



Hosts, parasites and their interactions respond to different climatic variables

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Abstract

Aim: Although there is a vast body of literature on the causes of variation in species composition in ecological communities, less effort has been invested in understanding how interactions between these species vary. Given that interactions are crucial to the structure and functioning of ecological communities, we need to develop a better understanding of their spatial distribution. Here, we investigate whether species interactions vary more in response to different climate variables than do individual species.

Location: Eurasia.

Time period: 2000s.

Major taxa: Animalia.

Methods: We used a measure of local contribution to β -diversity to evaluate the compositional uniqueness of 51 host–parasite communities of rodents and their ectoparasitic fleas across Eurasia, using publicly available data. We measured uniqueness based on the species composition and based on potential and realized biotic interactions (here, host–parasite interactions).

Results: We show that species interactions vary more, across space, than do species. In particular, we show that species interactions respond to some climatic variables that have no effect on species distributions or dissimilarity.

Main conclusions: Species interactions capture some degree of variation that is not apparent when looking at species occurrences only. In this system, this appeared as hosts and parasites interacting in different ways as a response to different environmental factors, especially the temperature and dryness. We discuss the implications of this finding for the amount of information that should be considered when measuring community dissimilarity.

KEYWORDS

β -diversity, climatic niches, data re-use, host–parasite interactions, species distribution models, species interaction networks

1 | INTRODUCTION

Ecological communities are made of species, but also of the interactions among them. Understanding how community structure changes requires addressing the variability of these two components. Biogeography has developed a very strong corpus of theory and empirical evidence pertaining to the fact that species occur at different locations

because of random chance, favourable habitat (environmental filtering) and the ability of these species to establish the biotic interactions they require to persist. Yet the spatial distribution of species interactions has been comparatively less studied. Poisot, Stouffer, and Gravel (2015) suggested that species interactions, at any given location, can vary because of neutral effects (species abundance), functional effects (species traits) or the effect of a suite of other unspecified factors.

Interactions are known to react in stochastic ways to population abundances (Canard et al., 2014) and are therefore inherently more variable. One of the possible factors that has not been evaluated is the effect of environmental conditions on the existence of interactions. If interactions respond to the environment, then ecological communities will vary across space. Here, we investigate whether knowledge of environmental conditions helps to explain species interaction variability across space.

One way to investigate the variability of ecological communities is through the use of spatially replicated systems. By describing the presence or absence of species originating from the same regional pool at different localities and documenting their interactions, it is possible to compare these communities to gain insights about why and how ecological communities vary. As this is done within a shared regional species pool, this approach reveals the role of factors other than species compositional changes on the variation of community structure. Information about both species and their interactions can be represented efficiently using ecological networks, and recent years have seen the development of approaches to quantify the variation of these networks (Poisot, Canard, Moullot, Mouquet, & Gravel, 2012). In parallel, advances in the quantification of β -diversity allow the identification of hot-spots of variation; that is, localities that, through their unique composition, have a high contribution to the overall β -diversity (Legendre, 2014; Legendre & De Cáceres, 2013; Legendre & Gauthier, 2014). In this manuscript, we bring these families of approaches together, describing the variation of community structure across space and identifying the mechanisms and environmental variables responsible for this variation.

Understanding species within their communities and, by extension, communities themselves can be done through a quantification of the species' Grinnellian and Eltonian niches (Devictor et al., 2010). To summarize, species contribute to community structure first by being present within the community (which assumes that the local environmental conditions are amenable to their persistence) and second by fulfilling a functional role within this community, which is defined in part by the way they interact with other species (Coux, Rader, Bartomeus, & Tylianakis, 2016). Communities therefore differ in two complementary ways: first, because they harbour different species; and second, because whichever species they share might interact in different ways (Poisot et al., 2015). This dissimilarity in both species composition and species interactions, over space and time, has attracted increasing empirical attention in recent years (Carstensen, Sabatino, Trøjelsgaard, & Morellato, 2014; Olito & Fox, 2015; Trøjelsgaard, Jordano, Carstensen, & Olesen, 2015; Vizentin-Bugoni, Maruyama, & Sazima, 2014). Although the drivers of species distributions have been well elucidated in the past decades, there is virtually no knowledge (nor hypotheses) regarding how species interactions should be distributed in space. Most of the earliest datasets on ecological interactions replicated across space (e.g., Havens, 1992) assume that interactions are constant; two species will always interact if they are found in the same location, environment, etc. Given that empirical data (most of which stem from host-parasite interactions, but also from mutualistic systems) contra-

dict this assumption, there is an urgent need to revise our understanding of how community structure should be defined at both the local and the global scale (Trøjelsgaard & Olesen, 2016). Specifically, we need to do so in a way that accounts simultaneously for the variability of species occurrences and for the variability of species interactions.

Building on the framework put forth by Poisot et al. (2015), it is possible to develop quantitative hypotheses regarding the variation of species presences, interactions and the relationship between them. An interaction between any pair of species is possible only when the two species of this pair co-occur. Therefore, because two species may co-occur but not interact (whereas the opposite is impossible), the dissimilarity of interactions is greater than or exactly equal to the dissimilarity of species composition; it cannot be smaller if the two dissimilarities are measured from the same community data (Poisot et al., 2012). There are two important consequences to this fact. First, species composition across multiple localities should be less dissimilar than species interactions. Second, interaction dissimilarity will produce a clearer picture of how communities differ, because the distribution of interactions is intrinsically more variable than that of species; even when two species are found together, they might not interact locally. Most importantly, species are nested within interactions (at least from a purely mathematical standpoint), whereby nodes/species are embedded into edges/interactions; therefore, describing interactions is a more informative way of depicting community structure, which not only includes but genuinely supersedes the usual approach of communities as lists of species.

In this study, we test whether species interactions respond to environmental conditions, using interaction data on rodents and fleas in Eurasia (Hadfield, Krasnov, Poulin, & Shinichi, 2013). Host-parasite systems offer an exceptional natural laboratory to investigate these questions. First, parasites require an interaction with their hosts to persist, so that we expect a strong matching in the distribution of parasites and their hosts (Poulin, Krasnov, & Moullot, 2011). Second, the effect of the parasite on its host's fitness is not strong enough to trigger large mortality events (Krasnov, Khokhlova, & Shenbrot, 2002), so we do not anticipate repulsion of parasites and hosts in space. Third, as fleas are macro-ectoparasites, they experience the same environment as their hosts, and therefore measuring their response to the same environmental conditions is justified. Finally, this rodents-fleas system has a very broad spatial distribution in Eurasia, covering a range of climatic regions. As a result of all of these characteristics, this is a perfectly suitable system in which we can test the effect of environmental conditions on species interactions in space.

This study demonstrates how, within a biogeographical perspective, species interactions reveal another dimension in which ecological communities vary; importantly, species interactions not only capture the variability of species distributions, but also capture another layer of environmental conditions. In particular, we show that (a) species interactions are more variable, and allow the identification of more sites with unique contributions to β -diversity, than species only; and (b) species interactions react to climatic variables of their own, in addition to capturing most of the climatic variables acting on species distributions.

We discuss these results in the light of our current definition of ecological communities and highlight ways of refining this definition in order to make more accurate and realistic predictions about community structure.

2 | MATERIALS AND METHODS

We measure how species composition and species interactions in host–parasite communities vary across environmental gradients. Specifically, we use novel methods to quantify the compositional uniqueness of localities based on different definitions of community structure (Legendre & De Cáceres, 2013), then identify the climatic variables involved in driving the dissimilarity between localities. An interactive document allowing the reproduction of all analyses (including downloading all data from their source) is available as supplementary information (Appendix S2 in Supporting Information); it requires the knitr package (Xie, 2014, 2015, 2016) to be compiled from R into HTML or into the PDF presented as Appendix S1 in Supporting Information.

2.1 | Species interaction data

The species interaction data were taken from Hadfield et al. (2013). They describe species interactions between rodents (121 species) and ectoparasites (206 species of fleas) at 51 locations throughout Eurasia. This system is species rich and probably originates from successive co-speciation events within pairs of interacting species (Hadfield, Krasnov, Poulin, & Nakagawa, 2014). These data were collected by combing a large number of individual rodents, then identifying the fleas that were collected (Krasnov, Mouillot, Shenbrot, Khokhlova, & Poulin, 2004). The data were downloaded from the *mangal* database (Poisot, Baiser, et al., 2016; <http://mangal.io/data/dataset/4/>). The communities (where ‘community’ is defined as the species and interactions detected at one location) have between 3 and 27 (median 11) hosts, between 7 and 40 (median 19) parasites and between 12 and 226 (median 63) interactions between them. Out of 326 species, 94 were observed only once, and 43 were observed at ≥ 10 locations.

For every location, we define two levels of analysis. First, the realized interactions; these correspond to interactions that were reported to occur within individual locations in the original data. Second, the potential interactions; these correspond to the interactions that could happen given the information contained in the entire dataset. For example, if parasite P_j and host H_k do not interact locally, but interact in at least one other location, there will be an interaction between them in the potential interaction network. These two levels encapsulate hypotheses about the filtering of species and interactions: potential interactions are what is expected if species vary in their spatial distribution, but interactions are entirely fixed; conversely, realized interactions are the interactions observed at each location, and therefore account for the spatial variability of interactions in addition to that of species (Poisot et al., 2012). For a given location, the realized and potential interaction networks have an equal number of species, but the realized network has as many or fewer interactions than the potential one.

This distinction between realized and potential interactions, as we detail below, is of great importance. There are many causes for the variation of species interactions, ranging from the distribution of functional traits (Olesen et al., 2011), expressed at macroecological scales, to neutral and random-chance events (Canard et al., 2014), expressed at the microecological scale. Rodents, in particular, are known to change part of their phenology in response to climate (Aars & Ims, 2002; Bozinovic & Rosenmann, 1989), and this in turn can affect the reproductive success of parasites. In short, we argue that potential interactions are more likely to reflect the evolutionary history of the species pairs (this is evidenced in this system by the strong phylogenetic signal in species interactions; Hadfield et al., 2014), whereas the realized interactions are more likely to reflect how the species pairs react to a set of local environmental conditions; specifically, changes in local conditions may, through their impact on species or on the interaction directly, prevent potential interactions from being realized. From the point of view of species distribution, potential interactions offer the possibility to look at the co-distribution of interacting pairs of species. Indeed, any effect of environmental variables on the spatial distribution of potential interactions is highly suggestive of the fact that species that interact also distribute non-randomly. Local factors affecting species interactions can blur whatever signal exists because of species co-distribution; this requires us to estimate this effect through the statistical signal of the models describing the effect of the environmental variables on both realized and potential interactions.

2.2 | Climatic variables data

We downloaded the 19 *bioClim* data (Hijmans, Cameron, Parra, Jones, & Jarvis, 2005) at a geographical resolution of 5 arc min. The data for each location were then extracted using the GPS coordinates of the sampling location. Given that the precise spatial extent that was sampled around each location is unknown, we deemed it more conservative to use a relatively coarse spatial resolution to capture the general environmental conditions around each site. Preliminary analyses revealed that the results are qualitatively unchanged when using different resolutions for the climatic data.

2.3 | Quantification of species and interactions variation

Our approach is represented in Figure 1. We use the method put forth by Legendre and De Cáceres (2013), where the overall β -diversity between sites of a spatially replicated sampling is measured as the variance of a community data matrix Y , and noted β_Y . Y is a binary matrix, with locations as rows and items on which to measure the dissimilarity as columns. These matrices are defined such that $Y(l, i)$; that is, the value in row l and column i of matrix Y is one if item i is found at location l , and zero otherwise. For site-by-species matrices, the row sums give the richness at the locations, and the column sums give the number of occurrences of the species. We define four community data matrices, which are special cases of Y . H has host species in columns, and P has parasite species in columns. These first two matrices will

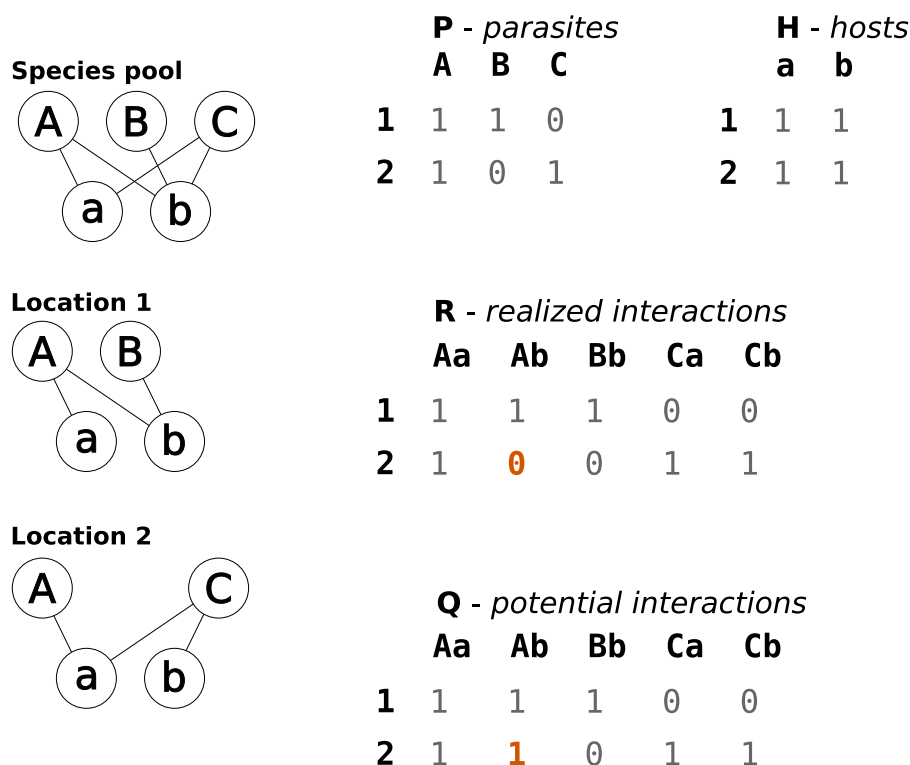


FIGURE 1 Illustration of the construction of the four community data matrices. The networks observed at every location are sampled from a regional pool of species (hosts a and b; parasites A, B and C), pictured at the top left. The two locations vary because they do not have the same species, nor do the shared species interact in the same way. The four matrices corresponding to this simplified situation are on the right side. Note that the interaction between host b and parasite A is 1 in the matrix of potential interactions for location 2; we know from the regional pool that it can happen, even though it was not observed at this location

generate a baseline estimate for the dissimilarity of localities based on species composition. Finally, we also define R , with realized interactions in columns, and Q with potential interactions in columns. The extent to which the realized and potential interactions differ is measured using β_{OS} (Poisot et al., 2012); in brief, this measure is, for every community, the pairwise dissimilarity between R and Q . Values close to zero mean that all potential interactions are realized (weak interaction filtering), whereas values close to unity indicate that most potential interactions have been filtered out (strong interaction filtering).

According to the recommendations of Legendre and Gallagher (2001), we applied the Hellinger transformation to the Y matrices. The dissimilarity matrices (D_H , D_P , D_R and D_Q) were computed using the Euclidean distance on transformed data (Legendre & De Cáceres, 2013). The transformed matrices were then analysed to measure the local contributions to β -diversity (LCBD). The LCBD is a quantification of how much every location (row of the Y matrix) contributes to the overall dissimilarity, presented as a vector, l_Y . We interpret this value as a measure of the originality of a location; a large contribution to β -diversity indicates that the location has a set of species or interactions that is different from the overall species pool. The values of LCBD are tested for statistical significance under a permutation scheme, specifically by redistributing species or interactions across locations. This tests whether LCBD values are larger than expected from random variations in species composition (H , P) or species interactions (Q , R). We assume that LCBDs larger than expected by chance

indicate that the locality is unique with regard to the species or interactions found therein. We used the default significance threshold of $\alpha = 0.05$ (after Holm–Bonferroni correction), with 9,999 random permutations of each column of the four community data matrices.

To summarize, for each community data matrix Y representing hosts, parasites, potential and realized interactions, we measure the β -diversity (β_Y), the matrix of pairwise distance between sites (D_Y), the extent to which each location contributes to β_Y (l_Y) and the statistical significance of each element $l_Y(i)$.

2.4 | Ordination and variable selection

We investigated the extent to which the dissimilarity among communities was driven by climatic variables using RDA (see Appendix S1 in Supporting Information; Legendre & Legendre, 2012). The 19 *bioClim* variables were put in a matrix E , with one location per row. To identify the most significant climatic variables, we used stepwise model building, with forward variable selection and AIC-like statistics over 9,999 replicate runs for each matrix. This approach yields four models of the form $RDA(Y \sim E)$ (i.e., we explain the structure of the Y matrix using the environmental conditions at each site as predictors), for which we extract e_Y , which is a vector of significant climatic variables to explain the structure of Y . In each model, we also record the rank of each climatic variable; variables selected early have a stronger contribution to the structure of community composition.

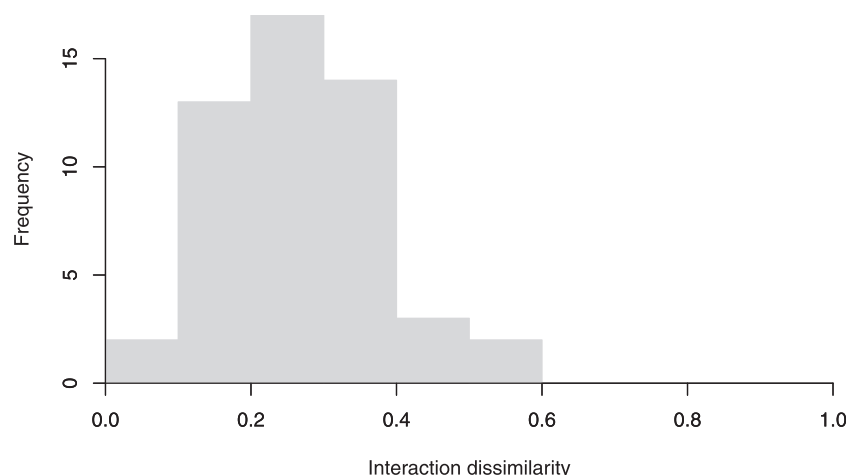


FIGURE 2 Distribution of β_{OS}' across the entire dataset. Local communities tend to keep a majority of the regional interactions, as evidenced by dissimilarity values going down to zero, with nevertheless some variability

2.5 | Interpretation

Taking a step back from the statistical analysis, we anticipate three possible outcomes with regard to the effect of environmental conditions on species and species interactions dissimilarity. In the first situation, environmental conditions have an impact on species, but not on their interactions; this would suggest that interactions are fully predicted by species traits or properties that are independent from environmental conditions (or that the suite of bioclimatic variables used did not contain the relevant information). In the second situation, species and interactions react to the same set of environmental predictors; this would suggest that species and interactions share environmental drivers and that interactions covary very strongly with species co-distribution. Finally, the alternative situation is one where interactions and species have some unique predictors; this would indicate that interactions are responding to aspects of the local environment that have no linear effect on species.

The difference in how realized and potential interactions respond to the environment will allow us to delineate the role of co-distribution versus local drivers of interactions. Specifically, unless the realized interactions are affected by drivers that have no impact on potential interactions, we will conclude that most of the variation in interactions over space is tied to species co-distribution rather than local variation driven by the environment. If drivers are shared between realized and potential interactions, especially if they are not shared with species, these environmental variables affect interactions at both scales.

3 | RESULTS

3.1 | Species and interactions vary across space

In Figure 2, we show that the 51 locations have realized interactions that are not the same as those found at the regional level. If interactions were perfectly conserved from the potential to realized steps, all locations would have $\beta_{OS}'=0$ (no dissimilarity). Although the strength of interactions varies across space, β_{OS}' measures whether their pres-

ence varies too (i.e., they are realized at one location, but not at another). Specifically, the mode of the distribution of β_{OS}' is c. 0.3, indicating that the filtering of interactions from the regional pool to local communities is comparable to other rodent-parasite communities (Poisot, Stanko, Miklisová, & Morand, 2013 reported values between 0.2 and 0.4 in a central European system). Using the Legendre and De Cáceres (2013) approach reveals that interactions vary more than species. Specifically, $\beta_R \approx 0.94$, $\beta_Q \approx 0.90$, $\beta_H \approx 0.80$ and $\beta_P \approx 0.79$. The finding that the variation in R is larger than that in Q is expected, as the realized interactions represent the action of additional ecological mechanisms involved in the filtering of potential interactions. The finding that localities in this dataset exhibit different interactions and have varying strengths of filtering from the potential to the realized interactions raises the question of identifying which of these localities have the strongest contribution to β -diversity.

3.2 | Species and interactions contribute differently to site uniqueness

Using the LCBP approach, we measure the extent to which each location contributes to the β -diversity of hosts, parasites, local and potential interactions. These results are presented in Figure 3. For each distance matrix, we clustered the locations using partition around the medoids (Kaufman & Rousseeuw, 1990) and selected the number of medoids that gave the smallest silhouette. This yielded three clusters for the hosts (east, north west and south west) and two for the parasites; the finding that hosts and parasites have different clusters suggests that they do not co-distribute perfectly. The same three clusters are found for the potential interactions and the realized interactions (north and south); this is expected because interactions are constrained by species composition, and so the finer division of host communities is also observed when using interactions as the proxy for community structure.

Overwhelmingly, the locations with significant contributions to β -diversity are found in the southernmost half of the dataset, in more desert areas (i.e., Turkestan, Taklamakan and Gobi deserts). These

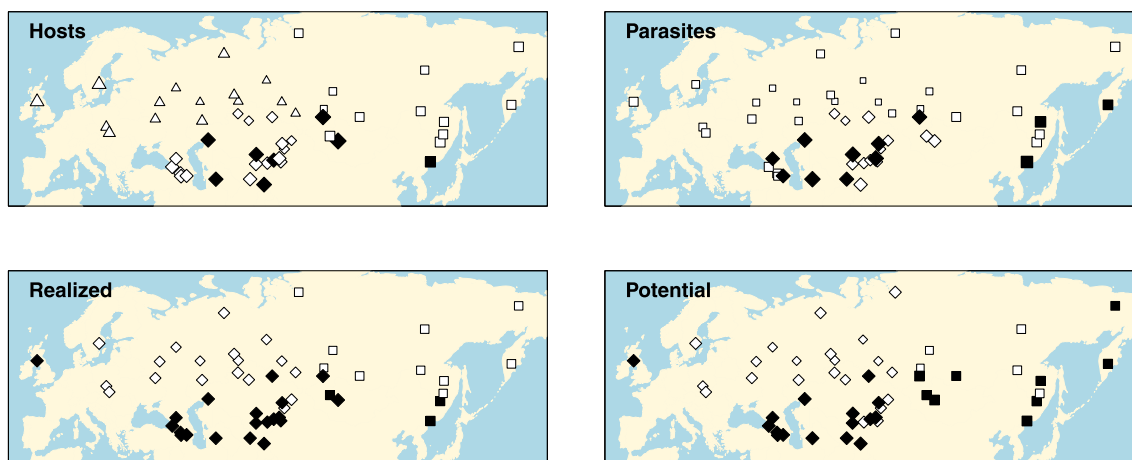


FIGURE 3 The β -diversity matrices were divided in clusters (indicated by different symbols). Black-filled symbols have significant local contribution to β -diversity (LCBD) values. The size of each symbol is proportional to its LCBD value. Note that there is no correspondence between the symbols used to denote cluster identity across the four panels

locations have environmental conditions that vary from what is found in the remainder of the dataset (see next section). There are eight host locations with significant contributions to β -diversity and 12 for parasites (i.e., locations that have unique species compositions). In contrast, there are 23 locations with unique realized interactions and 25 for potential interactions. With six exceptions, realized and potential interactions agree on which locations are unique.

3.3 | β -Diversity of species and interactions is structured by the environment

In Figure 4, we present the results of db-RDA analyses of the four β -diversity matrices, using the *bioClim* variables as predictors (additional information, such as number of constrained axes and inertia, are given in the Supporting Information). As in Figure 3 (and because environmental variables tend to be spatially autocorrelated), locations from the same cluster, and locations with significant contributions to β -diversity, tend to occupy the same area of the space defined by the canonical axes. This suggests two things. First, clustering of communities as a function of their species or interaction composition does indeed reflect that some species or interaction assemblages are specific to some combinations of environmental conditions. Second, locations with significant contributions to β -diversity are unique because their local environmental conditions select unique species and, within this local species pool, promote unique interaction assemblages.

Finally, in Table 1, we present the variables that have been retained during stepwise model building. The most important variables are the minimal temperature during the coldest month (bio06) and the minimal temperature during the wettest and coldest quarters (bio08 and bio10). Some precipitation-related variables were associated with hosts and interactions, but not parasites. Meanwhile, a number of variables were associated with parasites only (and, surprisingly, not with interactions). Finally, two variables (bio04, seasonality of temperature; bio11, mean temperature of warmest quarter) were associated with interactions only. From Table 1, it appears that the locations with the most

significant unique assemblages (in all matrices) also have low values of temperature during the wettest quarter (bio08), whereas locations with non-significant assemblages have high values of this variable.

4 | DISCUSSION

The β -diversity of interactions was higher than the β -diversity of species. Although notable, this is not a surprising result for at least two reasons. First, in any community there are many more interactions than species, and therefore interactions capture more of the variation of community structure than do species. Second, interactions are probabilistic events (Poisot, Cirtwill, et al., 2016) that can vary even if the two species involved are consistently co-occurring. Here, we show that part of this variability happens independently of species traits or attributes and is instead driven by local environmental conditions. Given that the information in species presence or absence is by definition included in the information about species interactions, it stands to reason that we would more adequately describe community structure and variation by systematically considering species interactions. This is emphasized by the fact that measuring LCBD based on interactions revealed over twice as many unique sites compared with measuring the LCBD of species (Figure 3). Quantifying the β -diversity of communities as based on their species composition (a) consistently underestimates β -diversity, and (b) consistently underestimates the uniqueness of localities in the dataset. A species-centred vision of diversity artificially homogenizes the structure of communities by missing important sources of variation.

An additional result of our study is that, although species and their interactions in this system do share a lot of their bioclimatic predictors, interactions (*R* and *Q*) responded to two variables that had no bearing on species (*P* and *H*) composition (Table 1). We recognize that this can, in part, stem from the mismatch of scales between the observation of species interactions and the observation of climatic variables; in itself, this is an intriguing question: what is the scale at which ecological interactions are affected by the environment? Methods such as Moran

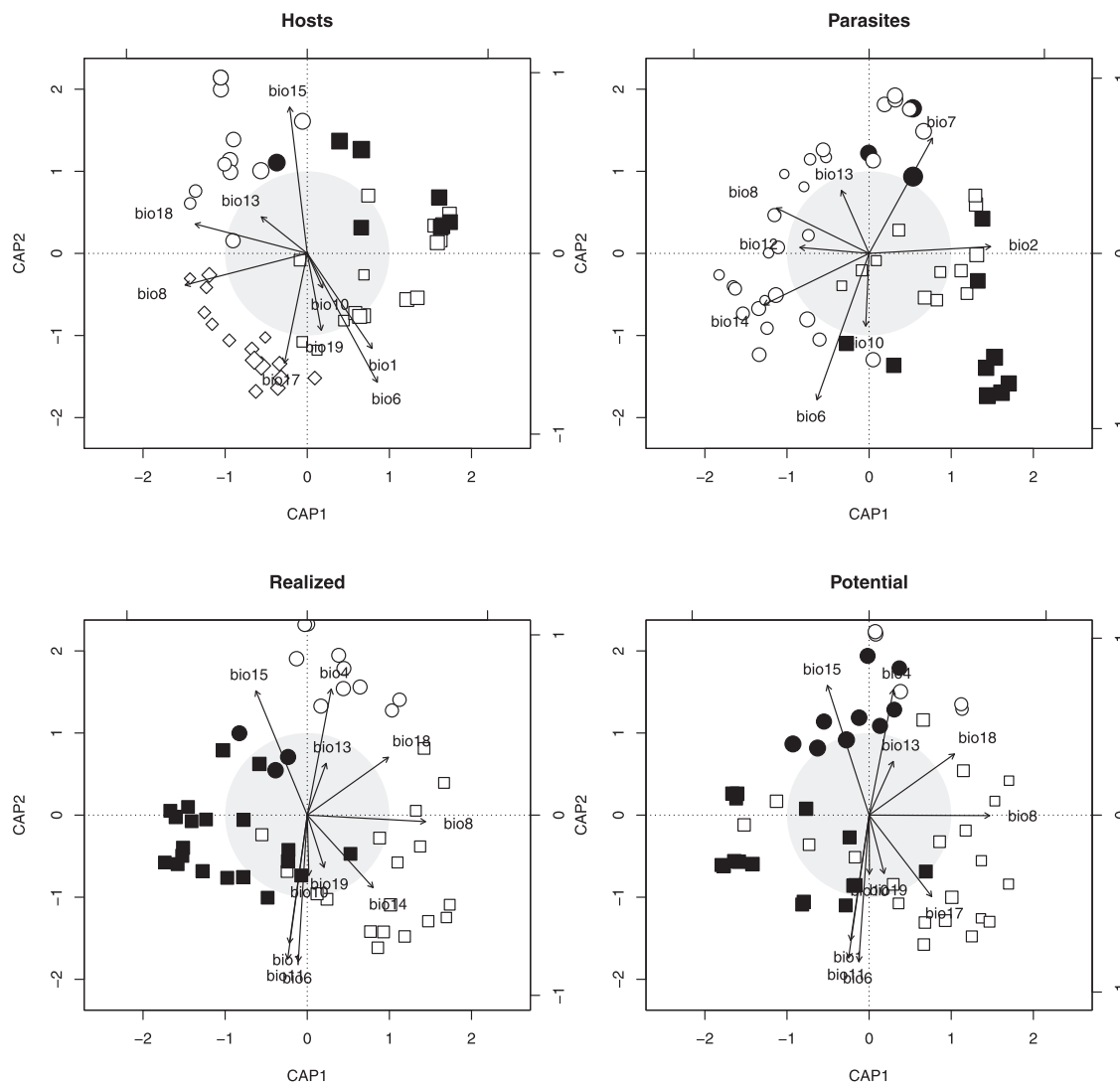


FIGURE 4 Ordinations of the four distance matrices, based on the scaled bioclimatic variables. Shape represents the cluster to which each location belongs, symbol size scales with local contribution to β -diversity (LCBD), and filled symbols have significant LCBD. The ordination diagrams were produced using the *capscale* function in *vegan*, hence the axes are labelled CAP1 and CAP2

eigenvector maps (Legendre & Gauthier, 2014) can help to pinpoint the right scale at which each environmental variable is acting on the system, but they are not applicable here because they would require the measurement of the environmental conditions that happen directly where and when interactions are sampled and cannot be extracted from interpolated bioclimatic data.

Nevertheless, our results highlight a challenge for current efforts to add biotic interactions to species distribution models (Araújo & Rozenfeld, 2014; Blois et al., 2014; Boulangeat, Gravel, & Thuiller, 2012). From additional predictors, species interactions become variables that must first be predicted themselves based on local environmental conditions. This is in addition to interactions requiring first and foremost the presence of both species (Gravel, Massol, Canard, Mouillot, & Mouquet, 2011) to be realized. Nevertheless, this problem may prove less complicated based on the observation that there is little difference (besides their relative importance) in the predictors of potential and realized interactions. What this suggests is that climatic variables act on the distribu-

tion of species pairs that can interact, and the signal at the level of realized interactions is inherited from the co-distribution of potential interaction partners; or in other words, the signal of potential interactions is a predictor of the co-occurrence of interacting species. As suggested by Poisot et al. (2015), interactions are filtered after the two species involved have co-distributed. It therefore stands to reason that environmental conditions that do not affect the co-distribution of the two species can affect the realization of the interaction locally.

4.1 | Biological implications

We report that communities close to desert areas tend to have unique interactions, more so than they have unique species assemblages. Rosenzweig and Winakur (1969) identified microhabitat features, such as resistance of soil to sheer stress, that modified the rules for competition between rodents species in deserts through their effect on vegetation cover. Likewise, because deserts are not environments as resource

TABLE 1 List of *bioClim* variables retained during the model selection (also used in Figure 4)

bioclim variable	Explanation	H	P	R	Q
6	Minimal temperature of coldest month	1	1	1	1
8	Mean temperature of wettest quarter	2	3	2	2
10	Mean temperature of coldest quarter	5	4	6	5
13	Precipitation of wettest month	4	5	9	4
1	Annual mean temperature	3		3	3
18	Precipitation of warmest quarter	6		5	6
15	Precipitation seasonality	7		4	8
19	Precipitation of coldest quarter	8		10	7
17	Precipitation of driest quarter	9			9
2	Mean diurnal range		2		
5	Maximal temperature of warmest month		6		
12	Annual precipitation		8		
14	Precipitation of wettest month		7		
4	Temperature seasonality			7	10
11	Mean temperature of warmest quarter			8	11
3	Isothermality				
7	Temperature annual range				
9	Mean temperature of driest quarter				
16	Precipitation of wettest quarter				

H = Hosts; P = parasites; Q = potential interactions; R = realized interactions. Note. Numbers identify the rank in the forward-selected model. Variables 4 and 11 (in bold) are unique to interactions.

rich as the usual rodent habitats, many rules of foraging are modified (Kotler & Brown, 1988). This includes increased among-patch mobility by hosts, and the overlap between diets of co-occurring species. Both intra- and interspecific transmission of ectoparasites can be modified as a result, creating unique communities in deserts. Kotler (1984) also identified an increase in predation risk in deserts, which are open habitats; this changes notably the times of the day when rodents are active. The joint constraints of finding food and avoiding predation can have cascading effects on the way hosts are exposed to parasites and the way parasites are transmitted. More recently, Bordes et al. (2015) examined the idea that habitat fragmentation (intensity of deforestation, switching from trees with a dense canopy to open shrubs and grasses unfavourable to rodents) can change the host–ectoparasite interaction networks. They found that increasingly open environments had increasingly different network structures (without looking into the detail of how interactions are organized within these networks). There is, in effect, a series of empirical evidence pointing to the fact that rodent–parasite interactions can be affected by the environment. Therefore, although our results should lead us to consider interaction networks as inherently dynamic objects that respond to the environment around them, they are not entirely surprising from an ecological point of view.

Across the entire spatial extent of the dataset, variables associated with interactions capture only warm temperatures (bio11) and temperature seasonality (bio04); this is an interesting contrast to the most important variables for all matrices that are associated with colder temperatures; this may suggest that interactions respond to modifications of host phenology when exposed to warmer temperatures, despite this aspect of the environment having no impact on host or parasite species occurrence. Temperature and precipitation have an impact on the specificity of fleas in this system (Krasnov et al., 2004). Our present study greatly expands this result. Not only does it pick up the contributions of variables to species distribution in addition to species interactions, but our approach works at the scale of each interaction (as opposed to the specificity, which aggregates across all interactions for a parasite). We are therefore confident that this method will allow researchers to study in greater detail, for a variety of systems, the way interactions respond to the environment.

4.2 | Conclusions

The results of this study highlight the following two important notions that we hope will be debated by biogeographers: (a) interactions contain intrinsically more information than species; and (b) interactions react to climatic variables in ways that seem to have no bearing on the concerned species. Taken together with the fact that the information on species occurrence is by definition nested within the information on species interactions, this points to the idea that we should be cautious when defining a ‘community’. In particular, we show that while describing the presence and absence of species at different locations is important, it misses a lot of information by systematically underestimating the variability and the uniqueness of these locations. For these reasons, we believe that the more desirable way of representing community structure is to describe not only the species, but also their interactions. Although this demands a much larger sampling effort, it is the correct approach not to neglect an entire aspect of what is community structure; species, not being independent entities, are organized in non-random ways through their interactions. Species interactions do also capture some information on species traits (Bartomeus et al., 2016; Coux et al., 2016), and adding traits as a key component of our definition of ‘community’ is indeed another obvious next step (McGill, Enquist, Weiher, & Westoby, 2006). As recent contributions expanded the framework of community dissimilarity to account for species traits (De Cáceres, Legendre, & He, 2013; Laliberté & Legendre, 2010), and the present contribution shows that the same work is possible with interactions, we hope that the method presented here is a solid enough methodological basis to start investigating whether, for any system, traits or interactions hold the most relevant information.

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AUTHOR CONTRIBUTION

T.P., D.G., P.L. and M.-J.F. designed the study; P.L. suggested how to compute LCBF from interaction matrices; C.G.-J. and T.P. performed the analyses and contributed equally to this study; T.P. and D.G. wrote the first draft of the manuscript. All authors provided comments and helped with the interpretation.

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BIOSKETCH

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