

HIGHLIGHT ON ECOLOGY AND EVOLUTION OF EXTRAFLORAL NECTARIES

Long-term temporal variation in the organization of an ant–plant network

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• **Background and Aims** Functional groups of species interact and coevolve in space and time, forming complex networks of interacting species. A long-term study of temporal variation of an ant–plant network is presented with the aims of: (1) depicting its structural changes over a 20-year period; (2) detailing temporal variation in network topology, as revealed by nestedness and modularity analysis and other parameters (i.e. connectance, niche overlap); and (3) identifying long-term turnover in taxonomic structure (i.e. switches in ant resource use or plant visitor assemblages according to taxa).

• **Methods** Fieldwork was carried out at La Mancha, Mexico, and ant–plant interactions were observed between 1989 and 1991, between 1998 and 2000, and between May 2010 and 2011. Occurrences of ants on extrafloral nectaries (EFNs) were recorded. The resulting ant–plant networks were constructed from qualitative presence–absence data determined by a species–species matrix defined by the frequency of occurrence of each pairwise ant–plant interaction.

• **Key Results** Network variation across time was stable and a persistent nested structure may have contributed to the maintenance of resilient and species-rich communities. Modularity was lower than expected, especially in the most recent networks, indicating that the community exhibited high overlap among interacting species (e.g. few species were hubs in the more recent network, being partly responsible for the nested pattern). Structurally, the connections created among modules by super-generalists gave cohesion to subsets of species that otherwise would remain unconnected. This may have allowed an increasing cascade-effect of evolutionary events among modules. Mutualistic ant–plant interactions were structured 20 years ago mainly by the subdominant nectarivorous ant species *Camponotus planatus* and *Crematogaster brevispinosa*, which monopolized the best extrafloral nectar resources and out-competed other species with broader feeding habits. Through time, these ants, which are still present, lost their position as network hubs and diminished in their importance in structuring the network; simultaneously, plants gained in importance.

• **Conclusions** The long-term network analysis reveals a decrease in attended plant species richness, a notable increase in plant species participation from 1990 to 2010 (sustained by less plant taxonomic similarity in the older 1990 network), an increase in the number of ant species and a diminishing dominance of super-generalist ants. The structure of the community has remained highly nested and connected with low modularity, suggesting overall a more participative, homogeneous, cohesive interaction network. Although previous studies have suggested that interactions between ants and EFN-bearing plants are susceptible to seasonality, abiotic factors and perturbation, this cohesive structure appears to be the key for biodiversity and community maintenance.

Key words: Ant–plant interaction interactive network, long-term variation, mutualism, nestedness, modularity, taxonomic structure.

INTRODUCTION

Functional groups of species interact and coevolve in space and time, forming complex networks of interacting species (Thompson, 2005; Bascompte, 2009). As a result, entire ecological communities can be analysed as a series of complex interaction networks (Burkle and Alarcón, 2011; Aizen *et al.*, 2012; Stouffer *et al.*, 2012). The analysis of interaction networks, such as plant–pollinator (e.g. Bascompte *et al.*, 2003; Bascompte, 2009; Burkle and Alarcón, 2011) and ant–plant networks (Guimarães *et al.*, 2006, 2007; Blüthgen *et al.*, 2007; Díaz-Castelazo *et al.*, 2010), has revealed several patterns

of interaction that are common to a wide range of free-living mutualisms. Mutualistic interactions are highly asymmetric (Vázquez and Aizen, 2004) and nested (Bascompte *et al.*, 2003; Díaz-Castelazo *et al.*, 2010), such that species with few partners primarily interact with hierarchical subsets of a generalized core group of partners. A few studies have explored the relationship between the architecture and functioning of ecological networks. They have found consistent relationships between the topology of the networks and their functioning, as variation in fitness attributes across populations has been correlated with higher nestedness, connectivity and clustering in mutualistic networks (but see also Rezende *et al.*, 2007;

Bastolla *et al.*, 2009; Fontaine *et al.*, 2011; Guimarães *et al.*, 2011). Bascompte *et al.* (2006) established that the architecture of quantitative mutualistic networks, characterized by a low number of strong dependencies, high asymmetry and heterogeneity in species strength (i.e. certain species interact very frequently or with high individual participation, in contrast to other species that rarely interact), may promote community coexistence, in turn favouring the long-term persistence of species-rich assemblages, i.e. biodiversity maintenance (Bascompte and Jordano, 2007; Ings *et al.*, 2009; Lewinsohn and Cagnolo, 2012).

Understanding the ecological and evolutionary causes and consequences of spatial and temporal variation in mutualistic interaction networks is important for addressing both basic and applied questions about community structure and function. However, the characterization of spatio-temporal variation of interactions has just begun (Olesen *et al.*, 2008; Díaz-Castelazo *et al.*, 2010; Burkle and Alarcón, 2011; Olesen *et al.*, 2011; Rico-Gray *et al.*, 2012). Ant–plant interactions are among the most temporally and spatially variable mutualistic interactions, especially regarding the species assemblages, although the outcomes of the interaction may vary as well (Rico-Gray, 1993; Bronstein, 1994; Oliveira *et al.*, 1999; Díaz-Castelazo *et al.*, 2004). In these interactions, plants produce food resources (e.g. extrafloral nectar) that attract and reward ants in exchange for their defence against natural enemies (Rico-Gray and Oliveira, 2007; Chamberlain and Holland, 2009). In recent years, several studies have described temporal patterns of mutualistic networks across 2–4 years of sampling (Burkle and Alarcón, 2011). To our knowledge, however, only two studies have addressed longer-term changes to network parameters in species interactions (Díaz-Castelazo *et al.*, 2010; Olesen *et al.*, 2011). Studies on more subtle temporal variation in mutualistic networks show a continuous switch in plant species used by animals in response to environmental disturbances and the availability of resources (Whittall and Hodges, 2007); plants can also adjust phenology and morphology to affect their visitors (Aizen and Vázquez, 2006; Kaiser-Bunbury *et al.*, 2010). Such rewiring of interspecific interactions could not only have ecological and environmental causes (e.g. changing resource availability), but could also reflect adaptive behaviour of species for enhancing the efficiency of resource utilization (Zhang *et al.*, 2011).

The redundant roles of species among cohesive subgroups (modules) that in turn are complementary to each other (Mello *et al.*, 2011) is relevant to robustness (i.e. the persistence of network structure and function even though certain species are lost at the community) because of the contributions of species to nestedness. Saavedra *et al.* (2011) have recently found that the extinction of stronger contributors to nested topology leads to a decrease in overall network persistence. With simulations they show that the removal of a strong contributor to nestedness tends to decrease overall network persistence more than the removal of a weak contributor. In addition, strong contributors to collective persistence did not gain individual survival benefits, but were in fact the nodes most vulnerable to extinction.

Temporal stability of a Neotropical ant–plant network over a 10-year period has been previously reported for La Mancha,

Veracruz, Mexico, despite changes in species composition which would potentially impact interaction structure (Díaz-Castelazo *et al.*, 2010). Here we extended this study to: (1) examine structural changes over an additional 10-year period, thus covering the 1990–2010 interval; (2) detail temporal variation of network topology, as revealed by nestedness and modularity analysis and other parameters such as connectance and niche overlap; and (3) identify long-term turnover in taxonomic structure (genus, subfamily/family, expressed as switches in ant resource use or plant visitor assemblages according to taxa.

METHODS

Study site

Fieldwork was carried out at Centro de Investigaciones Costeras La Mancha (CICOLMA), located on the coast of the state of Veracruz, Mexico (19°36'N, 96°22'W; elevation <100 m). The climate is warm and sub-humid; a rainy season occurs between June and September, total annual precipitation is approx. 1500 mm, and mean annual temperature is 22–26 °C. The major vegetation types in the study area are tropical deciduous forest, tropical dry forest, sand dune scrub, mangrove forest, freshwater marsh and flooded deciduous forest (Moreno-Casasola, 2006).

Sampling

Ant–plant interactions were observed monthly, first between May 1989 and April 1991 (Rico-Gray, 1993), then between October 1998 and September 2000 (Díaz-Castelazo *et al.*, 2004), and finally between May 2010 and April 2011 (hereafter 1990, 2000 and 2010, respectively) (Díaz-Castelazo *et al.*, 2010; Sánchez-Galván *et al.*, 2012). The study site is highly seasonal and it is important to mention that a category-3 hurricane impacted this coastal environment on September 2010, having an effect on the 2010 census (Sánchez-Galván *et al.*, 2012).

Censuses were conducted along six trails that represent main vegetation types in the area, such as sand dune pioneers, sand dune scrub, deciduous forest, dry forest, freshwater lagoon, flooded forest and mangrove forest (Rico-Gray, 1993; Díaz-Castelazo *et al.*, 2010). The transects varied in length, according to specific vegetation types per trial: sand dune pioneers, 638.5 m; sand dune scrub, 161.96 m; deciduous forest, 370.37 m; dry forest, 225.77 m; ecotone of freshwater marsh, deciduous forest and dune scrub, 457.66 m; mangrove forest, 171.03 m. All transects were 3 m wide. Although transect length was not equal among vegetation types, given their physiognomy (i.e. sand dune pioneer vegetation along the beach is quite sparse compared with other vegetation physiognomies such as dry forest), we performed equal numbers and seasonal repetitions of ant–plant interaction censuses within transects. We observed all plants (below 2.5 m. height, thus excluding large trees, canopy, and their associated epiphytes and vines), followed ants collecting liquids from plants, and subsequently recorded occurrences of ants on extrafloral nectaries (EFNs) strictly [based on published lists of EFN distribution within plant taxa and our knowledge of nectary morphology and position; whenever there were

doubts on nectar secretion, glucose test strips (Clinistix) were used]. On each visit we noted ant species, plant species and the plant organ on which the EFNs mediating the ant–plant interaction were located. Once an individual plant was marked as visited by ants, it was subsequently re-checked throughout the study. We considered extrafloral nectar produced either on the surface of reproductive structures such as the spike, pedicel, bud, calyx or fruit, or secreted by special structures on vegetative parts, such as leaves, shoots, petioles, bracts or stems (Koptur, 1992). Ants were considered to be feeding on nectar when they were immobile, with mouthparts in contact with nectar-secreting tissues, for periods of up to several minutes. Nectar-feeding ants often showed obviously distended gasters (Rico-Gray, 1993).

Statistics and network metrics

The resulting ant–plant networks analysed here (1990, 2000 and 2010) were constructed from qualitative presence–absence data determined by a species–species matrix defined by the frequency of occurrence of each pairwise ant–plant interaction. The following analyses were performed on these three network matrices.

Network-level estimates. We estimated nestedness for each network using the metric NODF (Almeida-Neto *et al.*, 2008). We used Aninhado 3.0 to compute NODF (Guimarães and Guimarães, 2006). The significance of NODF was estimated with a null model generating random matrices from the original matrix, null model 2 (Bascompte *et al.*, 2003), in which the probability of interaction between any pair of species is proportional to their total number of interactions (i.e. their degree). The *P*-value was obtained from the proportion of random matrices that had an NODF value equal or higher than the value obtained for the real matrix. Other structural features analysed for each network included degree (number of interactions among ant and plant species) and network connectance [proportion of realized links of the total possible in each network ($C = I/(P \times A)$), where *I* is the total number of interactions recorded for the network, *P* is plant species richness and *A* is ant species richness]. Using the ‘Bipartite’ in software ‘R’ (Dorman *et al.*, 2009; Dormann and Gruber, 2011), we obtained niche overlap for each trophic level using Horn’s index estimated with network-level function. To provide information on the within-year variation, for the 1990 network we reported the monthly peaks of nestedness and their corresponding *z*-score values. The *z*-score values are measures of relative nestedness, the standard deviation of the 1000 replicates obtained from the null model analysis minus the nestedness value of the analysed network. We then used these values to compare within-year variation with between-year variation. Thus, we are comparing each sampling date within and between the other sampling dates.

To test whether there were ant foraging preferences which produced guilds in the community, we assessed modularity within each network (Mello *et al.*, 2011). Modularity is a measure of how much the network is structured in cohesive subgroups of vertices (modules), in which the density of interactions is higher within than among modules. Modularity was calculated with the index *M* (from 0, no

subgroups, to 1, totally separated subgroups) estimated with Netcarto (Guimerà *et al.*, 2004; Guimerà and Amaral, 2005).

Temporal taxonomic turnover

We described changes in taxonomic composition of network nodes over time to depict temporal variation in ant resource use or plant visitor assemblages underlying changes in network structure. The network’s temporal taxonomic turnover was described at two levels. First, we summed all interspecific interactions observed between any given pair of ant/plant genera (e.g. *Camponotus/Crotalaria*). The same procedure was repeated for pairs of ant/plant subfamilies and families (e.g. Formicinae/Fabaceae). Taxonomically aggregated ecological interactions were then plotted as temperature matrices describing the number of interspecific interactions observed at the two taxonomic levels mentioned above, providing a quantitative visualization for relative roles of diverse taxa in the network over time.

We described the temporal stability of taxonomic composition by comparing Jaccard’s similarity index at both genus and family (for plants) or subfamily (for ants) levels between different time periods (1990, 2000 and 2010). Jaccard’s similarity index between the time periods *A* and *B* was computed as follows:

$$S_J(AB) = a/(a + b + c) \quad (1)$$

where *a* is the number of genera (family or subfamily) shared between the two periods, *b* is the number of genera (family or subfamily) present only in the first period and *c* is the number of genera (family or subfamily) present only in the last period.

RESULTS

Considering all data gathered at La Mancha throughout the two decades of study (1990–2010), we recorded 54 ant species (*A*) in 20 genera and five subfamilies attending 76 EFN-bearing plant species (*P*) in 61 genera and 29 families (Supplementary Data Table S1), adding up to 564 distinct ant–plant associations (*I*) and thus leading to an overall connectance (*C*) of 13.7%. Even though attended plant species richness (*P*) decreased from 1990 to 2010 while the number of ant species (*A*) increased, and turnover in species and interactions was high, connectance was relatively stable throughout the period (Fig. 1).

Like connectance, other network structure parameters were relatively stable across years. All ant–plant networks were significantly nested, $N = 47.4$ in 1990 ($N_{\text{random}} = 23.02$, $P < 0.001$), reaching a peak of $N = 50.5$ ($N_{\text{nullmodel}} = 26.92$, $P < 0.001$) in 2000, and decreasing to $N = 39.4$ ($N_{\text{nullmodel}} = 21.22$, $P < 0.001$) in 2010. The observed modularity of the 1990 network ($M = 0.426$) did not differ from the average modularity of random runs ($M_{\text{nullmodel}} = 0.4$, $P > 0.1$), while the modularity values recorded for both the 2000 ($M = 0.299$) and 2010 ($M = 0.33$) networks were significantly lower than that expected from null model simulations ($M_{\text{nullmodel}} = 0.329$, $P < 0.01$ for 2000 and $M_{\text{nullmodel}} = 0.368$, $P < 0.05$ for 2010). Thus, nestedness was higher than and modularity was often lower than or as expected by the heterogeneity of interactions across the years. Although our

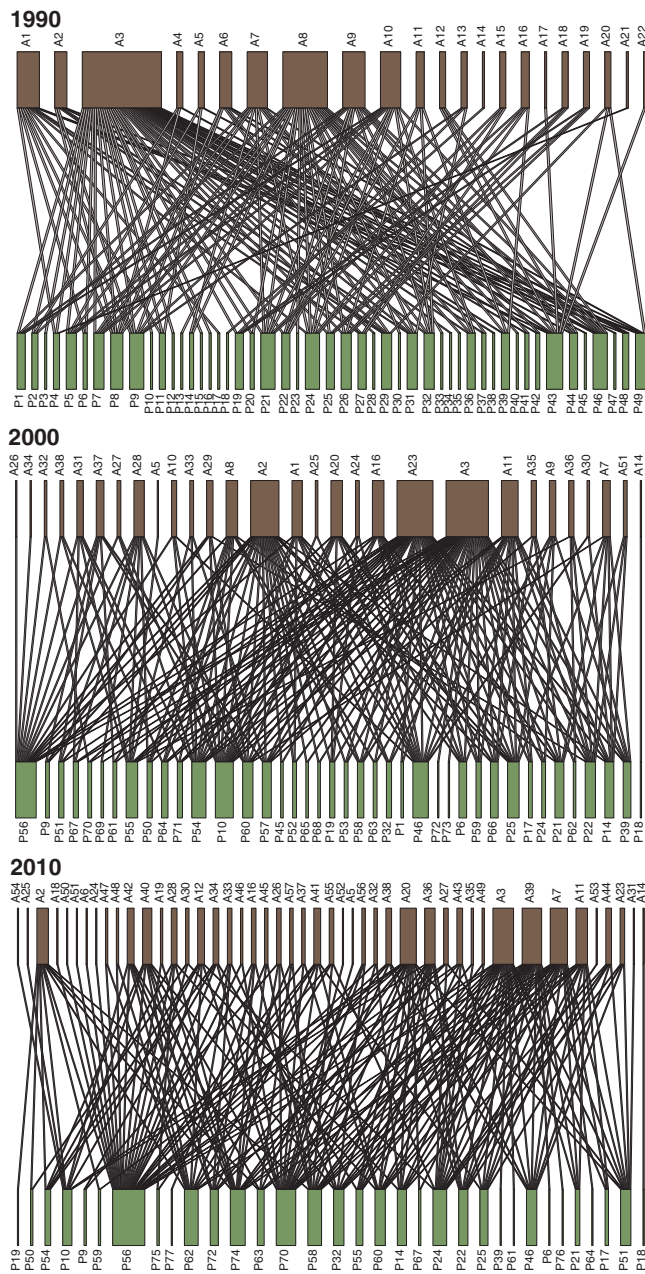


FIG. 1. Mutualistic bipartite networks of ants and extrafloral nectary-bearing plants over a 20-years period at La Mancha, Mexico. A, ants (brown); P, plants (green). 1990: S_P (plant species richness) = 49, S_A (ant species richness) = 22, C (connectance) = 0.14. 2000: S_P = 40, S_A = 29, C = 0.175. 2010: S_P = 33, S_A = 46, C = 0.138. Each label is species-specific and identifies an ant (e.g. A01) or a plant (e.g. P42).

long-term analysis is for presence/absence data, we also provide information regarding the frequency of interactions for networks in 1990, 2000 and 2010 (Supplementary Data Tables S2–4). These data indicate whether an individual link in the network is supported by a single observation or by several. Overall, the higher interaction frequencies are recorded for network 2000 relative to the other years.

For the first year of our study (1990 network) all months exhibited a significantly nested network topology ($P < 0.01$,

with NODF values ranging from 27.4 to 36.3 ($P < 0.0001$) from January to December, with peaks in March and June that, respectively, correspond to z -score values of 6.8 and 5.72 (Rico-Gray *et al.*, 2012). Our whole 1990 network (year-level) exhibited a significant NODF value of 47.43 ($P < 0.0001$), with a z -score of 8.95; this means that even when controlling for differences in species richness, number of interactions and connectance (as z -score values are measures of relative nestedness), our whole-year nestedness is considerably higher than their monthly counterparts. We also suggest that at our study site variation in ant–plant interaction networks is lower within a year than between years (i.e. non-contiguous period of time studied).

Although network structure was relatively constant through time, the relative importance of component species varied greatly (Fig. 2). The 1990 network had a plant species niche overlap of 0.345, characterized by the prevalence of interactions between Fabaceae species, which were present in 38 % of the links. This network exhibited the lowest ant species niche overlap (0.128) with ants in all subfamilies present (the most frequent being Formicinae and Myrmicinae) (Fig. 2A). Formicine ants, mostly *Camponotus* spp., were the dominant group in that period, participating in 40.5 % of the links and interacting with plants across 33 genera in 19 families. Myrmicine species, predominantly in the genera *Crematogaster*, *Forelius* and *Monomorium*, were responsible for 32.5 % of the links, interacting with plants in 27 genera and 13 families. Dolichoderinae species in the genera *Azteca* and *Dorymyrmex* occurred in 14 % of the interactions, being associated with 16 plant genera belonging to ten families. *Pseudomyrmex* species (Pseudomyrmecinae) were found in 12 % of the links, distributed in ten genera and four families (Fig. 2B). *Camponotus planatus* (Formicinae) and *Crematogaster brevispinosa* (Myrmicinae) were the species with the broadest range of interactions, acting as network hubs (Fig. 3).

The 2000 network exhibited a plant species niche overlap of 0.365, marked by the dominance peak of Formicine species (present in 51 % of the links), and on the plant side, a decrease in the relative participation of the Fabaceae (only 19 % of interactions). The period was also characterized by intermediate ant species niche overlap (0.195), with a high diversity of interactions with the Formicinae, including links between *Camponotus*, *Paratrechina* and *Brachymyrmex* and 36 plant genera in 23 families (the most frequent being Verbenaceae, Convolvulaceae, Boraginaceae and Bignoniaceae) (Fig. 2C). The only species acting as a network hub was a Formicine species (*Camponotus planatus*, Fig. 3). Myrmicinae were recorded in 25.6 % of the links, widespread through 24 genera in 15 plant families (the most common being Fabaceae and Bignoniaceae). Although the proportion of links involving the Dolichoderinae (8 %) and Pseudomyrmecinae (10 %) remained relatively constant, both groups expanded their range of attended plant families to 13 genera in ten families and 13 genera in eight families, respectively. Another major change in the 2000 network was the presence of the Ponerinae in 5 % of the links, comprising nine plant genera from eight plant families.

Finally, the 2010 network had the highest ant species niche overlap (0.215) and a prevalence of interactions involving Myrmicine ants (45 % of the interactions). The contributions of *Forelius*, *Crematogaster*, *Cephalotes* and *Monomorium*

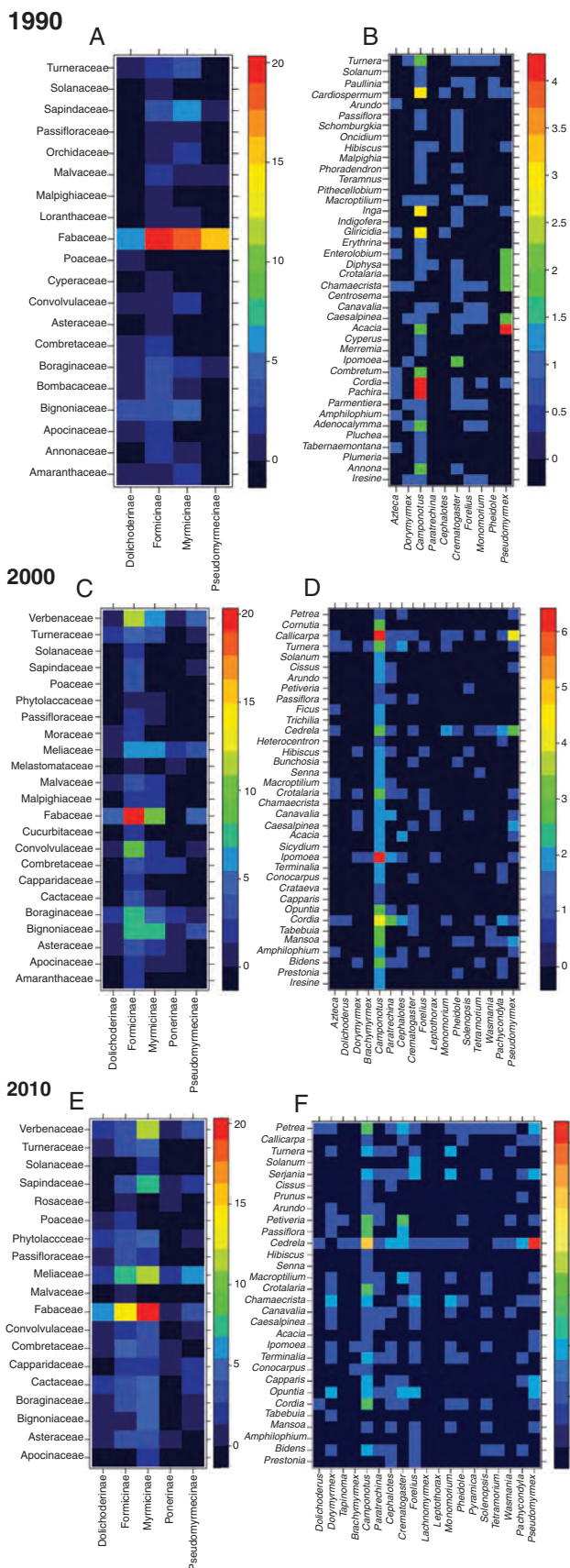


FIG. 2. Distribution of links between ants and extrafloral nectary-bearing plants at La Mancha, Mexico. The hotter the colour, the higher is the

species were roughly equal; they interacted with 23 genera in 16 plant families, the most common being Fabaceae, Meliaceae, Sapindaceae and Verbenaceae (Fig. 2E). Although there was a decrease in the proportion of Formicinae species in these links (29%), the subfamily remained as the one with the most taxonomically widespread interactions, embracing 26 plant genera in 17 plant families. The participation of Dolichoderinae, Pseudomyrmecinae and Ponerinae in the network links remained almost constant (approx. 10% for the former two groups and 4.3% for the latter), as well as their host plant ranges. The 2010 network also exhibited the lowest plant niche overlap (0.192), and plant species were more evenly visited by ants; the only species acting as a network hub was a plant (*Cedrela odorata*, Meliaceae; Fig. 3).

Taxonomic similarity at the genus and family level is shown in Table 1 for both plants and ants. The less similar network in terms of plant composition (at both the genus and the family level) is the 1990 network. This network is also the less similar in terms of the taxonomic composition of the ant fauna visiting EFNs.

DISCUSSION

This study of ant–plant network variation across time revealed a stable connectance and a persistent nested structure. These features have been demonstrated to maintain species richness and provide community resilience to disturbance (Bascompte *et al.*, 2006). Modularity was lower than expected, especially within the most recent networks, indicating that the networks show high overlap among interacting species. Thus, only a few species act as hubs in the network, and are partly responsible for the nested pattern (Olesen *et al.*, 2007). Network hubs are super-generalist species which have the broadest range of interactions; they are characterized not only by their high within-module degree, but by their high among-module connectivity (Olesen *et al.*, 2007), thus adjoining the whole community. In the 1990 network, two ant species were network hubs. In the 2000 network just one of those ant species remained as a network hub, while in the 2010 network a plant rather than ant species acted as a network hub or super-generalist. The connections that super-generalists create among modules produce cohesive subsets of species that otherwise would remain unconnected (Olesen *et al.*, 2007; Guimarães *et al.*, 2011). In general, this cohesive structure appears to be the key for biodiversity and community maintenance in mutualistic networks (Bascompte *et al.*, 2003).

This study provides one of the very few cases comparing the same communities over long periods of time focusing on the dynamics of interspecific interactions. Overall, our long-term analysis has documented a decrease in the richness of EFN-bearing plant species but a significant increase in the participation of those species (more homogeneous interactions of plant species visited by ants), concomitantly with an increase in the number of ant species and a diminishing dominance of super-generalist ants through time. The structure of the

number of species interactions between ant subfamilies and plant families (A, C, E) and between ant and plant genera (B, D, F). (A, B) 1990 network, (C, D) 2000 network, (E, F) 2010 network.

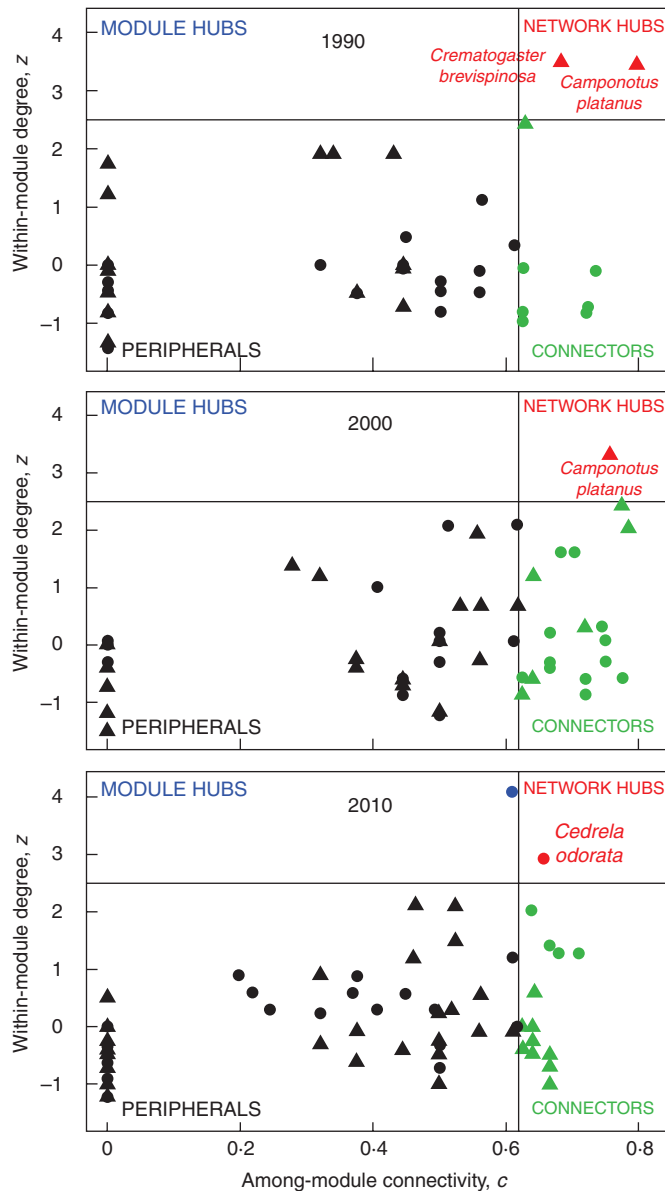


FIG. 3. Species roles in the three networks. Among-module connectivity describes how even is the distribution of links of a given species across species in different modules. Within-module degree quantifies how large is the number of interactions of a species compared with other species in the modules.

community remains highly nested and connected with low modularity. This suggests that a more participative, homogeneous and cohesive interaction network has arisen. Previous studies (Díaz-Castelazo *et al.*, 2010; Rico-Gray *et al.*, 2012) suggest that ant–plant interactions mediated by extrafloral nectaries at the study site are susceptible to seasonality, abiotic factors and perturbation, which may explain the important change in specific links (pairwise interactions) during our three sampling periods. Complementarily, seasonality, and natural and human perturbation do not seem to affect general network structure, nestedness or connectance. These findings are in accordance with Díaz-Castelazo *et al.* 2010 (in the context of ant–plant interactions) and in accordance with

TABLE 1. Taxonomic similarity of the extrafloral nectary-bearing plants assemblage and of ant species visiting extrafloral nectaries at La Mancha, Mexico, across a 20-year period; values above the diagonals describe Jaccard's similarities at the genus level whereas values below the diagonal describe the same index at the family level

	1990	2000	2010
Plants			
1990	–	0.24	0.25
2000	0.54	–	0.74
2010	0.5	0.75	–
Ants			
1990	–	0.59	0.45
2000	0.8	–	0.8
2010	0.8	1	–

Martínez *et al.* 2006 (in the context of ecosystem vulnerability), who state that relative to other sites along the Gulf of Mexico, La Mancha and the central region coasts would be considered as the least vulnerable and most resilient both to natural and to anthropogenic perturbation.

The 1990 network exhibited high numbers of interactions by the subdominant nectarivorous ant species *Camponotus platanus* and *Crematogaster brevispinosa*, which monopolized almost all the best EFN resources and which out-competed other species with broader feeding habits. Both ant species are frequent visitors to EFNs (Díaz-Castelazo *et al.*, 2004) and are considered as solitary leaf foragers that cover foliar area at greater rates than trophobiont-tender ants, in addition to their ability to rapidly take up large loads of nectar (Davidson *et al.*, 2004).

The presence of ruderal plants in the 2000 network, along with the increase in the richness of ant species visiting EFN-bearing plants and the arrival of some invasive ants (Díaz-Castelazo *et al.*, 2010), established new interactions. More coexisting ant species and the presence of generalist invasive ants may help to explain the increase in ant species niche overlap (more ant species use the same plant resources). Only one of the super-generalist ant species remained as a network hub in 2000. Within the most recent network no super-generalist dominant ants were noticed to monopolize plant resources. As a result, ant species such as *Cephalotes minutus* (Myrmicinae) visited noticeably more plant species in 2010 than they did in 1990. This ant is normally timid, is considered a ‘generalized Myrmicinae’ in relation to resource/habitat use (Andersen, 2000) and is less dependent on nectar food (Davidson *et al.*, 2003) than subdominant ants. These habits explain why *Cephalotes minutus* forages on more plants when no other ants are dominant.

Plant species niche overlap decreased noticeably as a consequence of being visited by distinct ant fauna. An extreme case is *Cedrela odorata*, a plant which has several features that makes it accessible to several ant species. It has numerous, small, dispersed EFNs, distributed in rachises (petioles) of compound leaves, non-lignified branches and stems and shoots. Modest amounts of nectar are secreted by these glands (N. Chavarro *et al.*, unpubl. data). However, their quantity and distribution across the plant favours simultaneous

visitation by ants, allowing them to avoid competition and perhaps facilitating their coexistence.

Overall, the 2010 network exhibited a more important contribution of plant species in structuring the mutualistic ant–plant community an increasing average degree (mean connections by node) of plant species occurring through time. This pattern may lead to an increase in convergence and complementarity for plant species as a consequence of the presence of super-generalist partners in the network (see also Guimarães *et al.*, 2011). Our results show a large shift through time in the contribution of ant subfamilies and plant families. Regarding plants, Fabaceae, a plant family where EFNs are most abundant and diverse (McKey, 1989; Koptur, 1992; Pascal *et al.*, 2000; Melo *et al.*, 2010) was particularly dominant. However, it decreased in relative importance over the 20 years of the study. In the 1990 network, 19 plant genera belonged to Fabaceae, in contrast to the seven recorded in the more recent networks. Besides a reduced number of genera, a reduced abundance of plant individuals may explain the lower degree for Fabaceae: plant cover for the 2000 network was 205.6 m, versus 349.7 m in 2010 (Sánchez-Galván *et al.*, 2012). Other plant families that interacted heavily with ants were Bignoniaceae, Verbenaceae, Convolvulaceae, Boraginaceae and Meliaceae, in increasing order of participation through time periods. These are also known as common EFN-bearing plant families that attract ant visitors (Schupp and Feener, 1991; Koptur, 1992).

With regard to ants, it is clear that the Formicinae, particularly the genus *Camponotus*, were associated with the majority of plant species. The extraordinary diversity of formicines foraging on plants reflects their diversification in feeding habits. Many species have proventricular adaptations that allow passive damming of sugary liquids, large crop capacities and seeping canals to nourish the midgut. These traits have turned foragers in several genera, such as *Camponotus*, into liquid ‘canteens’ that have allowed them to exhibit active and prolonged exploration of several environments, including open, hot habitats (Davidson *et al.*, 2004).

Although Formicinae species are important components of the three networks, across time periods the trend was towards a decrease in their interactions with EFN-bearing plants, and an increase in the relative importance of the ant subfamily Myrmicinae. Myrmicinae (mainly the cosmopolitan genera *Pheidole*, *Monomorium* and *Crematogaster*) are ubiquitous and often abundant; they have generalized habits in relation to nesting and dietary requirements, and are competitive at rich food sources, including nectar (Andersen, 1995). Generalized Myrmicinae (Andersen, 2000) are considered competitive species and have a higher tolerance to a variety of stresses and disturbances; this may explain their increased importance at the more recent network, which is also the most disturbed as the presence of some invasive/tramp ant species and ruderal plants reveals, but most importantly, because important abiotic perturbation occurred during this period of time, given a high-category hurricane impacting the study site (Sánchez-Galván *et al.*, 2012). Relative to 1990, in 2000 and 2010 an increased participation of other ant subfamilies, such as Dolichoderinae, Pseudomyrmecinae

and Ponerinae, was noted. Although the proportion of links involving these three subfamilies remained constant in the most recent network, all expanded their range of attended plant taxa, perhaps because no super-generalist species dominated interactions or monopolized nectar sources. Although *Pseudomyrmex* was a well-represented genus in the networks, its associations tend to change from legumes in the scrub and pioneer vegetation (Fabaceae, Caesalpinoideae) to shrubs or trees in the subdeciduous forests, such as the network hub *Cedrela odorata* (Meliaceae).

Our study provides long-term evidence of a nested, highly cohesive structure, suggestive of a community resilient to minor disturbance, as well as to the effects of abiotic factors (i.e. hurricanes, seasonality) on specific links: plant cover loss caused by abiotic phenomena resulted in higher redundancy of ant use of EFN resources and a decrease in the number of connector species. Importantly, super-generalist species contribute to overall network cohesion (Guimarães *et al.*, 2011) and the biological mechanisms behind this deserve future study. To understand how network structure and dynamics have implications for biodiversity and community maintenance, it is critical to continue the study of complex networks over long periods of time.

The community studied here shares structural and temporal properties with other mutualistic networks such as plant–pollinator networks which showed important daily and seasonal variation in species composition (Olesen *et al.*, 2008), where the new species entering the network tend to interact with already well-connected species, while deviations from the trend were explained by morphological mismatching and non-overlapping phenophases. Here as well, the new species entering the community tend to interact with generalists and the taxonomic turnover is higher during the first decade, as the 1990 network exhibited the smallest taxonomic similarities especially for plant genera and families. Thus, the mechanism behind species and taxonomic shifts in ant–plant networks among decades may be vegetation changes through both human and abiotic disturbance (Díaz-Castelazo *et al.*, 2010; Rico-Gray *et al.*, 2012; Sánchez-Galván *et al.*, 2012).

Furthermore, specialized mutualistic ant–plant networks, such as the symbiotic mutualisms between myrmecophilous plants and ants, are less prone to temporal variation and differ in having a modular, less generalistic structure (Guimarães *et al.*, 2007). It is the non-generalized, mainly facultative nature of the interactions we studied that reflects the importance of seasonal variation and disturbance (convergence) more than specific adaptations or complementarity, but provides the overall resilience and richer array of community members that may maintain biodiversity.

SUPPLEMENTARY DATA

Supplementary data are available online at www.aob.oxfordjournals.org and consist of the following. Table S1: list of species codes used for the plants and ants observed in the study. Table S2: interaction frequencies of the 1990 network. Table S3: interaction frequencies of the 2000 network. Table S4: interaction frequencies of the 2010 network.

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