplementary material, appendix S1) was recorded for at least one, and up to three of the largest individuals of each species based on four previously defined categories, from smooth (category 1) with lack of ornamentation (e.g. spines, lenticels,
fissures) and including smooth exfoliating barks (e.g., some *Bursera* species), to rough (category 4) with coarse ornamentation (large lenticels and/or fissures) and wood texture (category 5) rugose and ornamented barks are expected to have a greater load of epiphytes [22,29]. Data on wood density (specific gravity) of all tree species at the plots was obtained from the literature (see the electronic supplementary material, appendix S1).

(b) Observed interaction network

Ninety per cent of the bromeliad epiphytes were found on woody individuals with DBH greater than or equal to 5 cm, and we only considered these individuals for the network analyses. We constructed an interaction matrix *Y* describing the epiphyte–phorophyte network, pooling the data from the three sites. In this matrix the *i* rows correspond to woody species (all potential phorophytes) and the *j* columns to bromeliad species (epiphytes), and a cell *y* \[ij\] is an integer that represents the number of interactions that occur between woody species *i* and bromeliad species *j*, i.e. the sum of all epiphyte individuals of species *i* at the three plots, recorded on individuals of woody species *j*. Any woody species *i* at the plots, whose individuals did not host bromeliads is present in this matrix with \( \sum_j y_{i \cdot} = 0 \).

We use network statistics that characterize several aspects of network structure [40] to describe the observed network and to compare observed values with the values obtained from models of network determinants using functions in R [12,40–42] (see the electronic supplementary material, appendix S2): (i) connectance \( C = \text{number of links/} \text{df} \), (ii) nestedness [4], two metrics: \( N = 100 − T \), where \( T \) is the matrix temperature [43], and nestedness based on overlap and decreasing fill (NODF) [44]; (iii) interaction evenness [45]; (iv) \( H_z \), a quantitative metric of specialization that controls for the interaction frequencies expected from the total observations per species, i.e. the effect of the differences in the abundance of species, that results in abundant species interacting more frequently and with more partners, is removed [46]; (v) generality and vulnerability, the weighted mean number of phorophyte species per epiphyte species, and epiphyte species per phorophyte species, respectively [47]; and (vi) the average interaction strength asymmetry for phorophytes and for epiphytes [8].

First two statistics are based on unweighted links, whereas the remaining are based on weighted links. To allow for comparison, we assessed the significance of nestedness [43,48] using the equivalent of null model 2 of Bascompte et al. [4].

(c) Models of network determinants

We follow the conceptual and methodological framework proposed by Vázquez et al. [12], in which the observed matrix is a function of multiple interaction probability matrices determined by different factors. We built probability matrices from models that consider epiphyte and woody species abundance, bark texture and wood density of woody species, size (DBH) of woody individuals, and presence/absence of woody species and epiphyte species at each site (spatial overlap). In addition, we develop models that consider the joint effect of two or more factors, thus, all possible combinations (plant size is always assessed together with abundance, see below for details) of two, three and four factors, and a full model with the five factors are considered.

Consider a matrix \( X \) of the same size as \( Y \), whose entries \( x_{ij} \) are given weights according to a particular model defined by the factors we are assessing. To calculate a probability matrix \( P \) from any matrix \( X \), these weights are converted to probabilities of occurrence of pairwise interactions by normalizing the matrix so that all its elements sum to one. For the different models the calculation of \( x_{ij} \) is described in table 1. In the abundance model (A) interactions are determined by the abundance of species. The spatial overlap model (S) is based on the presence and absence of species on a local (site) scale, considering two species cannot interact if they do not co-occur at a site. The wood density model (W) weighs each species by its specific gravity. The model that considers higher probabilities of interaction to woody species with rougher bark texture. Model AD considers the joint influence of plant size and abundance. Because plant size is a trait of the individual and networks depict interactions among species, when considering the DBH of all individuals, the resulting probability matrix necessarily incorporates information on the abundance of each woody species. Models that consider the joint influence of spatial overlap and abundance are calculated so that not only presence/absence data are considered for each site, but the abundance of each species as well, thus incorporating the fine information on local abundance of this system. The performance of all models was compared to the performance of a null model that assumes that all interactions have the same probability to occur.

Two approaches are used to compare the performance of models in explaining the observed interaction network [12] (see the electronic supplementary material, appendix S2).

- The likelihood that a probability matrix explains the observed matrix is calculated assuming a multinomial distribution [12] and compared among models using the Akaike’s information criterion (AIC) [49].

\[
\text{AIC} = −2 \ln(L) + 2k
\]

where \( k \) is the number of parameters used to generate a probability matrix, which is the number of factors involved in the calculation of each probability matrix, except for plant size that involved two parameters (probability of presence and expected abundance). The model with the lowest \( \Delta \text{AIC} \), the difference in AIC between a given model and the AIC of matrix \( Y \) fitted to itself, indicates the model that better fits the data.

- A randomization algorithm [12] that assigns the total number of observed interactions (sum of elements in \( Y \)) to cells of a matrix of size \( Y \) with probabilities defined by a probability matrix is used to generate 1000 quantitative networks for each model; all network statistics are calculated for each generated network to obtain the 95% confidence intervals (CI) of the distribution of values of the statistics, against which the statistics of the observed network are compared. The only constraint of this algorithm [12] is to assign at least one interaction per species. To allow for empty rows, we did not use the constraint for woody species, though we left the constraint for epiphyte species. This simulates a scenario in which all woody species can potentially be colonized by epiphytes, and allows testing if the factors considered explain the absence of epiphytes on certain woody species. Network statistics are excluded calculating non-interacting woody species.

We identified the species with the predicted number of interactions that deviate the most from the observed values by assessing whether the observed value of each pairwise interaction falls outside the lower and upper 95% CI of 1000 runs of the algorithm.

(d) Phylogenetic signal in species associations

We evaluated the influence of evolutionary history on network patterns by assessing the presence of phylogenetic signal (i) on species degree (number of links per species); (ii) on species strength, a measure of the importance of a species to the partner’s set that considers the relative frequency of the species on each partner [50]; and (iii) on the assemblage of interacting partners of the species (ecological similarity) [13]. Phylogenetic signal is assessed separately for epiphytes and for phorophytes, and was evaluated for the regional species pool (data from the three sites) and for the assemblages at each sampled plot. Phylogenetic hypotheses were constructed using PHYLOMATIC software [51] for host species, and MESQUITE v. 2.75 [52] for epiphyte assemblages, based on Barfuss et al. [53] and Chew et al. [54].
Table 1. Model calculation of pairwise interaction weights \( x_{ij} \). (Letters composing the model name indicate the factors that it considers: A, abundance; B, bark texture of woody species; D, plant size (DBH); S, spatial overlap of species; W, wood density of woody species.)

<table>
<thead>
<tr>
<th>models</th>
<th>( x_{ij} ) calculation</th>
<th>variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>null</td>
<td>( x_{ij} = 1/\bar{I} )</td>
<td>( \bar{I} )—total number of woody species</td>
</tr>
<tr>
<td>A</td>
<td>( x_{ij} = a_i )</td>
<td>( a_i )—number of individuals of species ( i )</td>
</tr>
<tr>
<td>B</td>
<td>( x_{ij} = P )</td>
<td>( P )—number of individuals of species ( j )</td>
</tr>
<tr>
<td>S</td>
<td>( x_{ij} = s )</td>
<td>( s )—number of sites in which species ( i ) and ( j ) co-occur</td>
</tr>
<tr>
<td>W</td>
<td>( x_{ij} = w_i )</td>
<td>( w_i )—woody species specific gravity</td>
</tr>
<tr>
<td>ADF; ABD</td>
<td>( x_{ij} = \frac{1}{\bar{I}} \sum_{m=1}^{M} P( \text{presence of epiphytes}_m</td>
<td>\text{DBH}_m ) (n_m) (a_i) )</td>
</tr>
<tr>
<td>AS; ADS; ABDS</td>
<td>( x_{ij} ) calculated as for A, AD and ABD, respectively, but calculated for each of the three sites separately, and summing the three weights of ( x_{ij} ) obtained for each site.</td>
<td></td>
</tr>
<tr>
<td>AB; BS; ABS</td>
<td>The respective probability matrices are calculated as the element-wise multiplication of the probability matrices ( P ) of models A and B, B and S, AS and B, respectively. The resulting matrices are normalized again to obtain ( P ).</td>
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<tr>
<td>AW; BW; SW; ADW; ABW; ASW; ABSW; ABDSW; ABW</td>
<td>The respective probability matrices are calculated as the element-wise multiplication of each of the probability matrices ( P ) of models above (except W), and the probability matrix of model W. The resulting matrices are normalized again to obtain ( P ).</td>
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\( e \) is the Napier’s constant; \( z \) is calculated from a logistic regression equation that describes the log odds of presence of epiphytes on a tree, \( z = \ln(\text{odds}(\text{presence of epiphytes})) \), in which bark category and DBH are the explaining variables. \( z = b_0 + b_1 v_1 + b_2 v_2 \), where \( b_0 \) is the intercept, \( v_1 \) is the DBH category of the tree, \( v_2 \) is the bark category of the species to which the tree belongs, and \( b_1 \) and \( b_2 \) are the regression coefficients (see the electronic supplementary material, appendix S2). In this model \( v_1 \) is held constant to five, the minimum DBH category size of the individuals considered. In this model \( v_2 \) is held constant to the intermediate category (two), the bark category to which most tree species belong (see the electronic supplementary material, appendix S1). \( n_m \) is the back-transformation of the log (expected number of epiphytes on a woody individual), calculated from the regression equation that describes the linear relation between log(DBH of phorophytes) and log(epiphyte abundance) (see the electronic supplementary material, appendix S2).

We transformed all branch lengths to 1, because branch length information is absent for the Tillandsia phylogeny. The amount of phylogenetic signal for (i) and (ii) was evaluated calculating the K-statistic [55], and for (iii) with Mantel tests comparing phylogenetic distance matrices with ecological distance matrices. Analyses were performed over 100 fully resolved trees. See full details in the electronic supplementary material, appendix S2.

### 3. Results

We recorded 363 woody individuals, of which 221 (DBH greater than or equal 5 cm) were included in the network analyses, belonging to 50 species and 20 plant families. The network included 1304 bromeliads recorded on 142 phorophytes (64% of woody individuals). We registered 151 links between 12 Tillandsia spp. and 36 (72%) woody species (figure 1). Five of these Tillandsia spp. are endemic to Mexico (see the electronic supplementary material, appendix S1). The observed network showed low levels of specialization, with \( H_2 = 0.24 \), connectivity of 0.35, and a significantly nested structure (\( N = 90.5 \), mean for null matrices = 62.21, \( p < 0.001 \); NODF = 62.89, mean for null matrices = 44.75, \( p < 0.01 \)). Epiphyte species interact on average with 8.8 phorophyte species (generality), and phorophyte species interact with 4.9 epiphyte species (vulnerability). Interaction evenness is high (0.82), indicating no dominance of few interactions. The average strength of interactions between epiphytes and their partners is close to symmetry (mean asymmetry for epiphytes is very low = 0.023), whereas phorophytes tend to experience stronger effects from their interaction partners (–0.389).

In the likelihood analysis all the models tested had a better performance than the null matrix, and partially explained the observed patterns of pairwise interactions (figure 2). The full model (ABDSW, figure 1) was the best performing model, with \( \Delta AIC = 1171.41 \), which is 73.14 units away from the second lowest \( \Delta AIC \) model (figure 2). Thus, the likelihood that the observed data are explained by model ABDSW is higher than for all other models. Abundance contributed the most to explain the observed patterns (figure 2, compare performance of models including one factor). The model that considers abundance and spatial overlap of woody species and epiphytes (i.e. local presence and abundance, model AS) is the model of two factors that better fits the observed data. Size of woody individuals is
Figure 1. (a) The bromeliad–phorophyte network and (b) the network yielded by one run of the best performing model of network determinants, ABDSW. Widths of rectangles represent the relative interaction frequencies of bromeliad (a) and phorophyte (b) species. Links among species and their relative frequency are represented by the lines connecting the rectangles and their width, respectively. Networks are drawn to the same scale. Species order and code is the same at both networks. Species identity is shown in the electronic supplementary material, appendix S1.

the third factor in importance for explaining interactions between epiphytes and phorophytes, being ADS the model of three factors with lowest \( \Delta AIC \). Finally, wood density followed by bark texture contributed the least to explain the observed data (see figure 2 dark bars).

The models jointly including abundance, size of woody individuals and spatial overlap in general performed well for predicting network indices (figure 3 and electronic supplementary material, figure S1). In particular, connectance, interaction evenness, nestedness (NODF) and phorophyte interaction strength asymmetry were predicted by these models (figure 3 and electronic supplementary material, figure S1). Epiphyte interaction strength asymmetry was predicted or extremely close to the lower CI of models that incorporate abundance (figure 3). Similarly, nestedness (N) was predicted by models that incorporate abundance, and by model S (see the electronic supplementary material, figure S1).

No probability matrix predicted the values observed for indexes related to specialization; model matrices were more generalized. Epiphyte and phorophyte species interact with less species than predicted by any model (generality and vulnerability, electronic supplementary material, figure S1). Abundance, and to a lesser extent spatial overlap, contribute to explain vulnerability, being the models that include these factors the closest to the observed value, while all factors contribute to explain generality. The observed \( H_2 \) specialization is higher than predicted by any probability matrix. Incorporating information on the spatial heterogeneity of species abundance (models including AS) leads to higher values, yet lower than the observed (figure 3). The number of phorophyte species was overestimated by all models (see the electronic supplementary material, figure S1); ABDW (CI = 43–49) and ABDSW (CI = 44–49) yielded the matrices with the lowest number of phorophytes, yet more than the 36 observed.

The frequency of most links fell within the 95% CI of the frequencies predicted by the best performing models, i.e. models including ADS (mean = 92.4% of the links; electronic supplementary material, table S1 and figure S2). When links fell outside the 95% CI, more often, observed links were stronger than predicted by these models (see the electronic supplementary material, table S1). An improvement in the performance of the models can be observed, as the frequency of a larger number of interactions get closer to the line of best fit between mean values of a model and observed frequencies, and to the observed network (see the electronic supplementary material, figures S3 and S2, respectively). Finally, no significant phylogenetic signal was detected for the traits considered (K-statistic range for species degree and strength: 0.32–0.86; \( p \)-values \( \geq 0.14 \) for the z-statistic for ecological similarity; electronic supplementary material, table S2 and figure S4).

4. Discussion

We found that individual traits (woody individual’s size) and species traits (bark texture and wood density), and the abundance and spatial overlap of species contributed differentially to predict network metrics and the frequency of pairwise interactions in epiphytic bromeliads—woody species commensalisms in a tropical dry forest. Consistent with the finding that these ecological factors largely determine species–species interactions, and with the low specialization of epiphytes on host species, we did not detect a phylogenetic signal in the network, i.e. specialization, species strength and interaction partners are not conserved, neither in the phylogeny of epiphytes nor in the phylogeny of phorophytes.
(a) Network structure

Our results are consistent with previous work which suggests that neutrality contributes to network structure [8,15,25]. Neutrality only partly accounted for the observed patterns (see performance of model A). Species abundance explains network patterns when neutrality determines the establishment of interactions, but this is a component of the community structure, which is influenced in a complex way by many biological factors (see causal model of Vázquez et al. [21]), among them, the spatial and temporal distribution of species. Once considering the constraints posed by the local spatial and temporal distribution of species (models including ADS, figure 3, and electronic supplementary material, figure S1), neutrality explained to a large extent network patterns. That is, network assembly, involving abundance, spatial overlap and phorophyte size (proxy for time of exposure to epiphyte seed rain) implies that networks are gradually built up by random encounters of individuals (regardless of the species to which they belong) concurring in space and time. However, it is clear that other biological factors also affect the establishment of interactions, including bark texture and wood density of hosts. The network specialization is comparable to ant-nectar plant (‘low intimacy’) and seed dispersal networks [56]. Ecological differences among epiphyte species may lead to differentiation in host use, e.g. if species differ in their susceptibility to nutrient, light or humidity levels and if substrate suitability varies among phorophyte species.

Clumped spatial distributions of woody species in tropical dry forests [57], and epiphyte communities (e.g. orchids and ferns [22]) may account for the unexplained network structure. Anemochory of \textit{Tillandsia} can limit their dispersal kernel to a few metres resulting in spatial aggregation [28]. Asexual reproduction contributes for aggregated patterns when ramets detach and establish in the same or a neighbour host. Our models partly account for a clumped distribution: first, models that consider spatial overlap and abundance take into account differences in epiphyte abundances among plots. Second, when \textit{Tillandsia} reproduces sexually or asexually, the probability of some progeny being established on the same host is large, influencing the epiphyte abundance–host size relation considered in the models (table 1).

Using a spatially explicit model, Morales & Vázquez [58] simulated the assembly of plant–frugivore networks using scenarios with varying levels of spatial autocorrelation of tree species and distances of bird foraging movements. Spatial structure and limited mobility affected the ‘degree of mixing’ in the system, imposing constraints to encounter probabilities [58]. A similar scenario could be found in our system with limited seed dispersal of \textit{Tillandsia} and aggregated woody species. In Morales & Vázquez [58], a scenario analogous to our system (limited animal mobility and random...
individual tree spatial distribution), they show that with increasing spatial autocorrelation of tree identity, nestedness (N) varies slightly, connectance and evenness decrease, strength asymmetry increases, and unexpected presences/absences of interactions occur. Our findings agree with their results (figure 3 compare model A against AS, ADS and observed values which putatively increase in spatial structure), and the lower number of links, the many stronger interactions (see the electronic supplementary material, table S1, links above upper CI) and the absence of interactions in some woody species are evident in figure S2 (see the electronic supplementary material). This suggests that spatial processes contributed to structure the network.

We did not find evidence of a phylogenetic signal in the network structural patterns analysed, and the results were consistent across sites. A phylogenetic signal is not uncommon in mutualistic and antagonistic interactions [13,14,59]. Similar to our findings, the commensalistic networks between orchid epiphytes and their host trees do not show a phylogenetic signal [26]. The lack of phylogenetic signal could be related to the small size of the phylogenies [55]. Conversely, a phylogenetic signal may be present in other ways (e.g., species roles [60], within or among compartments [61], within clades [59]). Evolutionary history may influence network structure at other scales and analysing a phylogenetically more diverse epiphyte community or a larger network might lead to the detection of a phylogenetic signal. Additionally, if traits important for species associations are phylogenetically conserved, then phylogeny will indirectly influence network structure [62,63]. Abundance does not show a phylogenetic signal in the epiphyte or phorophyte assemblage (see the electronic supplementary material, table S2 and figure S4). Wood density is highly conserved across the entire seed plant phylogeny [64]. Here our results showed a weak signal, and the contribution of wood density on the network structure is low.

(b) Pairwise interactions
Our results are consistent with previous epiphyte community reports that have found stronger observed interactions when compared with results from null models that consider phorophyte abundance [32,36]. When accounting for the effect of other factors, a higher percentage of the interactions is explained (see the electronic supplementary material, table S1; model ABDSW versus A). Nevertheless, several observed interactions remain stronger (more frequent) than expected by the full model. This is consistent with a higher observed than model yielded network specialization ($H_s$). In our data set, the most extreme case was that of Caesalpinia slerocarpa, the only species in which all 12 Tillandsia species were registered. This is a timber species with dense wood (see the electronic supplementary material, appendix S1) and, therefore, more stable branches. A slower growth rate expected from dense wood also implies that size in this species represents older individuals than in other species. Despite considering all factors (ABDSW), five out of the 12 interactions were stronger than predicted. Caesalpinia slerocarpus’s architecture, with ramifications high on the trunk and fairly horizontal branches may favour the establishment of epiphytes. Species with dense wood are capable of growing taller [65], and this is one of the tallest species in the forest; Tillandsia are light demanding [66] and may be more successful high in the canopy. By contrast, the most abundant woody species, Apopilaxus paniculata, interacted with 10 Tillandsia spp., but was found interacting less frequently than expected by its abundance, bark texture, wood density, spatial overlap and tree sizes, with two Tillandsia species.

Despite several weaker links than predicted by ABDSW (11 cases, electronic supplementary material, table S1), we found little evidence for axenic species (free of epiphytes). From the 14 species recorded with no epiphytes, only Guazuma ulmifolia and Jacquina pungens were expected to interact with the most abundant epiphyte, T. usneoides. Overall, epiphytes tended to be absent or were less frequent on woody individuals that were young, on woody species with low abundance, present at a single site, and with smooth bark and/or soft wood. In the same way, less abundant epiphyte species tended to be hosted by a lower number of woody species.

(c) Conservation implications
Understanding community assembly will contribute to the conservation of interactions. Consequences of network structure for commensalistic interactions might be different than for other interactions where coevolution is involved. As other epiphyte–phorophyte networks [3,5,25], the bromeliad–phorophyte network showed high values of nestedness (N and NODF), higher than other types of interactions [5]. Several processes may lead to a nested pattern [63,67,68]. The specialization asymmetry implied in nestedness, suggested to arise through a coevolutionary process in other systems, with implications for extinction risk [4,63,69,70], seems to be here a consequence of the abundance and temporal and spatial distribution of species of a mostly generalized system. The one-way specialization of these bromeliads on an epiphytic life form but with low host specificity shows that many species can be adequate hosts, which in regard to host availability makes them less vulnerable to disturbance. However, hosts with suitable traits will give epiphytes higher probabilities of survival, with important implications for the conservation and management of this group.

Our analysis highlights the importance of time for the establishment of epiphyte communities. It takes at least one decade to the first reproduction of bromeliads [22,37], and the recruitment of species with different reproductive strategies (monocarpic versus polycarpic) may be affected differentially by disturbances. Epiphytes provide important resources for many taxa [22] and their disappearance will cascade through the ecosystem. Tillandsia is the richest genus of the Bromeliaceae, and Mexico is a center of diversification for this genus, where 43 per cent of its species, most of which are endemic [37,71], occur. Our study contributes to understanding the processes that assemble ecological interaction networks in general and commensalistic networks in particular. Vascular epiphytes are the most vulnerable plant life form, and this study highlights ecological factors that shape their communities.

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