Food web reconstruction through phylogenetic transfer of low-rank network representation

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Despite their importance in many ecological processes, collecting data and information on ecological interactions is an exceedingly challenging task. For this reason, large parts of the world have a data deficit when it comes to species interactions, and how the resulting networks are structured. As data collection alone is unlikely to be sufficient, community ecologists must adopt predictive methods. Here we develop such a method, relying on graph embedding and transfer learning to assemble a predicted list of trophic interactions between Canadian mammals. This interaction list is derived from the European food web, despite sharing only 5% of common species with Canada. The results of the predictive model are compared against databases of recorded pairwise interactions, showing that we correctly recover over 95% of known interactions. We provide guidance on how this method can be adapted by substituting some approaches or predictors in order to make it more generally applicable.

Introduction

- 2 There are two core challenges we are faced with in furthering our understanding of ecological networks
- across space, particularly at macro-ecologically relevant scales (e.g. Trøjelsgaard & Olesen 2016). First,
- 4 networks within a location are difficult to sample properly (Jordano 2016a, b), resulting in a widespread
- ⁵ "Eltonian shortfall" (Hortal et al. 2015), i.e. a lack of knowledge about inter and intra specific
- 6 relationships. This first challenge has been, in large part, addressed by the recent emergence of a suite of
- methods aiming to predict interactions within existing networks, many of which are reviewed in Strydom
- 8 et al. (2021a). Second, recent analyses based on collected data (Poisot et al. 2021a) or metadata (Cameron
- 9 et al. 2019) highlight that ecological networks are currently studied in a biased subset of space and
- bioclimates, which impedes our ability to generalize any local understanding of network structure.
- 11 Meaning that, although the framework to address incompleteness within networks exists, there would still
- be regions for which, due to a *lack* of local interaction data, we are unable to infer potential species
- interactions. Having a general solution for inferring *plausible* interactions (despite the unavailability of
- interaction data) could be the catalyst for significant breakthroughs in our ability to start thinking about
- species interaction networks over large spatial scales. In a recent overview of the field of ecological
- network prediction, Strydom et al. (2021a) identified two challenges of interest to the prediction
- of interactions at large scales. First, there is a relative scarcity of relevant data in most places
- 18 globally paradoxically, this restricts our ability to infer interactions to locations where
- inference is perhaps the least required; second, accurate predictions often demand accurate
- 20 predictors, and the lack of methods that can leverage small amount of data is a serious
- 21 impediment to our predictive ability globally.
- 22 Here, we present a general method to recommend possible trophic interactions, relying on the
- transfer learning of network representations, **specifically by using** similarities of species in a
- biologically/ecologically relevant proxy space (e.g. shared morphology or ancestry). Transfer learning is a
- 25 machine learning methodology that uses the knowledge gained from solving one problem and applying it
- to a related (destination) problem (Pan & Yang 2010; Torrey & Shavlik 2010). In this instance, we solve the
- 27 problem of predicting trophic interactions between species, based on knowledge extracted from another
- species pool for which interactions are known by using phylogenetic structure as a medium for transfer.
- There is a plurality of measures of species similarities that can be used for metaweb reconstruction (see e.g.

Morales-Castilla et al. 2015); however, phylogenetic proximity has several desirable properties when working at large scales. Gerhold et al. (2015) made the point that phylogenetic signal captures 31 diversification of characters (large macro-evolutionary process), but not necessarily community assembly 32 (fine ecological process); Dormann et al. (2010) previously found very similar conclusions. Interactions tend reflect a phylogenetic signal because they have a conserved pattern of evolutionary convergence that 34 encompasses a wide range of ecological and evolutionary mechanisms (Cavender-Bares et al. 2009; 35 Mouquet et al. 2012), and - most importantly - retain this signal even when it is not detectable at the 36 community scale (Hutchinson et al. 2017; Poisot & Stouffer 2018). Finally, species interactions at 37 macro-ecological scales seem to respond mostly to macro-evolutionary processes (Price 2003); which is 38 evidenced by the presence of conserved backbones in food webs (Dalla Riva & Stouffer 2016; Mora et al. **2018**), strong evolutionary signature on prey choice (Stouffer et al. 2012), and strong phylogenetic signature in food web intervality (Eklöf & Stouffer 2016). Phylogenetic reconstruction has also previously been used within the context of ecological networks, namely understanding ancestral plant-insect interactions (Braga et al. 2021). Taken together, these considerations suggest that phylogenies can reliably be used to transfer knowledge on species interactions.

[Figure 1 about here.]

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our methodology is outlined in fig. 1, where we provide an illustration based on learning the
embedding of a metaweb of trophic interactions for European mammals (known interactions;

Maiorano et al. 2020b, a) and, based on phylogenetic relationships between mammals globally
(i.e., phylogenetic tree Upham et al. 2019), infer a metaweb for the Canadian mammalian
species pool (interactions are treated as unknown in this instance). Following the definition of
Dunne (2006), a metaweb is a network analogue to the concept of a regional species pool.
Specifically, a metaweb is an inventory of all possible interactions within species likely to occurr
within a spatially delimited area (the network γ-diversity, in a sense). The metaweb is,
therefore, not a prediction of the food web at any specific locale within the frontiers of the
spatial area it recovers, and will in fact have a different structure (notably by having a larger
connectance; see e.g. Wood et al. 2015). These local food webs are expected to be a subset of both
the species and the interactions of their metaweb, and have been called "metaweb realizations"
(Poisot et al. 2015). The difference between a food web at a specific location and the metaweb

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has to do with a variety of mechanisms, including species co-occurrence, local environmental
   conditions, and local distribution of functional traits. Nevertheless, the metaweb represents the
   total of functional, phylogenetic, and macroecological processes (Morales-Castilla et al. 2015),
   and therefore still holds valuable ecological information. Because the metaweb can be
   down-sampled given appropriate knowledge of local species composition (the network
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   \alpha-diversity, in a sense), it is possible to infer what may drive the structure of food webs at finer
   spatial scales. This has been done for example for tree-gallers-parasitoid systems (Gravel et al.
   2018), fish trophic interactions (Albouy et al. 2019), tetrapods trophic interactions (O'Connor et
   al. 2020), and crop-pests networks (Grünig et al. 2020). Whereas the original metaweb definition
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   was based on presence/absence, we focus on probabilistic metawebs; not only does our method
   recommed interactions that may exist, it gives each interaction a score that is mathematically
   equivalent to the chance of success of a Bernoulli trial (see e.g. Poisot et al. 2016), which allows
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   properly weigh interactions as a function of how likely they are.
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   Our case study shows that phylogenetic transfer learning is an effective approach to the generation of
   probabilistic metawebs. This showcases that although the components (species) that make up the
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   Canadian and European communities may be minimally shared (the overall species overlap is less
   than 4%), if the medium (proxy space) selected in the transfer step is biologically plausible, we can still
   effectively learn from the known network and make biologically relevant predictions of interactions.
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   Indeed, as we detail in the result, when validated against known but fracational data of trophic
   interactions between Canadian mammals, our model achieves a predictive accuracy of approx.
   91%. It should be reiterated that the framework presented in fig. 1 is amenable to changes; notably, the
   measure of similarity may not be phylogeny, and can be replaced by information on foraging (Beckerman
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   et al. 2006), cell-level mechanisms (Boeckaerts et al. 2021), or a combination of traits and phylogenetic
   structure (Stock 2021). Most importantly, although we focus on a trophic system, it is an
   established fact that different types of interactions do themselves interact (Kéfi et al. 2012; see
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   e.g. Kawatsu et al. 2021). Future development of metaweb inference techniques should cover
   the prediction of multiple interaction types.
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Data used for the case study

We use data from the European metaweb assembled by Maiorano et al. (2020b).

88 TANYA TODO description of the data

- We induced the subgraph corresponding to all mammals by matching species names in the original
- 90 network to the GBIF taxonomic backbone (GBIF Secretariat 2021) and retaining all those who matched to
- mammals. This serves a dual purpose 1) to extract only mammals from the European network and 2) to
- match and standardize species names when aggregating the different data sources further downstream
- (which is an important consideration when combining datasets (Grenié et al. 2021)). All nodes had valid
- matches to GBIF at this step, and so this backbone is used for all name reconciliation steps as outlined
- 95 below.
- The European metaweb represents the knowledge we want to learn and transfer; the phylogenetic
- 97 similarity of mammals here represents the information for transfer. We used the mammalian consensus
- supertree by Upham et al. (2019), for which all approximatively 6000 names have been similarly matched
- to their GBIF valid names. This step allows us to place each node of the mammalian European metaweb
- in the phylogeny.
- The destination problem to which we want to transfer knowledge is the trophic interactions between
- mammals in Canada. We obtained the list of extant species from the IUCN checklist, and selected the
- terrestrial and semi-aquatic species (this corresponds to the same selection that was applied by Maiorano
- et al. (2020b) in the European metaweb). The IUCN names were, as previously, reconciled against GBIF to
- have an exact match to the taxonomy.
- After taxonomic cleaning and reconciliation as outlined in the following sections, the mammalian
- European metaweb has 260 species, and the Canadian species pool has 163; of these, 17 (about 4% of the
- total) are shared, and 89 species from Canada (54%) had at least one congeneric species in Europe. The
- similarity for both species pools predictably increases with higher taxonomic order, with 19% of shared
- genera, 47% of shared families, and 75% of shared orders; for the last point, Canada and Europe each had a
- single unique order (Didelphimorphia for Canada, Erinaceomorpha for Europe).
- In the following sections, we describe the representational learning step applied to European data, the
- transfer step through phylogenetic similarity, and the generation of a probabilistic metaweb for the

destination species pool.

Method description

The core point of our method is the transfer of knowledge of a known ecological network, in order to predict interactions between species from another location at which the network is unknown (or 117 partially known). In fig. 1, we give a high-level overview of the approach; in the example around which 118 this manuscript is built (leveraging detailed knowledge about binary trophic interactions between Mammalia in Europe to predict the less known trophic interactions between closely phylogenetically 120 related Mammalia in Canada), we use a series of specific steps for network embedding, trait inference, 121 network prediction and thresholding. Specifically, our approach can be summarized as follows: from the known network in Europe, we use a 123 truncated Singular Value Decomposition (t-SVD; Halko et al. 2011) to generate latent traits representing a 124 low-dimensional embedding of the network; these traits give an unbiased estimate of the node's position 125 in the latent feature spaces. Then, we map these latent traits onto a reference phylogeny (other 126 distance-based measures of species proximity that allow for the inference of features in the latent space 127 can be used, for example the dissimilarity in functional traits). Based on the reconstructed latent traits for species in the destination species pool, a Random Dot Product Graph model (hereafter RDPG; Young & 129 Scheinerman 2007) predicts the interaction between species through a function of the nodes' features 130 through matrix multiplication. Thus, from latent traits and node position, we can infer interactions. The method we develop is, ecologically speaking, a "black box," i.e. an algorithm that can be 132 understood mathematically, but whose component parts are not always directly tied to ecological processes. There is a growing realization in machine learning that (unintentional) 134 black box algorithms are not necessarily a bad thing Holm (2019), as long as their constituent 135 parts can be examined (which is the case with our method). But more importantly, data hold 136 more information that we may thought; as such, even algorithms that are disconnected from the model can make correct guesses most of the time (Halevy et al. 2009); in fact, in an instance 138 of ecological forecasting of spatio-temporal systems, model-free approaches (i.e. drawing all of 139 their information from the data) outperformed model-informed ones (Perretti et al. 2013).

141 Implementation and code availability

The entire pipeline is implemented in Julia 1.6 (Bezanson et al. 2017) and is available under the 142 permissive MIT License at https://osf.io/2zwgm/. The taxonomic cleanup steps are done using GBIF.jl 143 (Dansereau & Poisot 2021). The network embedding and analysis is done using EcologicalNetworks.jl (Poisot et al. 2019; Banville et al. 2021). The phylogenetic simulations are done using PhyloNetworks.jl 145 (Solís-Lemus et al. 2017) and Phylo. jl (Reeve et al. 2016). A complete Project.toml file specifying the 146 full tree of dependencies is available alongside the code. This material also includes a fully annotated copy of the entire code required to run this project (describing both the intent of the code and discussing some 148 technical implementation details), a vignette for every step of the process, and a series of Jupyter notebooks with the text and code. The pipeline can be executed on a laptop in a matter of minutes, and therefore does not require extensive computational power. 151

Step 1: Learning the origin network representation

The first step in transfer learning is to learn the structure of the original dataset. In order to do so, we rely 153 on an approach inspired from representational learning, where we learn a representation of the metaweb 154 (in the form of the latent subspaces), rather than a list of interactions (species a eats b). This approach is 155 conceptually different from other metaweb-scale predictions (e.g. Albouy et al. 2019), in that the metaweb 156 representation is easily transferable. Specifically, we use RDPG to create a number of latent variables that 157 can be combined into an approximation of the network adjacency matrix. RDPG results are known to 158 have strong phylogenetic signal, and to capture the evolutionary backbone of food webs (Dalla Riva & 159 Stouffer 2016); in other words, the latent variables of an RDPG can be mapped onto a 160 phylogenetic tree, and phylogenetically close predators should share phylogenetically close 161 **preys.** In addition, recent advances show that the latent variables produced this way can be used to 162 predict de novo network edges. Interstingly, the latent variables do not need to be prouced by decomposing the network itself; in a recent contribution, Runghen et al. (2021) show that deep 164 artificial neural networks are able to reconstruct the left and right subspaces of an RDPG, in 165 order to predict human movement networks from individual/location metadata. This is an 166 exciting opportunity, as it opens up the possibility of using additional predictors. 167

The latent variables are created by performing a truncated Singular Value Decomposition (t-SVD) on the

adjacency matrix. SVD is an appropriate embedding of ecological networks, which has recently been shown to both capture their complex, emerging properties (Strydom et al. 2021b) and to allow highly 170 accurate prediction of the interactions within a single network (Poisot et al. 2021b). Under SVD, an 171 adjacency matrix **A** (where $\mathbf{A}_{m,n} \in \mathbb{B}$ where 1 indicates predation and 0 an absence thereof) is decomposed into three components resulting in $A = U\Sigma V'$. Here, Σ is a $m \times n$ diagonal matrix and 173 contains only singular (σ) values along its diagonal, **U** is a $m \times m$ unitary matrix, and **V**' a $n \times n$ unitary 174 matrix. Truncating the SVD removes additional noise in the dataset by omitting non-zero and/or smaller 175 σ values from Σ using the rank of the matrix. Under a t-SVD $\mathbf{A}_{m,n}$ is decomposed so that Σ is a square $r \times r$ 176 diagonal matrix (whith $1 \le r \le r_{full}$ where r_{full} is the full rank of **A** and r the rank at which we truncate 177 the matrix) containing only non-zero σ values. Additionally, **U** is now a $m \times r$ semi unitary matrix and **V'** 178 a $n \times r$ semi-unitary matrix. As an aside, most ecologists are indirectly familiar with SVD: Principal Component Analysis is a special case of SVD, which is more sensitive to numerical 180 instabilities (see notably Shlens 2014). 181 The specific rank at which the SVD ought to be truncated is a difficult question. The purpose of SVD is to 182 remove the noise (expressed at high dimensions) and to focus on the signal, (expressed at low dimensions). 183 In datasets with a clear signal/noise demarcation, a scree plot of Σ can show a sharp drop at the rank where 184 noise starts (Zhu & Ghodsi 2006). Because the European metaweb is almost entirely known, the amount 185 of noise (uncertainty) is low; this is reflected in fig. 2 (left), where the scree plot shows no important drop, 186 and in fig. 2 (right) where the proportion of variance explained increases smoothly at higher dimensions. 187 For this reason, we default back to a threshold that explains 60% of the variance in the underlying data, corresponding to 12 dimensions - i.e. a tradeoff between accuracy and a reduced number of features. 189 An RDPG estimates the probability of observing interactions between nodes (species) as a function of the 190 nodes' latent variables, and is a way to turn a SVD (which decompose a matrix intro three) into 191 two matrices that can be multiplied to provide an approximation of the network. The latent 192 variables used for the RDPG, called the left and right subspaces, are defined as $\mathcal{L} = \mathbf{U}\sqrt{\Sigma}$, and $\mathcal{R} = \sqrt{\Sigma}\mathbf{V}'$ 193 - using the full rank of A, $\mathcal{LR} = A$, and using any smaller rank results in $\mathcal{LR} \approx A$. Using a rank of 1 for 194 the t-SVD provides a first-order approximation of the network. One advantage of using a RDPG rather 195 than a SVD is that the number of components to estimate decreases; notably, one does not have to estimate the eigenvalues of the SVD. Furthermore, the two subspaces can be directly multiplied to yield a network.

Because RDPG relies on matrix multiplication, the higher dimensions essentially serve to make specific 200 interactions converge towards 0 or 1; therefore, for reasonably low ranks, there is no guarantee that the 201 values in the reconstructed network will be within the unit range. In order to determine what constitutes 202 an appropriate threshold for probability, we performed the RDPG approach on the European metaweb, 203 and evaluated the probability threshold by treating this as a binary classification problem, specifically 204 assuming that both 0 and 1 in the European metaweb are all true. Given the methodological details given 205 in Maiorano et al. (2020b) and O'Connor et al. (2020), this seems like a reasonable assumption, although 206 one that does not hold for all metawebs. We used the thresholding approach presented in Poisot et al. 207 (2021b), and picked a cutoff that maximized Youden's J statistic (a measure of the informedness (trust) of 208 predictions; Youden (1950)); the resulting cutoff was 0.22, and gave an accuracy above 0.99. In Supp. 209 Mat. 1, we provide several lines of evidence that using the entire network to estimate the 210 threshold does not lead to overfitting; that using a subset of species would yield the same 211 threshold; that decreasing the quality of the original data by adding of removing interactions 212 would minimally affect the predicive accuracy of RDPG applied to the European metaweb; and that the networks reconstructed from artifically modified data are reconstructured with the 214 correct ecological properties. 215 The left and right subspaces for the European metaweb, accompanied by the threshold for prediction, 216 represent the knowledge we seek to transfer. In the next section, we explain how we rely on phylogenetic 217 similarity to do so.

Steps 2 and 3: Transfer learning through phylogenetic relatedness

In order to transfer the knowledge from the European metaweb to the Canadian species pool, we
performed ancestral character estimation using a Brownian motion model, which is a conservative
approach in the absence of strong hypotheses about the nature of phylogenetic signal in the network
decomposition (Litsios & Salamin 2012). This uses the estimated feature vectors for the European
mammals to create a state reconstruction for all species (conceptually something akin to a trait-based
mammalian phylogeny using generality and vulnerability traits) and allows us to impute the missing
(latent) trait data for the Canadian species that are not already in the European network; as we are focused

on predicting contemporary interactions, we only retained the values for the tips of the tree. We assumed that all traits (i.e. the feature vectors for the left and right subspaces) were independent, which is a 228 reasonable assumption as every trait/dimension added to the t-SVD has an additive effect to the one before 229 it. Note that the Upham et al. (2019) tree itself has some uncertainty associated to inner nodes of the phylogeny. In this case study, we have decided to not propagate this uncertainty, as it would complexify 231 the process. The Brownian motion algorithm returns the average value of the trait, and its upper and 232 lower bounds. Because we do not estimate other parameters of the traits' distributions, we considered that 233 every species trait is represented as a uniform distribution between these bounds. The choice of the 234 uniform distribution was made because the algorithm returns a minimum and maximum point 235 estimate for the value, and given this information, the uniform distribution is the one with 236 maximum entropy. Had all mean parameters estimates been positive, the exponential distribution would have been an alternative, but this is not the case for the subspaces of an 238 RDPG. In order to examine the consequences of the choice of distribution, we estimated the 239 variance per latent variable per node to use a normal distribution; as we show in Supp. Mat. 2, this decision results in dramatically over-estimating the number and probability of interactions, 241 and therefore we keep the discussions in the main text to the uniform case. The inferred left and 242 right sub-spaces for the Canadian species pool ($\hat{\mathcal{L}}$ and $\hat{\mathcal{R}}$) have entries that are distributions, representing the range of values for a given species at a given dimension. 244 These objects represent the transferred knowledge, which we can use for prediction of the Canadian 245 metaweb.

Step 4: Probabilistic prediction of the destination network

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The phylogenetic reconstruction of \hat{Z} and \hat{R} has an associated uncertainty, represented by the breadth of the uniform distribution associated to each of their entries. Therefore, we can use this information to assemble a *probabilistic* metaweb in the sense of Poisot *et al.* (2016), *i.e.* in which every interaction is represented as a single, independent, Bernoulli event of probability p.

[Figure 3 about here.]

Specifically, we have adopted the following approach. For every entry in $\hat{\mathcal{L}}$ and $\hat{\mathcal{R}}$, we draw a value from its distribution. This results in one instance of the possible left $(\hat{\ell})$ and right (\hat{r}) subspaces for the

Canadian metaweb. These can be multiplied, to produce one matrix of real values. Because the entries in $\hat{\ell}$ and \hat{r} are in the same space where ${\mathscr L}$ and ${\mathscr R}$ were originally predicted, it follows that the threshold ho256 estimated for the European metaweb also applies. We use this information to produce one random 257 Canadian metaweb, $N = \hat{\mathcal{L}}\hat{\mathcal{R}}' \geq \rho$. As we can see in (fig. 3) the European and Canadian metawebs are structurally similar (as would be expected given the biogeographic similarities) and the two (left and right) 259 subspaces are distinct i.e. capturing predation (generality) and prey (vulnerability) traits. 260 Because the intervals around some trait values can be broad (in fact, probably broader than what they would actually be, see e.g. Garland et al. 1999), we repeat the above process 2×10^5 times, which results in 262 a probabilistic metaweb P, where the probability of an interaction (here conveying our degree of trust that 263 it exists given the inferred trait distributions) is given by the number of times where it appears across all random draws N, divided by the number of samples. An interaction with $P_{i,j}=1$ means that these two 265 species were predicted to interact in all 2×10^5 random draws. 266 It must be noted that despite bringing in a large amount of information from the European 267 species pool and interactions, the Canadian metaweb has distinct structural properties. 268 Following an approach similar to Vermaat et al. (2009), we show in Supp. Mat. 3 that not only can we observed differences in a multivariate space between the European and Canadian metaweb, we can also observe differences in the same space between random subgraphs from 271 these networks. These results line up with the studies spatializing metawebs that have been 272 discussed in the introduction: changes in the species pool are driving local structural changes in the networks.

Data cleanup, discovery, validation, and thresholding

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Once the probabilistic metaweb for Canada has been produced, we followed a number of data inflation steps to finalize it. This step is external to the actual transfer learning framework but rather serves as a way to augment and validate the predicted metaweb.

[Figure 4 about here.]

First, we extracted the subgraph corresponding to the 17 species shared between the European and
Canadian pools and replaced these interactions with a probability of 0 (non-interaction) or 1 (interaction),

according to their value in the European metaweb. This represents a minute modification of the inferred network (about 0.8% of all species pairs from the Canadian web), but ensures that we are directly re-using 283 knowledge from Europe. 284 Second, we looked for all species in the Canadian pool known to the Global Biotic Interactions (GLoBI) 285 database (Poelen et al. 2014), and extracted their known interactions. Because GLOBI aggregates observed 286 interactions, it is not a networks data source, and therefore the only information we can reliably extract 287 from it is that a species pair was reported to interact at least once. This last statement should yet be taken with caution, as some sources in GLOBI (e.g. Thessen & Parr 2014) are produced through text analysis, 289 and therefore may not document direct evidence of the interaction. Nevertheless, should the predictive 290 model work, we would expect that a majority of interactions known to GLOBI would also be predicted. 29 We retrieved 366 interactions between mammals from the Canadian species pool from GLOBI, 292 33 of which were not predicted by the model; this results in a success rate of 91%. After performing 293 this check, we set the probability of all interactions known to GLOBI to 1. Finally, we downloaded the data from Strong & Leroux (2014), who mined various literature sources to 295 identify trophic interactions in Newfoundland. This dataset documented 25 interactions between mammals, only two of which were not part of our (Canada-level) predictions, resulting in a success rate of 297 92%. These two interactions were added to our predicted metaweb with a probability of 1. A table listing 298 all interactions in the predicted Canadian metaweb can be found in the supplementary material. 299

[Figure 5 about here.]

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Because the confidence intervals on the inferred trait space are probably over-estimates, we decided to apply a thresholding step to the interactions after the data inflation (fig. 5). Cirtwill & Hambäck (2021) 302 proposed a number of strategies to threshold probabilistic networks. Their methods assume the 303 underlying data to be tag-based sequencing, which represents interactions as co-occurrences of predator 304 and prey within the same tags; this is conceptually identical to our Bernoulli-trial based reconstruction of 305 a probabilistic network. We performed a full analysis of the effect of various cutoffs, and as they either 306 resulted in removing too few interactions, or removing enough interactions that species started to be 307 disconnected from the network, we set this threshold for a probability equivalent to 0 to the largest possible value that still allowed all species to have at least one interaction with a non-zero probability. The 309

need for this slight deviation from the Cirtwill & Hambäck (2021) method highlights the need for additional development on network thresholding.

Results and discussion of the case study

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In fig. 5, we examine the effect of varying the cutoff on $P(i \rightarrow j)$ on the number of links, species, and connectance. Determining a cutoff using the maximum curvature, or central difference approximation of the second order partial derivative, as suggested by *e.g.* Cirtwill & Hambäck (2021), results in species being lost, or almost all links being kept. We therefore settled on the value that allowed all species to remain with at least one interaction. This result, in and of itself, suggests that additional methodological developments for the thresholding of probabilistic networks are required.

[Figure 6 about here.]

The t-SVD embedding is able to learn relevant ecological features for the network. fig. 6 shows that the first rank correlates linearly with generality and vulnerability (Schoener 1989), i.e. the number of preys 321 and predators. Importantly, this implies that a rank 1 approximation represents the configuration model 322 for the metaweb, i.e. a set of random networks generated from a given degree sequence (Park & Newman 323 2004). Accounting for the probabilistic nature of the degrees, the rank 1 approximation also represents the 324 soft configuration model (van der Hoorn et al. 2018). Both models are maximum entropy graph models 325 (Garlaschelli et al. 2018), with sharp (all network realizations satisfy the specified degree sequence) and 326 soft (network realizations satisfy the degree sequence on average) local constraints, respectively. The (soft) 327 configuration model is an unbiased random graph model widely used by ecologists in the context of null 328 hypothesis significance testing of network structure (e.g. Bascompte et al. 2003) and can provide 329 informative priors for Bayesian inference of network structure (e.g. Young et al. 2021). It is noteworthy 330 that for this metaweb, the relevant information was extracted at the first rank. Because the first rank 331 corresponds to the leading singular value of the system, the results of fig. 6 have a straightforward 332 interpretation: degree-based processes are the most important in structuring the mammalian food web.

34 Discussion

One important aspect in which Europe and Canada differ (despite their comparable bioclimatic conditions) is the degree of the legacy of human impacts, which have been much longer in Europe. 336 Nenzén et al. (2014) showed that even at small scales (the Iberian peninsula), mammal food webs retain 337 the signal of both climate change and human activity, even when this human activity was orders of 338 magnitude less important than it is now. Similarly, Yeakel et al. (2014) showed that changes in human 339 occupation over several centuries can lead to food web collapse. Megafauna in particular seems to be very 340 sensitive to human arrival (Pires et al. 2015). In short, there is well-substantiated support for the idea that human footprint affects more than the risk of species extinction (Marco et al. 2018), and can lead to 342 changes in interaction structure. Yet, owing to the inherent plasticity of interactions, there have been 343 documented instances of food webs undergoing rapid collapse/recovery cycles over short periods of time 344 (Pedersen et al. 2017). The embedding of a network, in a sense, embeds its macro-evolutionary history, 345 especially as RDPG captures ecological signal (Dalla Riva & Stouffer 2016); at this point, it is important to 346 recall that a metaweb is intended as a catalogue of all possible interactions, which should then be filtered 347 (Morales-Castilla et al. 2015). In practice (and in this instance) the reconstructed metaweb will predict interactions that are plausible based on the species' evolutionary history, however some interactions 349 would not be realized due to human impact. 350 Dallas et al. (2017) suggest that most links in ecological networks may be cryptic, i.e. uncommon 351 or otherwise hard to observe. This argument essentially echoes Jordano (2016b): the sampling of 352 ecological interactions is difficult because it requires first the joint observation of two species, 353 and then the observation of their interaction. In addition, it is generally expected that weak or 354 rare links would be more common in networks (Csermely 2004), compared to strong, persistent 355 links; this is notably the case in food chains, wherein many weaker links are key to the stability 356 of a system (Neutel et al. 2002). In the light of these observations, the results in fig. 4 are not particularly surprising: we expect to see a surge in these low-probability interactions under a 358 model that has a good predictive accuracy. Because the predictions we generate are by design 359 probabilistic, then one can weigh these rare links appropriately. In a sense, that most ecological interactions are elusive can call for a slightly different approach to sampling: once the common 36 interactions are documented, the effort required in documenting each rare interaction may

increase exponentially. Recent proposals suggest that machine learning algorithms, in these situations, can act as data generators (Hoffmann et al. 2019): in this perspective, high quality 364 observational data can be supplemented with synthetic data coming from predictive models, 365 which increases the volume of information available for inference. Indeed, Strydom et al. (2021a) suggest that knowing the metaweb may render the prediction of local networks easier, 367 because it fixes an "upper bound" on which interactions can exist; indeed, with a probabilistic 368 metaweb, we can consider that the metaweb represents an aggregation of informative priors on 369 the interactions. 370 Related to the last point, Cirtwill et al. (2019) show that network inference techniques based on 371 Bayesian approaches would perform far better in the presence of an interaction-level informative prior; 372 the desirable properties of such a prior would be that it is expressed as a probability, preferably 373 representing a Bernoulli event, the value of which would be representative of relevant biological processes 374 (probability of predation in this case). We argue that the probability returned at the very last step of our 375 framework may serve as this informative prior; indeed, the output of our analysis can be used in 376 subsequent steps, also possibly involving expert elicitation to validate some of the most strongly 377 recommended interactions. One important caveat to keep in mind when working with interaction 378 inference is that interactions can never really be true negatives (in the current state of our methodological 379 framework and data collection limitations); this renders the task of validating a model through the usual 380 application of binary classification statistics very difficult (although see Strydom et al. 2021a for a 381 discussion of alternative suggestions). The other way through which our framework can be improved is by substituting the predictors that are used for transfer. For example, in the presence of information on 383 species traits that are known to be predictive of species interactions, one might want to rely on functional 384 rather than phylogenetic distances - in food webs, body size (and allometrically related variables) has been established as such a variable (Brose et al. 2006); the identification of relevant functional traits is 386 facilitated by recent methodological developments (Rosado et al. 2013). It should be noted that Xing & 387 Fayle (2021) highlight phylogenetic relatedness as one of the core components of network comparison at 388 the global scale. In this case study, we have embedded the original metaweb using t-SVD, because it lends 389 itself to an RDPG reconstruction, which is known to capture the consequences of evolutionary processes 390 (Dalla Riva & Stouffer 2016); this being said, there are others ways to embed graphs (Cai et al. 2017; Arsov 391 & Mirceva 2019; Cao et al. 2019), which can be used as alternatives.

As Herbert (1965) rightfully pointed out, "[y]ou can't draw neat lines around planet-wide problems"; in this regard, our approach (and indeed, any inference of a metaweb at large scales) must contend 394 with several interesting and interwoven families of problems. The first is the limit of the metaweb to 395 embed and transfer. If the initial metaweb is too narrow in scope, notably from a taxonomic point of view, the chances of finding another area with enough related species to make a reliable inference **decreases**; 397 this would likely be indicated by large confidence intervals during ancestral character 398 estimation, but the lack of well documented metawebs is currently preventing from developing 399 more concrete guidelines. The question of phylogenetic relatedness and dispersal is notably true 400 if the metaweb is assembled in an area with mostly endemic species, and as with every predictive 401 algorithm, there is room for the application of our best ecological judgement. Conversely, the 402 metaweb should be reliably filled, which assumes that the S^2 interactions in a pool of S species have been examined, either through literature surveys or expert elicitation. Supp. Mat. 1 provides some guidance 404 as to the type of sampling effort that should be prioritized. Although RDPG was able to 405 maintain very high predictive power when interactions where missing, the addition of false positive interactions was immediately detected; this suggests that it may be appropriate to err 407 on the side of "too many" interactions when constructing the initial metaweb to be transfered. 408 The second series of problems are related to determining which area should be used to infer the new metaweb in, as this determines the species pool that must be used. In our application, we focused on the mammals of Canada. The upside of this approach is that information at the country level is likely to be 411 required by policy makers and stakeholders for their biodiversity assessment, as each country tends to set 412 goals at the national level (Buxton et al. 2021) for which quantitative instruments are designed (Turak et al. 2017), with specific strategies often enacted at smaller scales (Ray et al. 2021). And yet, we do not 414 really have a satisfying answer to the question of "where does a food web stops?"; the current 415 most satisfying solutions involve examining the spatial consistency of network area relationships (see e.g. Galiana et al. 2018; Galiana et al. 2019, 2021), which is of course impossible 417 in the absence of enough information about the network itse;f. This suggests that an a posteriori 418 refinement of the results may be required, based on a downscaling of the metaweb. The final family of problems relates less to the availability of data or quantitative tools, and more to the praxis of spatial ecology. Operating under the context of national divisions, in large parts of the 421 world, reflects nothing more than the legacy of settler colonialism. Indeed, the use of ecological

data is not an apolitical act (Nost & Goldstein 2021), as data infrastructures tend to be designed to answer questions within national boundaries, and their use both draws upon and reinforces 424 territorial statecraft; as per Machen & Nost (2021), this is particularly true when the output of 425 "algorithmic thinking" (e.g. relying on machine learning to generate knowledge) can be re-used for governance (e.g. enacting conservation decisions at the national scale). We therefore 427 recognize that methods such as we propose operate under the framework that contributed to the 428 ongoing biodiversity crisis (Adam 2014), reinforced environmental injustice (Choudry 2013; Domínguez 429 & Luoma 2020), and on Turtle Island especially, should be replaced by Indigenous principles of land 430 management (Eichhorn et al. 2019; No'kmaq et al. 2021). As we see AI/ML being increasingly 431 mobilized to generate knowledge that is lacking for conservation decisions (e.g. Lamba et al. 432 2019; Mosebo Fernandes et al. 2020), our discussion of these tools need to go beyond the technical, and into the governance consequences they can have. 434 Acknowledgements: We acknowledge that this study was conducted on land within the traditional 435 unceded territory of the Saint Lawrence Iroquoian, Anishinabewaki, Mohawk, Huron-Wendat, and 436 Omàmiwininiwak nations. TP, TS, DC, and LP received funding from the Canadian Institue for Ecology & 437 Evolution. FB is funded by the Institut de Valorisation des Données. TS, SB, and TP are funded by a 438 donation from the Courtois Foundation. CB was awarded a Mitacs Elevate Fellowship no. IT12391, in 439 partnership with fRI Research, and also acknowledges funding from Alberta Innovates and the Forest 440 Resources Improvement Association of Alberta. M-JF acknowledges funding from NSERC Discovery 441 Grant and NSERC CRC. RR is funded by New Zealand's Biological Heritage Ngā Koiora Tuku Iho National Science Challenge, administered by New Zealand Ministry of Business, Innovation, and 443 Employment. BM is funded by the NSERC Alexander Graham Bell Canada Graduate Scholarship and the FRQNT master's scholarship. LP acknowledges funding from NSERC Discovery Grant (NSERC RGPIN-2019-05771). TP acknowledges financial support from NSERC through the Discovery Grants and 446 Discovery Accelerator Supplement programs.

References

- Adam, R. (2014). Elephant treaties: The Colonial legacy of the biodiversity crisis. UPNE.
- Albouy, C., Archambault, P., Appeltans, W., Araújo, M.B., Beauchesne, D., Cazelles, K., et al. (2019). The

- marine fish food web is globally connected. *Nature Ecology & Evolution*, 3, 1153–1161.
- 452 Arsov, N. & Mirceva, G. (2019). Network Embedding: An Overview. Available at:
- http://arxiv.org/abs/1911.11726. Last accessed.
- 454 Banville, F., Vissault, S. & Poisot, T. (2021). Mangal.jl and EcologicalNetworks.jl: Two complementary
- packages for analyzing ecological networks in Julia. *Journal of Open Source Software*, 6, 2721.
- ⁴⁵⁶ Bascompte, J., Jordano, P., Melian, C.J. & Olesen, J.M. (2003). The nested assembly of plant-animal
- mutualistic networks. *Proceedings of the National Academy of Sciences*, 100, 9383–9387.
- Beckerman, A.P., Petchey, O.L. & Warren, P.H. (2006). Foraging biology predicts food web complexity.
- *Proceedings of the National Academy of Sciences*, 103, 13745–13749.
- Bezanson, J., Edelman, A., Karpinski, S. & Shah, V. (2017). Julia: A Fresh Approach to Numerical
- 461 Computing. SIAM Review, 59, 65–98.
- Boeckaerts, D., Stock, M., Criel, B., Gerstmans, H., De Baets, B. & Briers, Y. (2021). Predicting
- bacteriophage hosts based on sequences of annotated receptor-binding proteins. Scientific Reports, 11,
- 464 1467.
- Braga, M.P., Janz, N., Nylin, S., Ronquist, F. & Landis, M.J. (2021). Phylogenetic reconstruction of ancestral
- ecological networks through time for pierid butterflies and their host plants. Ecology Letters, n/a.
- Brose, U., Jonsson, T., Berlow, E.L., Warren, P., Banasek-Richter, C., Bersier, L.-F., et al. (2006).
- 468 ConsumerResource Body-Size Relationships in Natural Food Webs. *Ecology*, 87, 2411–2417.
- Buxton, R.T., Bennett, J.R., Reid, A.J., Shulman, C., Cooke, S.J., Francis, C.M., et al. (2021). Key
- information needs to move from knowledge to action for biodiversity conservation in Canada.
- Biological Conservation, 256, 108983.
- ⁴⁷² Cai, H., Zheng, V.W. & Chang, K.C.-C. (2017). A Comprehensive Survey of Graph Embedding:
- 473 **Problems, Techniques and Applications. Available at:** http://arxiv.org/abs/1709.07604. Last
- accessed.
- ⁴⁷⁵ Cameron, E.K., Sundqvist, M.K., Keith, S.A., CaraDonna, P.J., Mousing, E.A., Nilsson, K.A., et al. (2019).
- Uneven global distribution of food web studies under climate change. *Ecosphere*, 10, e02645.
- 477 Cao, R.-M., Liu, S.-Y. & Xu, X.-K. (2019). Network embedding for link prediction: The pitfall and
- improvement. Chaos: An Interdisciplinary Journal of Nonlinear Science, 29, 103102.

- Cavender-Bares, J., Kozak, K.H., Fine, P.V.A. & Kembel, S.W. (2009). The merging of community ecology and phylogenetic biology. *Ecology Letters*, 12, 693–715.
- ⁴⁸¹ Choudry, A. (2013). Saving biodiversity, for whom and for what? Conservation NGOs, complicity,
- colonialism and conquest in an era of capitalist globalization. In: NGOization: Complicity,
- contradictions and prospects. Bloomsbury Publishing, pp. 24–44.
- Cirtwill, A.R., Ekl, A., Roslin, T., Wootton, K. & Gravel, D. (2019). A quantitative framework for
 investigating the reliability of empirical network construction. *Methods in Ecology and Evolution*, 0.
- Cirtwill, A.R. & Hambäck, P. (2021). Building food networks from molecular data: Bayesian or fixed-number thresholds for including links. *Basic and Applied Ecology*, 50, 67–76.
- Csermely, P. (2004). Strong links are important, but weak links stabilize them. *Trends in Biochemical Sciences*, 29, 331–334.
- Dalla Riva, G.V. & Stouffer, D.B. (2016). Exploring the evolutionary signature of food webs' backbones using functional traits. *Oikos*, 125, 446–456.
- Dallas, T., Park, A.W. & Drake, J.M. (2017). Predicting cryptic links in host-parasite networks.

 PLOS Computational Biology, 13, e1005557.
- Dansereau, G. & Poisot, T. (2021). SimpleSDMLayers.jl and GBIF.jl: A Framework for Species Distribution

 Modeling in Julia. *Journal of Open Source Software*, 6, 2872.
- Domínguez, L. & Luoma, C. (2020). Decolonising Conservation Policy: How Colonial Land and
- Conservation Ideologies Persist and Perpetuate Indigenous Injustices at the Expense of the
- Environment. Land, 9, 65.
- Dormann, C.F., Gruber, B., Winter, M. & Herrmann, D. (2010). Evolution of climate niches in European mammals? *Biology Letters*, 6, 229–232.
- Dunne, J.A. (2006). The Network Structure of Food Webs. In: *Ecological networks: Linking structure and dynamics* (eds. Dunne, J.A. & Pascual, M.). Oxford University Press, pp. 27–86.
- Eichhorn, M.P., Baker, K. & Griffiths, M. (2019). Steps towards decolonising biogeography. *Frontiers of Biogeography*, 12, 1–7.
- Eklöf, A. & Stouffer, D.B. (2016). The phylogenetic component of food web structure and intervality.
- Theoretical Ecology, 9, 107–115.

- Galiana, N., Barros, C., Braga, J., Ficetola, G.F., Maiorano, L., Thuiller, W., et al. (2021). The spatial
- scaling of food web structure across European biogeographical regions. *Ecography*, n/a.
- 6509 Galiana, N., Hawkins, B.A. & Montoya, J.M. (2019). The geographical variation of network
- structure is scale dependent: Understanding the biotic specialization of hostparasitoid
- networks. *Ecography*, 42, 1175–1187.
- Galiana, N., Lurgi, M., Claramunt-López, B., Fortin, M.-J., Leroux, S., Cazelles, K., et al. (2018).
- The spatial scaling of species interaction networks. *Nature Ecology & Evolution*, 2, 782–790.
- ⁵¹⁴ Garland, T., JR., Midford, P.E. & Ives, A.R. (1999). An Introduction to Phylogenetically Based Statistical
- Methods, with a New Method for Confidence Intervals on Ancestral Values 1. American Zoologist, 39,
- 516 374-388.
- Garlaschelli, D., Hollander, F. den & Roccaverde, A. (2018). Covariance structure behind breaking of
- ensemble equivalence in random graphs. *Journal of Statistical Physics*, 173, 644–662.
- GBIF Secretariat. (2021). GBIF Backbone Taxonomy.
- Gerhold, P., Cahill, J.F., Winter, M., Bartish, I.V. & Prinzing, A. (2015). Phylogenetic patterns are not
- proxies of community assembly mechanisms (they are far better). Functional Ecology, 29, 600–614.
- Gravel, D., Baiser, B., Dunne, J.A., Kopelke, J.-P., Martinez, N.D., Nyman, T., et al. (2018).
- Bringing Elton and Grinnell together: A quantitative framework to represent the
- biogeography of ecological interaction networks. *Ecography*, 0.
- 6525 Grenié, M., Berti, E., Carvajal-Quintero, J.D., Winter, M. & Sagouis, A. (2021). Harmonizing taxon names
- in biodiversity data: A review of tools, databases, and best practices.
- 6527 Grünig, M., Mazzi, D., Calanca, P., Karger, D.N. & Pellissier, L. (2020). Crop and forest pest
- metawebs shift towards increased linkage and suitability overlap under climate change.
- 529 Communications Biology, 3, 1–10.
- Halevy, A., Norvig, P. & Pereira, F. (2009). The Unreasonable Effectiveness of Data. IEEE
- 531 Intelligent Systems, 24, 8–12.
- Halko, N., Martinsson, P.G. & Tropp, J.A. (2011). Finding Structure with Randomness: Probabilistic
- Algorithms for Constructing Approximate Matrix Decompositions. SIAM Review, 53, 217–288.
- Herbert, F. (1965). *Dune*. **1st edn**. Chilton Book Company, Philadelphia.

- Hoffmann, J., Bar-Sinai, Y., Lee, L.M., Andrejevic, J., Mishra, S., Rubinstein, S.M., et al. (2019).
- Machine learning in a data-limited regime: Augmenting experiments with synthetic data
- uncovers order in crumpled sheets. Science Advances, 5, eaau6792.
- 538 Holm, E.A. (2019). In defense of the black box. *Science*, 364, 26–27.
- Hortal, J., de Bello, F., Diniz-Filho, J.A.F., Lewinsohn, T.M., Lobo, J.M. & Ladle, R.J. (2015). Seven
- Shortfalls that Beset Large-Scale Knowledge of Biodiversity. Annual Review of Ecology, Evolution, and
- *Systematics*, 46, 523–549.
- Hutchinson, M.C., Cagua, E.F. & Stouffer, D.B. (2017). Cophylogenetic signal is detectable in pollination
- interactions across ecological scales. *Ecology*, n/a-n/a.
- Jordano, P. (2016a). Chasing Ecological Interactions. *PLOS Biol*, 14, e1002559.
- Jordano, P. (2016b). Sampling networks of ecological interactions. Functional Ecology, 30, 1883–1893.
- Kawatsu, K., Ushio, M., van Veen, F.J.F. & Kondoh, M. (2021). Are networks of trophic
- interactions sufficient for understanding the dynamics of multi-trophic communities?
- Analysis of a tri-trophic insect food-web time-series. *Ecology Letters*, 24, 543–552.
- Kéfi, S., Berlow, E.L., Wieters, E.A., Navarrete, S.A., Petchey, O.L., Wood, S.A., et al. (2012). More
- than a meal... integrating non-feeding interactions into food webs: More than a meal....
- 551 *Ecology Letters*, 15, 291–300.
- Lamba, A., Cassey, P., Segaran, R.R. & Koh, L.P. (2019). Deep learning for environmental
- conservation. Current Biology, 29, R977-R982.
- Litsios, G. & Salamin, N. (2012). Effects of Phylogenetic Signal on Ancestral State Reconstruction.
- *Systematic Biology*, 61, 533–538.
- 556 Machen, R. & Nost, E. (2021). Thinking algorithmically: The making of hegemonic knowledge
- in climate governance. Transactions of the Institute of British Geographers, 46, 555–569.
- Maiorano, L., Montemaggiori, A., Ficetola, G.F., O'Connor, L. & Thuiller, W. (2020a). Data from: Tetra-EU
- 1.0: A species-level trophic meta-web of European tetrapods.
- Maiorano, L., Montemaggiori, A., Ficetola, G.F., O'Connor, L. & Thuiller, W. (2020b). TETRA-EU 1.0: A
- species-level trophic metaweb of European tetrapods. Global Ecology and Biogeography, 29, 1452–1457.

- Marco, M.D., Venter, O., Possingham, H.P. & Watson, J.E.M. (2018). Changes in human footprint drive changes in species extinction risk. *Nature Communications*, 9, 4621.
- Mora, B.B., Gravel, D., Gilarranz, L.J., Poisot, T. & Stouffer, D.B. (2018). Identifying a common
- backbone of interactions underlying food webs from different ecosystems. Nature
- 566 Communications, 9, 2603.
- Morales-Castilla, I., Matias, M.G., Gravel, D. & Araújo, M.B. (2015). Inferring biotic interactions from proxies. *Trends in Ecology & Evolution*, 30, 347–356.
- Mosebo Fernandes, A.C., Quintero Gonzalez, R., Lenihan-Clarke, M.A., Leslie Trotter, E.F. &
- Jokar Arsanjani, J. (2020). Machine Learning for Conservation Planning in a Changing
- Climate. Sustainability, 12, 7657.
- Mouquet, N., Devictor, V., Meynard, C.N., Munoz, F., Bersier, L.-F., Chave, J., et al. (2012).
- Ecophylogenetics: Advances and perspectives. *Biological Reviews*, 87, 769–785.
- Nenzén, H.K., Montoya, D. & Varela, S. (2014). The Impact of 850,000 Years of Climate Changes on the
- Structure and Dynamics of Mammal Food Webs. *PLOS ONE*, 9, e106651.
- Neutel, A.-M., Heesterbeek, J.A.P. & de Ruiter, P.C. (2002). Stability in Real Food Webs: Weak
 Links in Long Loops. Science, 296, 1120–1123.
- No'kmaq, M., Marshall, A., Beazley, K.F., Hum, J., joudry, shalan, Papadopoulos, A., et al. (2021).
- "Awakening the sleeping giant": Re-Indigenization principles for transforming biodiversity
- conservation in Canada and beyond. *FACETS*, 6, 839–869.
- Nost, E. & Goldstein, J.E. (2021). A political ecology of data. *Environment and Planning E: Nature*and Space, 25148486211043503.
- O'Connor, L.M.J., Pollock, L.J., Braga, J., Ficetola, G.F., Maiorano, L., Martinez-Almoyna, C., et al. (2020).
- Unveiling the food webs of tetrapods across Europe through the prism of the Eltonian niche. *Journal of*
- 585 Biogeography, 47, 181–192.
- Pan, S.J. & Yang, Q. (2010). A Survey on Transfer Learning. *IEEE Transactions on Knowledge and Data*Engineering, 22, 1345–1359.
- Park, J. & Newman, M.E.J. (2004). Statistical mechanics of networks. *Physical Review E*, 70, 066117.

- Pedersen, E.J., Thompson, P.L., Ball, R.A., Fortin, M.-J., Gouhier, T.C., Link, H., *et al.* (2017). Signatures of the collapse and incipient recovery of an overexploited marine ecosystem. *Royal Society Open Science*,
- 591 4, 170215.
- Perretti, C.T., Munch, S.B. & Sugihara, G. (2013). Model-free forecasting outperforms the correct
- mechanistic model for simulated and experimental data. Proceedings of the National
- 594 Academy of Sciences, 110, 5253-5257.
- Pires, M.M., Koch, P.L., Fariña, R.A., de Aguiar, M.A.M., dos Reis, S.F. & Guimarães, P.R. (2015).
- Pleistocene megafaunal interaction networks became more vulnerable after human arrival.
- 597 Proceedings of the Royal Society B: Biological Sciences, 282, 20151367.
- Poelen, J.H., Simons, J.D. & Mungall, C.J. (2014). Global biotic interactions: An open infrastructure to
- share and analyze species-interaction datasets. *Ecological Informatics*, 24, 148–159.
- Poisot, T., Belisle, Z., Hoebeke, L., Stock, M. & Szefer, P. (2019). EcologicalNetworks.jl analysing ecological networks. *Ecography*.
- Poisot, T., Bergeron, G., Cazelles, K., Dallas, T., Gravel, D., MacDonald, A., et al. (2021a). Global
- knowledge gaps in species interaction networks data. Journal of Biogeography, n/a.
- Poisot, T., Cirtwill, A.R., Cazelles, K., Gravel, D., Fortin, M.-J. & Stouffer, D.B. (2016). The structure of probabilistic networks. *Methods in Ecology and Evolution*, 7, 303–312.
- Poisot, T., Ouellet, M.-A., Mollentze, N., Farrell, M.J., Becker, D.J., Albery, G.F., et al. (2021b). *Imputing*
- the mammalian virome with linear filtering and singular value decomposition. Available at:
- 609 http://arxiv.org/abs/2105.14973. Last accessed.
- Poisot, T. & Stouffer, D.B. (2018). Interactions retain the co-phylogenetic matching that communities lost.
- 611 Oikos, 127, 230–238.
- Poisot, T., Stouffer, D.B. & Gravel, D. (2015). Beyond species: Why ecological interaction
- networks vary through space and time. *Oikos*, 124, 243–251.
- Price, P.W. (2003). Macroevolutionary theory on macroecological patterns. Cambridge University Press.
- Ray, J.C., Grimm, J. & Olive, A. (2021). The biodiversity crisis in Canada: Failures and challenges of
- federal and sub-national strategic and legal frameworks. FACETS, 6, 1044–1068.

- Reeve, R., Leinster, T., Cobbold, C.A., Thompson, J., Brummitt, N., Mitchell, S.N., et al. (2016). How to
- partition diversity. Available at: http://arxiv.org/abs/1404.6520. Last accessed. 618
- Rosado, B.H.P., Dias, A. & de Mattos, E. (2013). Going Back to Basics: Importance of Ecophysiology when 619
- Choosing Functional Traits for Studying Communities and Ecosystems. Natureza & conservaç~ao 620
- revista brasileira de conservaç~ao da natureza, 11, 15-22. 621
- Runghen, R., Stouffer, D.B. & Dalla Riva, G.V. (2021). Exploiting node metadata to predict interactions in 622 large networks using graph embedding and neural networks.
- Schoener, T.W. (1989). Food webs from the small to the large. Ecology, 70, 1559-1589.
- Shlens, J. (2014). A Tutorial on Principal Component Analysis. Available at: 625
- http://arxiv.org/abs/1404.1100. Last accessed. 626
- Solís-Lemus, C., Bastide, P. & Ané, C. (2017). PhyloNetworks: A Package for Phylogenetic Networks. 627
- Molecular Biology and Evolution, 34, 3292-3298. 628
- Stock, M. (2021). Pairwise learning for predicting pollination interactions based on traits and phylogeny. 629
- Ecological Modelling, 14. 630

623

- Stouffer, D.B., Sales-Pardo, M., Sirer, M.I. & Bascompte, J. (2012). Evolutionary Conservation of Species' 631
- Roles in Food Webs. *Science*, 335, 1489–1492. 632
- Strong, J.S. & Leroux, S.J. (2014). Impact of Non-Native Terrestrial Mammals on the Structure of the
- Terrestrial Mammal Food Web of Newfoundland, Canada. PLOS ONE, 9, e106264. 634
- Strydom, T., Catchen, M.D., Banville, F., Caron, D., Dansereau, G., Desjardins-Proulx, P., et al. (2021a). A 635
- roadmap towards predicting species interaction networks (across space and time). 636
- Philosophical Transactions of the Royal Society B: Biological Sciences, 376, 20210063. 637
- Strydom, T., Dalla Riva, G.V. & Poisot, T. (2021b). SVD Entropy Reveals the High Complexity of Ecological 638
- Networks. Frontiers in Ecology and Evolution, 9. 639
- Thessen, A.E. & Parr, C.S. (2014). Knowledge extraction and semantic annotation of text from the
- encyclopedia of life. PloS one, 9, e89550. 641
- Torrey, L. & Shavlik, J. (2010). Transfer learning. In: Handbook of research on machine learning 642
- applications and trends: Algorithms, methods, and techniques. IGI global, pp. 242–264. 643

- Trøjelsgaard, K. & Olesen, J.M. (2016). Ecological networks in motion: Micro- and macroscopic variability across scales. *Functional Ecology*, 30, 1926–1935.
- Turak, E., Brazill-Boast, J., Cooney, T., Drielsma, M., DelaCruz, J., Dunkerley, G., *et al.* (2017). Using the
 essential biodiversity variables framework to measure biodiversity change at national scale. *Biological Conservation*, SI:measures of biodiversity, 213, 264–271.
- Upham, N.S., Esselstyn, J.A. & Jetz, W. (2019). Inferring the mammal tree: Species-level sets of
 phylogenies for questions in ecology, evolution, and conservation. *PLOS Biology*, 17, e3000494.
- van der Hoorn, P., Lippner, G. & Krioukov, D. (2018). Sparse Maximum-Entropy Random Graphs with a

 Given Power-Law Degree Distribution. *Journal of Statistical Physics*, 173, 806–844.
- Vermaat, J.E., Dunne, J.A. & Gilbert, A.J. (2009). Major dimensions in food-web structure
 properties. *Ecology*, 90, 278–282.
- Wood, S.A., Russell, R., Hanson, D., Williams, R.J. & Dunne, J.A. (2015). Effects of spatial scale of
 sampling on food web structure. *Ecology and Evolution*, 5, 3769–3782.
- Xing, S. & Fayle, T.M. (2021). The rise of ecological network meta-analyses: Problems and prospects.
 Global Ecology and Conservation, 30, e01805.
- Yeakel, J.D., Pires, M.M., Rudolf, L., Dominy, N.J., Koch, P.L., Guimarães, P.R., *et al.* (2014). Collapse of an ecological network in Ancient Egypt. *PNAS*, 111, 14472–14477.
- Youden, W.J. (1950). Index for rating diagnostic tests. *Cancer*, 3, 32–35.
- Young, J.-G., Cantwell, G.T. & Newman, M.E.J. (2021). Bayesian inference of network structure from unreliable data. *Journal of Complex Networks*, 8.
- Young, S.J. & Scheinerman, E.R. (2007). Random Dot Product Graph Models for Social Networks. In:
- Algorithms and Models for the Web-Graph, Lecture Notes in Computer Science (eds. Bonato, A. &
- 666 Chung, F.R.K.). Springer, Berlin, Heidelberg, pp. 138–149.
- Zhu, M. & Ghodsi, A. (2006). Automatic dimensionality selection from the scree plot via the use of profile likelihood. *Computational Statistics & Data Analysis*, 51, 918–930.

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Figure 1: Overview of the phylogenetic transfer learning (and prediction) of species interactions networks. Starting from an initial, known, network, we learn its representation through a graph embedding step (here, a truncated Singular Value Decomposition; Step 1), yielding a series of latent traits (vulnerability traits representing species at the lower trophic-level and generality traits representing species at higher trophic-levels; *sensu* Schoener (1989)); second, for the destination species pool, we perform ancestral character estimation using a phylogeny (here, using a Brownian model for the latent traits; Step 2); we then sample from the reconstructed distribution of latent traits (Step 3) to generate a probabilistic metaweb at the destination (here, assuming a uniform distribution of traits), and threshold it to yield the final list of interactions (Step 4).

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Figure 2: Left: representation of the screeplot of the singular values from the t-SVD on the European metaweb. The screeplot shows no obvious drop in the singular values that may be leveraged to automatically detect a minimal dimension for embedding, after *e.g.* Zhu & Ghodsi (2006). Right: cumulative fraction of variance explained by each dimension up to the rank of the European metaweb. The grey lines represent cutoffs at 50, 60... 90% of variance explained. For the rest of the analysis, we reverted to an arbitrary threshold of 60% of variance explained, which represented a good tradeoff between accuracy and reduced number of features.

figures/figure-subspaces.png

Figure 3: Visual representation of the left (green/purple) and right (green/brown) subspaces, alongside the adjacency matrix of the food web they encode (greyscale). The European metaweb is on the left, and the imputed Canadian metaweb (before data inflation) on the right. This figure illustrates how much structure the left sub-space captures. As we show in fig. 6, the species with a value of 0 in the left subspace are species without any prey.

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Figure 4: Left, comparison of the probabilities of interactions assigned by the model to all interactions (grey curve), the subset of interactions found in GLOBI (red), and in the Strong & Leroux (2014) Newfoundland dataset (blue). The model recovers more interaction with a low probability compared to data mining, which can suggest that collected datasets are biased towards more common or easy to identify interactions. Right, distribution of the in-degree and out-degree of the mammals from Canada in the reconstructed metaweb. This figure describes a flat, relatively short food web, in which there are few predators but a large number of preys.

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Figure 5: Left: effect of varying the cutoff for probabilities to be considered non-zero on the number of unique links and on \hat{L} , the probabilistic estimate of the number of links assuming that all interactions are independent. Right: effect of varying the cutoff on the number of disconnected species, and on network connectance. In both panels, the grey line indicates the cutoff $P(i \rightarrow j) \approx 0.08$ that resulted in the first species losing all of its interactions.

figures/figure-degree.png

Figure 6: Top: biological significance of the first dimension. Left: there is a linear relationship between the values on the first dimension of the left subspace and the generality, *i.e.* the relative number of preys, *sensu* Schoener (1989). Species with a value of 0 in this subspace are at the bottom-most trophic level. Right: there is, similarly, a linear relationship between the position of a species on the first dimension of the right subspace and its vulnerability, *i.e.* the relative number of predators. Taken together, these two figures show that the first-order representation of this network would capture its degree distribution. Bottom: topological consequences of the first dimension. Left: differences in the *z*-score of the actual configuration model for the reconstructed network, and the prediction based only on the first dimension. Right: distribution of the differences in the left panel.