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32 Abstract

Current global changes make it important to be able to predict which interactions will 33 occur in emerging ecosystems. Most of the current methods to infer the existence 34 of interactions between two species require a good knowledge of their behaviour or 35 a direct observation of interactions. In this paper, we overcome these limitations by 36 developing a method, inspired from the niche model of food web structure, using 37 the statistical relationship between predator and prey body size to infer the matrix 38 of potential interactions among a pool of species. The novelty of our approach is to 39 infer, for any species of a given species pool, the three species specific parameters of the niche model. The method applies to both local and metaweb scales. It allows 41 one to evaluate the feeding interactions of a new species entering the community. We 42 find that this method gives robust predictions of the structure of food webs, and that its efficiency is increased when the strength of the body- size relationship between predators and preys increases. We finally illustrate the potential of the method to infer the metaweb structure of pelagic fishes of the Meditarrannean sea under different global change scenarios.

1 Introduction

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One of the current challenges in ecology is to predict the emergence and re-assembly 49 of communities following species responses to global changes. Understanding this is 50 made all the more important as these novel ecosystems will be more common with 51 increasing human pressure. We know that species invasions, biomass harvesting, 52 ranges shifts, disturbances and changes in land use are important drivers of biodi-53 versity turnover. How they affect species composition is now well described (Pereira 54 et al. 2010), but forecasting their impact on community structure and functioning 55 requires a priori knowledge of potential interactions among species. Predicting inter-56 actions among species that never co-occurred proves challenging, as traditional em-57 pirical methods of food web sampling such as stomacal content analysis cannot be 58 applied. While for competitive communities this task could be achieved using predictive modeling based on functional traits (Albouy et al. 2010; McGill et al. 2006) or phylogeny (Cavender-Bares et al. 2009; Mouquet et al. 2012), it is much more difficult 61 to forecast trophic interactions in a food web (Ings et al. 2009; Montoya & Raffaelli 2010; Tylianakis et al. 2008). 63

Inferring potential interactions among species of an arbitrary defined pool is a major step to predict the structure of emergent communities and their functioning. We call these potential interactions among a given set of species, wether at the local or regional scales, the *metaweb* (Dunne 2006). A metaweb takes the form of an adjacency matrix \mathbf{MW} , of size $s \times s$ for a pool of s species, and in which $\mathbf{MW_{ij}} = 1$ if species i can consume species j, and 0 otherwise. This matrix aggregates the trophic interactions among all species from the pool that are susceptible to both co-occur and interact at the regional scale (Dunne 2006). With a metaweb in hands, one could analyze the impacts of global changes, such as range shifts or species invasions, on the potential community structure, by extracting the relevant species and examining the proper-

ties of the community they define. While this concept is progressively finding its way
through theoretical spatial food web ecology (Gravel et al. 2011a,b; Lafferty & Dunne
2010; Pillai et al. 2009), it is still limited by data availability and predictive accuracy.
Published metaweb data rely on literature surveys (Baiser et al. 2012; Havens 1992;
Piechnik et al. 2008), or a compilation of several local food webs (Poisot et al. 2012;
Stanko et al. 2002), making them resolved only for species that co-occurred a large enough number of times.

Development of predictive models of trophic interactions could greatly improve 81 our understanding of large scale food web structure and our capacity to anticipate major changes in ecosystem functionning. Theory should provide some guidance about 83 the general rules underpinning interactions among a set of species and thus help us 84 to infer the metaweb. The actual food web theory is largely derived from the niche 85 model (Williams & Martinez 2000). This model simply and intuitively poses that each 86 species in a food web has a niche position n_i , a feeding niche optimum c_i and a range 87 r_i of suitable preys around that optimum (Fig. 1). These simples rules are sufficient 88 to generate realistic food web structures that fit most of published food webs (Dunne 89 2006). The niche model was a substantial improvement of the previous cascade model 90 (Cohen1990) and subsequent models (e.g. the nested hierarchy model (Cattin et al. 91 2004), the minimum potential model (Allesina et al. 2008), the probabilistic niche 92 model (Williams et al. 2010) are somehow derived from these rules (Stouffer et al. 93 2005). Although it is based on different assumptions, the adaptive foraging theory of 94 food web structure (Petchey et al. 2008) provides comparable predictions (Williams et 95 al. 2010). One major recent breakthrough in food web theory have been the attempts 96 to parameterize the niche model and other food web models from field data and to compare their fit through likelihood methods (Allesina et al. 2008; Williams 2011; 98 Williams et al. 2010). These methods provide, for each species of the food web, the

optimal parameters to fit the empirical web structures, given the hypothesized underlying rules of the niche model.

Despite their theoretical interest, these methods however come with several draw-backs when comes the time to perform biodiversity scenarios. Firstly, they are difficult to apply at large scale because of the technical and logistical requirements of metaweb data collection. Secondly, once the model is parameterized, it could only be used to infer feeding interactions between species with already documented interactions (i.e. it is impossible to infer potential interactions among species that do never co-occurred). Finally, the model optimization is a serious challenge for large datasets with a large number of parameters to evaluate and rough likelihood surfaces. There is consequently an urgent need for a method that could rapidly and easily provide an estimate of potential interactions in a metaweb based on incomplete data.

In this paper, we present a method designed to infer the potential interactions between all pairs of species of a species pool based on observations of body size of predators and their prey. The method applies to both the local and the metaweb scales. We do so by a parameterization of the niche model, based on the well-documented allometric scaling relationship between predator and prey (Brose et al. 2006; Cohen et al. 2003; Riede et al. 2010). We first develop the method and apply it to food webs from various environments. We find the method accurately predicts the interactions (and lack thereof), and that the accuracy increases with the strength of the predator-prey body size relationship. We then analyze the sensitivity of the method to incomplete data (missing links) and find that it is robust to sampling effort. We finally illustrate the potential of the method to infer the metaweb structure of pelagic fishes of the Meditarrannean sea and the consequences of alteration of body size distribution by global changes or anthropic forcings. The method is best suited for strongly size-structured food webs and will likely not hold for other types of non-body-size structured interac-

tions. We therefore conclude on the future issues to generalize this approach to other types of trait matching models.

2 Model description

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- The method aims to infer the potential interactions among a pool of species from a subset of observations of predator-prey interactions. The method follows the following steps, with details provided below:
- Step 1: Log transformation of the body size data;
- Step 2: Statistical analysis of the predator-prey body size relationship;
- Step 3: Inferrence of the niche model parameters for all species from the species pool;
- 135 **Step 4:** Interpretation of the parameters and computation of the metaweb.
- We also provide an example of R code (R Core Development Team) and data in the Supplementary Material, detailling the step by step procedure and the format of the input data.

2.1 Inferring parameters from the niche model and building the metaweb

The niche model predicts the food web structure from a set of three species-specific parameters (Williams & Martinez 2000): the niche position n_i , the feeding niche optimum c_i (called the centroid), and the feeding range r_i . A species i will predate all species j whose niche position n_j lies within the interval $[c_i - r_i/2, c_i + r_i/2]$ (Fig. 1). We evaluate all of these parameters from the predator-prey body size relationship, enabling us to parameterize the metaweb from *observed* interactions only. The parameterization is robust to the sampling effort, as it is much easier to document interactions

than their absence (Martinez et al. 1999). Our approach is however mostly limited to predatory interactions since the body size relationship between herbivores and primary producers is not as general (Riede et al. 2010), and obviously do not hold for parasitic, mutualistic, or competitive networks.

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Assuming that body size is the main niche axis structuring trophic interactions, 152 the parameter n_i corresponds simply to the log of body size (in mass or length) of 153 species i. Though only the relative position of all species along the body size gradi-154 ent needs to be respected, it is possible to standardize log body size values between 0 155 (minimum size in the regional species pool) and 1 (maximal size). We then consider 156 a linear relationship between the decimal logarithm of body size and the centroid of 157 the niche (the dark line at Fig. 1). This relationship is obtained by fitting the linear 158 model $c = \log_{10}(M_{prey}) = \alpha_0 + \alpha_1 \times \log_{10}(M_{pred})$ to the data, where M_{prey} and M_{pred} are 159 the prey and predator body size respectively. The lower and upper boundaries of the 160 feeding range are easily obtained by fitting the 5% and 95% quantile regressions be-161 tween $\log_{10}(M_{prev})$ and $\log_{10}(M_{pred})$ (the dotted lines at Fig. 1, see the example at Fig. 162 2). We note them as $r_{low,i}$ and $r_{high,j}$ respectively, and the corresponding parameters of 163 the linear quantile regresions are $\beta_{0,low/high}$ and $\beta_{1,low/high}$. In sum, the parameter n_i 164 for any species of the metaweb is given by the standardized value of the log body size 165 M_i , c_i is estimated from the linear regression between predator and prey log body size 166 and r_i from the quantile regressions. Type I regressions were used because there is no 167 equivalence of type II models for quantile regressions. 168

The next step of the methodology is to reconstruct the metaweb. Once these parameters are calculated from a subsample of species from the regional pool, the coefficients of the different linear models are used to infer the niche parameters of each species of the species pool. Again, the niche parameter for any species i, n_i , is given by the log of body size. The centroid of the niche is obtained by the relationship $c_i = \alpha_0 + \alpha_1 \times n_i$,

the lower boundary of the niche is $r_{low,i} = \beta_{0,low} + \alpha_{1,low} \times n_i$ and the upper boundary ary $r_{high,i} = \beta_{0,high} + \alpha_{1,high} \times n_i$. A feeding link from species j to species i occurs if $n_j > r_{low,i} \& n_j < r_{high,i}$. We provide an example at Fig. 2.

77 3 Method accuracy

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178 3.1 Predictive performance

We illustrate the method with the food web datasets of Brose et al. (Brose et al. 2005). 179 The meta-analysis of Brose (Brose et al. 2006) was conducted on this dataset to test 180 the generality of the predator-prey body size relationship across different systems (ter-181 restrial, aquatic and marine). The relationship was found to be very strong across all 182 systems despite exhibiting variability from one to another. Each web has between 26 183 and 380 species and 18 and 1466 feeding links. Several of these webs are repetitions 184 over time at a single location, in which cases we pooled the data for each of the 15 185 different locations to calculate the predator-prey body size relationship. We removed 186 4 datasets that had a non-significant predator-prey relationship and were thus useless 187 with our approach. The links are not systematically sampled, meaning that any ab-188 sence of a link between two species for a given dataset could either be a real absence 189 or due to insufficient sampling or lack of information. While the predator-prey body 190 size relationship is very strong over all datasets (Brose2006), there is quite substan-191 tial variation among them, enabling us to assess the sensitivity of the method to the 192 strength of this relationship. 193

We assessed the performance of our method using the True Skill Statistic (TSS). The TSS is based on the partition of events (the prediction of a trophic interaction) between four components: the component a reports the number of links that are both predicted and observed, b reports predicted links with no corresponding observation,

c reports the number of observed links that are predicted absent, and d reports the number of predicted and observed absences of links. The TSS is then calculated as TSS = (ad-bc)/[(a+c)(b+d)]. The TSS quantifies the proportion of prediction success relative to false predictions and returns values ranging between 1 (perfect predictions) and -1 (inverted forecast) (Allouche et al. 2006).

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We calculated the TSS for each of the 11 different webs and related it to the 203 strength of the predator-prey body size relationship, measured by the R^2 of the lin-204 ear model. We find that the TSS is positive for all webs, ranging from 0.13 to 0.76 205 (Fig. 3A). We find a positive relationship between the R^2 of the linear model and the 206 TSS ($R^2 = 0.50$, p = 0.016). When we decompose the different components of predictions and observations, we find that the fraction of prediction match is high, with an 208 average of $(\overline{a} + \overline{d})/S^2 = 0.58$, Fig. 3B). The fraction of wrong predictions is lower, at 209 $\overline{b+c}/S^2 = 0.40$, and decreases with the R^2 of the predator-prey body size relationship. The parameterized niche model tends to overestimate the number of links in a 211 web (see the example at Fig. 2). This result is significant but not surprising, given 212 that these datasets do not necessarily contain all links, as they were not designed with 213 this purpose, and thus some of the links might have been missed. This interpretation 214 is also reminiscent of previous debates on the difficulty to sample all links in a web 215 (Martinez et al. 1999). It is also well known that the niche model predicts a continuous 216 diet along the niche axis (the webs are said to be interval (Cohen et al. 1990; Stouffer 217 et al. 2006)), while real food webs do not have this characteristic. We thus might over 218 predict link density within the niche of a given species. Previous studies (Allesina et al. 219 2008) and the Application 1 however show this problem is easily circumvented when 220 a second niche axis, e.g. an environmental niche, comes into play. A recent study on 221 dimensionality of networks shown that most webs have between 3 and 6 dimensions 222 (Eklof et al. 2013), meaning that adding a few more niche axes can greatly improve the 223

Sensitivity to sampling effort

We subsequently explored the impact of sampling effort on the accuracy of the model 226 predictions. We assumed a random subsampling of all the interactions occurring in 227 the food web. To do so, we selected a species rich food web (S = 67) from the Brose 228 et al. dataset, with 601 observed feeding links and a good TSS (0.51). We randomly 229 removed from 0 to 90% of the observed links to do the evaluation of the parameters of 230 the linear regressions (i.e. e decrease the quantity of information used to calibrate the 231 model). We then after compared the empirical web (with all species) to the inferred 232 web with these parameters (all species also). The comparison was done again with the 233 TSS. We performed 100 randomizations per number of removed links. This numerical 234 simulation reproduces incomplete sampling in the process of building the food web. 235 We find that the *TSS* remains constant up until 80% of the observed links are removed 236 (Fig. 3). At this level, the *TSS* starts to decline drastically and its variance increases. 237 This result shows that our method of parameterization of the niche model is robust 238 with regard to sampling effort. Note however that a biased sampling with respect to 239 body size (e.g. sampling of the largest species) might be more likely to reduce the fit 240 of the predator-prey body size relationship. The aggregation intro trophic species that 241 do have the same body size and the same diet should not impact the prediction since it 242 will not affect parameter evaluation. It will however reduce the fit of the predator-prey

variation in body size (see the Discussion). 246

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body size relationship and therfore the accuracy of the method if the species do have

different body sizes. The same artifact will be observed if there is strong intraspecific

4 Application: Mediterranean food web structure under fishing pressure

9 4.1 Dataset

We now present an application of the method to infer the metaweb of interactions 250 among fishes of the Mediterranean sea. The Mediterranean is known as a hotspot of 251 fish diversity that is severely threatened by climate change and overfishing (Mouillot 252 et al. 2011). There are 557 fish species in the regional pool, with a maximum body 253 size ranging from 2.3 cm to 1100 cm (Froese & Pauly 2011; Louisy 2005; Whithead et 254 al. 1986). Chondrichthyans, mammals and turtles were under represented in the two 255 trophic networks and were removed from the analyses. We parameterized the niche 256 model with a subset of species, from two different highly resolved food webs from 257 the Catalan area (Coll et al. 2006) (82 species) and Corsica (Albouy et al. 2010) (58 258 species). 259

4.2 Inferring the metaweb for Mediterrannean fishes

We estimated parameters *n*, *c* and *r* for each of the 557 species and inferred the poten-261 tial interactions among all of them (the example in the Supplementariy code is based 262 on this dataset). The metaweb has a total of 126 501 links, for a connectance of 0.41 263 (Fig. 5A). The metaweb is also highly nested (specialist species feed on a subset of 264 prey of the most generalist species. Fig. 5A). We also considered a second niche axis 265 related to species spatial distribution. Most fish species have restricted geographic 266 range within the Mediterranean sea because of specific response to temperature and 267 other environmental variables (Albouy et al. 2012). We therefore removed from the metaweb all links between species having no range overlap. Data on the extent of oc-

currence of fish species were compiled from a published atlas of fishes of the northern Atlantic and the Mediterranean (Whitehead & Unesco 1986). This atlas is based on 271 regional data sets and expert knowledge and was edited between 1984 and 1986. It 272 currently provides the only available basin-wide information on the extent of occur-273 rence of all Mediterranean Sea fish species. The above mentioned atlas do not account 274 for the bathymetric distribution of Mediterranean fish species, yet bathymetry is con-275 sidered as one of the main factors accounting for marine fish distributions (Louisy 276 2005). We therefore refined the extent of occurrence maps by clipping off areas with 277 depths that fall outside the minimum or maximum known for the species. Species' 278 bathymetric ranges were obtained from FishBase (Froese 2010; Louisy 2005). The resulting metaweb has a total of 95 989 links, for a connectance of C = 0.31. Connectance 280 decreases because links are removed by incomptabilities in bathymetry but the total 281 number of species stays constant. This metaweb is clearly less interval (Fig. 5B). Con-282 tiguous gaps in the diet are likely to emerge from modularity in the co-occurrence 283 matrix (Araújo et al. 2011). Future studies should explore how co-occurrence is con-284 straining the topological structure of metawebs. 285

4.3 Impact of global changes

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Our method to parameterize the niche model has a unique feature relative to the original inal niche model (Williams & Martinez 2000): the network properties of the original model, such as the number of links, food chain length and degree distribution (the number of in and out feeding links per species), are controlled by the input parameters (species richness and connectance), wherea all properties of our parameterized niche model are emergent features of the predator-prey body size relationship and the frequency distribution of body size. The connectance in the niche model is fixed by the user; in our case it strongly depends on the scaling of the feeding range with body

size and the frequency of larger body species.

This feature is particularly important to understand global change impacts on com-296 munity structure. For instance, a common prediction of the impact of fisheries on the 297 body size distribution is the reduction in the average and variance of body size (Jack-298 son et al. 2001). Similarly, global warming is also expected to alter fish body size distri-299 bution towards smaller species (Cheung et al. 2012). We explored by simulations the 300 impact of these changes on the degree distribution of the Mediterranean metaweb. Re-301 sults are illustrated at Fig. 6. The shape of the cumulative degree distribution provides 302 a visual assessment of the distribution of diet specificity in a network. The curves be-303 comes steeper and the fall moves to the left with increasing specialization (indicative of a larger proportion of species with a low degree). Even if the simulated scenarios are 305 crude representations of the predicted alterations of community structure, they both 306 show that fisheries impacts on body size will substantially alter the network proper-307 ties. The results at Fig. 6 show that reduction in the average body size will decrease 308 the average number of preys per predator (there is an initial sharper decline of the 309 cumulative distribution), but that super generalist species will also appear (because of 310 the shift of the tail toward the right). The removal of the 40% largest species reduces 311 much more the average generality which is expected as larger species are predators 312 with a large feeding niche in the niche model. The two scenarios consequently in-313 creases substantially the relative abundance of species with a smaller degree, even for 314 the average size reduction scenario where the total number of species is held constant. 315 The change in the degree distribution is likely to reduce substantially the expected 316 persistence and stability of these communities (Gravel et al. 2011a,b). 317

5 Discussion

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In this paper, we presented a method to infer potential interactions among an arbitrary 319 pool of species. The data required to perform this methodology are simple to obtain, 320 as the body size of a large number of species are available from reference databases, 321 or easy to measure. Because we rely on a robust allometric relationship, applying 322 this method requires neither complex statistical techniques, nor an exhaustive knowl-323 edge of the realized interactions within the metaweb of interest. In addition to the 324 opportunity to simulate biodiversity scenarios, our method can also help generating 325 baseline expectations about the food web structure for environments which are no-326 toriously difficult to sample, such as soils, deep-sea environments, or fossil records. 327 However, because this allometric relationship is characteristic of predatory interac-328 tions, our method will likely not hold for other types of non-body-size structured in-329 teractions such as herbivory and parasitism. 330

It appears at Fig. 2 that the feeding range increases with the body size of the preda-331 tor. This result was quite general across datasets with a good fit of the predator-prey 332 body size relationship, although with considerable variation in the slope of the re-333 lationship. It offers the possibility to compare empirical based estimates of the pa-334 rameters to the original niche model of Williams & Martinez (2000). For this specific 335 dataset that the model for the centroid is $c_i = -1.34 + 0.66n_i$ and the average range 336 $(\overline{r}_i = (r_{high,i} - r_{low,i})/2)$ is $\overline{r}_i = 0.63 + 0.78n_i$. The original niche models assumed that r_i 337 scales linearly with connectance C and the niche position, such that $r_i = 2Cn_i$. Even 338 though the parameterization suggests a linear increase of generality with body size, 339 the final number of preys in the diet for a given size will depend on the frequency dis-340 tribution of body size. A large predator for instance could have a large feeding range, 341 but only few species to find on and effectively appears as a specialist. This relation-342 ship will have to be investigated further, among several food webs (e.g. Digel2011), because the scaling of generality with trophic rank has several consequences on persistence (Gravel et al. 2011b) and food web dynamics (**Berlow2009**; Brose et al. 2006).

Ontogenic shifts in diet are common in many size-structured populations and could be accounted for in two ways with this methodology. First, they will enter the model parameterization by simply including a link from species A (the largest) to species B (the smallest), and the opposite. This would yield in a data point for species B figuring above the 1:1 relationship. Such a data point will obviously influence the model calibration and enlarge the regime of all species. It is not obvious however that ontogenic shifts will be found in the metaweb as it would require an upper limit of the range above the 1:1 relationship. The alternative approach would be to distinguish "sub-species" by size categories in the model calibration and then in the reconstruction of the metaweb. This approach would force ontogenic shifts and perhaps more precisely represent the interaction matrix where the within-population size structure is important.

The method should be completed with other sources of informations to better predict gaps in the interaction matrix. The original niche model was definitely inspired by the predator-prey body size relationship, but was intented to be more general and eventually deal with several niche axes (Williams & Martinez 2000). Allesina et al. (2008) indeed found that adding a second axis, creating gaps into the first axis interaction matrix, increases the fit of the model to empirical data. It also makes the network less interval (a continuous succession of species diets along the niche axis), a structural issue of the niche model that was reported by Cattin et al. (2004) and Bersier et al. (2006). There are numerous sources of information that could be used to improve the model, such as co-occurrence and functional traits.

Our application with the Mediterannean pelagic fish food web provides an example of how straightforward it is to add additional information so as to improve the parameterization. A similar approach could also build on the comptability of other traits such as phenology, location in the water column or hunting modes. While the approach we describe here is based on very simple statistics, the next methodological efforts will also have to take into account more various and heterogeneous sources of data such as phylogeny (Eklöf et al. 2012) and expert knowledge. Bayesian inference appears a good candidate to achieve goal. It offers the possibility to constrain the prior distribution of the likelihood of any pairwise interaction, based on trait matching, using alternative source of data. The method we employed here was based on the match between predator and prey traits. This method could be generalized by looking at the match between all potential traits. Eklof et al. (2013) indeed found that the match between only three traits is enough to encompass a vast majority of interactions. But when such data is not always available, then it might be possible to use phylogeny as a proxy for a high dimensional trait species, provided that traits are conserved along the phylogeny (Mouquet et al. 2012). A bayesian addition to the model would be to weight the likelihood that species *j* is a predator of species *i*, considering that the closest relative of this predator is also a predator of i. It could account for part of the unexplained variation we found in our analysis, for instance by constraining the range of the diet for some large predators or by creating holes within the feeding range (Rall2011).

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Adding this method to the ecologist's toolbox is a timely issue because there is an urgent need to include interactions into future biodiversity scenarios (Thuiller et al. 2013). There has been impressive progress in biogeography over the last fifteen years to predict the impact of global changes on range shifts, extinction risks, and the future distribution of biodiversity, but we now face the realization that emerging ecosystems are unlikely to be a spatial displacement of the ones we currently know. This method will help assessing global change impacts of community structure, moving the field of biogeography closer to ecosystem functioning.

6 Acknowledgements

- We thank I. Gounand for preparing the food web data. We also thank N. Martinez, U.
- ³⁹⁸ Brose and two anonymous reviewers for their constructive comments and stimulating
- 399 discussion. DG received financial support from NSERC and Canada Research Chair
- program. TP is funded by a MELS-FQRNT post-doctoral fellowship.

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506 Figure legends

Figure 1

Schematic representation of the niche model and its application to the predator-prey 508 body size relationship. Three consumer species are represented. The white triangles 500 correspond to the niche position of predators (their position on the X axis is trans-510 posed along the Y axis). For most cases, prey size is smaller that the predator size and 511 thus fall under the 1:1 relationship. In some occasions, however, the feeding range can 512 overlay the 1:1 relationship, for instance in presence of ontogenic shifts in diet. For a 513 given predator, we consider that the niche centroid (c_i) is the average prey size, given by the linear relationship between predator and prey size (black line). The boundaries of the feeding range (r_i) are given by the 5% and 95% quantile regression lines (dot-516 ted lines). A predator feeds on all prey species which their niche lies between these 517 boundaries. In this specific example, the largest predator feeds on the two smaller 518 species (the white triangles depicting the niche lies between range boundaries) and 519 the intermediate size predator feeds on the smallest species. 520

Figure 2

Example of predator-prey body size relationship and predictions from the parameterized niche model. Open dots represent observed links while black dots are predicted
interactions based on the parameterization of the niche model. The bullseye symbols
thus correspond to a match between predicted occurrence of a link and the real data
(the "a" classification in the calculation of the TSS, see main text), a black dot a predicted link with not observation ("b") and an open circle a non-predicted and observed
link ("c"). Data from a creek bed located in Australia (Brose et al. 2005).

Performance of the parameterized niche model to predict food web structure for the 11 food webs. A) Relationship between the TSS and the R^2 of the predator-prey body size relationship. B) Partitionning of the different components of the TSS.

533 Figure 4

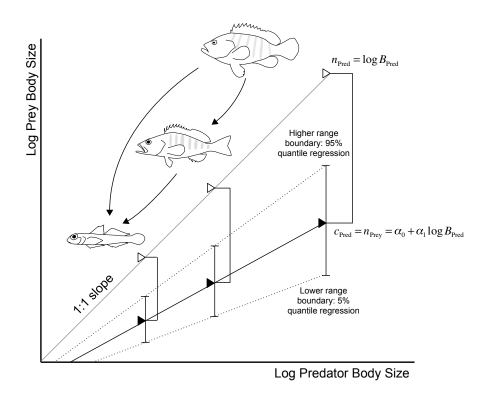
Effect of the sampling effort on the accuracy of the parameterized niche model to predict food web structure. The simulation was conducted with the same data as Fig.

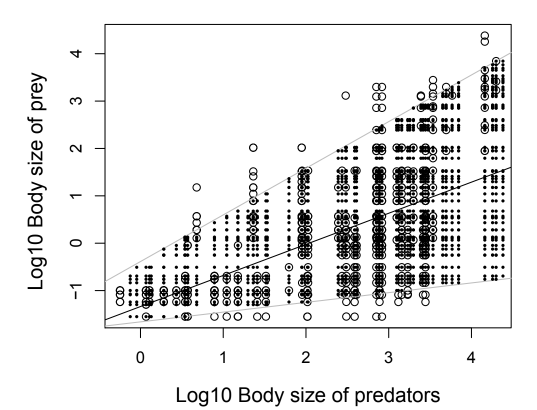
537 Figure 5

Illustration of the metaweb of the Mediterannean pelagic fishes (interactions in black).
Panel A) represents the full metaweb and B) the same metaweb for which links between species not co-occurring together were removed (their range overlap for less than 0.1% of the Mediterannean sea).

Figure 6

Impacts of altering the body size distribution of Meditarrennean fishes on the degree distribution of the metaweb. The black line corresponds to the original metaweb degree distribution. For the first scenario, we simulated a 40% reduction in average body size and recalculated the metaweb properties, keeping the predator-prey body size relationship constant (dotted line). For the second scenario we eliminated the top 40% largest species and recalculated the metaweb properties (dashed line).





26

