Spatially explicit predictions of food web structure from regional level data

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Gabriel Dansereau Ceres Barros Timothée Poisot

Abstract

3

Knowledge about how ecological networks vary across global scales is currently limited given the complexity of acquiring repeated spatial data for species interactions. Yet, recent developments of metawebs highlight efficient ways to first document possible interactions within regional species pools. Downscaling metawebs towards local network predictions is a promising approach to use current data to investigate the variation of networks across space. However, issues remain in how to represent the spatial 10 variability and uncertainty of species interactions, especially for large scale large-scale food webs. Here, we present a probabilistic framework to downscale a metaweb based on the Canadian mammal metaweb and species occurrences from GBIF. We investigate global databases. We investigated how our approach 13 can be used to represent the variability of networks and communities between ecoregions in Canada. 14 Our results highlighted mismatches in the distribution of species Species richness and interactions, especially in their within-ecoregion variability, indicating that interactions vary differently than species distributions over continental-scale gradients. Results summarized by ecoregion showed non-constant 17 variation within and between ecologically meaningful biogeographic boundaries and followed a similar latitudinal gradient across ecoregions but at the same time identified contrasting diversity hotspots. Network motifs revealed additional areas of variation in network structure compared to species richness

and number of links. Our method offers the potential to bring global predictions down to a more actionable local scale, and increases the diversity of ecological networks that can be projected in space.

Introduction

Because species interactions vary in time and space, and because species show high turnover over larger spatial extents, adequately capturing the diversity of ecological networks is a challenging task 25 (Jordano 2016). Most studies on food webs have previously focused on local networks limited in size and extent, and are rarely replicated in space or time (Mestre et al. 2022). Interactions can show important variations in space (Poisot et al. 2015; Zarnetske et al. 2017), yet available network data also show important geographical bias by focusing sampling efforts in a few areas or biomes, limiting our ability to 29 answer questions in many biomes and over broad spatial extents (Poisot et al. 2021). Moreover, global monitoring of biotic interactions is insufficient to properly describe and understand how ecosystems are reacting to global change (Windsor et al. 2023). Approaches to predict species interactions (e.g., 32 Morales-Castilla et al. 2015; Desjardins-Proulx et al. 2017) are increasingly used as an alternative to 33 determine potential interactions; they can handle limited data to circumvent data scarcity (Strydom et al. 2021), but are still rarely used to make explicitly spatial predictions. As a result, there have been repeated calls for globally distributed interaction and network data coupled to repeated sampling in time 36 and space (Mestre et al. 2022; Windsor et al. 2023), which will help understand the macroecological 37 variations of food webs (Baiser et al. 2019). Despite these limitations, food web ecologists often can infer a reasonable approximation of the network 39

Despite these limitations, food web ecologists often can infer a reasonable approximation of the network existing within a region. This representation, called a metaweb, contains all possible interactions between species in a given regional species pool (Dunne 2006), and provides a solid foundation to develop approaches to estimate the structure of networks at finer spatial scales.

When assembled by integrating different data sources (and potentially with additional predictive steps), the metaweb allows to overcome sampling limitations and to raise network data to a global scale. For example, Albouy et al. (2019) coupled data on fish distributions with a statistical model of trophic 45 interactions to provide estimates of the potential food web structure at the global scale. Recent studies have focused on assembling metawebs for various taxa through literature surveys and expert elicitation (European terrestrial tetrapods, Maiorano et al. 2020) or using predictive tools (marine fishes, Albouy et al. 2019; Canadian mammals, Strydom et al. 2022). At a finer spatial scale, the local food webs (i.e. the local "realization" of the metaweb when combined with species distributions, Poisot et al. 2012) reflect local environmental conditions but still retain the signal of the metaweb to which they belong 51 (Saravia et al. 2022). Given this, Strydom et al. (2023) defended that predicting the metaweb's structure should be the core goal of predictive network ecology, as if. Assuming there is a strong link between the metaweb and its local realizations, more accurante accurate predictions of the metaweb will have the potential to bring us closer to producing accurate local (downscaled) predictions (Strydom et al. 2023). Therefore, establishing or predicting the metaweb should be the first target in systems lacking information about local realizations. This is not the same as using interactions to improve predictions 57 of species distributions, as recent studies have done (Moens et al. 2022; Poggiato et al. 2022; Lucas et al. 2023), although these are incredibly relevant and answer long-standing calls to include interactions within such models (Wisz et al. 2013). Instead, predicting networks in space is a different task, and it serves a different goal: focusing first on the distribution of network structures and its drivers rather than 61 on the distribution of species.

Explicit spatial predictions (such as downscaled metaweb predictions) are essential as they will allow comparisons with extant work for species-rich communities. Recent approaches to metaweb downscaling combined a regional metaweb with species distribution maps to generate local assemblages for European tetrapods (Braga *et al.* 2019; O'Connor *et al.* 2020; Galiana *et al.* 2021; Gaüzère *et al.* 2022), Barents

Sea marine taxa (Kortsch et al. 2019), and North Sea demersal fishes and benthic epifauna (Frelat et al. 2022). These downscaled assemblages allowed studying network structures in novel ways, for instance, assessing changes in food web structure across space (Braga et al. 2019), describing the scaling of network 69 area relationships (Galiana et al. 2021). Other examples have shown that the metaweb can be used to 70 investigate large-scale variation in food web structure, indicating high geographical connections and heterogeneous robustness against species extinctions (Albouy et al. 2019), which are only apparent when the local and global networks are both available. Further comparisons between network structure and other community properties are relevant as they may highlight new and surprising elements regarding network biogeography. For instance, Frelat et al. (2022) found a strong spatial coupling between community composition and food web structure, but a temporal mismatch depending on the spatial scale. Poisot et al. (2017) found that interaction uniqueness captures more composition variability than community uniqueness, and that sites with exceptional compositions might differ for networks and communities, because species distributions and species interactions had different bioclimatic drivers. Spatialized network data will allow these comparisons, identifying important conservation targets for networks and whether they differ geographically from areas currently prioritized for biodiversity 81 conservation. A key challenge remains in how to downscale a regional metaweb towards local network predictions 83 that reflect the spatial variability of interactions. Even when the metaweb is known, local networks may vary substantially and differ both amongst themselves and from the metaweb (McLeod et al. 2021), emphasizing the need for methods to generate local, downscaled network predictions. A potential limitation to previous downscaling approaches is that they assume interactions are constant across 87 space, which ignores well-documented interaction variability, and masks the effect of environmental conditions on interaction realization (Braga et al. 2019). In contrast, recent studies argued that seeing interactions as probabilistic (rather than binary) events allows us to account for their variability in space

(Poisot et al. 2016) and that this should also be reflected at the metaweb level (Strydom et al. 2023). Gravel et al. (2019) introduced a probabilistic framework describing how the metaweb can generate local realizations and showed how it could be used for modelling interaction distributions. This approach to downscaling is relevant when combined with in situ observations of interactions and local networks to train interaction models (in this case, with willow-galler-parasitoid networks). However, such data is rarely available across broad spatial extents (Hortal et al. 2015; Poisot et al. 2021; Windsor et al. 2023). Spatially replicated interaction data required for such models are especially challenging to document 97 with large food web systems such as European tetrapod and Canadian mammal metawebs (Maiorano et al. 2020; Strydom et al. 2022), where hundreds of species result in tens of thousands of species pairs that may potentially interact. We currently lack a downscaling framework that is both probabilistic and 100 can be trained without replicated in situ interaction data. Additionally, a probabilistic view can allow 101 propagating uncertainty, which can play a key role in evaluating the quality of the predictions. Assessing 102 model uncertainty would enable us to determine to which degree we should trust our predictions and to 103 identify what to do to improve the current knowledge. 104 Here, we present a workflow to downscale a metaweb in space, and illustrate it by spatially reconstructing 105 local instances of a probabilistic metaweb of Canadian mammals. We do so using a probabilistic approach 106 to both species distributions and interactions in a system without spatially replicated interaction data. 107 We then explore how the spatial structure of the downscaled metaweb varies in space and how the 108 uncertainty of interactions can be made spatially explicit. We further show that the downscaled metaweb can highlight important biodiversity areas and bring novel ecological insight compared to traditional 110 community measures like species richness. 111

112 Methods

Fig. 1 shows a conceptual overview of the methodological workflow leading to the downscaled metaweb.

Its components were grouped as non-spatial and spatial data, localized site steps (divided into singlespecies-level, two-species-level, and network-level steps), and the final downscaled and spatialized
metaweb. Throughout these steps, we highlight the importance of presenting the uncertainty of interactions and their distribution in space. We argue that this requires adopting a probabilistic view and
incorporating variation between scales.

19 Data

120 Metaweb

The main source of interaction data was the metaweb for Canadian mammals from Strydom et al. (2022), which is a-spatial, i.e., it represents interactions between mammals that can occur anywhere 122 in Canada. It contains 163 species and 3280 links with a probability of interaction. The species list for 123 the Canadian metaweb was extracted from the International Union for the Conservation of Nature 124 (IUCN) checklist (Strydom et al. 2022). Briefly, the metaweb was developed using graph embedding and phylogenetic transfer learning based on the metaweb of European terrestrial mammals, which is itself 126 based on a comprehensive survey of interactions reported in the scientific literature (Maiorano et al. 127 2020). The Canadian metaweb showed a 91% success rate for known interactions between Canadian 128 mammals recorded in global databases (Strydom et al. 2022). This metaweb is probabilistic, which has 129 the advantage of reflecting the likelihood of an interaction taking place given the phylogenetic and trait 130 match between two species. This allows incorporating interaction variability between species (i.e., taking 131 into account that two species may not always interact whenever or wherever they occur); however, we highlight that other factors beyond trait and phylogenetic matching (e.g., population densities) will also

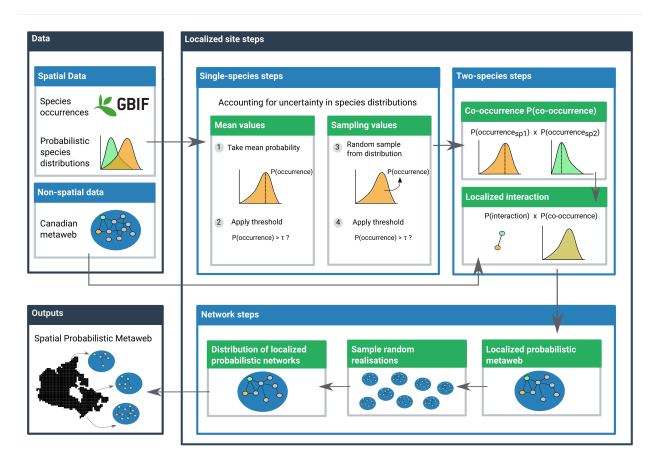


Figure 1: Conceptual figure of the proposed workflow used to downscale the probabilistic metaweb in space. The workflow has three components: the data, the localized steps, and the final spatial output. The data are composed of spatial data (with information in every cell) and non-spatial data (constant for all of Canada). The localized steps use these data and are performed separately in every cell, first at a single-species level (using distribution data), then for every species pair (adding interaction data from the metaweb), and finally at the network level by combining the results of all species pairs. The final output of the network-level steps contains a downscaled probabilistic metaweb for every cell across the study extent.

contribute to observed interaction frequencies.

135 Species occurrences

The downscaling of the metaweb involved combining it with species occurrence and environmental 136 data. First, we extracted species occurrences from the Global Biodiversity Information Facility (GBIF; 137 www.gbif.org) for the Canadian mammals after reconciling species names between the Canadian metaweb and GBIF using the GBIF Backbone Taxonomy (GBIF Secretariat 2021). This step removed 139 potential duplicates by combining species listed in the Canadian metaweb which were considered as 140 a single entity by GBIF. We collected occurrences for the updated species list (159 species) using the 141 GBIF download API on October 21st 2022 (GBIF.org 2022). We restricted our query to occurrences with coordinates between longitudes 175°W to 45°W and latitudes 10°N to 90°N. This was meant to 143 collect training data covering a broader range than our prediction target (Canada only) and include 144 observations in similar environments. Then, since GBIF observations represent presence-only data and 145 most predictive models require absence data, we generated the same number of pseudo-absence data 146 using the surface range envelope method, which selects random non-observed sites within the spatial 147 rangedelimited by the presence data (as occurrences for every species (Barbet-Massin et al. 2012). We 148 weighted candidate sites by their distance to a known observation (separately for each species) using the DistanceToEvent method from the Julia package SpeciesDistributionToolkit (https://github 150 .com/PoisotLab/SpeciesDistributionToolkit.jl), making it more likely to select sites further away from 151 an observation and the known species range. This is because our intent was to model the potential 152 distribution of species, capturing wider responses to the environment, as the downscaled metaweb we 153 aimed to produce is potential in nature (see Downscaled metaweb section below).

155 Environmental data

We used species distribution models (SDMs, Guisan & Thuiller 2005) to project Canadian mammal habitat suitability across the country, which we treated as information on potential distribution. For 157 each species, we related occurrences and pseudo-absences with 19 bioclimatic variables from CHELSA 158 (Karger et al. 2017) and 12 consensus land-cover variables from EarthEnv (Tuanmu & Jetz 2014). The 159 CHELSA bioclimatic variables (bio1-bio19) represent various measures of temperature and precipitation (e.g., annual averages, monthly maximum or minimum, seasonality) and are available for land areas 161 across the globe. We used the most recent version, the CHELSA v2.1 dataset (Karger et al. 2021), 162 and subsetted it to land surfaces only using the CHELSA v1.2 (Karger et al. 2018), which does not 163 cover open water. The EarthEnv land-cover variables represent classes such as Evergreen broadleaf trees, Cultivated and managed vegetation, Urban/Built-up, and Open Water. Values range between 0 165 and 100 and represent the consensus prevalence of each class in percentage within a pixel (hereafter 166 called sites). We coarsened both the CHELSA and EarthEnv data from their original 30 arc-second 167 resolution to a 2.5 arc-minute one (around 4.5 km at the Equator) using GDAL (GDAL/OGR contributors 168 2021). This resolution compromised capturing both local variations and broad-scale patterns while 169 limiting computation costs to a manageable level as memory requirements rapidly increase with spatial 170 resolution. 171

172 Analyses

173 Species distribution models

Our selection criteria for choosing an SDM algorithm was to have a method that generated probabilistic results (similar to Gravel *et al.* 2019), including both a probability of occurrence for a species in a specific site and the uncertainty associated with the prediction. These were crucial to obtaining a probabilistic version of the metaweb as they were used to create spatial variations in the localized

interaction probabilities (see next section). One suitable method for this is Gradient Boosted Trees with a Gaussian maximum likelihood from the EvoTrees.jl Julia package (https://github.com/E 179 vovest/EvoTrees.jl). This method returns a prediction for every site with an average value and a 180 standard deviation, which we used considered as a measure of uncertainty. We used the average value 181 and standard deviation to build a Normal distribution for the probability of occurrence of a given species 182 at all sites each site (represented as probability distributions on Fig. 1). We trained models across the 183 extent chosen for occurrences (longitudes 175°W to 45°W and latitudes 10°N to 90°N), then predicted 184 species distributions only for Canada. We used the 2021 Census Boundary Files from Statistics Canada 185 (Statistics Canada 2022) to set the boundaries for our predictions, which gave us 970,698 sites in total. 186 Performance evaluation for the single species SDMs are available on GitHub.

188 Building site-level instances of the metaweb

The next part of the workflow was to produce local metawebs for every site (*Localized steps* box on Fig. 1).

This component was divided into single-species, two-species, and network-level steps.

The single-species steps represented four possible ways to account for uncertainty in the species distributions and bring variation to the spatial metaweb. We explored four different options to select a 192 value (*P(occurrence)*; Fig. 1) from the occurrence distributions obtained in the previous steps: 1) taking 193 the mean from the distribution as the probability of occurrence (option 1 in Fig. 1); 2) converting the 194 mean value to a binary one using a specific threshold per species (option 2); 3) sampling a random value within the Normal distribution (option 3); or 4) converting a random value into a binary result (option 4, 196 using a separate draw from option 3 and the same threshold as in option 2). The threshold (τ in Fig. 1) 197 used was the value that maximized Youden's J informedness statistic (Youden 1950), the same metric 198 used by Strydom et al. (2022) at an intermediate step while building the metaweb. The four sampling options were intended to explore how uncertainty and variation in the species distributions can affect 200

the metaweb result. We expected thresholding to have a more pronounced effect on network structure
as it should reduce the number of links by removing many of the rare interactions (Poisot *et al.* 2016).
On the other hand, we expected random sampling to create higher spatial heterogeneity compared to
the mean probabilities, as including some extreme values should confound the potential effects of main
trends promoted by environmental gradients. We chose option 1 to present our results as it is intuitive
and essentially represents the result of a probabilistic SDM (as in Gravel *et al.* 2019), but results obtained
with other sampling strategies are available in Supplementary Material, Fig. S1.

Next, the two-species steps were aimed at assigning a probability of observing an interaction between two species in a given site. For each species pair, we multiplied the product of the two species' occurrence probabilities (*P(co-occurrence)*; Fig. 1) (obtained using one of the sampling options above) by their interaction probability in the Canadian metaweb. For cases where species in the Canadian metaweb were considered as the same species by the GBIF Backbone Taxonomy (the reconciliation step mentioned earlier), we used the highest interaction probabilities involving the duplicated species.

The network-level steps then created the probabilistic metaweb for the site. We assembled all the local interaction probabilities (from the two-species steps) into a probabilistic network (Poisot *et al.* 2016). We then sampled several random network realizations to represent the potential local realization process (Poisot *et al.* 2015). This resulted in a distribution of localized networks, which we averaged over the number of simulations to obtain a single probabilistic network for the site.

Downscaled metaweb

The final output of our workflow was the downscaled metaweb, which contains a localized probabilistic metaweb in every site across the study area (*Outputs* box in Fig. 1). The metaweb sets an upper bound on the potential interactions (Strydom *et al.* 2023), therefore, the downscaled metaweb is a refined upper boundary at the local scale that takes into account co-occurrences. It is still potential in nature and

differs from a local realization, from which it should have a different structure. Nonetheless, from the downscaled metaweb, we can create maps of network properties (e.g. number of links, connectance) 225 measured on the local probabilities of species interactions and occurrences, and compute some traditional 226 community-level measures such as species richness. We chose to compute and display the expected 227 number of links (measured on probabilistic networks following Poisot et al. 2016; see Gravel et al. 228 2019 for a similar example) as its relationship with species richness has been highlighted in a spatial context in recent studies (Galiana et al. 2021, 2022). We also computed the uncertainty associated with 230 the community and network measurements (richness variance and link variance, respectively) and 231 compared their spatial distribution (see Supplementary Material). 232

Analyses of results by ecoregions

Since both species composition and network summary values display a high spatial variation and complex 234 patterns, we simplified the representation of their distribution by grouping sites by ecoregion, as species 235 and interaction composition have been shown to differ between ecoregions across large spatial scales 236 (Martins et al. 2022). To do so, we rasterized the Canadian subset of the global map of ecoregions from 237 (Dinerstein et al. 2017; also used by Martins et al. 2022), which resulted in 44 different ecoregions. 238 For every measure we report (e.g. species richness, number of links), we calculated the median site value for each ecoregion, as a way to avoid bias due to long tails in the distributions. We also measured 240 within-ecoregion variation as the 89% interquantile range of the site values in each ecoregion (threshold 241 chosen to avoid confusion with conventional significance tests; McElreath 2020).

43 Analyses of ecological uniqueness

We compared the compositional uniqueness of the networks and the communities to assess whether
they indicated areas of exceptional composition. We measured uniqueness using the local contributions

to beta diversity (LCBD, Legendre & De Cáceres 2013), which identify sites with exceptional composition by quantifying how much one site contributes to the total variance in the community composition. 247 While many studies used LCBD values to evaluate uniqueness on local scales or few study sites (for 248 example, da Silva & Hernández 2014; Heino & Grönroos 2017), recent studies used the measure on 249 predicted species compositions over broad spatial extents and a large number of sites (Vasconcelos et 250 al. 2018; Dansereau et al. 2022). LCBD values can also be used to measure uniqueness for networks by computing the values over the adjacency matrix, which has been shown to capture more unique sites 252 and uniqueness variability than through species composition (Poisot et al. 2017). Here, we measured and 253 compared the uniqueness of our localized community and network predictions. For species composition, 254 we assembled a site-by-species community matrix (970,698 sites by 159 species) with the probability of 255 occurrence of each species at every site from obtained in the species distribution models. For network 256 composition, we assembled a site-by-interaction matrix with the localized interaction values from the 257 spatial probabilistic metaweb probability of interaction at every site given by the downscaled metaweb 258 (therefore, 970,698 sites by 3,108 interactions with defined probabilities in the metaweb). We applied 259 the Hellinger transformation on both matrices and computed the LCBD values from the total variance 260 in the matrices (Legendre & De Cáceres 2013). High LCBD values indicate a high contribution to the 261 overall variance and a unique species or interaction composition compared to other sites. Since the 262 values themselves are very low given our high number of sites (as in Dansereau et al. 2022), what 263 matters primarily is the magnitude of the difference between the sites. Given this, we divided values 264 by the maximum value in each matrix (species or network) and suggest that these should be viewed as 265 relative contributions compared to the highest observed contribution. As with other measures, we then summarized the local uniqueness values by ecoregion by taking the median LCBD value and measuring 267 the 89% interquantile range. 268

269 Analyses of network motifs

To further explore network structure in space, we investigated the distribution of network motifs across 270 space. Motifs are defined sets of interaction between species (Milo et al. 2002; Stouffer et al. 2007), for 271 instance two predators sharing one prey, which are repeated within larger and more complex food webs. 272 Motifs are linked to community persistence (Stouffer & Bascompte 2010) and community structure 273 (Cirtwill & Stouffer 2015; Simmons et al. 2019), are conserved across scales (Baker et al. 2015; Baiser 274 et al. 2016), and are part of a common backbone of interactions among all food webs (Mora et al. 275 2018). We focused on four of the most studied three-species motifs (Stouffer et al. 2007; Stouffer 276 & Bascompte 2010; Baiser et al. 2016): S1 (tri-trophic food chains), S2 (omnivory), S4 (exploitative 277 competition) and S5 (apparent competition). These motifs can be grouped into two pairs according to their ecological information: S1 and S2 highlight different trophic structures, while S4 and S5 indicate 279 different competition types. Therefore, we compared the spatial distribution of the motifs in each pair 280 to see which ones were dominant across all our sites. First, we computed the expected motif count 281 for each of the four motifs for all sites using the localized probabilistic networks from the downscaled 282 metaweb (following Poisot et al. 2016). Then, we compared the expected counts of the motifs within 283 the two pairs. To do so, we used a normalized difference measure similar to the Normalized Difference 284 Vegetation Index (NDVI), where we compute the difference between the two motif counts over their 285 sum. We called the index comparing the two trophic motifs (S1 and S2) the Normalized Difference 286 Trophic Index (NDTI) and the one comparing the competition motifs (S4 and S5) the Normalized 287 Difference Competition Index. We defined both indexes as: 288

$$NDTI = \frac{(S1 - S2)}{(S1 + S2)}$$

289

$$NDCI = \frac{(S4 - S5)}{(S4 + S5)}$$

Values for both indexes are bounded between -1 and 1. A value of 0 indicates that both motifs have

the same expected counts. Positive values indicate that the first motif in each index (S1 and S4) is 291 dominant and has a higher expected count, while negative values indicate that the second motif (S2 and 292 S5) is dominant. As with previous measures, we then summarized both index values by ecoregion by 293 taking the ecoregion median and measuring its within-ecoregion variation with the 89% interquantile 294 range. Ecoregion values therefore indicate if one type of trophic structure (for NDTI) and one type of 295 competition (for NDCI) is dominant in the ecoregion, while the interquantile range values measure 296 whether the dominant type varies within the ecoregion. We used Julia v1.9.0 (Bezanson et al. 2017) to implement all our analyses. We used packages 298 GBIF. j1 (Dansereau & Poisot 2021) to reconcile species names using the GBIF Backbone Taxonomy, SpeciesDistributionToolkit.jl (https://github.com/PoisotLab/SpeciesDistributionToolkit.jl) 300 to handle raster layers, species occurrences and generate pseudoabsences pseudo-absences (using the 301 DistanceToEvent method), EvoTrees.jl (https://github.com/Evovest/EvoTrees.jl) to perform the 302 Gradient Boosted Trees, EcologicalNetworks.jl (Poisot et al. 2019) to analyze network and metaweb 303 structure, and Makie.jl (Danisch & Krumbiegel 2021) to produce figures. Our data sources (CHELSA, 304 EarthEnv, Ecoregions) were all unprojected, and we did not use a projection in our analyses. However, 305 we displayed the results using a Lambert conformal conic projection more appropriate for Canada 306 using GeoMakie.jl (https://github.com/MakieOrg/GeoMakie.jl). All the code used to implement our 307 analyses is archived on Zenodo (https://doi.org/10.5281/zenodo.8350065; Dansereau & Poisot 2023) 308 and includes instructions on how to run a smaller example at a coarser resolution. Note that running 309 our analyses at full scale is resource and memory-intensive and required the use of computer clusters provided by Calcul Québec and the Digital Research Alliance of Canada. Full-scale computations 311

(excluding motifs) required 900 CPU core-hours and peaked at 500 GB of RAM. Computations for network motifs were even more intensive and required 12 CPU core-years.

Results

Our workflow allowed us to display the spatial distribution of ecoregion-level community measures (here, expected species richness) and network measures (expected number of links; Fig. 2). We highlight that the 316 community and network-level measures presented here are not actual predictions first computed over 317 the predicted communities and networks obtained when downscaling the metaweb, then summarized 318 across the ecoregions (taking the median within each ecoregion). They are not a direct prediction of 319 the measure itself (e.g., we do not present a prediction of actual species richness at each location). 320 Instead, they are the reflection of these metrics from the localized predictions of the communities and 321 networks obtained from the downscaling of the metaweb, then summarized for the ecoregions (using the median value). Expected ecoregion richness (Fig. 2A) and expected number of links (Fig. 2B) 323 displayed similar distributions with a latitudinal gradient and higher values in the south. However, 324 within-ecoregion Within-ecoregion variability was distributed differently, as some ecoregions along the 325 coast displayed higher interquantile ranges, while ecoregions around slightly differently with a less constant latitudinal gradient, notably lower interquantile ranges near the southern border displayed 327 narrower ones ((for example, near Vancouver Island and the Rockies on the West Coast, and near 328 the Ontario Peninsula, the Saint-Lawrence Valley, and Central New-Brunswick in the East; Fig. 2C-329 D). Bivariate comparison of the distributions of species richness and expected number of links and of 330 their respective within-ecoregion variability further shows some areas of mismatches, indicating that 331 richness and links do not co-vary completely although they may show similar distributions for median 332 values (see Supplementary Material, Fig. S1). All results shown are based on the first sampling strategy 333 (option 1) mentioned in the Building site-level instances of the metaweb section, where we used the mean

value of the species distributions as the species occurrence probabilities (results for other sampling strategies are shown in Supplementary Material, Fig. S1S2). Site-level results (before summarizing by 336 ecoregion) are also provided in Supplementary Material (Figs. \$2-\$5). 337 Direct comparison of the spatial distributions of species richness and expected number of links showed some areas with mismatches, both regarding the median estimates and regarding the within-ecoregion 339 variability (Fig. ??). Median values for the ecoregions showed a similar bivariate distribution, with 340 ecoregions in the south mostly displaying high species richness and a high number of links (Fig. ??A). 341 The northernmost ecoregions (Canadian High Artic Tundra and Davis Highlands Tundra) displayed higher richness (based on the quantile rank) compared to the number of links. Inversely, ecoregions 343 further south (Canadian Low Artic Tundra, Northern Canadian Shield Taiga, Southern Hudson 344 Bay Taiga) ranked higher for the number of links than for species richness. On the other hand, 345 within-ecoregion variability showed different bivariate relationships and a less constant latitudinal 346 gradient (Fig. ??B). This indicates that richness and links do not co-vary completely (i.e. their variability 347 is not highly correlated) although they may show similar distributions for median values S3-S6). 348 Our results also indicate a mismatch between the uniqueness of communities and networks (Fig. 3). 349 Uniqueness was higher mostly in the north and along the south border for communities, but only mainly 350 in the north for networks (Fig. 3A-B). Consequently, ecoregions with both unique community composi-351 tion and unique network composition were mostly in the north (Fig. 3C). Meanwhile, some areas were 352 unique for one element but not the other. For instance, the New England-Acadian forests ecoregion 353 (south-east, near 70°W and 48°N) had a highly ecoregions along the south border had a unique species 354 composition but a more common network composition (Fig. 3C). Opposite areas with Two ecoregions 355 showed the opposite (unique network compositions onlywere observed at higher between latitudes 52°N 356 and) at higher latitudes (Davis Highlands tundra, near 70°N(Eastern Canadian Shield Taiga, Northern 357 Canadian Shield Taiga, Canadian Low Artic Tundra). Also, and Southern Hudson Bay taiga, near 54°N). 358

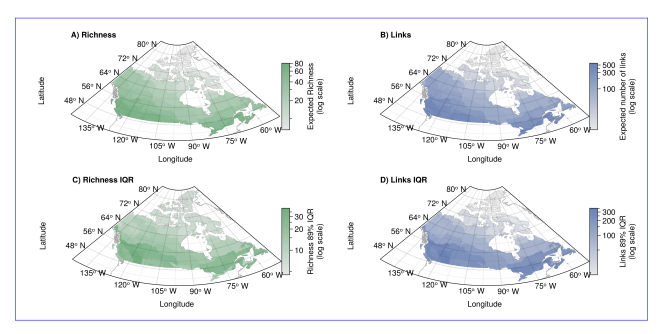


Figure 2: (A-B) Example of a community measure (A, expected species richness) and a network one (B, expected number of links). Both measures are assembled from the predicted probabilistic communities and networks, respectively. Values are first measured separately for all sites; then, the median value within each ecoregion was taken to represent the ecoregion-level value. (C-B) Representation of the 89% interquantile range of values within the ecoregion for expected richness (C) and expected number of links (D). All colour bars follow a log scale with tick marks representing even intervals. Real values (non-log transformed) are shown beside major tick marks while minor ticks represent half increments.

Moreover, network uniqueness values for ecoregions spanned a narrower range between the 44 ecoregions than species LCBD values (Fig. 3D, left). Within-ecoregion variation was also lower for network
values with generally lower 89% interquantile ranges among the site-level LCBD values (Fig. 3D, right).

Moreover, mismatched sites (unique for only one element)formed two distinct groups when evaluating
the relationship between species richness and the number of links (see Supplementary Material, Fig.S5).

The areas only unique for their species composition had both a high richness and number of links

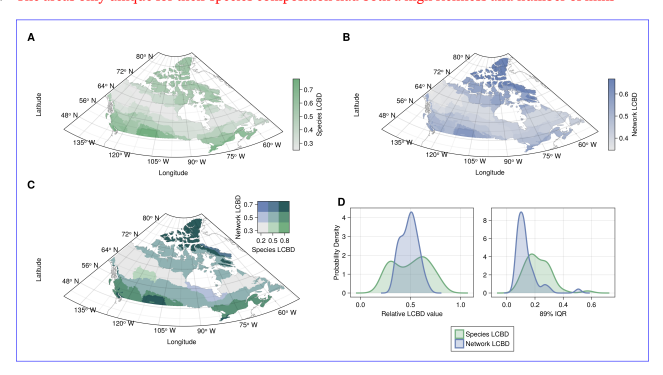


Figure 3: (A-B) Representation of the ecoregion uniqueness values based on species composition (A) and network composition (B). LCBD values were first computed across all sites and scaled relative to the maximum value observed. The ecoregion LCBD value is the median value for the sites in the ecoregion. (C) Bivariate representation of species and network composition LCBD. Values are grouped into three quantiles separately for each variable. The colour combinations represent the nine possible combinations of quantiles. The species uniqueness (horizontal axis) goes left to right from low uniqueness (light grey, bottom left) to high uniqueness (green, bottom right). The network composition uniqueness goes bottom-up from low uniqueness (light grey, bottom left) to high uniqueness (blue, top left). (D) Probability densities for the ecoregion LCBD values for species and network LCBD (left), highlighting the variability of LCBD values between ecoregions, and the 89% interquartile range of the values within each ecoregion (right), highlighting the variability within the ecoregions.

Comparing the distribution of dominant network motifs revealed additional areas of variation in network structure (Fig. 4). NDTI displayed a latitudinal gradient between the trophic motifs.

Northern ecoregions showed positive NDTI values and high dominance of S1 (tri-trophic chains) 367 expected counts compared to S2 (omnivory) but ecoregions along the south border showed a reduced 368 dominance (Fig. 4A). Ecoregions near the Ontario Peninsula and Saint-Lawrence Valley showed values 369 close to zero, indicating a balance between two motifs, while Central New Brunswick had slightly 370 negative values, indicating a low dominance of S2. In comparison, NDCI values showed an evenly 371 high dominance of S5 (apparent competition) over S4 (exploitative competition) across all ecoregions (Fig. 4B). Meanwhile, within-ecoregion variance displayed a different spatial distribution from the 373 median values. NDTI interquantile ranges spanned a wide range of values and were higher both in the 374 north and in the south (although not in the ecoregions with higher NDTI median values) (Fig. 4C). On 375 the other hand, the sites only unique for their networks had both lower richness and a lower number of links, although they were not the sites with the lowest values for both NDCI interquantile ranges 377 showed lower within-ecoregion variance in most ecoregions except in the northernmost one (Canadian 378 High Arctic tundra), which has a notably higher value (Fig. 4D). Although this higher variance does 379 not reflect in the NDCI median values, it does appear when looking at the site-level values, where this 380 ecoregion is the only one with patches with high positive NDCI values (indicating a dominance of S4) 381 surrounded by highly negative values (indicating a dominance of S5) as in other ecoregions (Fig. S6B). 382

83 Discussion

Our approach presents a way to downscale a metaweb, produce localized predictions using probabilistic networks as inputs and outputs, and incorporate uncertainty, as called for by Strydom *et al.* (2023). It gives us an idea of what local metawebs or networks could look like in space, given species distributions and their variability, as well as the uncertainty around species interactions. We also provide the first spatial representation of the metaweb of Canadian mammals (Strydom *et al.* 2022) and a probabilistic equivalent to how the European tetrapod metaweb (Maiorano *et al.* 2020) was used to predict localized

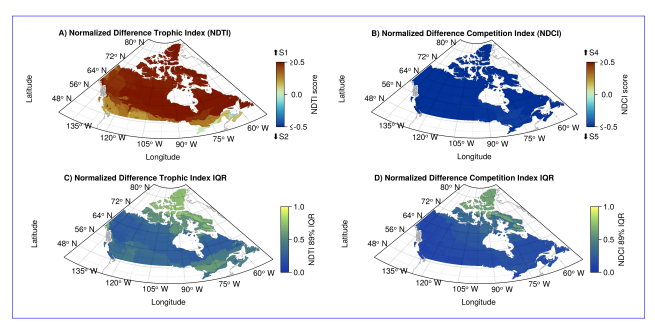


Figure 4: Comparison of the dominant ecological motifs across ecoregions. A) Normalized Difference Index (NDTI) comparing the trophic motifs S1 (tri-trophic food chains) and S2 (omnivory). Positive values indicate a dominance of S1, while negative values indicate a dominance of S2. Values equal or superior to |0.5| are shown with the same colour as they indicate a high dominance of one motif. B) Normalized Difference Index (NDCI) comparing the competition motifs S4 (exploitative competition) and S5 (apparent competition). Positive values indicate a dominance of S4, while negative values indicate a dominance of S5. (C-D) Representation of the 89% interquantile range of values within the ecoregion for the trophic motifs index (NDTI, C) and competition motifs index (NDCI, D).

networks in Europe (Braga et al. 2019; O'Connor et al. 2020; Galiana et al. 2021; Gaüzère et al. 2022; Botella et al. 2023). Therefore, our approach could open similar possibilities of investigations on the 391 variation of structure in space (Braga et al. 2019) and on the effect of land-use intensification (Botella et 392 al. 2023) on North American food webs, particularly Canadian mammal food webs. Other interesting 393 research applications include assessing climate change impacts on network structure (e.g., Kortsch et al. 394 2015) or investigating linkages between network structure and stability (Windsor et al. 2023). As our approach is probabilistic, it does not assume species interact whenever they co-occur and incorpo-396 rates variability based on environmental conditions (via projected species distributions), which could lead 397 to different results by introducing a different association between species richness and network properties. 398 Galiana et al. (2021) found that species richness had a large explanatory power over network properties, 390 but mentioned this could potentially be due to interactions between species being constant across space. 400 Here, we found mismatches in the distribution of that potential species richness and interactions that 401 were especially apparent in their number of links displayed similar distributions following a latitudinal 402 gradient, but that the within-ecoregion variability variance was lower along the southern border than 403 the measures themselves (Fig. ??), highlighting that interactions might vary differently than species 404 distributions even over continental-scale gradients. Network-2). Why the relationship between the 405 median ecoregion values and the ecoregion variance is not constant across space could be verified 406 through more tests in future studies (for instance, testing the effect of higher urban density in the south). 407 Examples of metaweb structures in space are rare and not ideal for comparison with our results. For 408 instance, we found that network density (links on Fig. ??A) were also 2B) was lower in the north, 409 contrarily which is contrary to what was observed for all European terrestrial tetrapods in Europe for 410 the terrestrial tetrapod metaweb (Braga et al. 2019; Galiana et al. 2021) and for willow-galler-parasitoid 411 networks (Gravel et al. 2019), where connectance was higher in northern regions. However, those 412 are systems with different numbers of species and environmental conditions (e.g. Europe and Canada 413

could differ due to varying climatic conditions at the same latitudes). Further research should investi-414 gate why these results might differ between the two continents and continents and ecological systems 415 and whether it is due to the methodology, data, or biogeographical processes. 416 Our LCBD and uniqueness results highlighted that areas with unique network composition differ from sites with unique species composition. In other words, the joint distribution of community and 418 network uniqueness highlights different diversity hotspots. Poisot et al. (2017) showed a similar result 419 with host-parasite communities of rodents and ectoparasitic fleas. Our results further show how that 420 these differences could be distributed across ecoregions and in over a broad spatial extent. Areas unique for only one element (species or network composition) differed in their combination of species 422 richness and number of links (Supplementary Material, Fig. S5), with species-unique sites displaying 423 high values of both measures, and network-unique sites displaying low values. Moreover, for mammal food webs. LCBD scores essentially highlight variability hotspots and are a measure of the variance of 425 community or network structure. Here, they also serve as an inter-ecoregion variation measure, which 426 can be compared to the within-ecoregion variation highlighted by the interquantile ranges. The narrower 427 range of values for network LCBD values and the lower IQR values indicate that both the inter-ecoregion 428 and within-ecoregion variation are lower for networks than for species (Fig. 3). Additionally, higher 429 values for network LCBD also indicate that most ecoregions can hold ecologically unique sites 430 Our analysis of the distribution of dominant network motifs revealed additional areas of variation in 431 network structure. Trophic motifs (S1 and S2, measured through NDTI values) showed a latitudinal 432 gradient different from the richness and links ones, with high dominance of tri-trophic chains (S1) in the 433 north and higher omnivory counts (S2) only in a few ecoregions in the south. These results did not seem 434 related to ecoregion variance, which once again showed a very different distribution from the median 435 values. Meanwhile, competition motifs (S4 and S5, measured through NDCI values) showed an even 436 dominance of apparent competition (S5) but high variance in a single ecoregion. Overall, our results

show that dominant motifs within a mammal food web vary between ecoregions and vary differently across space.

When to use the workflow we presented here will depend on the availability of interaction data or 440 existing metawebs, and on the intent to incorporate interaction variability, as well as ecoregion-level variability. In systems where in situ interaction and complete network data are available, the approach 442 put forward by Gravel et al. (2019) achieves a similar purpose as we attempted here, but is more rigourous 443 rigorous and allows modelling the effect of the environment on the interactions themselves. Without such data, establishing or predicting the metaweb (e.g. using transfer learning) should be the first step toward producing localized predictions ().-Strydom et al. 2023). Our framework then downscales the 446 metaweb towards the localized predictions, here using the probabilistic Canadian mammal one, but it 447 can also use other metawebs generated through various means. Well-documented binary metawebs 448 ones such as the European tetrapod metaweb could be partly combined with our approach if used with 449 probabilistic SDMs and summarized by ecoregions (as they would only lack an initial probabilistic 450 metaweb, but would still obtain a more probabilistic output). Our approach will essentially differ from 451 previous attempts in how it perceives uncertainty and variability. For instance, rare interactions should not be over-represented (Poisot et al. 2016) and should have lesser effects over computed network 453 measures. Furthermore, summarizing results by ecoregion allows for showing variation within and 454 between ecologically meaningful biogeographic boundaries (Martins et al. 2022), which, as our results 455 showed, is not constant across space and can help identify contrasting diversity hotspots.

Adding validation to our framework will be highly important for the predictions to be actionable. We

see it as the next step since data availability currently remains an issue. Developing a way to generate

actionable information when information is initially scarce, as we present here, is highly important in

itself. Our manuscript highlights the application of a predictive pipeline in a case where interactions

are almost entirely predicted, but it can equally be applied when there are additional data about the

interaction network. Moreover, the predictions we make are already actionable, as any future sampling
of food web structure can be used to validate them, and fed into the model to iterate these results again.

As Strydom *et al.* (2023) point out, validation of metaweb predictions, empirical sampling, and method
design should all proceed jointly, and making conceptual progress in one of these areas helps all the
others.

The recent shift in focus towards building metawebs opens many opportunities for projections of networks
in space through probabilistic downscaling, as we suggested here. Metawebs have been documented in
many systems, allowing us to build new ones from predictions. How the European tetrapod metaweb

many systems, allowing us to build new ones from predictions. How the European tetrapod metaweb 469 (Maiorano et al. 2020) was used to predict the Canadian mammal metaweb (Strydom et al. 2022) is one 470 such case, but recent examples also extend to other systems. Metawebs have been compiled for many 471 marine food webs (e.g., Barents Sea, Kortsch et al. 2019; North Scotia Sea, López-López et al. 2022; Gulf of Riga, Kortsch et al. 2021) and used to predict the probability of novel interactions (Artic food web of the Barents sea, Pecuchet et al. 2020). Olivier et al. (2019) built a temporally resolved metaweb of demersal 474 fish and benthic epifauna but also suggested combining their approach with techniques estimating 475 the probability of occurrence of trophic relationships to describe spatial and temporal variability more accurately. Lurgi et al. (2020) built a metaweb and probabilistic (occurrence-based) networks for rocky 477 intertidal communities (and in doing so, they also showed that environmental factors do not affect the 478 structure of binary and probabilistic networks in different ways). Albouy et al. (2019) predicted the global marine fish food web using a probabilistic model, showing the potential to describe networks across broad spatial scales. Similarly, predictive approaches are also increasingly used with other interaction 481 types to highlight interactions interaction hotspots on global scales (e.g. -mapping geographical hotspots 482 of predicted host-virus interactions between bats and betacoronaviruses, Becker et al. 2022; predicting 483 the distribution of hidden interactions in the mammalian virome, Poisot et al. 2023). Our workflow offers the potential to bring these global predictions down to the local scale where they can be made 485

more actionable, and vastly increases the diversity of ecological networks that can be projected in space.

487 Acknowledgements

We acknowledge that this study was conducted on land within the traditional unceded territory of the
Saint Lawrence Iroquoian, Anishinabewaki, Mohawk, Huron-Wendat, and Omàmiwininiwak nations.
GD is funded by the NSERC Postgraduate Scholarship – Doctoral (grant ES D – 558643), the FRQNT
doctoral scholarship (grant no. 301750), and the NSERC CREATE BIOS² program. TP is funded by
the Wellcome Trust (223764/Z/21/Z), NSERC through the Discovery Grant and Discovery Accelerator
Supplements programs, and the Courtois Foundation. This research was enabled in part by support
provided by Calcul Québec (calculquebec.ca) and the Digital Research Alliance of Canada (alliancecan
.ca) through the Narval general purpose cluster.

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