# Modeling Food Webs: Exploring Unexplained Structure Using Latent Traits

Rudolf Philippe Rohr,<sup>1,\*</sup> Heike Scherer,<sup>2</sup> Patrik Kehrli,<sup>1,†</sup> Christian Mazza,<sup>3</sup> and Louis-Félix Bersier<sup>1</sup>

Unit of Ecology and Evolution, University of Fribourg, Chemin du Musée 10, 1700 Fribourg, Switzerland;
Department of Mathematics, University of Geneva, 2–4 rue du Lièvre, 1211 Geneva, Switzerland;
Department of Mathematics, University of Fribourg, Chemin du Musée 23, 1700 Fribourg, Switzerland

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ABSTRACT: Several stochastic models have tried to capture the architecture of food webs. This approach is interesting, but it is limited by the fact that different assumptions can yield similar results. To overcome this limitation, we develop a purely statistical approach. Body size in terms of an optimal ratio between prey and predator is used as explanatory variable. In 12 observed food webs, this model predicts, on average, 20% of interactions. To analyze the unexplained part, we introduce a latent term: each species is described by two latent traits, foraging and vulnerability, that represent nonmeasured characteristics of species once the optimal body size has been accounted for. The model now correctly predicts an average of 73% of links. The key features of our approach are that latent traits quantify the structure that is left unexplained by the explanatory variable and that this quantification allows a test of whether independent biological information, such as microhabitat use, camouflage, or phylogeny, explains this structure. We illustrate this method with phylogeny and find that it is linked to one or both latent traits in nine of 12 food webs. Our approach opens the door to the formulation of more complex models that can be applied to any kind of biological network.

*Keywords:* biological network, latent variable, statistical model, body size, phylogeny, community structure.

#### Introduction

Understanding food-web structure remains one of the central questions in ecology, and it is of both fundamental and practical relevance. Cohen (1977) pioneered this research by assembling a collection of webs and analyzing regularities in their structures. This statistical approach was based on derived descriptors (e.g., proportions of top, intermediate, and basal species, connectance, intervality; Sugihara et al. 1989; Cohen et al. 1990; Bersier et al. 2002), which were typically regressed with regard to food-web size as measured by number of species. Stochastic and probabilistic models of food-web architecture were proposed to account for the observed relationships (Cohen and Newman 1985; Williams and Martinez 2000; Drossel et al. 2001; Cattin et al. 2004; Rossberg et al. 2006; Allesina et al. 2008; Petchey et al. 2008). The goodnesses of fit of these models were evaluated by comparing observed and predicted values of derived descriptors (with the notable exception of the model by Allesina et al. [2008], who used a likelihood approach to compute the probability of reproducing the entire web).

The aim of these models was to uncover the major factors underlying food-web organization. In almost all cases, a hierarchy was assumed to exist between the species that was described by a rank or a niche value. Two nonexclusive factors have been proposed to generate the hierarchy: body size, and phylogeny. In Cohen and Newman's (1985) pioneering cascade model, a species with a given rank is allowed to eat any species of lower rank, with a given probability. The other models are built on this general framework, usually with the number of species and the total number of trophic links as parameters. Earlier models (cascade: Cohen and Newman 1985; niche: Williams and Martinez 2000; nested hierarchy: Cattin et al. 2004) differ only in the rules used to assign the trophic links between consumers and resources. Later approaches involved more parameters to improve the quality of the models. Rossberg et al. (2006) introduced a dynamical stochastic model for food-web structure that had five free parameters. In that model, each species is characterized by the log of its body size and two binary vectors: the foraging traits and the vulnerability traits. These behavioral traits do not correspond directly to measurable quantities; they are latent parameters. These traits and the number of species evolve over time by speciation, extinction, and adaptation. Drossel et al. (2001) developed a model where

<sup>\*</sup> Corresponding author; e-mail: rudolf.rohr@unifr.ch.

<sup>&</sup>lt;sup>†</sup> Present address: Station de Recherche Agroscope Changins-Wädenswil, 1260 Nyon, Switzerland.

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population dynamics are explicitly modeled and determine extinctions. Again, they introduced for each species a latent trait ("features of species") that controls trophic interactions. Recently, Petchey et al. (2008) introduced a mechanistic model. This was the first attempt to predict the actual links in real trophic matrices as opposed to simply generating plausible webs with a stochastic model. The body-size distribution was used to predict feeding interactions on the basis of allometric rules and optimal foraging theory. It showed that body size plays an important role in trophic relations, but the authors also pointed out that a second trait would be required to better explain the structure of the predation matrix.

In this article, we propose a purely statistical approach to explore the structure of food webs. This approach is inspired by the methods used in social-network studies (Hoff 2009). We define and investigate two models: the body-size model and the latent-trait model. The former uses the prey-over-predator body-size ratio as the explanatory variable. We show that, with this model, a large part of the variability remains unexplained. In the second model, our aim is to quantify the part that is not explained by the first model. For this purpose, we introduce latent variables-one vulnerability trait and one foraging traitfor each species (Rossberg et al. 2006). Even though these abstract latent traits are not directly observable, we are able to estimate them from the predation matrix and the body sizes of the species by using Bayesian statistics and Markov Chain Monte Carlo (MCMC) techniques. Finally, we show that these latent variables are very useful to test whether biological information other than body-size ratios plays a role in determining the architecture of the food web.

#### Material and Methods

The predation matrix is the matrix  $\{a_{ij}\}$ , which is defined by  $a_{ij} = 1$  if *j* consumes *i* and  $a_{ij} = 0$  otherwise. The aim of our statistical models is to fit the predation matrix of a food web. We achieve this in the following manner:

1. For each pair of species (i, j), we estimate the linking probability  $Pr(a_{ij} = 1)$  using the model.

2. We compute the expected number of trophic links  $\hat{L} = \sum_{ij} \Pr(a_{ij} = 1)$ .

3. The fitted predation matrix is obtained by attributing trophic links to the  $\hat{L}$  pairs of species having the highest linking probabilities.

From a statistical point of view, the attribution of trophic links is a binary classification problem. In this situation, a frequently applied method is the logistic regression. Thus, the equations for the models are given for the logit of  $\Pr(a_{ij} = 1)$ , that is,  $\log \left[\Pr(a_{ij} = 1) / \Pr(a_{ij} = 0)\right]$ .

#### Body-Size Model

Our body-size model uses the log ratio of the body sizes of resources over those of consumers (equivalently, the difference in the log body sizes) as the explanatory variable (Brose et al. 2006; Petchey et al. 2008). In order to implement the idea of an optimal size range for the prey, we model the logit of the linking probability in the following manner:

$$\log\left[\frac{\Pr\left(a_{ij}=1\right)}{\Pr\left(a_{ij}=0\right)}\right] = \alpha + \beta \log\left(\frac{m_i}{m_j}\right) + \gamma \log^2\left(\frac{m_i}{m_j}\right)$$

The parameters to be estimated are  $\alpha$ ,  $\beta$ , and  $\gamma$ . The quadratic polynomial in the log body-size ratio for the logit creates a curve for the linking probabilities whose general aspect resembles a Gaussian distribution function. The bell shape reflects the idea of an optimal range for the prey, consistent with niche theory. In this respect, the optimum (in difference of log) is given by  $-\beta/2\gamma$ , and the ecological range (standard deviation in difference of log) is approximated by  $(-1/2\gamma)^{1/2}$ . As a typical example, figure 1*a* shows the estimated probabilities for the body-size model in red and the values of the observed predation matrix in blue as functions of the log body-size ratios. The dashed line indicates the threshold above which we attribute the trophic links.

#### Latent-Trait Model

Since the body-size model typically predicts only a low percentage of trophic links, we introduce latent terms to quantify the part that is not explained by the optimal body-size ratio. The most general way to do this is to summarize the unobserved part into a matrix **M**, that is,

$$\log\left[\frac{\Pr\left(a_{ij}=1\right)}{\Pr\left(a_{ij}=0\right)}\right] = \alpha + \beta \log\left(\frac{m_i}{m_j}\right) + \gamma \log^2\left(\frac{m_i}{m_j}\right) + M_{ij}.$$

The matrix  $\mathbf{M}$  can be understood as an analog of the residuals in a Gaussian framework. At this point, one could be tempted to try to estimate the components of  $\mathbf{M}$  directly, but since the number of parameters exceeds the number of observations, this would not be reasonable. Note also that each component of  $\mathbf{M}$  refers to a trophic interaction (present or absent) between two species. A better approach is to find a structure where the latent terms refer not to pairwise interactions but to species, which allows a biological interpretation. This can be achieved



Figure 1: Comparison of the body-size model and the latent trait model for the Tuesday Lake food web. In *a* and *d*, with respect to log body-size ratio, the values of the observed predation matrix are shown in blue and those of the fitted linking probabilities are shown in red (note that these are individual points corresponding to the  $66^2$  possible pairs of species). The black curve illustrates the part of the fitted linking probability contributed by the body-size term of the corresponding model. The dashed line indicates the threshold above which the fitted linking probabilities are interpreted as trophic links. *b* and *e* show the observed predation matrix (with the 66 species ordered from smallest to largest on both axes), where the black circles represent trophic links. The background color illustrates the magnitudes of the corresponding predicted linking probabilities, increasing from pale yellow to red. *c* and *f* show again the magnitudes of the fitted linking probabilities, now overlaid with the fitted predation matrix.

using singular-value decomposition (SVD) and concentrating only on the most explicative singular value (see "Singular-Value Decomposition (SVD)" in the online edition of the *American Naturalist*). With this approach, the matrix **M** has the structure  $M_{ij} = v_i \delta f_j$ , where  $\delta$  is the maximal singular value,  $v_i$  (where  $i = 1 \dots S$ ) refers to a species as prey, and  $f_j$  (where  $j = 1 \dots S$ ) refers to a species as predator. Accordingly, our latent-trait model is

$$\log \left[ \frac{\Pr(a_{ij} = 1)}{\Pr(a_{ij} = 0)} \right] =$$
  
$$\alpha + \beta \log \left( \frac{m_i}{m_j} \right) + \gamma \log^2 \left( \frac{m_i}{m_j} \right) + v_i \delta f_j.$$

Each species *i* is now characterized by three numbers: the known body size  $m_i$  and the unknown latent traits  $v_i$  and  $f_i$ , which we call vulnerability and foraging, respectively. These are estimated from the observed predation matrix during the fitting of the model. The introduction of the latent-trait term allows noncontiguous diet ranges so that, as in nature, predators are not constrained to feed on all prey within a continuous region of the body-mass axis. In this case, a large positive latent-trait term (the product  $v_i \delta f_j$ ) can compensate for an unfavorable body-size ratio, whereas a negative latent-trait term can produce a low linking probability even if the body-size ratio is favorable. In figure 1*d*, the estimated linking probabilities are depicted in red as a function of the log ratio of body size.

The black curve represents the effect of body size, corresponding to the quadratic polynomial in the logit function. It is clear that the contribution of the latent traits to the linking probability is very important.

#### Variants of the Latent-Trait Model

We have also considered the following subcases of the latent-trait model: (1) a model with latent traits only, that is,  $\log [\Pr(a_{ij} = 1) / \Pr(a_{ij} = 0)] = v_i \delta f_i$ ; (2) a model with the latent trait of vulnerability only, that is,  $M_{ij} = v_i \delta$ ; and (3) a model with the latent trait of foraging only, that is,  $M_{ij} = \delta f_i$ .

#### Parameter Estimation

The body-size model is a generalized linear model (GLM) with explanatory variables  $\log (m_i/m_j)$  and  $\log^2 (m_i/m_j)$ . Thus, we estimate  $\alpha$ ,  $\beta$ , and  $\gamma$  by using a GLM-fitted algorithm based on maximum likelihood with logit linking function and binomial distribution. For the latent-traits model we estimate the parameters  $\alpha$ ,  $\beta$ ,  $\gamma$ , and  $\delta$ , and for each species we estimate the two latent traits. It is important to understand that the latent traits and  $\delta$  cannot be estimated from the residuals of the body-size model but instead must be fitted simultaneously with  $\alpha$ ,  $\beta$ , and  $\gamma$ . We achieve this using a Bayesian approach and an MCMC technique (see "Estimation of the Parameters with Markov Chain Monte Carlo (MCMC) Techniques" in the online edition of the *American Naturalist*).

#### Interpretation of Latent Traits

A priori, we can interpret the latent traits in two ways: mathematically, as their product, or biologically, as their distance. Latent traits are introduced in the model in terms of their product,  $v_i \delta f_i$ . Thus, a positive term (implying that  $v_i$  and  $f_i$  are both of the same sign) increases the probability of the existence of a trophic interaction between *i* and *j* (relative to the optimal ratio part), and vice versa. Reasoning in terms of distance allows a separate interpretation of foraging (or vulnerability) traits: if two species i and khave similar foraging traits—more precisely, if  $|f_i - f_k|$  is small—then for any given potential prey *i*, the term  $v_i \delta f_i$ is close to the term  $v_i \delta f_k$ . For consumers j and k, the contribution of the latent term to the linking probability is of similar magnitude, that is, there should be similarities in their predation behavior. The same reasoning applies to the vulnerability traits. Thus, our latent traits can be used to quantify the similarities between species in their roles of predator and prey, with the key characteristic that the effect of the optimal body-size ratio is removed. A

posteriori, we can of course try to understand whether the values of latent traits can be interpreted directly.

#### How to Relate Latent Traits to Biological Information

As latent traits quantify the similarity between species, a sensible approach to relate them to external information is to compare similarity matrices. A standard method is the Mantel test (Sokal and Rohlf 2001), which can be used to detect whether there is a correlation between similarity matrices on the basis of latent traits and the biological descriptors of interest. Here we demonstrate this technique with (1) trophic structure, (2) body size, and (3) phylogeny. The first comparisons are used as a test to assess whether our latent traits capture the corresponding information of the food-web matrix. The second comparisons test whether there remains an effect of body size that is not accounted for by the optimal ratio. Finally, the third comparison uses the latent traits to test whether evolutionary history contributes to food-web architecture when the optimal ratio has been removed.

For the foraging latent traits, we first compute the distances  $d_{ij} = |f_i - f_j|$ . The coefficients of the corresponding similarity matrix are  $1 - d_{ij} / \max_{ij} d_{ij}$ . The similarity matrix with respect to the vulnerability traits is computed in the same manner. We use the Jaccard index (Jaccard 1908) to describe trophic similarity, separating foraging, and vulnerability. Trophic similarity with respect to foraging between two species *i* and *j* (Jaccard foraging) is the number of common prey divided by the total number of prey of the two species. Similarly, the Jaccard index with respect to vulnerability between species i and j (Jaccard vulnerability) is the number of common predators divided by the total number of predators of the two species. Similarity in body size is computed in the same way as for latent traits. The phylogenetic similarity between two species *i* and *j* is computed by assigning values to taxonomic levels from 1 (phylum) to 17 (genus) and dividing the value of most precise common taxonomic level by 1 plus the value of the most precise level to which either of the two species was determined (Cattin et al. 2004).

We use Spearman's rank correlation coefficient in the Mantel tests since it is less sensitive to outliers than Pearson's coefficient. When comparing phylogenetic similarity and similarity with respect to latent traits, it is important to note that the two types of similarity matrices contain body-size information (see "Results and Discussion"). In order to compare them independently of common bodysize components, we use a partial Mantel test; that is, before comparison, we first remove the linear correlation of body size from both the latent-trait and the phylogenetic-similarity matrices.

#### Data

We used the published data set of Brose et al. (2005), from which we extracted 12 highly resolved food webs: (1) Silwood Park (Cohen et al. 2005), (2) grassland (Dawah et al. 1995), (3) Sierra Lakes (Harper-Smith et al. 2005), (4) Tuesday Lakes (Jonsson et al. 2005), (5) Mill Stream (M. E. Ledger, F. Edwards, and G. Woodward, unpublished data), (6) Broom (Memmott et al. 2000), (7) Celtic Sea (Pinnegar et al. 2003), (8) Mulgrave River (T. S. Rayner, unpublished data), (9) Goettingen (Ulrich 1999), (10) Skipwith Pond (Warren 1989), (11) Sheffield (P. H. Warren, unpublished data), and (12) Broadstone Stream (Woodward et al. 2005). We discarded webs for which trophic interactions were not determined from direct observations, as well as, for computational reasons, one very large web. Detailed information is presented in table A1, which is available in a zip file in the online edition of the American Naturalist.

#### **Results and Discussion**

#### Goodness of Fit

We fitted the body-size model and the latent-trait model to 12 high-quality food webs. In order to compare the performance of the two models, we computed as a first criterion the proportion of correctly predicted trophic links (table 1; Petchey et al. 2008). It is clear that the latenttrait model performs much better than the body-size model. The observed and predicted connectance values for the 12 food webs are listed in table 1. Both models performed almost equally as well with respect to the fitted connectance, and in only one case did the body-size model predict a slightly different value. That the latent-trait model outperforms the body-size model is not surprising, as it involves 2S + 1 more parameters. However, in 10 of the 12 food webs, the Akaike Information Criterion (AIC) was lower for the latent-trait model than it was for the body-size model (table 1). This confirms that the addition of the 2S latent traits is justified by the considerable gain in accuracy.

Figure 1 allows for visual comparison of the quality of the fit of the two models for the Tuesday Lake food web. One can easily see that the body-size model poorly reproduces the structure of the food web. Figure 1*f* shows that the magnitudes of the fitted probabilities from the latent-trait model closely follow the structures of the real predation matrix and that noncontiguous diets are permitted. The corresponding figures for the 11 other food webs are available in a zip file in the online edition.

We computed variants of our latent-trait model: one with latent traits only (without information on body size), one with body size and the vulnerability trait only, and one with body size and the foraging trait only. Table 2 gives the results for these variants and reveals that they never outperformed the original latent-trait model.

Additionally, we computed the predicted optimal  $\log_e$  body-size difference and the ecological range representing an optimal diet range for both the body-size and the latent-trait models (table 1). We point out that for all food webs except 1, 2, and 9, the predicted optimal  $\log_e$  body-size difference is negative, implying that predators are larger than their prey. Webs 1, 2, and 9 are parasitoid networks, and positive estimates are thus expected (Cohen et al. 2005). Web 2 is unusual in that the body-size ratio fails

Table 1: Results and goodness of fit of the body-size and latent-trait models

					Body-size model				Latent-trait model				
Food web	S	С	Body-size measure	Ĉ	%	AIC	Optimum difference	Ecological range	Ĉ	%	AIC	Optimum difference	Ecological range
1	34	4.8	Length	9.2	16	439	.1	.4	4.8	71	374	.0	.3
2	25	3.1	Length	3.1	2.9	913			3.1	44	944		
3	37	14.0	Length	13.7	29	960	-4.3	3.2	14.0	93	337	-2.3	1.2
4	66	8.7	Mass	8.7	46	1,643	-9.9	3.3	8.7	87	873	-10.2	2.1
5	79	5.9	Mass	5.9	36	1,626	-20.8	5.6	5.9	74	1,358	-24.4	6.3
6	55	3.3	Length	3.3	12	873	-4.1	3.8	3.3	66	657	-4.5	14.6
7	57	6.3	Mass	6.6	19	1,223	-4.4	2.0	6.3	65	1,018	-5.2	2.1
8	62	5.5	Length	5.5	11	1,307	-6.0	2.6	5.5	64	1,010	-6.7	2.7
9	24	4.2	Mass	4.3	13	173	1.9	1.0	4.2	83	209	1.6	.7
10	67	7.8	Length	7.3	13	2,269	-1.4	1.2	7.8	67	1,491	-1.7	1.1
11	58	4.9	Length	5.1	6	1,276	-1.4	1.5	4.9	72	762	-3.9	1.8
12	29	18.6	Mass	18.5	36	736	-11.6	6.4	18.6	92	285	-5.9	3.1

Note: We report for each food web the number of species (*S*), the connectance (*C*), the body-size measure, the fitted connectance ( $\hat{C}$ ), the proportion of correctly fitted links (%), the Akaike Information Criterion (AIC), the optimum difference in log body size, and the ecological range. The identifiers in the first column refer to those listed in "Data."

Table 2: Model comparison

Food	B.S	. only	B.S.	+ L.T.	L.T	only.	B.S.	+ vul.	B.S.	+ for.
web	%	AIC	%	AIC	%	AIC	%	AIC	%	AIC
1	16	439	71	374	52	405	25	462	38	397
2	2.9	913	44	944	29	924	18	927	23	926
3	29	960	93	337	68	623	47	918	66	530
4	46	1,643	87	875	37	1,607	60	1,429	46	1,633
5	36	1,626	74	1,358	57	1,552	56	1,502	55	1,558
6	12	873	66	657	41	761	38	687	15	953
7	19	1,223	65	1,018	56	1,086	43	1,128	47	1,123
8	11	1,307	64	1,010	57	1,105	31	1,226	48	1,143
9	13	173	83	209	25	251	29	220	29	202
10	13	2,269	67	1,491	52	1,619	31	2,154	41	1,774
11	6	1,276	72	762	55	811	25	1,255	46	802
12	36	736	92	285	87	410	37	791	83	307

Note: For each model, we report the proportion (%) of correctly predicted trophic links and the Akaike Information Criterion (AIC). We consider the following models: the body-size model (B.S. only), the body-size model and the "original" latent-traits model (B.S. + L.T.), the model with the latent traits only (L.T. only), the model with body size and the latent trait of vulnerability only (B.S. + vul.), and the model with body size and the latent traits of foraging only (B.S. + for.). For each food web, the best value is bolded.

to totally explain interaction structure and no optimum can be estimated (see fig. A2, available in a zip file in the online edition).

#### Latent Traits

At this point, the key question is whether the addition of latent traits, which quantify the part of the variation not explained by the optimal body-size ratio, just introduces purely artificial terms or whether it represents a biologically meaningful and justified model improvement. Because the optimal ratio leaves a large part of the variation unexplained, one may expect that some correlation is left between the latent traits and the original information on vulnerability and foraging. The latter can be measured with Jaccard similarity.

We compared the similarity in latent traits to the Jaccard similarities with Mantel tests and found highly significant correlations in 21 out of 24 cases (see table A2, available in a zip file in the online edition). These simple tests confirm that the vulnerability and foraging latent traits are not just fitting parameters but also carry biologically relevant information, with the crucial feature that they are independent of an optimal body-size ratio. We can now use our latent traits to explore which independent biological factors should be added to the optimal ratio to better understand food-web structure.

*Comparison with Body Size.* Does the optimal-ratio term of the model capture all of the influence of body size on trophic behavior? In other words, is there a correlation

between latent traits and body size? We performed Mantel tests between the similarity in body size and against the similarity in latent vulnerability and foraging traits, respectively. Table 3 shows that the optimal body-size ratio does not carry all of the information related to body size. It is interesting that the latent foraging traits are, in general, more closely related to body size than the vulnerability traits are.

Comparison with Phylogeny. Phylogeny has been suggested as a major factor underlying food-web architecture (Cattin et al. 2004; Bersier and Kehrli 2008). Mantel tests between phylogeny and latent-traits similarities resulted in six significant cases out of 12 food webs for the vulnerability traits and seven cases for the foraging traits (table 3). In all, latent traits were not correlated with phylogeny in only three webs, all of which are from freshwater habitats. Phylogeny and body size are not independent, so that the optimal body-size ratio may carry some phylogenetic signal. Finding a significant correlation indicates that phylogeny intervenes in the network in a manner independent from the optimal ratio. Apart from body size and phylogeny, other species characteristics could determine the probability of trophic interactions and could be analyzed in future work using latent traits, for example, microhabitat use, color and camouflage patterns, chemical and other defensive traits, or circadian activity.

Direct Interpretation of the Values of Latent Traits. In order to explore whether we can directly interpret the values of the latent traits, we plotted the estimated foraging traits versus vulnerability traits in figure 2 for the Goettingen food web (for the other webs, see figs. A1-A24 in the zip file in the online edition). We observed three general features: (1) all basal species have very similar foraging traits, whereas their vulnerability traits are more variable: the top predators show the opposite pattern, and the intermediate species generally have a high variability in both traits; (2) the higher the absolute value of the foraging trait, the higher the number of prey, and similarly for the number of predators and the vulnerability trait (see the threedimensional figures in the zip file available in the online edition); (3) for almost all trophic interactions, the sign of the vulnerability trait of the prey is the same as the sign of the foraging trait of the predator. Their product is thus positive, yielding a positive contribution to the overall probability for a link. This indicates that a direct biological interpretation of latent traits must be performed with caution. For example, two prey characterized by a similar number of predators can have vulnerability traits of opposite sign (e.g., see species 30 and 35 in fig. A12 in the zip file available in the online edition).

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	Body size versus latent vulnerability		Body size versus latent foraging		Phylogeny latent vulne	versus rability	Phylogeny versus latent foraging	
Food web	Correlation coefficient	Р	Correlation coefficient	Р	Correlation coefficient	Р	Correlation coefficient	Р
1	.092	.075	.132	.014	.484	<.001	.660	<.001
2	.178	.007	.052	.143	043	.830	.222	<.001
3	.139	.078	.324	<.001	.224	<.001	018	.583
4	.124	.052	.409	<.001	.358	<.001	.399	<.001
5	.020	.340	.243	<.001	.75	.037	.289	<.001
6	.222	<.001	154	.974	094	.890	.153	.001
7	115	.974	.373	<.001	.012	.403	084	.967
8	.030	.285	.291	<.001	.225	<.001	.374	<.001
9	.221	.089	015	.522	.214	.015	095	.839
10	.000	.473	.168	.002	.042	.174	.076	.019
11	.033	.107	.021	.261	036	.700	.025	.338
12	.093	.116	181	.960	.005	.455	056	.790

Table 3: Results of the Mantel tests

Note: We compare body-size similarity with similarity of latent vulnerability and with latent foraging traits and phylogenetic similarity with similarity of latent vulnerability and with latent foraging traits. P values <.05 are indicated in bold; P values from .05 to .1 are indicated in italic type.

#### Conclusions

The body-size model is, to our knowledge, the first purely statistical model of food-web structure using an independent variable (body-size ratio) as a predictor of trophic interactions. We show how the inclusion of latent traits can dramatically improve the fit of the model and, more importantly, that these parameters can be used to shed light on the factors that structure trophic interactions in natural communities.

Our analyses reveal that the optimal body-size ratio does not capture all the trophic information of a food web and that the latent traits quantify a large part of the structure. First, the interpretation of latent traits shows that the effect of body size is not fully embodied in the optimal ratio, since for more than half of the food webs, body size is also correlated with the latent traits. More importantly, this additional effect of body size is found predominantly for the foraging traits. It might be speculated that this asymmetry is due to a more opportunistic behavior of predators, which pushes them to consume outside of the optimal body-size ecological range (Rossberg et al. 2006; Bersier and Kehrli 2008). Second, latent traits are related to phylogeny in most food webs, which confirms the importance of long-term evolution on community structure. However, the signal of phylogeny was not found in three webs for which latent traits nonetheless greatly improved the proportion of correctly predicted interactions. Without additional information on these systems, it is difficult to speculate on the biological factors that could account for this unexplained variation. The very value of latent traits is that they allow an exploration of such nontrivial structures.

Latent traits can also be used in meta-analyses to highlight general properties of systems. For example, we found that the increase of accuracy with the latent-trait model is more important for terrestrial webs (on average, by a factor of 7.8) than for aquatic webs (by an average factor of 4.5). This result indicates that the optimal ratio plays a larger role in aquatic environments than in terrestrial environments, which supports the finding that body size is a major driver of trophic interactions in the former systems (Shurin et al. 2006). Finally, latent traits as im-



Figure 2: Estimated latent traits for the Goettingen food web. The lines indicate the trophic interactions.

plemented here (one trait assigned to outflows and one to inflows) could easily be applied to analyze networks in various domains of biology, from pollinators to protein networks.

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## Appendix from R. P. Rohr et al., "Modeling Food Webs: Exploring Unexplained Structure Using Latent Traits"

(Am. Nat., vol. 176, no. 2, p. 170)

### **Supplementary Materials and Methods**

### Singular-Value Decomposition (SVD)

Let **M** be an  $S \times S$  square matrix (we explain the SVD for square matrices, but it can be applied to any rectangular matrix). The singular value decomposition of **M** (see chap. 7 of Watkins 1991) is a factorization into a product of three matrices,  $\mathbf{M} = \mathbf{VDF}$ , where

$$\mathbf{V} = \begin{pmatrix} \begin{vmatrix} & & & \\ \mathbf{v}^{1} & \cdots & \mathbf{v}^{S} \\ \mid & & \mid \end{pmatrix},$$
$$\mathbf{D} = \begin{pmatrix} \delta_{1} & & \\ & \ddots & \\ & & \delta_{S} \end{pmatrix},$$
$$\mathbf{F} = \begin{pmatrix} - & \mathbf{f}^{1} & - \\ & \vdots & \\ - & \mathbf{f}^{S} & - \end{pmatrix}.$$
(A1)

The matrix **D** is a diagonal matrix with *S* nonnegative numbers  $\delta_1, \ldots, \delta_s$  called the singular value of **M**; **V** is a matrix of *S* orthogonal unitary (of length 1) column vectors  $\mathbf{v}^1, \ldots, \mathbf{v}^s$ ; **F** is a matrix of *S* orthogonal unitary row vectors  $\mathbf{f}^1, \ldots, \mathbf{f}^s$ . Without loss of generality, is it always possible to reorder the singular values from the largest to the smallest, that is,  $\delta_1 \ge \delta_2 \ge \ldots \ge \delta_s$ ; the columns of **V** and the rows of **F** must be reordered accordingly. Note that SVD is unique; that is, the singular values are unique up to their ordering, and the vectors are also unique up to their orientation in the space.

We can approximate the matrix **M** using the SVD in the following ways: The simplest approximation consists of considering only the first singular value (the largest) and setting all others to 0, that is,  $\delta_2 = ... = \delta_s = 0$ . This gives us  $\mathbf{M} \approx \mathbf{v}^1 \delta_1 \mathbf{f}^1$ , which corresponds exactly to the latent term in our latent-trait model. This approximation is called the first-rank approximation of **M**.

The second-simplest approximation consists of considering only the two first singular values, which gives us  $\mathbf{M} \approx \mathbf{v}^1 \delta_1 \mathbf{f}^1 + \mathbf{v}^2 \delta_2 \mathbf{f}^2$ . This approximation is called the second-rank approximation of  $\mathbf{M}$ . We can continue until we consider all nonzero singular values; in the last case, we obtain exactly  $\mathbf{M}$ .

Note that SVD is closely related to principal component analysis. The singular values of the data matrix correspond to the eigenvalues of the covariance matrix. The columns of the matrix  $\mathbf{V}$  are the eigenvectors of the covariance matrix.

Finally, note that the condition on the vectors  $\mathbf{f}^i$  and  $\mathbf{v}^i$  to be of unit length is not important for the computation of the similarity matrices (see "How to Relate Latent Traits to Biological Information"). Similarity matrices are independent of the choice of scale.

#### Estimation of the Parameters with Markov Chain Monte Carlo (MCMC) Techniques

To fit the parameters of the latent-trait model, maximum likelihood estimation could not be used because the likelihood function is too complex to maximize. Instead, we employed a Bayesian approach. Contrary to maximum likelihood, the parameters in the Bayesian framework are considered to be random variables. In this

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context, we have to specify a prior distribution  $p(\theta)$  for the parameter  $\theta$  and determine the likelihood  $p(y|\theta)$  of the model (y denotes the data). Using Bayes's theorem, we then obtain the posterior distribution of the parameters, that is, their distribution when the data are known:

$$p(\theta|y) = \frac{p(y|\theta)}{p(y)}p(\theta).$$
 (A2)

These posterior distributions of the parameters are too complex to be computed analytically, and a common method is to "sample" these using an MCMC technique. The main idea is to take samples from a discrete-time Markov chain  $(X_n)_{n\geq 0}$  whose stationary distribution is the posterior distribution of the parameters  $\theta$  we are looking for and to let it run for a long time. Under certain not very restrictive conditions, the Markov chain will converge to its steady state and the  $X_n$  for large n can be considered as (certainly not independent) samples from the posterior distribution of the parameters. Moreover, it can be shown that

$$\frac{1}{N-k}\sum_{n=k}^{N}X_{n} \to E(\theta), \tag{A3}$$

as  $N \rightarrow \infty$ . In other words, the mean of the Markov chain will converge to the expected values of the parameters. Since this is true for any *k*, it can be useful to choose *k* such that the first and most imprecise values of the Markov chain (the so-called burn-in phase) are not taken into account. This will accelerate the convergence. For a good introduction to MCMC techniques and their application to Bayesian statistics, see Gilks et al. (1996) and Yang (2005). More precise and technical information can be found in Robert and Casella (2004).

#### **Details of the Algorithm**

The parameters to be estimated in the latent-trait model are  $\alpha$ ,  $\beta$ ,  $\gamma$ , and  $\delta$  and, for each species *i* (where  $1 \le i \le S$ ), the latent traits  $\mathbf{v}_i$  and  $\mathbf{f}_i$ . The data are the values (i.e., 0 or 1) of the food-web matrix **A**. To construct the Markov chain in which stationary distribution converges to the posterior distribution of the parameters, we use the Metropolis-Hastings algorithm (sec. 3.4 of Yang 2005). In the next section, we describe the algorithm used. Finally we give some technical details concerning the distributions.

#### **Metropolis-Hastings** Algorithm

Algorithm: suppose the Markov chain to be constructed up to state *n*; more precisely, suppose  $\alpha_n$ ,  $\beta_n$ ,  $\gamma_n$ ,  $\delta_n$ ,  $\mathbf{v}_n$ , and  $\mathbf{f}_n$  are known (note that  $\mathbf{v}_n$  and  $\mathbf{f}_n$  denote the vectors at the *n*th state of the Markov chain and not the *n*th components of the vectors).

Propose a new value  $\alpha^*$  by sampling from  $N(\alpha_n, \sigma_\alpha)$ . Accept this new value and put  $\alpha_{n+1} = \alpha^*$  with probability

$$\min\left\{1, \frac{p(\alpha^*|\beta_n, \gamma_n, \delta_n, \mathbf{v}_n, \mathbf{f}_n, \mathbf{A})}{p(\alpha_n|\beta_n, \gamma_n, \delta_n, \mathbf{v}_n, \mathbf{f}_n, \mathbf{A})}\right\},\tag{A4}$$

else  $\alpha_{n+1} = \alpha_n$ .

Propose a new value  $\beta^*$  by sampling from  $N(\beta_n, \sigma_\beta)$ . Accept this new value and put  $\beta_{n+1} = \beta^*$  with probability

$$\min\left\{1, \frac{p(\beta^*|\alpha_{n+1}, \gamma_n, \delta_n, \mathbf{v}_n, \mathbf{f}_n, \mathbf{A})}{p(\beta_n | \alpha_{n+1}, \gamma_n, \delta_n, \mathbf{v}_n, \mathbf{f}_n, \mathbf{A})}\right\},\tag{A5}$$

else  $\beta_{n+1} = \beta_n$ .

Propose a new value  $\gamma^*$  by sampling from  $N(\gamma_n, \sigma_\gamma)$ . Accept this new value and put  $\gamma_{n+1} = \gamma^*$  with probability

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$$\min\left\{1, \frac{p(\gamma^*|\alpha_{n+1}, \beta_{n+1}, \delta_n, \mathbf{v}_n, \mathbf{f}_n, \mathbf{A})}{p(\gamma_n | \alpha_{n+1}, \beta_{n+1}, \delta_n, \mathbf{v}_n, \mathbf{f}_n, \mathbf{A})}\right\},\tag{A6}$$

else  $\gamma_{n+1} = \gamma_n$ .

Propose a new value  $\delta^*$  by sampling from  $N(\delta_n, \sigma_{\delta})$ . Accept this new value and put  $\delta_{n+1} = \delta^*$  with probability

$$\min\left\{1, \frac{p(\delta^*|\alpha_{n+1}, \beta_{n+1}, \mathbf{\gamma}_{n+1}, \mathbf{v}_n, \mathbf{f}_n, \mathbf{A})}{p(\delta_n|\alpha_{n+1}, \beta_{n+1}, \gamma_{n+1}, \mathbf{v}_n, \mathbf{f}_n, \mathbf{A})}\right\},\tag{A7}$$

else  $\delta_{n+1} = \delta_n$ .

Propose a new value  $\mathbf{v}^*$  by sampling from vMF( $\mathbf{v}_n, \boldsymbol{\kappa}_v$ ). Accept his new value and put  $\mathbf{v}_{n+1} = \mathbf{v}^*$  with probability

$$\min\left\{1, \frac{p(\mathbf{v}^*|\boldsymbol{\alpha}_{n+1}, \boldsymbol{\beta}_{n+1}, \boldsymbol{\gamma}_{n+1}, \boldsymbol{\delta}_{n+1}, \mathbf{f}_n, \mathbf{A})}{p(\mathbf{v}_n|\boldsymbol{\alpha}_{n+1}, \boldsymbol{\beta}_{n+1}, \boldsymbol{\gamma}_{n+1}, \boldsymbol{\delta}_{n+1}, \mathbf{f}_n, \mathbf{A})}\right\},\tag{A8}$$

else  $\mathbf{v}_{n+1} = \mathbf{v}_n$ .

Propose a new value  $\mathbf{f}^*$  by sampling from vMF( $\mathbf{f}_n, \mathbf{\kappa}_f$ ). Accept his new value and put  $\mathbf{f}_{n+1} = \mathbf{f}^*$  with probability

$$\min\left\{1, \frac{p(\mathbf{f}^*|\boldsymbol{\alpha}_{n+1}, \boldsymbol{\beta}_{n+1}, \boldsymbol{\gamma}_{n+1}, \boldsymbol{\delta}_{n+1}, \mathbf{v}_{n+1}, \mathbf{A})}{p(\mathbf{f}_n|\boldsymbol{\alpha}_{n+1}, \boldsymbol{\beta}_{n+1}, \boldsymbol{\gamma}_{n+1}, \boldsymbol{\delta}_{n+1}, \mathbf{v}_{n+1}, \mathbf{A})}\right\},\tag{A9}$$

else  $\mathbf{f}_{n+1} = \mathbf{f}_n$ . Here vMF( $\mu$ ,  $\kappa$ ) denotes the von Mises-Fisher distribution, which is a distribution on the unit sphere in  $\mathbf{R}^s$ , parameterized by its mean direction  $\mu$  and concentration parameter  $\kappa$ . The sampling technique for a von Mises-Fisher distribution is explained in Wood (1994).

#### Likelihood, Prior and Posterior Distribution

The prior distributions of the parameters  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$ ,  $\mathbf{v}$ , and  $\mathbf{f}$  are chosen to be independent and of the following types: normal for  $\alpha$ ,  $\beta$ ,  $\gamma$ , and  $\delta$  and uniform on the unit sphere in  $\mathbf{R}^s$  for  $\mathbf{v}$  and  $\mathbf{f}$  (recall once again that  $\mathbf{v}$  and  $\mathbf{f}$  are of length 1). As explained in the preceding section, it is necessary to compute the acceptance probabilities for the proposal values of the parameters. These probabilities are expressed as quotients of posterior probabilities of the parameters. As an illustration of how these probabilities can be simplified, consider the case of the parameter  $\alpha$ : since the prior distributions of the parameters  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$ ,  $\mathbf{v}$ , and  $\mathbf{f}$  are assumed to be independent, Bayes's theorem yields

$$p(\alpha^*|\beta,\gamma,\delta,\mathbf{v},\mathbf{f},\mathbf{A}) = \frac{p(\mathbf{A}|\alpha^*,\beta,\gamma,\delta,\mathbf{v},\mathbf{f})}{p(\beta,\gamma,\delta,\mathbf{v},\mathbf{f},\mathbf{A})}p(\alpha^*),$$
$$p(\alpha_n|\beta,\gamma,\delta,\mathbf{v},\mathbf{f},\mathbf{A}) = \frac{p(\mathbf{A}|\alpha_n,\beta,\gamma,\delta,\mathbf{v},\mathbf{f})}{p(\beta,\gamma,\delta,\mathbf{v},\mathbf{f},\mathbf{A})}p(\alpha_n).$$

Hence, the only function needed to compute the acceptance probabilities are the likelihood function and the prior distribution, which are both known:

$$\frac{p(\alpha^*|\beta,\gamma,\delta,\mathbf{v},\mathbf{f},\mathbf{A})}{p(\alpha_n|\beta,\gamma,\delta,\mathbf{v},\mathbf{f},\mathbf{A})} = \frac{p(\mathbf{A}|\alpha^*,\beta,\gamma,\delta,\mathbf{v},\mathbf{f})p(\alpha^*)}{p(\mathbf{A}|\alpha_n,\beta,\gamma,\delta,\mathbf{v},\mathbf{f})p(\alpha_n)}.$$

By definition of the model, the likelihood function is

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$$p(\mathbf{A}|\alpha,\beta,\gamma,\delta,\mathbf{v},\mathbf{f}) = \prod_{i,j=1}^{S} \Pr(a_{ij}=1)^{a_{ij}} \Pr(a_{ij}=0)^{1-a_{ij}},$$

where  $\Pr(a_{ij} = 1) = e^{\theta_{ij}}/(1 + e^{\theta_{ij}})$  for  $\theta_{ij} = \log [\Pr(a_{ij} = 1)/\Pr(a_{ij} = 0)]$ , the logit of the linking probability, is given by the equation of the latent-trait model (see "Latent-Trait Model").

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N	Common name	Geographic location	General habitat	Specific habitat	pw/fw *
1	Silwood Park	United Kingdom; UTM: 51.24'N, 0.34'W; Silwood Park, Berkshire	terrestrial	abandoned field	pw
2	Grassland	United Kingdom	terrestrial	grasslands	fw
3	Sierra	United States of America; John	freshwater	small subalpine and	fw
	Lakes	Muir Wilderness Area and Kings		alpine lakes, less than	
		Canyon National Park, Sierra Nevada mountains, California		3m deep	
4	Tuesday Lakes	United States of America; UTM: 89.32'W, 46.13'N; Tuesday Lake, Michigan	freshwater	small, mildly acidic lake, 1984	fw
5	Mill	United Kingdom: the River Frome	freshwater	lowland chalk stream	fw
0	Stream	Dorset: UTM: 50 40'N 2 11'W	ii esii watei		1.00
6	Broom	United Kingdom; UTM: 51.24'N, 0.34'W; Silwood Park, Berkshire	terrestrial	<i>Cytisis scoparius</i> (Scotch broom) patch, source web on broom	fw
7	Celtic Sea	Europe, Celtic Sea ecosystem	marine	demersal food web	pw
8	Mulgrave River	Australia; Mulgrave River; UTM: 17.08'S, 145.52'E	freshwater	lowland coastal river	pw
9	Goettingen	Germany; Goettingen; UTM: 51.31'N, 09.56'E;	terrestrial	Beech forest	pw
10	Skipwith Pond	United Kingdom; UTM: 53.40'N, 0.59'W; Skipwith Common, North Yorkshire	freshwater	acidic pond, up to 1 m deep, 0.25 ha	fw
11	Sheffield	United Kingdom; Sheffield;	freshwater	laboratory study	pw
12	Broadstone Stream	United Kingdom; UTM: 51.05'N, 0.03'E; Broadstone Stream in Sussex	freshwater	spring-fed acidic head- water stream, 120m ele- vation	fw

\*fw indicates a complete food web, pw a partial web

	Vul. traits vs.	Jaccard vul.	For. traits vs	Jaccard for.
Food web	corr. coeff.	p-value	corr. coeff.	p-value for
1	0.502	<0.001	0.196	0.004
2	0.304	<0.001	0.042	0.183
3	0.540	<0.001	0.061	0.224
4	0.669	<0.001	0.424	<0.001
5	0.227	<0.001	0.596	<0.001
6	-0.006	0.525	0.465	<0.001
7	0.273	<0.001	0.352	<0.001
8	0.397	<0.001	0.224	<0.001
9	0.404	<0.001	0.193	<0.001
10	0.320	<0.001	0.280	<0.001
11	0.379	<0.001	0.465	<0.001
12	0.286	0.008	0.419	<0.001

Table A2: Results of the Mantel tests comparing the similarity in latent traits of vulnerability (foraging) with the Jaccard index of vulnerability (foraging).

Boldface type indicates p-values  $\leq 0.01$ .

For each of the 12 studied food webs, there are three pages.

- **1st page.** This figure is the same as Fig. 1 in the main article: comparison of the Body Size Model and the Latent Trait Model. Figures a and d show, with respect to log body size ratio, the coefficients of the observed predation matrix in blue and the fitted linking probabilities in red (note that these are individual points corresponding to all possible pairs of species). The black curve illustrates the part of the fitted linking probability contributed by the body size term of the corresponding model. The dashed line indicates the threshold above which the fitted linking probabilities are interpreted as trophic links. Figures b and e show the observed predation matrix, where black circles represent trophic links. The background color illustrates the magnitudes of the corresponding predicted linking probabilities, increasing from pale yellow to red. Figures c and f show again the magnitudes of the fitted linking probabilities, now overlaid with the fitted predation matrix.
- **2nd page.** Upper figure : graph of the estimated foraging against the estimated vulnerability traits Symbols indicate trophic level and lines represent trophic interactions. Lower left and right : 3-dimensional graph of the number of consumers and number of resources, respectively, against. the estimated latent traits.
- **3nd page.** Names of the species in the food web. The numbering is the same as in the preceding graphs.



Figure A1: Silwood Park



Figure A2: Silwood Park

	Silwood Park
Sp. no.	Species name
1	Aphelinus varipes adults
2	Aphelinus abdominalis adults
3	Capitophorus carduinis larvae
4	Aphis larvae
5	Binodoxys acalephe adults
6	Capitophorus carduinis adults
7	Praon abjectum adults
8	Aphidius matricariae adults
9	Aphis adults
10	Megoura viciae larvae
11	Sitobion larvae
12	Sitobion ptericolens larvae
13	Metopolophium albidum larvae
14	Praon volucre adults
15	Ephedrus plagiator adults
16	Aphidius rhopalosiphi adults
17	Amphorophora rubi larvae
18	Sitobion adults
19	Praon dorsale adults
20	Macrosiphum funestum larvae
21	Metopolophium albidum adults
22	Aphidius ervi adults
23	Acyrthosiphon pisum larvae
24	Aphidius eadyi adults
25	Amphorophora rubi adults
26	Aphidius picipes adults
27	Microlophium carnosum larvae
28	Aphidius urticae adults
29	Aphidius microlophii adults
30	Sitobion ptericolens adults
31	Macrosiphum funestum adults
32	Megoura viciae adults
33	Acyrthosiphon pisum adults
34	Microlophium carnosum adults



Figure A3: Grassland



Figure A4: Grassland

Grassland						
Sp. no.	Species name	Sp. no.	Species name			
1	Pediobius festucae larvae	41	Tetramesa hyalipennis larvae			
2	Pediobius claridgei larvae	42	Pediobius dactylicola larvae			
3	Pediobius planiventris larvae	43	Tetramesa longicornis larvae			
4	Pediobius alaspharus larvae	44	Chlorocytus larvae			
5	Mesopolobus graminum larvae	45	Eurytoma tapio larvae			
6	Eurytoma roseni larvae	46	Eurytoma appendigaster larvae			
7	Sycophila mellea larvae	47	Tetramesa petiolata larvae			
8	Eurytoma flavimana larvae	48	Ahtola atra larvae			
9	Tetramesa brevicornis larvae	49	Eurytoma pollux larvae			
10	Chlorocytus agropyri larvae	50	Tetramesa angustipennis larvae			
11	Chlorocytus formosus larvae	51	Chlorocytus harmolitae larvae			
12	Tetramesa cornuta larvae	52	Eupelmus atropurpureus larvae			
13	Tetramesa linearis larvae	53	Eurytoma danuvica larvae			
14	Homoporus fulviventris larvae	54	Tetramesa calamagrostis larvae			
15	Pediobius larvae	55	Macroneura vesicularis larvae			
16	Pediobius deschampiae larvae	56	Tetramesa eximia larvae			
17	Eurytoma phalaridis larvae	57	Endromopoda larvae			
18	Tetramesa fulvicollis larvae					
19	Chlorocytus deschampiae larvae					
20	Chlorocytus pulchripes larvae					
21	Eurytoma larvae					
22	Pediobius eubius larvae					
23	Eurytoma erdoesi larvae					
24	Chlorocytus phalaridis larvae					
25	Homoporus febriculosus larvae					
26	Homoporus luniger larvae					
27	Torymus baudysi larvae					
28	Bracon larvae					
29	Bracon erythrostictus larvae					
30	Chlorocytus ulticonus larvae					
31	Homoporus larvae					
32	Eurytoma collaris larvae					
33	Tetramesa airae larvae					
34	Tetramesa brevicollis larvae					
35	Tetramesa longula larvae					
36	Tetramesa albomaculata larvae					
37	Pediobius calamagrostidis larvae					
38	Tetramesa phleicola larvae					
39	Pediobius phalaridis larvae					
40	Eurytoma castor larvae					

Table A4



Figure A5: Sierra Lakes



Figure A6: Sierra Lakes

	Sierra Lakes
Sp. no.	Species name
1	Keratella
2	Keratella quadrata
3	Polyarthra
4	Kellicottia
5	Chydorus sphaericus
6	Daphnia rosea
7	Leptodiaptomus signicauda
8	Cyclopoda
9	Daphnia middendorffiana
10	Hesperodiaptomus shoshone
11	Hydroporini adults
12	Hydroporini larvae; adults
13	Pisidium
14	Culex
15	Corixidae
16	Psychoglypha
17	Sialis
18	Agabus
19	Polycentropus
20	Limnephilus
21	Desmona
22	Callibaetis ferrugineus
23	Chironomidae
24	Ameletus
25	Hesperophylax
26	Oligochaeta
27	Rana muscosa larvae
28	Rana muscosa adults
29	Thamnophis elegans elegans
30	Salvelinus fontinalis
31	Salvelinus fontinalis adults
32	Oncorhynchus mykiss aguabonita
33	Oncorhynchus mykiss aguabonita adults
34	Oncorhynchus mykiss
35	Oncorhynchus mykiss adults
36	Salmo trutta
37	Salmo trutta adults



Figure A7: Tuesday Lake



Figure A8: Tuesday Lake

Tuesday Lake						
Sp. no.	Species name	Sp. no.	Species name			
1	Chromulina	41	Ploesoma			
2	Unclassified microflagellates	42	Gastropus stylifer			
3	Ankistrodesmus spiralis	43	Gastropus hyptopus			
4	Ankyra judai	44	Conochiloides dossuarius			
5	Trachelomonas	45	Trichocerca multicrinis			
6	Closteriopsis longissima	46	Filinia longispina			
7	Chroococcus dispersus	47	Ascomorpha eucadis			
8	Selenastrum minutum	48	Polyarthra vulgaris			
9	Unclassified flagellates	49	Trichocerca cylindrica			
10	Dictyosphaerium pulchellum	50	Synchaeta			
11	Cryptomonas	51	Bosmina longirostris			
12	Schroederia setigera	52	Diaphanosoma leuchtenbergianum			
13	Dinobryon sociale	53	Conochilus			
14	Nostoc	54	Tropocyclops prasinus			
15	Quadrigula	55	Leptodiaptomus siciloides			
16	Mallomonas	56	Daphnia rosea			
17	Chroococcus limneticus	57	Skistodiaptomus oregonensis			
18	Arthrodesmus	58	Cyclops varicans rubellus			
19	Peridinium pusillum	59	Orthocyclops modestus			
20	Oscillatoria	60	Daphnia pulex			
21	Dinobryon cylindricum	61	Holopedium gibberum			
22	Cosmarium	62	Chaoborus punctipennis			
23	Dinobryon bavaricum	63	Phoxinus eos			
24	Oocystis	64	Phoxinus neogaeus			
25	Glenodinium pulvisulcus	65	Umbra limi			
26	Quadrigula lacustris	66	Micropterus salmoides			
27	Glenodinium quadridens					
28	Dinobryon sertularia					
29	Keratella cochlearis					
30	Sphaerocystis schroeteri					
31	Keratella testudo					
32	Microcystis aeruginosa					
33	Kellicottia					
34	Kellicottia bostoniensis					
35	Peridinium wisconsinense					
36	Peridinium cinctum					
37	Kellicottia longispina					
38	Synedra					
39	Peridinium limbatum					
40	Gloeocystis					

Table A6



Figure A9: Mill Stream



Figure A10: Mill Stream

	Mill Stream					
Sp. no.	Species name	Sp. no.	Species name			
1	Cyanobacteria	41	Pericoma trivialis larvae			
2	Fragilaria elliptica	42	Procladius larvae			
3	Achnanthes elliptica	43	Pisidium adults; juveniles			
4	Gomphonema olivaceum	44	Macropelopia larvae			
5	Amphora pediculus	45	Limnophila larvae			
6	Fragilaria leptostauron	46	Limonia larvae			
7	Achnanthes lanceolata	47	Antocha vitripennis larvae			
8	Gongrosira incrustans	48	Tinodes waeneri larvae			
9	Cymbella minuta	49	Microtendipes			
10	Melosira varians	50	Microtendipes larvae			
11	Cocconeis placentula	51	Athripsodes cinereus larvae			
12	Navicula menisculus	52	Aphelocheirus aestivalis larvae			
13	Surirella ovalis	53	Lumbriculidae adults; juveniles			
14	Navicula gregaria	54	Eiseniella tetraedra adults; juveniles			
15	Nitzschia dissipata	55	Valvata piscinalis adults; juveniles			
16	Nitzschia perminuta	56	Polycentropus flavomaculatus larvae			
17	Fragilaria vaucheriae	57	Gammarus pulex adults; juveniles			
18	Rhoicosphenia curvata	58	Simuliidae			
19	Ostracoda	59	Simuliidae larvae			
20	Diatoma vulgare	60	Asellus aquaticus adults; juveniles			
21	Navicula tripunctata	61	Limnophora larvae			
22	Amphora ovalis	62	Potamopyrgus jenkinsi adults; juveniles			
23	Navicula lanceolata	63	Limnius larvae			
24	Gyrosigma obtusatum	64	Theodoxus fluviatilis adults; juveniles			
25	Ciliophora	65	Anabolia nervosa larvae			
26	Cymatopleura solea	66	Brychius elevatus larvae			
27	Nematoda	67	Potamophylax latipennis larvae			
28	Synorthocladius larvae	68	Hydropsyche larvae			
29	Cricotopus	69	Halesus radiatus larvae			
30	Cricotopus larvae	70	Lymnea peregra adults; juveniles			
31	Hydroptila larvae	71	Ancylus fluviatilis adults; juveniles			
32	Naididae adults; juveniles	72	Brachycentrus subnubilus larvae			
33	Heterotrissocladius	73	Erpobdella octoculata adults; juveniles			
34	Heterotrissocladius larvae	74	Ephemera danica			
35	Pentaneura	75	Ephemera danica larvae			
36	Rhyacophila dorsalis larvae	76	Sialis lutaria larvae			
37	Limnephilus larvae	77	Sericostoma personatum larvae			
38	Elmis aenea larvae	78	Tipula montium			
39	Bezzia larvae	79	Tipula montium larvae			
40	Tubificidae adults; juveniles					



Figure A11: Broom



Figure A12: Broom

Broom						
Sp. no.	Species name	Sp. no.	Species name			
1	Beauveria bassiana	41	Meta segmentata			
2	Paecilomyces	42	Coccinella septempunctata			
3	Anystis baccarum	43	Agonopteryx assimilella			
4	Hylastinus obscurus	44	Xantholinus linearis			
5	Mesopolobus mediterraneus	45	Anatis ocellata			
6	Phloeophthorus rhododactylus	46	Chesias legatella			
7	Necremnus metalarus	47	Harpalus rubripes			
8	Aprostocetus tibialis	48	Forficula auricularia			
9	Acyrthosiphon spartii	49	Araneus diadematus			
10	Chrysocharis gemma	50	Philonthus politus			
11	Pringalio soemias	51	Pisaura mirabilis			
12	Microctonus	52	Abax parallelepipedus			
13	Habrocytus sequester	53	Parus caeruleus			
14	Apion fuscirostre	54	Parus major			
15	Centistes excrucians	55	Fringilla coelebs			
16	Leucoptera spartifoliella					
17	Apion immune					
18	Arytaina genistae					
19	Arytaina spartii					
20	Aphis sarathamni					
21	Bruchidius ater					
22	Apanteles fulvipes					
23	Apanteles vitripennis					
24	Cheiropachys colon					
25	Dinotiscus bidentulus					
26	Braconid					
27	Anthocoris sarothamni					
28	Ilyobates nigricollis					
29	Gonioctena olivacea					
30	Phytodecta olivacea					
31	Exochomus quadripustulatus					
32	Propylaea quatuordecimpunctata					
33	Adalia decempunctata					
34	Adalia bipunctata					
35	Sitona regensteinensis					
36	Heterocordylus tibialis					
37	Philodromus aurelolus caespiticolis					
38	Linyphia triangularis					
39	Evarcha arcuata					
40	Xysticus cristatus					



Figure A13: Celtic Sea



Figure A14: Celtic Sea

Celtic Sea				
Sp. no.	Species name	Sp. no.	Species name	
1	Glyptocephalus cynoglossus	41	Trisopterus luscus	
2	Fish-Larvae	42	Leucoraja naevus	
3	Sandeel	43	Lophius budegassa	
4	Goby	44	Zeus faber	
5	Agonus cataphractus	45	Melanogrammus aeglefinus	
6	Capros aper	46	Lepidorhombus boscii	
7	Myctophid	47	Raja undulata	
8	Gaidropsarus vulgaris	48	Squalus acanthias	
9	Gadiculus argenteus	49	Leucoraja fullonica	
10	Cepola macrophthalma	50	Galeorhinus galeus	
11	Rockling	51	Lophius piscatorius	
12	Hatchet-fish	52	Gadus morhua	
13	Scaldfish	53	Raja montagui	
14	Sprattus sprattus	54	Raja clavata	
15	Buglossidium luteum	55	Molva molva	
16	Trisopterus esmarkii	56	Pollachius virens	
17	Clupeoid	57	Pollachius pollachius	
18	Gadoid			
19	Callionymus lyra			
20	Hippoglossoides platessoides			
21	Microchirus variegatus			
22	Arnoglossus imperialis			
23	Argentina sphyraena			
24	Flatfish			
25	Engraulis encrasicolus			
26	Trisopterus			
27	Fish-unidentified			
28	Trisopterus minutus			
29	Micromesistius poutassou			
30	Trachurus trachurus			
31	Sardina pilchardus			
32	Microstomus kitt			
33	Scomber scombrus			
34	Clupea harengus			
35	Eutrigla gurnardus			
36	Merlangius merlangus			
37	Lepidorhombus whiffiagonis			
38	Chelidonichthys cuculus			
39	Scyliorhinus canicula			
40	Merluccius merluccius			



Figure A15: Mulgrave River



Figure A16: Mulgrave River

Mulgrave River			
Sp. no.	Species name	Sp. no.	Species name
1	Diatoms and desmids	41	Awaous acritosus
2	Filamentous algae	42	Gerres filamentosus
3	Detritus	43	Herring
4	Testate amoeba adults	44	Fish
5	Collembola juveniles	45	Giuris margaritacea
6	Diptera (not Chironomidae) larvae	46	Leiognathus equulus
7	Aquatic macrophytes	47	Caranx ignobilis
8	Elmidae larvae	48	Kuhlia rupestris
9	Hemiptera adults	49	Notesthes robusta
10	Arachnida adults	50	Microphis brachyurus
11	Diptera larvae	51	Hephaestus tulliensis
12	Ostracoda adults	52	Acanthopagrus berda
13	Chironomidae larvae	53	Arrhamphus sclerolepis
14	Corixidae and Notonectidae - incl. Plea adults	54	Lutjanus argentimaculatus
15	Ephemeroptera larvae	55	Nematalosa erebi
16	Small terrestrial inverts adults	56	Bunaka gyrinoides
17	Planktonic inverts - Cladocera etc adults	57	Anguilla reinhardtii
18	Hymenoptera adults	58	Neosilurus ater
19	Lepidoptera larvae	59	Tandanus tandanus
20	Trichoptera larvae	60	Mugil cephalus
21	Terrestrial vegetation	61	Anguilla australis
22	Gerridae and Mesovelioidea adults	62	Lates calcarifer
23	Large terrestrial inverts adults		
24	Fruit		
25	Decapoda adults		
26	Orthoptera adults		
27	Odonata larvae		
28	Mollusca adults		
29	Toxotes chatareus		
30	Redigobius bikolanus		
31	Xiphophorus maculatus		
32	Pseudomugil signifer		
33	Hypseleotris compressa		
34	Macrobrachium adults		
35	Ambassis agassizii		
36	Craterocephalus stercusmuscarum		
37	Glossamia aprion		
38	Melanotaenia splendida		
39	Tilapia mariae		
40	Glossogobius		



Figure A17: Goettingen



Figure A18: Goettingen

Goettingen		
Sp. no.	Species name	
1	Basalys parva adults	
2	Kleidotoma psiloides adults	
3	Pentapleura adults	
4	Idiotypa nigriceps adults	
5	Aspilota (Species 2) adults	
6	Alloxysta victrix adults	
7	Aspilota (Species 5) adults	
8	Ismarus dorsiger adults	
9	Aspilota adults	
10	Dendrocerus carpenteri adults	
11	Pediobius foliorum adults	
12	Aspilota (Species 3) adults	
13	Aphidius ervi adults	
14	Aphelopus holomelas adults	
15	Limosina sp. pupae	
16	Aphelopus melaleucus adults	
17	Aphelopus serratus adults	
18	Orthostigma adults	
19	Megaselia pupae	
20	Megaselia ruficornis pupae	
21	Eulophus larvarum adults	
22	Atractodes adults	
23	Gymnophora arcuata pupae	
24	Pegomya pupae	

Table A11



Figure A19: Skipwith



Figure A20: Skipwith

Skipwith			
Sp. no.	Species name	Sp. no.	Species name
1	Chydorus latus adults	41	Argyroneta aquatica larvae
2	Chydorus latus larvae	42	Agabus sturmii adults
3	Scapholeberis mucronata adults	43	Agabus sturmii larvae
4	Scapholeberis mucronata juveniles	44	Agabus bipustulatus adults
5	Scapholeberis mucronata larvae	45	Agabus bipustulatus larvae
6	Oribatei sp adults	46	Illybius fuliginosus adults
7	Acanthocyclops vernalis adults	47	Illybius fuliginosus larvae
8	Acanthocyclops vernalis larvae	48	Sialis lutaria larvae
9	Enchytraeidae adults	49	Lestes sponsa adults
10	Chironomidae larvae	50	Lestes sponsa larvae
11	Enchytraeidae larvae	51	Sialis lutaria adults
12	Corynoneura scutellata adults	52	Limnephilus marmoratus adults
13	Corynoneura scutellata larvae	53	Limnephilus marmoratus larvae
14	Tanytarsus adults	54	Sympetrum scoticum adults
15	Tanytarsus larvae	55	Sympetrum scoticum larvae
16	Chironomidae adults	56	Corixa punctata adults
17	Procladius sagittalis adults	57	Corixa punctata larvae
18	Procladius sagittalis larvae	58	Corixa dentipes adults
19	Hydroporus erythrocephalus adults	59	Corixa dentipes larvae
20	Hydroporus erythrocephalus larvae	60	Notonecta glauca larvae
21	Polycelis tenuis larvae	61	Notonecta glauca adults
22	Enallagma cyathigerum adults	62	Lumbriculus variegatus adults
23	Enallagma cyathigerum larvae	63	Lumbriculus variegatus larvae
24	Glyptotendipes pallens adults	64	Aeshna juncea adults
25	Glyptotendipes pallens larvae	65	Aeshna juncea larvae
26	Holocentropus picicornis adults	66	Dytiscus marginalis adults
27	Holocentropus picicornis larvae	67	Dytiscus marginalis larvae
28	Sigara semistriata adults		
29	Sigara semistriata larvae		
30	Chironomus dorsalis adults		
31	Chironomus dorsalis juveniles		
32	Chironomus dorsalis larvae		
33	Callicorixa praeusta adults		
34	Callicorixa praeusta larvae		
35	Hesperocorixa linnei adults		
36	Hesperocorixa linnei larvae		
37	Hesperocorixa sahlbergi adults		
38	Hesperocorixa sahlbergi larvae		
39	Arctocorisa germari larvae		
40	Argyroneta aquatica adults		



Figure A21: Sheffield



Figure A22: Sheffield

Sheffield				
Sp. no.	Species name	Sp. no.	Species name	
1	Chydoridae (Species 1)	41	Ceratopogonidae	
2	Lymnea	42	Limnephilus centralis	
3	Acari (Species 1)	43	Limnephilus flavicornis	
4	Cyclopoidea	44	Corixa punctata	
5	Simocephalus vetulus	45	Sialis lutaria adults	
6	Daphnia pulex	46	Chaoborus obscuripes	
7	Acari (Species 2)	47	Agabus larvae	
8	Ostracoda	48	Chironomidae (Species 1)	
9	Caenis horaria	49	Chaoborus crystallinus larvae	
10	Potamopyrgus jenkinsi	50	Chaoborus crystallinus pupae	
11	Hygrotus inaequalis	51	Sialis lutaria	
12	Haliplus ruficollis	52	Notonecta glauca adults	
13	Polycelis tenuis	53	Tubificidae	
14	Unionicola Pentatax	54	Cyrnus flavidus	
15	Daphnia magna	55	Aeshna cyanea larvae	
16	Hesperocorixa castanea	56	Sigara dorsalis larvae	
17	Cymatia coleoptrata	57	Sigara nigrolineata	
18	Cloeon dipterum	58	Tubificidae (Species 1)	
19	Noterus clavicornis			
20	Laccophilus minutus			
21	Caenis luctuosa			
22	Helobdella stagnalis			
23	Helobdella stagnalis adults			
24	Nemoura cinerea			
25	Crangonyx pseudogracilis			
26	Lymnea peregra			
27	Chironomidae (Species 2)			
28	Culicidae			
29	Sigara dorsalis			
30	Polycelis tenuis adults			
31	Asellus aquaticus			
32	Ischnura elegans			
33	Agabus nebulosus			
34	Chironomidae (Species 3)			
35	Chironomidae (Species 5)			
36	Chironomidae (Species 4)			
37	Gammarus pulex			
38	Coenagrion puella larvae			
39	Piscicola geometra			
40	Agabus bipustulatus adults			

Table A13



Figure A23: Broadstone Stream



Figure A24: Broadstone Stream

Broadstone Stream				
Sp. no.	Species name	Sp. no.	Species name	
1	Corynoneura lobata larvae	41	Argyroneta aquatica larvae	
2	Micropsectra bidentata larvae	42	Agabus sturmii adults	
3	Polypedilum albicorne larvae	43	Agabus sturmii larvae	
4	Heterotrissocladius marcidus larvae	44	Agabus bipustulatus adults	
5	Zavrelimyia barbatipes larvae	45	Agabus bipustulatus larvae	
6	Brillia modesta larvae	46	Illybius fuliginosus adults	
7	Paraleptophlebia submarginata larvae	47	Illybius fuliginosus larvae	
8	Trissopelopia longimana larvae	48	Sialis lutaria larvae	
9	Leuctra nigra larvae	49	Lestes sponsa adults	
10	Simulium larvae	50	Lestes sponsa larvae	
11	Nemurella pictetii larvae	51	Sialis lutaria adults	
12	Macropelopia nebulosa larvae	52	Limnephilus marmoratus adults	
13	Prodiamesa olivacea larvae	53	Limnephilus marmoratus larvae	
14	Leuctra hippopus larvae	54	Sympetrum scoticum adults	
15	Siphonoperla torrentium larvae	55	Sympetrum scoticum larvae	
16	Niphargus aquilex adults	56	Corixa punctata adults	
17	Diptera larvae	57	Corixa punctata larvae	
18	Dicranota larvae	58	Corixa dentipes adults	
19	Adicella reducta larvae	59	Corixa dentipes larvae	
20	Oligochaeta larvae; adults	60	Notonecta glauca larvae	
21	Plectrocnemia conspersa larvae	61	Notonecta glauca adults	
22	Asellus meridianus adults	62	Lumbriculus variegatus adults	
23	Tipulidae larvae	63	Lumbriculus variegatus larvae	
24	Oligochaeta	64	Aeshna juncea adults	
25	Scirtidae (previous Helodidae) sp. larvae	65	Aeshna juncea larvae	
26	Sialis fuliginosa larvae	66	Dytiscus marginalis adults	
27	Potamophylax cingulatus larvae	67	Dytiscus marginalis larvae	
28	Pedicia larvae			
29	Cordulegaster boltonii larvae			
30	Chironomus dorsalis adults			
31	Chironomus dorsalis juveniles			
32	Chironomus dorsalis larvae			
33	Callicorixa praeusta adults			
34	Callicorixa praeusta larvae			
35	Hesperocorixa linnei adults			
36	Hesperocorixa linnei larvae			
37	Hesperocorixa sahlbergi adults			
38	Hesperocorixa sahlbergi larvae			
39	Arctocorisa germari larvae			
40	Argyroneta aquatica adults			