Beta and phylogenetic diversities tell complementary stories about ecological networks biogeography

Abstract: The beta-diversity of interactions between communities does not necessarily correspond to the differences related to their species composition because interactions show greater variability than species co-occurrence. Additionally, the structure of species interaction networks can itself vary over spatial gradients, thereby adding constraints on the dissimilarity of communities in space. We used published data on the parasitism interaction between fleas and small mammals in 51 regions of the Palearctic to investigate how beta-diversity of networks and phylogenetic diversity are related. The networks could be separated in groups based on the metrics that best described the differences between them, and these groups were also geographically structured. We also found that each network beta-diversity index relates in a particular way with phylogenetically community dissimilarity, reinforcing that some of these indexes have a strong phylogenetic component. Our results clarify important aspects of the biogeography of hosts and parasites communities in Eurasia, while suggesting that networks beta-diversity and phylogenetic dissimilarity interact with the environment in different ways.

Last revision: February 10, 2021

Authors

Gracielle Higino ^{1,2,*} Timothée Poisot ^{3,2,*}

Affiliations

- ¹ Universidade Federal de Goiás
- ² Québec Centre for Biodiversity Sciences
 ³ Université de Montréal

Correspondance

 * graciellehigino@gmail.com * timothee.poisot@umontreal.ca

Introduction

Ecological networks are complex units that incorporate many threads of the fabric of biodiversity, namely species identity, interactions, and shared coevolutionary history. Investigating the structure and the biogeography of communities through species interactions can therefore be highly informative. Local networks carry a record of both biogeographical and historical features of the regional pool of species and interactions, as they are subsets of a regional metaweb (Holt 2002), and so result from the fact that species both co-occur and interact. However, some of these characteristics can be lost over time, in response to local ecological pressure or random change. This is notably true of the co-phylogenetic signal of interactions (Desdevises et al. 2015; Boris R. Krasnov, Morand, and Poulin 2015), which can be eroded by environmental filtering during community assembly. This would result in a non-correlative variation of ecological networks components (Poisot and Stouffer 2018; Poisot et al. 2016). Dissimilarity of species interactions are always equal to, or greater than, the differences in species composition, because there cannot be an interaction without the presence of both partners. Therefore, interactions can be more informative than the species richness or functional diversity alone (Poisot et al. 2017). For instance, the probability of interaction may be modified by environmental changes that affect the metabolic rate of organisms (Rall et al. 2012), by changes in their habitats (Tylianakis and Morris 2017) or by community's phylogenetic structure (Coelho, Rodrigues, and 20 Rangel 2017) - which, in turn, varies with the abundance and specialization of species involved (Canard et al. 2014; Tylianakis and Morris 2017), but is not captured by looking solely at species composition. Environmental conditions also have direct effects over species fitness. In this sense, environmental gradients can change the frequency of interactions through direct influence on species' characteristics and population abundance, which, on the other

the environment can affect the production of secondary metabolites that exert selective pressure on the organisms that interact with certain plants (Muola et al. 2010), 29 how the geographical variation of functional characteristics generates changes in the interaction network and in species composition (König, Wiklund, and Ehrlén 2014; Cha et al. 2015), as well as the substitution of species along environmental gradients, variation in reproductive success and in the trophic network, or, yet, how the population density regulated by the environment can change the sign of an interaction (Bruder et al. 2017; Doxford, Ooi, and Freckleton 2013; Kaplan and Eubanks 2005). Because of all independent factors that can determine their occurrence, the differences between communities related to interactions may be, but not necessarily are, correspondent to those related to their species composition (Poisot, Stouffer, and Gravel 2014), and therefore the indexes that measure characteristics of ecological networks can also respond to environmental gradients in space and time (Dalsgaard et al. 2013; Baiser et al. 2019; Gravel et al. 2019). One of these indexes that carries important historical information is the phylogenetic diversity, measured as the sum of the lengths of the phylogeny branches that include all the species that interact in a community. Dispersion and speciation events are the main factors that affect the phylogenetic diversity of a network of ecological interactions (Coelho, Rodrigues, and Rangel 2017; Sebastián-González et al. 2015; Trøjelsgaard and Olesen 2013). Moreover, phylogenetic diversity is very sensitive to addition of species and may indicate, for example, the extent of impacts caused by an invasive species in a community (Davies and Buckley 2011). Therefore, beta diversity (the difference in the composition of communities) and the phylogenetic diversity of interaction networks are related, and both can respond to environmental variation in different ways. 51 Based on a parasite-host system distributed over a vast biogeographic region (Eurasia),

hand, are also affected by interactions (Poisot, Stouffer, and Gravel 2014). For example,

we identified similar numerical and geographical clusters between the phylogenetic

diversity and the dissimilarity of species composition and interactions of ecological networks. This result adds to our previous understanding of biodiversity distribution and help us tell a more complete story on the biogeography of ecological communities.

Specifically, we have found that local networks are characterized by beta-diversity metrics in different ways across the metaweb, and these metrics covary with networks' phylogenetic diversity.

Methods

- We used the Hadfield et al. (2014) data on the parasitism interaction between fleas
 and small mammals (Soricomorpha and Rodentia) in 51 regions of the Palearctic to
 investigate how beta-diversity of networks and phylogenetic diversity are related.
 This publication gathers occurrence records of 536,000 mammal individuals of 121
 species, 1,692,000 individuals from 206 flea species that occurred in those mammals,
 and the interactions between them (Hadfield et al. 2013). Original data is available
 at Data Dryad (http://dx.doi.org/10.5061/dryad.jf3tj) and interaction data is
 available at mangal database (http://mangal.io).
- The authors also used molecular and morphological traits of species to retrieve the phylogenetic relationships between species. We used the resulting trees to measure the phylogenetic community dissimilarity (PCD) of both hosts and parasites meta-communities using the function pcd of the package phyr, in R (Li et al. 2020; R Core Team 2018). To do that, we discarded sites with no correspondents taxa in the phylogenetic trees. The output of the pcd function can be divided in compositional (PCDc) and phylogenetic (PCDp) aspects of beta-diversity, which were summarized through a Principal Component Analysis (PCA) and grouped by their own *k-means* for both parasites and hosts.
- ⁷⁸ Because of particular characteristics such as communities' species composition and

- relationship with local environment, the differences in ecological networks can be
- ⁸⁰ due to species turnover, links established by shared species or a combination of both.
- In this sense, networks beta-diversity indexes are composed by their characteristics
- on species composition and interactions both on local and regional networks (Poisot
- et al. 2012). Here we assessed three indexes that summarize these information in dif-
- 84 ferent ways:
- 1. β s: this index corresponds to the differences on species composition between net-
- works. A high β s means solely a high species turnover (Koleff, Gaston, and Lennon
- 87 2003).
- 2. β os: this index represents the differences on interactions between shared species.
- 89 It is the component of networks dissimilarity only related to interactions, not species
- identity (Canard et al. 2014).
- 3. β wn: this summarizes the global differences between all networks in a metaweb,
- calculated as $\beta_w n = \beta_o s + \beta_s t$. It has two components: the difference in interactions
- between shared species (β os) and the difference in interactions due to species turnover
- β st. Therefore, β os can not assume values higher than β wn (Canard et al. 2014).
- These measures were calculated using the EcologicalNetworks.jl and Mangal.jl
- modules in Julia (Poisot et al. 2020; Poisot, Banville, and Dansereau 2020; Bezanson
- et al. 2017) and summarized with the KGL11 function, which calculates the Sørensen
- index of beta-diversity (Koleff, Gaston, and Lennon 2003). β s was the only metric cal-
- 99 culated separately for hosts and parasites because it represents their taxonomic diver-
- sities. The dissimilarity matrices resulting from this analysis represented, therefore,
- the differences between networks considering each of the indexes described above.
- In order to use these matrices on the following analyses as a single variable, we per-
- formed a PCA on each matrix and selected the first component of each. A subsequent
- PCA and *k-means* analysis on a combined matrix of these variables allowed us to in-
- vestigate how they co-vary among networks.

Results & Discussion

107 Communities can be grouped according to network beta-diversity.

The beta-diversity indexes described the dissimilarity of local networks in different ways across the metacommunity. In our case these differences were very prominent, making it possible to group communities by their interactions dissimilarity decomposition.

The first two axes of the Principal Component Analysis performed on the network 112 beta-diversity indexes, which explain 95.5% of the variation of the data, separates the 113 50 networks (those with corresponding species in the phylogenetic trees) in those that 114 have more similar βs , βos and βwn values (fig. 1). This separation is more explicit be-115 tween β s and β wn, and more diffuse for β os, which is aligned to the assumption that β s and β wn are only indirectly related, while β os have a more proximate relationship both with βwn and βs . The fact that the networks grouped by βs values are so different from those grouped by βwn may suggest that the turnover of species in the first group causes loss of links through loss of co-occurrence, while in the former group this turnover is translated into new connections. The β os group, however, would be 121 composed by communities that change less in species composition, but more in eco-122 logical interactions. 123

Because βs and the species composition of the phylogenetic community dissimilarity (PCDc) can be interpreted in the same way, a Principal Component Analysis of PCDc would provide a closer look to the βs metric. Our results suggest that, from a species composition point of view, parasites communities are much more similar across the metaweb than hosts, that can be more easily described in three main groups (fig. 1 D and fig. 1 E, respectively). The diversity of fleas can be much more uniform in space because it is common that a single host interacts with more than one species of parasites. In this particular case, only a few fleas communities have a distinguished

species composition and can be grouped together.

139

On the other hand, the three groups of the phylogenetic component of PCD (PCDp)
for both hosts and parasites are distinct: the diffuse group that appears in parasites'
PCDc does not repeat on PCDp. Additionally, both clades are arranged similarly in
the Principal Components space, with groups 2 and 3 being more alike than group 1.
This may be a reflex of the biogeographic history of communities, where one group is ancestral to the other two.

[Figure 1 about here.]

Each beta-diversity index relates in a particular way with phylogenetically community dissimilarity (PCD).

As expected, βs and PCDc are proxies for each other both for hosts and parasites, while PCDc is inversely correlated with βwn (fig. 2). Communities with a high βs value are very different from those around them, and the change in species composition could also represent a shift in the links inside these networks either because new species will probably explore different ranges of ecological niche or because the loss of species would also represent a loss of interaction. These changes in links inside networks are represented by βos , and its relationship with both PCDc and PCDp is highly variable (fig. 3 and fig. 5).

Because any change in species composition highly affects phylogenetic diversity, βs is also positively correlated with PCDp (fig. 3). Communities with high values for any of those metrics are located in regions with expected higher biodiversity (fig. 4 and fig. 5), and this may indicate that the biogeographical history of these communities are more related to migration than diversification of local lineages (Davies and Buckley 2011). Therefore, networks with high PCDp also represent communities with lower

ecological redundancy and higher functional diversity because it indicates that the species turnover is happening between species phylogenetically distant.

On the other hand, networks that are better represented by βwn - i.e., those which differences between them are significantly smaller than the differences in relation to the metaweb - are also phylogenetically similar, varying always inside a limited range of small dissimilarity (both with PCDc and PCDp). Because these communities also have low values of βs , indicating less frequent species turnover, this dissimilarity is due to different links between shared species. This result may reflect two possible scenarios:

1. In similar communities with low phylogenetic diversity (shorter branch lengths) the turnover of species could be adding very ecologically similar lineages, which leads to different interactions to prevent local extinction through competition.

2. In similar communities with high phylogenetic diversity (longer branch lengths) the species turnover may have been a result of invasion and migration, which may lead to opportunistic interactions.

This is also illustrated in fig. 2 and fig. 3 on scatterplots of β os vs. PCD: networks that differ little in phylogenies have a broader range of values of β os, while highly phylogenetically distinct networks only have very low values of β os - meaning that, for communities with high values of PCD, the few species that are shared interact in the same way. Additionally, because those same communities also have low values of β wn (i.e., they are very similar to the overall metaweb) and high values of β s (i.e., high species turnover), the interactions are probably being conserved also when species are replaced, like when two species that are phylogenetically distant replace each other in the same ecological function.

[Figure 2 about here.]

[Figure 3 about here.]

180

181

The separation of communities by components of beta-diversity was also observed geographically

There is a gradual transition between networks that were better described by turnover of species, clustered in central south Eurasia, to those more unique compared to the metaweb, spread in the north (fig. 4). The regional species pool is expected to be more diverse towards the tropics, and therefore local networks have a higher chance to have different species composition, which results in a strong contribution of β s for net-188 works beta-diversity. Because of the high diversity, species are functionally "packed," 189 and although some species could have more generalist interactions, they would rarely 190 do so, in order to avoid competition. Heading north, species turnover would be less 191 frequent due to a decrease in regional species richness, and now networks have more 192 shared species. They start to "unpack" and establish interactions with other remain-193 ing species, and therefore the β os component of beta-diversity explains better why 194 networks are different. The third group of networks, characterized by a high value 195 of β wn, is also composed by phylogenetically similar communities (as seen in fig. 3). 196 Because the species richness is even lower, any change in composition can have a high 197 impact on interactions. Therefore, the β os component is still very important, but now 198 differences in interactions due to species turnover contribute much more to networks' beta-diversity.

The phylogenetic community dissimilarity of networks was also geographically grouped, and in the region where βs was more important, there was a very distinguished group for both fleas' and mammals' phylogenetic dissimilarity (fig. 5). The two other groups are differently arranged in space: PCDc groups have a similar latitudinal distribution, but different longitudinal ranges, while PCDp groups are the opposite. This distribution of phylogenetic groups highlight the uniqueness of the southern-central set of communities, which suggests historical isolation of species. Additionally, the purely phylogenetic component of PCD reinforces the geographic distribution of betadiversity metrics as seen in fig. 4, with one group largely spread in the north - occupying a diverse range of environments - and two other groups restricted to latitudes under 60° (fig. 5).

[Figure 4 about here.]

[Figure 5 about here.]

4 Conclusion

212

213

The conspicuous association - both numerical and geographical - between the evolu-215 tionary history of species and networks' beta-diversity properties clarifies key aspects 216 of the biogeography of hosts and parasites communities in Eurasia. For example, the 217 longitudinal PCDc clusters separation roughly coincide with the presence of the Ural Mountains. From this point of view, considering the longitudinal spread of PCDp, the history of both hosts and parasites seems to follow a path of migration and diversification from south-central Eurasia towards the north. This history is also sustained 221 by the metaweb beta-diversity metrics: with a distinctive β s group at the south of the Ural Mountains suggesting higher species richness and common origin, followed towards north by gradual changes in interactions and composition, they sum up to the information unveiled by PCDp to describe a very likely biogeographic history. By describing how the phylogenetic differences between networks vary in the same 226 way within groups, this result seems to reinforce previous findings that there is no 227 co-phylogenetic matching between regional and local networks (Poisot and Stouffer 228 2018). If networks co-varied in continental scale in the same way they co-vary in local 229 scale, our analyses would not detect the groups illustrated in fig. 5. 230 Finally, this paper highlights how beta-diversity and phylogenetic dissimilarity are 231 related to each other, and sheds light on the possibility that they interact with the

environment in different ways. While βs seems to be connected to environmental uniqueness and geographical barriers, βos and βwn better reflect migration processes and evolutive trajectories. As stated at the beginning of this text, ecological networks are valuable, multidimensional lenses through which we can investigate biodiversity and its history. Although we did not account for properties such as phenology and natural history aspects of species, we did find that small scale processes such as species interactions can be integrated in large scale investigations and can have a stamp in macroecological processes.

Interaction networks between parasites and hosts have great potential to be used as 241 study systems in the geographic variation of interactions (Proulx, Promislow, and 242 Phillips 2005; Poulin 2010). Because of the particular type of association between 243 parasites and hosts, the dissimilarity of these interactions networks reflect not only 244 the environmental differences, but also the replacement of the host species (Eriksson 245 et al. 2019; Boris R. Krasnov et al. 2005; Poulin and Krasnov 2010). Nevertheless, the 246 association between parasites and hosts is often the result of the evolutionary his-247 tory of the groups, and this history can result in a non-neutral contribution of these 248 species to the beta diversity of these communities (Poisot et al. 2012). The underlying logic of our approach pertains to a wide diversity of systems; not only do rodents act as reservoirs for zoonotic diseases, (2020) show that understanding the global-scale structure of host-virus interactions requires a joint understanding of the geographical and evolutionary mechanisms involved in shaping them. We argue that when the data are available, there is even more information to be gained by looking at the way interactions vary.

References

Albery, Gregory F., Evan A. Eskew, Noam Ross, and Kevin J. Olival. 2020. "Predict-

- ing the Global Mammalian Viral Sharing Network Using Phylogeography." *Nature*
- 259 Communications 11 (1): 2260. https://doi.org/10.1038/s41467-020-16153-4.
- Baiser, Benjamin, Dominique Gravel, Alyssa R. Cirtwill, Jennifer A. Dunne, Ashkaan
- K. Fahimipour, Luis J. Gilarranz, Joshua A. Grochow, et al. 2019. "Ecogeographical
- Rules and the Macroecology of Food Webs." Global Ecology and Biogeography 28
- 263 (9): 1204–18. https://doi.org/10.1111/geb.12925.
- Bezanson, Jeff, Alan Edelman, Stefan Karpinski, and Viral B Shah. 2017. "Julia: A
- Fresh Approach to Numerical Computing." *SIAM Review* 59 (1): 65–98.
- ²⁶⁶ Bruder, Andreas, Romana K Salis, Peter E Jones, and Christoph D Matthaei. 2017. "Bi-
- otic Interactions Modify Multiple-Stressor Effects on Juvenile Brown Trout in an
- Experimental Stream Food Web." Global Change Biology 23 (9): 3882–94. https:
- //doi.org/10.1111/gcb.13696.
- ²⁷⁰ Canard, E F, N Mouquet, D Mouillot, M Stanko, D Miklisova, and D Gravel. 2014. "Em-
- pirical Evaluation of Neutral Interactions in Host-Parasite Networks." The Ameri-
- *can Naturalist* 183 (4): 468–79.
- ²⁷³ Cha, Muha, Xiaodong Wu, Heping Fu, Shuai Yuan, Yunga Wu, and Xiaodong Zhang.
- 2015. "An Empirical Research of Rodent Metacommunities in Alashan Desert."
- 275 Acta Ecologica Sinica 35 (17). https://doi.org/10.5846/stxb201312092913.
- Coelho, Marco Túlio Pacheco, João Fabrício Mota Rodrigues, and Thiago F Rangel.
- 2017. "Neutral Biogeography of Phylogenetically Structured Interaction Networks."
- Ecography 40 (12): 1467-74.
- Dalsgaard, Bo, Kristian Trøjelsgaard, Ana M Martín González, David Nogués-Bravo,
- Jeff Ollerton, Theodora Petanidou, Brody Sandel, et al. 2013. "Historical Climate-
- 281 Change Influences Modularity and Nestedness of Pollination Networks." Ecogra-
- *phy* 36 (12): 1331–40.
- Davies, T Jonathan, and Lauren B Buckley. 2011. "Phylogenetic Diversity as a Win-

- dow into the Evolutionary and Biogeographic Histories of Present-Day Richness
 Gradients for Mammals." *Philos. Trans. R. Soc. Lond. B Biol. Sci.* 366 (1576):
 2414–25.
- Desdevises, Yves, Serge Morand, Boris R. Krasnov, and Julien Claude. 2015. "Comparative Analysis: Recent Developments and Uses with Parasites." In *Parasite Diver-*sity and Diversification: Evolutionary Ecology Meets Phylogenetics, edited by Serge
 Morand, Boris R. Krasnov, and D. Timothy J. Littlewood, 337–50. Cambridge:
 Cambridge University Press. https://doi.org/10.1017/CB09781139794749.
- Doxford, Simon W., Mark K. J. Ooi, and Robert P. Freckleton. 2013. "Spatial and Temporal Variability in Positive and Negative Plant-Bryophyte Interactions Along a Latitudinal Gradient." *Journal of Ecology* 101 (2): 465–74. https://doi.org/10.
- Eriksson, Alan, Jean-françois Doherty, Erich Fischer, Gustavo Graciolli, and Robert
 Poulin. 2019. "Hosts and Environment Overshadow Spatial Distance as Drivers of
 Bat Fly Species Composition in the Neotropics." *Journal of Biogeography*.
- Gravel, Dominique, Benjamin Baiser, Jennifer A. Dunne, Jens-Peter Kopelke, Neo D.

 Martinez, Tommi Nyman, Timothée Poisot, et al. 2019. "Bringing Elton and Grinnell Together: A Quantitative Framework to Represent the Biogeography of Ecological Interaction Networks." *Ecography* 42 (3): 401–15. https://doi.org/10.

 1111/ecog.04006.
- Hadfield, Jarrod D, Boris R Krasnov, Robert Poulin, and Shinichi Nakagawa. 2014. "A
 Tale of Two Phylogenies: Comparative Analyses of Ecological Interactions." The
 American Naturalist 183 (2): 174–87.
- Hadfield, Jarrod D, Boris R Krasnov, Robert Poulin, and Nakagawa Shinichi. 2013.

 "Data from: A Tale of Two Phylogenies: Comparative Analyses of Ecological In-

```
Holt, Robert D. 2002. "Food Webs in Space: On the Interplay of Dynamic Instability
31
       and Spatial Processes." Ecological Research 17 (2): 261-73. https://doi.org/10.
       1046/j.1440-1703.2002.00485.x.
   Kaplan, Ian, and Micky D. Eubanks. 2005. "Aphids Alter the Community-Wide Impact
314
       of Fire Ants." Ecology 86 (6): 1640–49. https://doi.org/10.1890/04-0016.
315
   Koleff, Patricia, Kevin J Gaston, and Jack J Lennon. 2003. "Measuring Beta Diversity
316
       for Presence-Absence Data." Journal of Animal Ecology 72 (3): 367-82.
317
   König, Malin A. E., Christer Wiklund, and Johan Ehrlén. 2014. "Context-Dependent
318
       Resistance Against Butterfly Herbivory in a Polyploid Herb." Oecologia 174 (4):
319
       1265-72. https://doi.org/10.1007/s00442-013-2831-4.
320
   Krasnov, Boris R., Serge Morand, and Robert Poulin. 2015. "Phylogenetic Signals in
321
       Ecological Properties of Parasites." In Parasite Diversity and Diversification: Evo-
322
       lutionary Ecology Meets Phylogenetics, edited by Serge Morand, Boris R. Krasnov,
323
       and D. Timothy J. Littlewood, 351–59. Cambridge: Cambridge University Press.
324
       https://doi.org/10.1017/CB09781139794749.024.
325
   Krasnov, Boris R, Georgy I Shenbrot, David Mouillot, Irina S Khokhlova, and Robert
326
       Poulin. 2005. "Spatial Variation in Species Diversity and Composition of Flea As-
327
       semblages in Small Mammalian Hosts: Geographical Distance or Faunal Similar-
328
       ity?" Journal of Biogeography 32 (4): 633-44.
329
   Li, Daijiang, Russell Dinnage, Lucas Nell, Matthew R. Helmus, and Anthony Ives. 2020.
330
       "Phyr: An R Package for Phylogenetic Species-Distribution Modelling in Ecologi-
331
       cal Communities." bioRxiv. https://doi.org/10.1101/2020.02.17.952317.
332
   Muola, Anne, Pia Mutikainen, Marianna Lilley, Liisa Laukkanen, Juha Pekka Salmi-
       nen, and Roosa Leimu. 2010. "Associations of Plant Fitness, Leaf Chemistry,
```

teractions." https://doi.org/10.5061/DRYAD.JF3TJ.

310

and Damage Suggest Selection Mosaic in Plant-Herbivore Interactions." Ecology. 335 https://doi.org/10.1890/09-0589.1. 336 Poisot, Timothée, Francis Banville, and Gabriel Dansereau. 2020. "EcoJulia/Mangal.jl: 337 V0.3.1." Zenodo. https://doi.org/10.5281/zenodo.4299306. 338 Poisot, Timothée, Elsa Canard, David Mouillot, Nicolas Mouquet, and Dominique 339 Gravel. 2012. "The Dissimilarity of Species Interaction Networks." Ecology Letters 15 (12): 1353-61. 341 Poisot, Timothée, Alyssa R Cirtwill, Kévin Cazelles, Dominique Gravel, Marie Josée Fortin, and Daniel B Stouffer. 2016. "The Structure of Probabilistic Networks." 343 Methods in Ecology and Evolution 7 (3): 303-12. https://doi.org/10.1111/ 344 2041-210X.12468. Poisot, Timothée, Cynthia Guéveneux-Julien, Marie Josée Fortin, Dominique Gravel, and Pierre Legendre. 2017. "Hosts, Parasites and Their Interactions Respond to Different Climatic Variables." Global Ecology and Biogeography 26 (8): 942-51. https://doi.org/10.1111/geb.12602. Poisot, Timothée, Michiel Stock, Laura Hoebeke, Piotr Szefer, Francis Banville, and Giulio V. Dalla Riva. 2020. "Ecological Networks Analyses in Julia." Zenodo. 351 https://doi.org/10.5281/zenodo.4302247. 352 Poisot, Timothée, and Daniel B. Stouffer. 2018. "Interactions Retain the Co-Phylogenetic 353 Matching That Communities Lost." Oikos 127 (2): 230–38. https://doi.org/10. 354 1111/oik.03788. 355 Poisot, Timothée, Daniel B. Stouffer, and Dominique Gravel. 2014. "Beyond Species: 356 Why Ecological Interaction Networks Vary Through Space and Time." Oikos 124 357 (3): 243-51. https://doi.org/10.1111/oik.01719. 358 Poulin, Robert. 2010. "Network Analysis Shining Light on Parasite Ecology and Di-359

360

versity." Trends in Parasitology 26 (10): 492-98. https://doi.org/10.1016/j.

```
pt.2010.05.008.
```

- Poulin, Robert, and Boris R Krasnov. 2010. "Similarity and Variability of Parasite

 Assemblage Across Geographical Space." In *The Biogeography of Host-Parasite In-*teractions, edited by Serge Morand and Boris R Krasnov, 115127. Great Claredon

 Street, Oxford: Oxford University Press.
- Proulx, S, D Promislow, and P Phillips. 2005. "Network Thinking in Ecology and Evolution." *Trends in Ecology & Evolution* 20 (6): 345-53. https://doi.org/10.

 1016/j.tree.2005.04.004.
- R Core Team. 2018. *R: A Language and Environment for Statistical Computing*. Manual.

 Vienna, Austria: R Foundation for Statistical Computing.
- Rall, B. C., U. Brose, M. Hartvig, G. Kalinkat, F. Schwarzmuller, O. Vucic-Pestic, and
 O. L. Petchey. 2012. "Universal Temperature and Body-Mass Scaling of Feeding
 Rates." *Philosophical Transactions of the Royal Society B: Biological Sciences* 367
 (1605): 2923–34. https://doi.org/10.1098/rstb.2012.0242.
- Sebastián-González, Esther, Bo Dalsgaard, Brody Sandel, and Paulo R Guimarães.

 2015. "Macroecological Trends in Nestedness and Modularity of Seed-Dispersal

 Networks: Human Impact Matters." *Global Ecology and Biogeography* 24 (3): 293–

 303.
- Trøjelsgaard, Kristian, and Jens M Olesen. 2013. "Macroecology of Pollination Networks." *Global Ecology and Biogeography* 22 (2): 149–62.
- Tylianakis, Jason M., and Rebecca J. Morris. 2017. "Ecological Networks Across Environmental Gradients." *Annual Review of Ecology, Evolution, and Systematics* 48 (1):

 annurev-ecolsys-110316-022821. https://doi.org/10.1146/annurev-ecolsys-110316-022821.

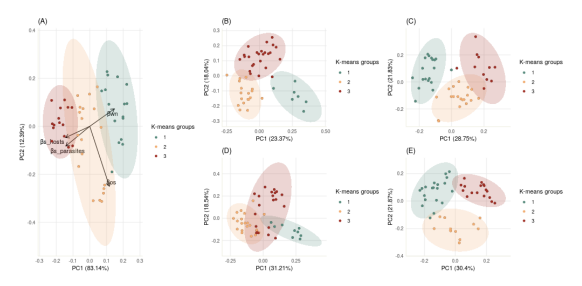


Figure 1: Principal Component Analysis of networks beta-diversity metrics and projection of local networks. For the dataset used here, networks are described by three dimensions of beta-diversity: while βs captures part of the variation that is complementary to that captured by βwn , βos describes a completely different dimension of the data. (A) PCA of networks beta-diversity metrics βs , βwn and βos ; PCA of the phylogenetic component of PCD (PCDp) for parasites (B) and hosts (C); PCA of the compositional component of PCD (PCDc) for parasites (D) and hosts (E).

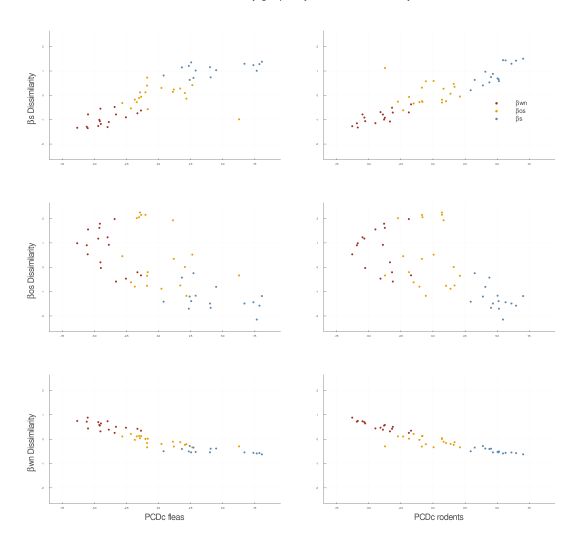


Figure 2: Effects of the compositional element of phylogenetic diversity dissimilarity on network beta-diversity for both parasites (left) and hosts (right). The colours correspond to the groups described on fig. 1. Networks with higher values of PCDc are taxonomically more distinct and therefore have higher values of βs and lower values of βos because they do not share many species.

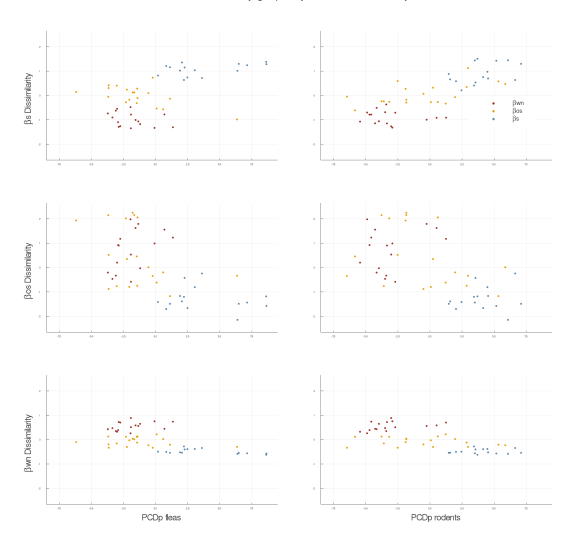


Figure 3: Effects of the phylogenetic component of the phylogenetic diversity dissimilarity on network beta-diversity for both parasites (left) and hosts (right). The colours correspond to the groups described on fig. 1. Networks with higher values of PCDp are phylogenetically more distinct, and therefore have lower values of β os (because they do not share many species). Networks better represented by β wn and β os are less distinguished on this aspect, but usually have lower values of PCDp.

Spatial clusters of networks by beta-diversity metrics

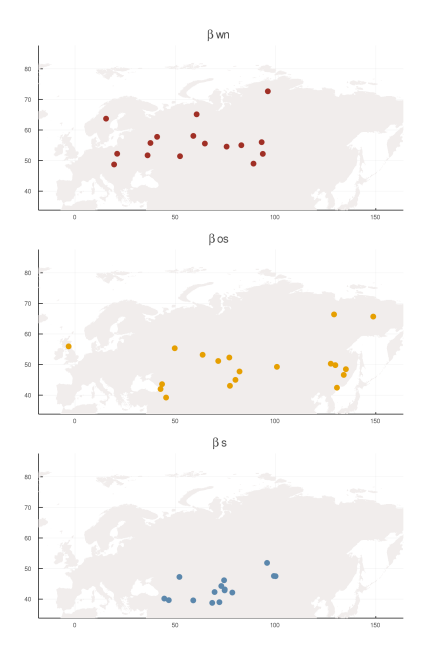


Figure 4: Spatial distribution of beta-diversity metrics. The groups detected in fig. 1 are also geographically distinguished

Spatial distribution of PCD components clusters for hosts and parasites



Figure 5: Spatial distribution of PCD components. Again, a distinct PCDc cluster (as seen on the third map of the left column) matches the cluster for which βs metric is more important.