



Food-web structure in relation to environmental gradients and predator-prey ratios in tank-bromeliad ecosystems

Olivier Dézerald, Céline Leroy, Bruno Corbara, Jean-François Carrias,
Laurent Pelozuelo, Alain Dejean, Régis Céréghino

► To cite this version:

Olivier Dézerald, Céline Leroy, Bruno Corbara, Jean-François Carrias, Laurent Pelozuelo, et al.. Food-web structure in relation to environmental gradients and predator-prey ratios in tank-bromeliad ecosystems. PLoS ONE, 2013, vol. 8 (8), pp. 1-8. 10.1371/journal.pone.0071735 . hal-00913931

HAL Id: hal-00913931

<https://hal.science/hal-00913931>

Submitted on 4 Dec 2013

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution 4.0 International License



Open Archive TOULOUSE Archive Ouverte (OATAO)

OATAO is an open access repository that collects the work of Toulouse researchers and makes it freely available over the web where possible.

This is an author-deposited version published in : <http://oatao.univ-toulouse.fr/>
Eprints ID : 10204

To link to this article : DOI:10.1371/journal.pone.0071735
URL : <http://dx.doi.org/10.1371/journal.pone.0071735>

To cite this version : Dézerald, Olivier and Leroy, Céline and Corbara, Bruno and Carrias, Jean-François and Pelozuelo, Laurent and Dejean, Alain and Céréghino, Régis. *Food-web structure in relation to environmental gradients and predator-prey ratios in tank-bromeliad ecosystems*. (2013) PLoS ONE, vol. 8 (n° 8). pp. 1-8. ISSN 1932-6203

Any correspondence concerning this service should be sent to the repository administrator: staff-oatao@listes-diff.inp-toulouse.fr

Food-Web Structure in Relation to Environmental Gradients and Predator-Prey Ratios in Tank-Bromeliad Ecosystems

Olivier Dézerald^{1*}, Céline Leroy², Bruno Corbara³, Jean-François Carrias³, Laurent Pélozuelo^{4,5}, Alain Dejean^{1,4}, Régis Céréghino^{4,5}

¹ EcoFoG, Ecologie des Forêts de Guyane, CNRS UMR 8172, Kourou, France, ² AMAP, botAnique et bioinforMatique de l'Architecture des Plantes, IRD UMR 123, Montpellier, France, ³ LMGE, Laboratoire Microorganismes: Génome et Environnement, Université Blaise Pascal, CNRS UMR 6023, Aubière, France, ⁴ EcoLab, Laboratoire Ecologie Fonctionnelle et Environnement, Université Paul Sabatier, Toulouse, France, ⁵ EcoLab, Laboratoire Ecologie Fonctionnelle et Environnement, CNRS UMR 5245, Toulouse, France

Abstract

Little is known of how linkage patterns between species change along environmental gradients. The small, spatially discrete food webs inhabiting tank-bromeliads provide an excellent opportunity to analyse patterns of community diversity and food-web topology (connectance, linkage density, nestedness) in relation to key environmental variables (habitat size, detrital resource, incident radiation) and predator:prey ratios. We sampled 365 bromeliads in a wide range of understorey environments in French Guiana and used gut contents of invertebrates to draw the corresponding 365 connectance webs. At the bromeliad scale, habitat size (water volume) determined the number of species that constitute food-web nodes, the proportion of predators, and food-web topology. The number of species as well as the proportion of predators within bromeliads declined from open to forested habitats, where the volume of water collected by bromeliads was generally lower because of rainfall interception by the canopy. A core group of microorganisms and generalist detritivores remained relatively constant across environments. This suggests that (i) a highly-connected core ensures food-web stability and key ecosystem functions across environments, and (ii) larger deviations in food-web structures can be expected following disturbance if detritivores share traits that determine responses to environmental changes. While linkage density and nestedness were lower in bromeliads in the forest than in open areas, experiments are needed to confirm a trend for lower food-web stability in the understorey of primary forests.

* E-mail: olivier.dezerald@gmail.com

Introduction

Biodiversity is not only the sum of coexisting species, it is also the diversity of interactions that connect these species. It is widely acknowledged that linkage patterns among species, rather than species richness alone, are a key component of the diversity-stability relationship [1]. For instance, for a fixed number of species, food-web persistence and robustness (two concepts linked to network stability) are believed to increase with connectance, the proportion of all possible interactions that are realized [2]. A rich body of literature has focused on the relationship between network structure and dynamics [3], and on how different types of interactions (either antagonistic or mutualistic) affect network properties [4]. Most studies however focused on network structure regardless of variation in the abiotic environment. Consequently, our understanding of the environmental determinants of network structure lags behind the increasingly vast knowledge of species richness- and interactions-mediated patterns [5,6]. Identifying

linkages between ecological networks and physical environments could be relevant to predict the extent to which entire communities will respond to increasing environmental disturbances (e.g., climate change, habitat fragmentation, overharvesting, introduction of invasive species, pollution).

On a local scale, both the spatial/temporal changes in habitat conditions and species traits regulate community structure [7], thereby affecting ecosystem functions. For instance, the nature and extent of the riparian vegetation determines food quality and availability in still [8] and running waters [9], thus influencing the functional feeding group composition of macroinvertebrate communities [10]. However, we know little about how linkage patterns among species (food-web topology) change against a background of environmental gradients over broad, regional scales. Most of what we know about environmental determinants of food-web structure comes from experiments in single locations [11], on a target fraction of the food web (e.g., vertebrates, macro-

and micro-invertebrates excluding components such as microorganisms, bacteria, flagellates, rotifers and viruses; [12]).

It has proven challenging to study the effects of both abiotic and biotic factors on food webs. High species diversity and population densities in large ecosystems preclude the accurate characterization of trophic links (notably in continuous habitats), and because it is often difficult to manipulate entire ecosystems. To tackle these issues, we focussed on small, spatially discrete food webs that naturally span a broad range of environmental gradients. Bromeliaceae are flowering plants represented by 59 genera and some 3140 species native mainly to the Neotropics [13]. The interlocking leaves of tank-forming bromeliads form wells that collect water, leaf litter and other organic detritus. The detritus that enter the tank constitute the main source of nutrients for the aquatic food web [14]. The aquatic communities inhabiting tank-bromeliads provide excellent opportunities to study food-web structure because they contain several trophic levels (from bacteria to predatory macroinvertebrates; [15]) and can be exhaustively sampled [16].

The aim of this study was to determine whether spatial patterns in food-web structure can be predicted from a small set of environmental factors and/or the richness of predators relative to their prey. To address this question, we sampled 365 tank-bromeliads in a wide range of environments in French Guiana (plantations, pioneers growths, rock savannah, primary forest) and drew the 365 corresponding food webs using the gut contents of invertebrates as well as observations of predator-prey encounters. Previous studies on tank-bromeliads concluded that detrital resources at the base of the food web, understorey light environments (energy available for algal production), habitat size and predation play key roles in shaping aquatic community composition [17,18]. Detritus constitute the main source of energy for aquatic bromeliad communities, however, the high algal biomass found in sun-exposed bromeliads [19] may provide a complementary non-detrital resource to the upper trophic levels. If (i) species richness and abundance increase with bromeliad (habitat) size [20–24], and (ii) bromeliads from open area benefit from both detrital and non-detrital resources and understorey bromeliad foodwebs are solely supported by detrital matter [15], then, for a given habitat size, the diversity of invertebrate functional feeding groups should increase from forest to open areas. One may expect shifts in food-web connectance, linkage density, and/or nestedness as species with particular traits are replaced or complemented by species with other traits when shifting from forest understorey to open areas. Conversely, if the resource that supports food webs does not differ from forest to open areas, then one should not observe significant shifts in food-web structure and functions. To test these hypotheses, we used Linear Mixed Effect modelling to analyse patterns of community diversity and food-web topology in relation to key environmental variables and predator:prey richness ratios.

Materials and Methods

Ethics Statement

This study was conducted according to relevant national and international guidelines. Sample collections necessary to scientific research were authorized by the French *Office National des Forêts* (ONF) provided that their impact upon the environment is considered negligible.

Study Area, Bromeliads and Environmental Variables

The study was conducted in French Guiana, from March 2006 to October 2011. The climate is tropical moist with 3,000 -

3,400 mm of yearly precipitation mainly distributed over 280 days. There is a major reduction in rainfall between September and November and another shorter and more irregular dry period in March. The maximum monthly temperature averages 33.5°C (32.1–35.8°C), and the monthly minimum averages 20.3°C (19.7–21°C).

We selected five sampling localities distributed across a south-east to north-west range (Fig. 1) and sampled 365 bromeliads (i.e. 365 food webs) in the understorey of primary and transitional forests, in pioneer growths, in a rock savannah, and in plantations. The main habitat characteristics of tank-bromeliad species found in five vegetation types at five localities (hereafter “vegetation types”, within localities) are summarized in Table 1. Further descriptions of the Nouragues (sampling period in April 2006), Petit-Saut (March 2009) and Kaw (Oct. 2008) localities and their bromeliads can be found in [15,22,25]. Saint-Elie (Oct. 2007) and Angoulême (April 2010) are *Citrus* plantations. Both epiphytic bromeliads and those that had taken root on the ground were included in the study.

In order to prevent seasonality effects (i.e., dry vs rainy season) on water volumes and species abundance, the sampling took place at the transition between rainy and dry seasons, where invertebrate abundances reach a peak. We also sampled tank-bromeliads that were full of water. Species occurrence was not an issue as bromeliad invertebrates have fast life cycles (<2 months) with overlapping cohorts. To characterize habitat size, we emptied the wells in each plant by sucking the water out (see invertebrate sampling) and recorded the corresponding volume of water (WV, mL). The amount of fine particulate organic matter (FPOM; 1000 to 0.45 µm in size) was expressed as preserved volume (mL) after decantation in graduated test-tubes [12]. Finally, percentages of total incident radiation (IR) above the bromeliads were calculated using hemispherical photographs and an image processing software (Gap Light Analyzer 2.0; [26]), as described in [23]. In this study, we consider that all bromeliads with an IR below 50% were located in partially shaded areas or forested environments while higher percentages defined sun-exposed areas.

Aquatic Invertebrate Communities

For both ethical (extensive sampling could destroy local populations) and legal reasons (the Nouragues Research Station is located in a nature reserve), we decided to use a non-destructive sampling technique for all studied bromeliads. To sample the water retained in the tanks, we used 5-mL and 10-mL automatic micropipettes with the end trimmed to widen the orifice. We homogenized the water within leaf wells by sucking in and out with a pipet, before sucking out the content. Although less efficient than plant dissection, we and other researchers have already successfully used this technique [24,27]. It was consistently used for all of the samples and most of the water (>95%) was collected. Both early and late instars of prey and predator invertebrates were captured, so we were confident that our technique was efficiently implemented. The samples were preserved in the field in 4% (final concentration) formalin. Invertebrates were sorted in the laboratory and preserved in 70% ethanol. They were identified to genus, species or morphospecies and enumerated (species lists in [21,24]). Species abundance data (individuals per plant) were used to calculate evenness (Simpson index) and entropy (Shannon-Weaver index) for each invertebrate community. The Simpson index was calculated as $D = \sum p_i^2$. The Shannon-Weaver index (hereafter Shannon index) was defined as $H = -\sum p_i \log(b) p_i$, where p_i is the proportional abundance of species i and b is the base of the logarithm (natural logarithm in this case). Species richness and abundance being components of these two indices, they were not

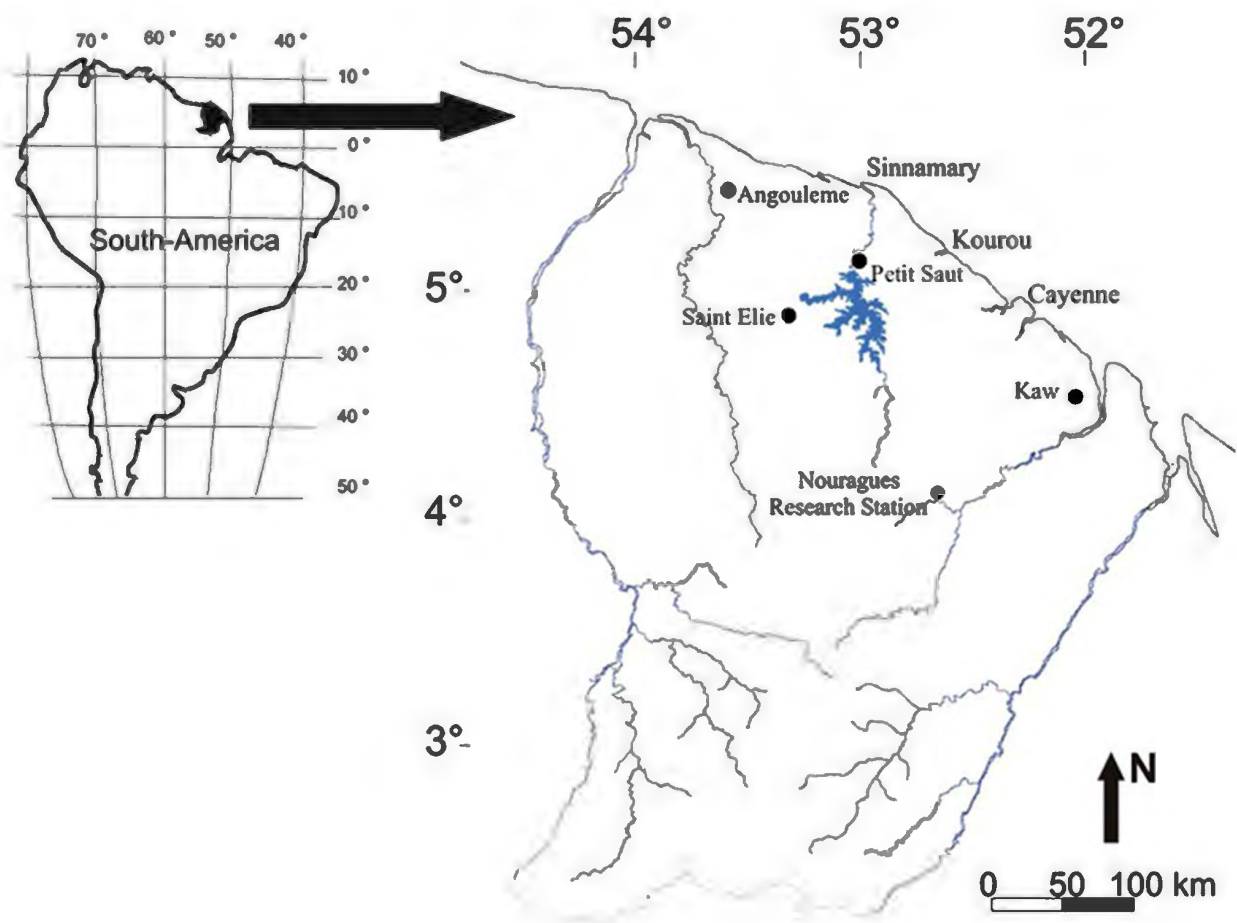


Figure 1. Distribution of the sampling locations in French Guiana (see Table 1).
doi:10.1371/journal.pone.0071735.g001

considered alone in subsequent modeling. Macroinvertebrate taxa were partitioned into predators (carnivorous species which attack and consume living macroinvertebrates) and prey (species which sift fine particulates and microorganisms from the water column

and/or gather FPOM and associated microorganisms from the accumulated debris), and these categories were used to calculate Predator:Prey Richness ratios (number of predatory taxa/number of prey taxa, hereafter PPR).

Table 1. Main habitat characteristics (mean $\pm$ standard error) of tank-bromeliad species found in five vegetation types.

Locality	Vegetation types	Species	N	IR	WV	FPOM	#Taxa
Nouragues	Rock savannah	CB	29	73.3 $\pm$ 2.4	40.62 $\pm$ 3.75	0.74 $\pm$ 0.08	10.55 $\pm$ 0.15
		AA	31	66.97 $\pm$ 2.73	949.23 $\pm$ 102.64	10.84 $\pm$ 1.26	15.87 $\pm$ 0.42
	Transitional forest	VP	30	25.12 $\pm$ 0.29	73.2 $\pm$ 9.11	1.7 $\pm$ 0.28	10.3 $\pm$ 0.35
		AB	26	25.69 $\pm$ 0.47	137.85 $\pm$ 21.1	1.08 $\pm$ 0.25	11.23 $\pm$ 0.32
	Primary forest	VS	26	18.75 $\pm$ 0.4	48.54 $\pm$ 5.03	4.29 $\pm$ 0.42	10.35 $\pm$ 0.25
Petit-Saut	Pioneer growth	GL	19	15.9 $\pm$ 0.6	17.46 $\pm$ 2.53	1.04 $\pm$ 0.16	9.21 $\pm$ 0.29
		AM	63	39.33 $\pm$ 2.63	84.38 $\pm$ 10.1	6.16 $\pm$ 1.26	12.49 $\pm$ 0.28
	Primary forest	VS	34	16.64 $\pm$ 0.38	26.18 $\pm$ 3.84	3.46 $\pm$ 0.71	10.68 $\pm$ 0.23
Kaw	Pioneer growth	AM	45	35.1 $\pm$ 2.98	92.46 $\pm$ 11.82	4.3 $\pm$ 0.68	11.78 $\pm$ 0.28
Angoulême	Citrus plantation	AM	35	33.03 $\pm$ 2.95	31.01 $\pm$ 4.54	2.32 $\pm$ 0.31	11.8 $\pm$ 0.34
Saint-Elie	Citrus plantation	AM	27	32.07 $\pm$ 1.3	56.41 $\pm$ 8.29	4.58 $\pm$ 0.86	14.22 $\pm$ 0.36

CB: *C. berteroniana*; AA: *A. aquilega*; VP: *V. pleiosticha*; AB: *A. bromeliifolia*; VS: *V. splendens*; GL: *G. lingulata*; AM: *A. mertensii*. N = number of plants sampled, IR = incident radiation (%), WV = water volume extracted (mL), FPOM = fine particulate organic matter (mL after decantation in test-tubes), #Taxa = number of taxa per food web.
doi:10.1371/journal.pone.0071735.t001

Food Web Characterization

The diet of the various invertebrate species that make up food webs was determined by dissecting the entire guts. Twenty to 50 individual guts from each taxon were dissected across both localities and vegetation types in order to encompass the variability in diets of omnivores and predators. The gut contents of predators were placed into a drop of water on a glass slide, spread out, and analysed using a binocular microscope (Leica® MZ 12.5) and an Optiphot-2 Nikon® microscope whenever necessary. Most of the prey items could be identified from the guts of predators by comparing the chitinous parts (head capsules or legs, setae...) with specimens of bromeliad invertebrates archived in our collection (Univ. of Toulouse III). Only the gut contents of piercers (Heteroptera Veliidae, Diptera Tabanidae) could not be identified visually; in this case, we relied on observations of arranged encounters in test tubes.

The microorganisms found in the guts of detritivores were identified to a coarse taxonomic level using an Optiphot-2 Nikon microscope at x600 magnification. Determinations were based on earlier descriptions ([28,29]), and on recent microbiological samples (Carrias J-F, unpublished data). The presence of particulate organic matter was also recorded in the gut contents, and we used an ocular micrometer to distinguish fine particulate organic matter (FPOM, 1000 to 0.45 µm in size) from coarse particulate organic matter (CPOM, >1000 µm). Gut contents and other observations were used to build 365 interaction matrices (one per bromeliad) that described 365 “connectance webs” *sensu* [30] (see Fig. 1).

Three topological descriptors were used to describe food webs: connectance, linkage density, and nestedness. In the literature, two measures of the connectance are widely used: connectance and normalized connectance. Both refer to the proportion of all possible interactions that are realized, but normalized connectance is rather used for food webs comprising less than 20 species [1]. However, in this study, the two measures were strongly correlated (Pearson’s correlation coefficient = 0.966, $p < 0.0001$) and we therefore used the connectance as defined by [31]:

$$C = \frac{2 \times \text{Links}}{\text{Species} \times (\text{Species} - 1)}$$

Linkage density was calculated as the ratio between the number of links and the number of species. Levels of nestedness were expressed as matrix temperatures, with values ranging from 0 (perfectly nested) to 100 (see [32]) were calculated from the 365 interaction matrices (binary data) using the BINMATNEST3 program [33]. Departure from random was tested against a number of 1000 random matrices using the most conservative null model in BINMATNEST3 (namely “model 3”). A matrix temperature is affected by both its structure (the fill) and its size (the number of rows and columns). The extent to which the observed temperature departs from random depends on how random matrices are built. Unlike the null models 1 and 2, the model 3 is a packing algorithm that generates random matrices that are less influenced by the structure of the input matrix, thus allowing for comparisons with other studies [33]. More specifically, the null model 3 generates random matrices where each cell has a probability of containing a “1” equal to the averaged probability of occupancy of rows and columns of the input matrix. Therefore, the fill of each cell depends on both the fraction of “1s” in rows and columns in which the cell is included.

Data Analysis

To analyze the relationships between food-web descriptors, environmental variables and PPRs, we used linear mixed effect modeling. All variables were log-transformed to fit a normal distribution. Because a given vegetation type was found at only one locality, we qualified this structure as the “vegetation type nested within locality”. In contrast, a given bromeliad species could be sampled at many localities or many vegetation types, therefore the variable “bromeliad species” was not nested within the former or the latter. Since “locality”, “vegetation type”, “vegetation type nested within locality” or “bromeliad species” could be included as random factors, we performed a model selection based on the more conservative Bayesian Information Criterion (BIC) of the full models (models considering all environmental variables) for each dependent variable. Then, for a selected random factor (the remaining random effects being dropped from the model), the relationship between dependent and independent variables were explored using a stepwise backwards removal procedure and only the final models containing significant variables were presented. Departures from homoscedasticity and the normality of the residual errors were evaluated graphically for each final model. All statistical analyses were evaluated under a 95% confidence level and were conducted using R software V. 2.14.1 [34] and the associated packages for Linear Mixed Effect Modeling (nlme, lmer4).

Results

Abiotic Environments and Food-web Composition

There were large differences in WV and FPOM among bromeliad species, even at a given vegetation type (Table 1). The two bromeliads growing in rock savannah (*A. aquilega* and *C. berteroniana*) showed the highest and lowest mean values for FPOM (10.84 mL vs. 0.74 mL, respectively), and the highest and one of the lower mean values for WV (949.23 mL vs. 40.62 mL, respectively) compared to the others tank-bromeliad species. Thus, even though the understorey of primary forests consistently received lower incident radiation (i.e. *V. splendens* and *G. lingulata*) and higher litter inputs than sun-exposed areas, the amount of FPOM inside the tank of bromeliads from primary forests was not necessarily higher than those from rock savannah. However, bromeliads from primary forests had higher FPOM:WV ratios (0.05–0.09 mL FPOM/mL WV) than bromeliads from open areas (0.01–0.02 mL FPOM/mL WV).

Food webs comprised 8 to 20 taxa (median = 11 taxa), including macroinvertebrates, rotifers, and miscellaneous microorganisms (bacteria, cyanobacteria, fungi, algae, heterotrophic flagellates, and ciliates) identified in the gut contents. The predators belonged to the Diptera Culicidae (*Toxorhynchites purpureus*), Corethrellidae (*Corethrella* sp.) and Tabanidae, and to the Odonata (one unidentified Coenagrionidae species). Detritivores mostly consisted in Diptera Culicidae (*Culex* spp. and *Wyeomyia* spp.), Limoniinae, Chironomidae (*Tanytarsus* sp.) and Psychodidae (*Telmatoctopus* spp.). The list of taxa and the corresponding functional groups is provided in Fig. 2.

Model Selection and Influence of Abiotic and Biotic variables on Food Webs

All random factors were selected in at least one model (Table 2), suggesting that different spatial scales could provide complementary explanations to the observed pattern. More specifically, “vegetation type” and “bromeliad species” were included as random factors in the Simpson index (BIC = -314.16) and connectance models (BIC = -1108.648), respectively. “Locality”

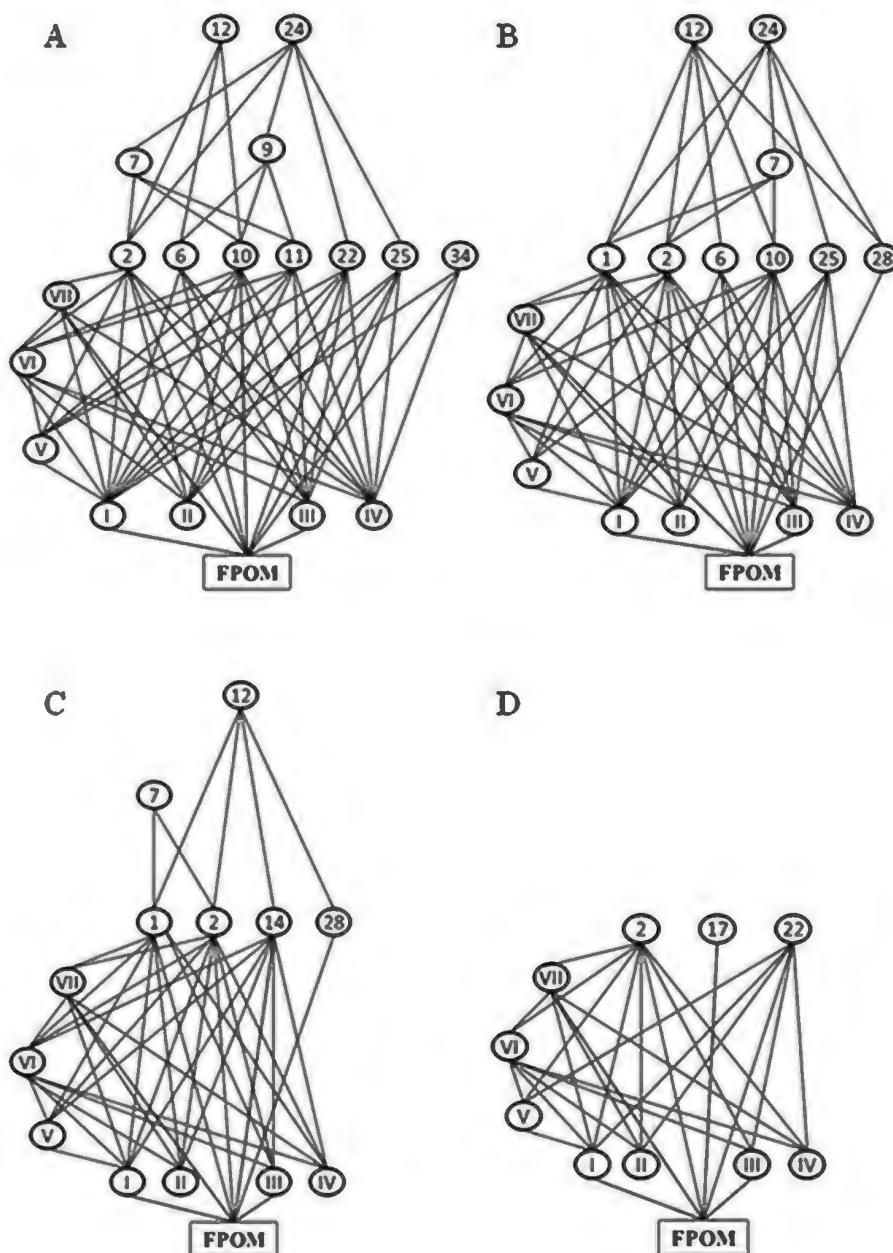


Figure 2. Examples of connectance webs under two contrasted environmental conditions: open areas (A, B) and forest areas (C, D). The upper trophic level (predators) is at the top of the graphs, and the lower (algae, detritus) at the bottom. Numbers and abbreviations are for: *Wyeomyia* spp. (1; filter-feeder); *Culex* spp. (2; filter-feeder); *Forcypomiinae* sp2 (6; filter-feeder); *Bezzia* sp. (7; predator); Tanypodinae (9; predator); Chironominae (10; detritivore); Tanytarsinii (11; detritivore); *Corethrella* sp. (12; predator); *Telmatoscopus* sp1 (14; detritivore); Limoniinae (17; detritivore-shredder); *Cyphon* sp. (22; shredder); Coenagrionidae (24; predator); *Aulophorus superterrenus* (25; detritivore); Hydracarina (28; detritivore); *Elpidium* sp. (34; detritivore-scraper); Bacteria (I); Cyanobacteria (II); Fungi (III); Algae (IV); Flagellate (V); Ciliate (VI); Rotifera (VII); Fine Particulate Organic Matter (FPOM).

doi:10.1371/journal.pone.0071735.g002

was selected in the Shannon index, linkage density and nestedness models (BIC = 21.83, BIC = -586.99, and BIC = -546.25, respectively). The subsequent stepwise backward removal procedures for each food-web descriptor were thus performed using their selected random factors. The computation of likelihood ratios demonstrated that we could not discriminate the model with "locality" from the one with "vegetation type" as random factor for the Shannon index model ($L = 0.034$, $p = 0.92$). Therefore, the

variable selection for the Shannon index model was also performed with "vegetation type" as random factor. In general, the inclusion of random factors with higher BIC values (e.g., "vegetation type", "vegetation type nested within locality" or "bromeliad species") only slightly affected the outcomes of the models in terms of significant and non-significant interactions among variables. More specifically, interactions that were barely

Table 2. Bayesian Information Criterion (BIC) of the full models with four different random effects for each food-web descriptor.

Models	Random effects			
	Vegetation type	Locality	Locality/Vegetation type	Bromeliad species
Shannon-Weaver's index	21.87	21.83	25.79	31.69
Simpson's index	-314.16	-310.39	-308.73	-310.19
Connectance	-1102.57	-1090.4	-1097.89	-1108.65
Linkage density	-584.88	-586.99	-582.41	-564.51
Nestedness	-544.09	-546.25	-541.21	-527.14

The random effect 'Locality/Vegetation type' means that the vegetation types are nested within the localities. Bold characters highlight the lowest BICs.
doi:10.1371/journal.pone.0071735.t002

significant with the selected random factor became non-significant when other random factors were included.

The Shannon and Simpson indices were both positively correlated with WV ($p < 0.0001$) and with PPR ($p = 0.0009$ and $p = 0.0027$, respectively; Table 3), showing that invertebrate community diversity increased with habitat size and the relative number of predators. Moreover, these two models showed a negative interaction between FPOM and PPR ($p = 0.036$ and $p = 0.047$, respectively). However, when "vegetation type" was included as random factor in the Shannon index model, both water volume and PPR remained significant but the FPOM:PPR interaction was not significant ($p = 0.081$). Last, the variable "IR" was not significantly correlated with the Shannon index, whatever the random factor included ($p = 0.6$ – 0.75). Finally, the two negative interactions IR:WV and PPR:FPOM become significant ($p = 0.046$ and $p = 0.014$, respectively) for the Simpson index when "Locality" was included as a random factor.

Connectance showed a negative and significant correlation with WV and PPR ($p = 0.0079$ and $p < 0.0001$, respectively), and there was a trend for decreasing connectance with increasing FPOM amount ($p = 0.039$; Table 3). Incident radiation was not significantly correlated with connectance ($p = 0.24$). When "Locality" was included, the two negative interactions IR:WV and WV:FPOM turned to be significant ($p = 0.0002$ and $p = 0.009$, respectively) in the connectance model. Both the linkage density and nestedness models showed a significant negative interaction between WV and IR ($p = 0.008$ and $p = 0.0071$, respectively). Linkage density and nestedness were positively correlated with WV and IR, and negatively correlated with PPR (Table 3). Linkage density was positively correlated with the IR:FPOM interaction ($p = 0.034$), but did not correlate with FPOM alone ($p = 0.064$). Finally, levels of nestedness (N) ranged from 21.54 to 49.79 (median $N = 39.98$). Overall, 44.4% of our interaction matrices were significantly different from the null matrices obtained at random ($p = 0.0001$ to 0.99 , median $p = 0.08$). Nevertheless, this percentage increased with the number of taxa; for instance, 66.1% and 88.8% of all of the interaction matrices comprising ≥ 13 and ≥ 17 taxa, respectively, were significantly different from random.

Discussion

Based on an unprecedented number of replicates under natural conditions, our models have proven informative in assessing whether environmental factors and biological interactions explain some patterns of food-web structure. Overall, variations in food-web structure across environments and spatial scales were mostly due to interactions between habitat size and the distribution of predators (Kendall's test, positive correlation between WV and

PPR, $\rho = 0.179$, $p < 0.0001$). There was however a trend for more feeding links in open areas compared to forest understorey. Finally, a core of generalists remained relatively constant across large regional scales, so that patterns of food-web connectance, linkage density and nestedness were mostly related to the addition or loss of predators.

Water volume was one of the most significant variables in all models, suggesting that habitat size was the primary factor controlling species composition and abundance patterns and, subsequently, food-web structure. Larger habitats are more easily colonized by immigrants, resulting in positive species-area relationships [35]. An increase in habitat area also fosters functional diversity [36], while allowing for a better partitioning of food resources by coexisting species [37]. Hence, at any given locality, both the number of taxa and individuals per plant increased with habitat size (there was a positive correlation between WV and PPR, Kendall's test = 0.179, $p < 0.0001$), which is consistent with previous findings on bromeliad ecosystems [21] and small wetlands in general [38]. However, there was a trend for bromeliad with higher incident radiation to accumulate more water (positive WV to IR relationship, $r = 0.32$, $p < 0.05$, log-transformed data) certainly because in open areas there are fewer overhanging trees to keep most of the rain from reaching the bromeliads. Indeed, based on measures made in the understorey of the primary forest in Petit-Saut, we estimated that 30–38% of the rain is intercepted by the canopy. Hence, for a given bromeliad species and size, the containers hold more water at sun-exposed areas. Nevertheless, the complex interaction between habitat size and incident radiation (significant interaction between IR and WV in the linkage density and nestedness models, but not in the connectance model) precludes a precise distinction of their relative impacts on food webs. We cautiously suggest an indirect effect where IR mediates food-web features through bromeliads' carrying capacity for aquatic invertebrates.

In aquatic ecosystems, detrital inputs form a strong trophic link between plant production, decomposer microorganisms, and larger metazoans [39]. In tank-bromeliads, detritus constitutes the main source of nutrients for the aquatic food web [14]. Debris-chewing invertebrates process incoming litter. Small particles, including faeces, are then washed into the plant pools where the FPOM is further processed in the gut of invertebrate collectors and filterers. Dead organisms, litter, and faecal particles, which collect in the leaf bases, are utilized by bacteria and other microorganisms. The nature and extent of the vegetation that surrounds these systems was therefore expected to have a strong influence on food-web structure through food quality and availability. On one hand, algae were found to account for more than 30% of the total microbial diversity and biomass in sun-exposed bromeliads

Table 3. Models evaluating the patterns of community diversity (Shannon's entropy and Simpson's evenness) and food-web structure (connectance, linkage density, nestedness) in relation to environmental variables and their interactions.

Fixed Effects	Estimate±SE	t-value	df	p	Random Effects
Shannon					Locality
Intercept	0.296±0.068	4.305	356	<0.0001	
Slope					
WV	0.091±0.011	7.72	356	<0.0001	
FPOM	0.044±0.025	1.773	356	0.077	
PPR	0.268±0.08	3.357	356	0.0009	
FPOM:PPR	-0.126±0.059	-2.103	356	0.036	
Simpson					Vegetation type
Intercept	0.226±0.036	6.157	352	<0.0001	
Slope					
WV	0.04±0.008	4.889	352	<0.0001	
FPOM	0.02±0.016	1.226	352	0.22	
PPR	0.15±0.049	3.025	352	0.0027	
FPOM:PPR	-0.074±0.037	-1.992	352	0.047	
Connectance					Bromeliad species
Intercept	0.60±0.017	35.018	355	<0.0001	
Slope					
WV	-0.007±0.002	-2.671	355	0.0079	
FPOM	-0.007±0.003	-2.07	355	0.039	
PPR	-0.14±0.009	-15.039	355	<0.0001	
Linkage Density					Locality
Intercept	1.136±0.102	11.117	354	<0.0001	
Slope					
IR	0.079±0.027	2.85	354	0.0046	
WV	0.141±0.029	4.77	354	<0.0001	
FPOM	-0.087±0.047	-1.854	354	0.064	
PPR	-0.186±0.019	-9.365	354	<0.0001	
IR:WV	-0.028±0.008	-3.4	354	0.008	
IR:FPOM	0.028±0.013	2.124	354	0.034	
Nestedness					Locality
Intercept	3.302±0.107	30.664	355	<0.0001	
Slope					
IR	0.087±0.029	3.014	355	0.0028	
WV	0.106±0.025	4.126	355	<0.0001	
FPOM	0.02±0.007	2.619	355	0.0092	
PPR	-0.237±0.021	-11.204		<0.0001	
IR:WV	-0.019±0.008	-2.708	355	0.0071	

IR = %incident radiation, WV = water volume (mL), FPOM = fine particulate organic matter (mL), PPR = Predator:Prey Ratio (see text). Only variables and interactions with $p < 0.05$ are interpreted as statistically significant and presented in the table.
doi:10.1371/journal.pone.0071735.t003

[17,19], which usually receive lower amounts of leaf litter [15]. We suggest that such a complementary “green” food web could contribute to reducing functional redundancy in FPOM-poor

systems. Overall, there was a positive correlation between WV and FPOM (Kendall's test = 0.349, $p < 0.0001$). However, higher FPOM concentrations (i.e. the amount of FPOM in relation to WV) decreased the amount of open water inside of the tank, which had a negative effect on aquatic organism diversity. High FPOM concentrations may have the effect of clogging habitats in small freshwater ecosystems and, to a certain extent, space availability [21]. Since FPOM was not consistently significant in our models, we assume that resource availability is not a limiting factor in bromeliad systems. However, the influence of resource quality (detritus *vs.* algae) on individual food apportionment should deserve more attention in future studies.

The PPR was a highly significant variable in all models. Connectance, linkage density and nestedness (matrix temperature) significantly decreased with increasing proportions of predators. A decrease in connectance and linkage density can be due to a gain of specialists and/or a loss of generalists [1]. In our study, predators were generalist species in that they fed on numerous prey species and multiple trophic levels, but they could be considered as “node specialists” since they interacted mainly with their prey while prey established links with both basal species (algae, rotifers, ciliates...) and predators. The extent to which predators are specialized on specific prey plays a great role in generating patterns of community diversity from site-to-site to large regional scales [40,41]. We found that the Shannon and Simpson indices (entropy and evenness, respectively) were positively correlated with the PPR at relatively small scales (from individual bromeliads to local scales), but overall, bromeliad food webs comprising less than 13 species were primarily composed of microorganisms and primary consumers, and were characterized by high levels of diffuse interactions where all species were more or less closely linked to each other. Indeed, a core of highly-connected species remained relatively constant at lower trophic levels (see also [42]), both in terms of species identity and ecological function, between individual bromeliads and from open to closed areas. For instance, the dominant generalist detritivores belonging to the Culicidae (*Wyeomyia* sp., *Culex* sp., and *Anopheles* sp.), Chironomidae and Oligochaeta were found in 310, 215 and 133 plants out of 365 respectively. The main top-predators, namely Corethrellidae, Ceratopogonidae and *Toxorhynchites* sp., occurred in 122, 105 and 83 plants respectively. It is likely that this highly-connected core ensures food-web stability and key ecosystem functions (e.g., decomposition, nutrient cycling) across environments. If detritivores share traits that determine responses to environmental changes in any given area (e.g., resistance to dessication, dispersal ability), then larger deviations in food-web structures can be expected following a major disturbance. Beyond 13 species, community diversity and food-web nestedness increased as a result of the addition of specialist predators and generalist detritivores. Our observations suggest that predators could enter the food webs at an average water volume of 49.9 mL, which corresponded to 1 predator species for 3 prey species on average. Regardless of the locality, larger bromeliad hosted a higher proportion of predators. In addition to their intrinsic value for biological diversity, predators (i.e., *Toxorhynchites* sp., *Corethrella* sp. and *Coenagrionidae*) thus played a key role in generating food-web patterns. This assumption is supported by the fact that, whilst predators are less frequent than detritivores, the predator:prey ratio was a significant variable in all three food-web topology models (connectance, linkage density and nestedness). We suggest that the addition of species at higher trophic levels contributes to departure from random interactions through the partition of prey resources between coexisting predators. Our results are also in line with studies which demonstrated that specialized species tend to be

dependent on a core of densely connected generalist species, and that natural communities display non-random interaction patterns [32,43–45]. Therefore, although we relied on unweighted trophic links to analyse spatial patterns in food-webs, this study provides strong empirical support on how food-web structure change along environmental and predator richness gradients. We conclude that at the bromeliad scale, habitat size determines the number of species that constitute food-web nodes and the proportion of predators (positive relationship), and subsequently, food-web topology. Hence, the number of species as well as the proportion of predators within assemblages globally decline from open to closed (forest) habitats, where the volume of water collected by bromeliads is generally lower because of rainfall interception by the canopy. The extent to which stability is related to linkage patterns (notably nestedness) in networks such as food webs remains however unclear and obviously deserves further research [46]. While linkage density and nestedness were lower in forested

areas, experimental research is now needed to confirm a trend for lower food-web stability in the understorey of primary forests.

Acknowledgments

We are grateful to the members of the *Laboratoire Environnement de Petit Saut* (HYDRECO) and to the members of the Nouragues Research Station for technical support and the use of their facilities. The English text was proofread by Andrea Yockey-Dejean and Kurtis Trzcinski. Two anonymous reviewers provided insightful comments on earlier version of this paper.

Author Contributions

Conceived and designed the experiments: RC BC CL AD J-FC. Performed the experiments: RC BC CL AD J-FC LP. Analyzed the data: OD. Contributed reagents/materials/analysis tools: RC BC J-FC AD CL LP. Wrote the paper: OD RC CL.

References

- Gilbert AJ (2009) Connectance indicates the robustness of food webs when subjected to species loss. *Ecol Indic* 9: 72–80.
- Rall BC, Guill C, Brose U (2008) Food-web connectance and predator interference dampen the paradox of enrichment. *Oikos* 117: 202–213.
- Kondoh M (2003) Foraging adaptation and the relationship between food-web complexity and stability. *Science* 299: 1388–1391.
- Fontaine C, Guimarães PR, Kéfi S, Loeuille N, Memmott J, et al. (2011) The ecological and evolutionary implications of merging different types of networks. *Ecol Lett* 14: 1170–1181.
- Strong DR, Lawton JH, Southwood TRE (1984) *Insects on Plants: Community Patterns and Mechanisms*. Oxford: Blackwell 313 pp.
- Dunne JA, Williams RJ, Martinez ND (2002) Food-web structure and network theory: the role of connectance and size. *Proc Natl Acad Sci USA* 99: 12917–12922.
- Townsend CR (1989) The patch dynamics concept of stream community ecology. *J N Am Benthol Soc* 8: 36–50.
- Declercq S, De Bie T, Ercken D, Hampel H, Schrijvers S, et al. (2006) Ecological characteristics of small farmland ponds: Associations with land use practices at multiple spatial scales. *Biol Conserv* 131: 523–532.
- Vannote RL, Minshall GW, Cummins KW, Sedell JR, Cushing CE (1980) The River Continuum Concept. *Can J Fish Aquat Sci* 37: 130–137.
- Compin A, Céréghino R (2007) Spatial patterns of macroinvertebrate functional feeding groups in streams in relation to physical variables and land-cover in Southwestern France. *Landscape Ecol* 22: 1215–1225.
- Srivastava DS, Bell T (2009) Reducing horizontal and vertical diversity in a food web triggers extinctions and impacts functions. *Ecol Lett* 12(10): 1016–1028.
- Paradise CJ (2004) Relationship of water and leaf litter variability to insects inhabiting treeholes. *J N Am Benthol Soc* 23: 793–805.
- Givnish TJ, Barfuss MH, Ee BV, Riina R, Schulte K, et al. (2011) Phylogeny, adaptive radiation, and historical biogeography in Bromeliaceae: Insights from an eight-locus plastid phylogeny. *Am J Bot* 98: 872–895.
- Benzing DH (2000) *Bromeliaceae: profile of an adaptive radiation*. Cambridge University Press, Cambridge.
- Brouard O, Céréghino R, Corbara B, Leroy C, Pélouzel L, et al. (2012) Understorey environments influence functional diversity in tank-bromeliad ecosystems. *Freshwater Biol* 57: 815–823.
- Romero GQ, Srivastava DS (2010) Food-web composition affects cross-ecosystem interactions and subsidies. *J Anim Ecol* 79: 1122–1131.
- Marino NAC, Guariento RB, Dib V, Azevedo FD, Farjalla VF (2011) Habitat size determine algae biomass in tank-bromeliads. *Hydrobiologia* 678: 191–199.
- Lounibos LP, Nishimura N, Escher RL (1993) Fitness of a treehole mosquito: influences of food type and predation. *Oikos* 66: 114–118.
- Brouard O, Lejeune A-H, Leroy C, Céréghino R, Roux O, et al. (2011) Are algae relevant to the detritus-based food web in tank-bromeliads? *PLoS One*, 6: e20129.
- Armbruster P, Hutchinson RA, Cotgreave P (2002) Factors influencing community structure in a South American tank bromeliad fauna. *Oikos* 96: 225–234.
- Jabiol J, Corbara B, Dejean A, Céréghino R (2009) Structure of aquatic insect communities in tank-bromeliads in a East-Amazonian rainforest in French Guiana. *Forest Ecol Manag* 257: 351–360.
- Céréghino R, Leroy C, Carrias J-F, Pélouzel L, Ségura C, et al. (2011) Ant-plant mutualisms promote functional diversity in phytotelm communities. *Funct Ecol* 25: 954–963.
- Leroy C, Corbara B, Dejean A, Céréghino R (2009) Ants mediate foliar structure and nitrogen acquisition in a tank-bromeliad. *New Phytol* 183: 1124–1133.
- Céréghino R, Leroy C, Dejean A, Corbara B (2010) Ants mediate the structure of phytotelm communities in an ant-garden bromeliad. *Ecology* 91: 1549–1556.
- Bongers F, Charles-Dominique P, Forget PM, Théry M (2001) *Dynamics and plant-animal interactions in a neotropical rainforest*. Kluwer Academic Publishers, Dordrecht, N.L.
- Frazer GW, Canham CD, Lertzman KP (1999) *Gap Light Analyzer (GLA) 2.0: Imaging software to extract canopy structure and gap light transmission indices from true-colour fisheye photographs: users manual and program documentation*. Simon Fraser University, Burnaby, British Columbia, and the Institute of Ecosystems Studies, Millbrook, New York.
- Jocqué M, Kernahan A, Nobes A, Williams C, Field R (2010) How effective are non-destructive sampling methods to assess aquatic invertebrate diversity in bromeliads? *Hydrobiologia* 649: 293–300.
- Carrias J-F, Cussac ME, Corbara B (2001) A preliminary study of freshwater protozoa in tank bromeliads. *J Trop Ecol* 17: 611–617.
- Carrias J-F, Brouard O, Leroy C, Céréghino R, Pélouzel L, et al. (2012) An ant-plant mutualism induces shifts in the protist community structure of a tank-bromeliad. *Basic Appl Ecol* 13: 698–705.
- Post DM (2002) The long and short food chain length. *Trends Ecol Evol* 17: 269–277.
- Morin PJ (1999) *Community ecology*. Blackwell Science, Inc. Malden, MA.
- Bascompte J, Jordano P, Melian CJ, Olesen JM (2003) The nested assembly of plant-animal mutualistic networks. *Proc Natl Acad Sci* 100: 9383–9387.
- Rodriguez-Gironés MA, Santamaría L (2006) A new algorithm to calculate the nestedness temperature of presence-absence matrices. *Journal of Biogeogr* 33: 924–935.
- R Development Core Team (2010) *R: A language and environment for statistical computing*. R Foundation for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0, URL <http://www.R-project.org/>.
- Drakare S, Lennon JJ, Hillebrand H (2006) The imprint of the geographical, evolutionary and ecological context on species-area relationships. *Ecol Lett* 9: 215–227.
- Takimoto G, Post DM (2012) Environmental determinants of food-chain length: a meta-analysis. *Ecol Res* 1–7.
- Richardson BA, Rogers C, Richardson MJ (2000) Nutrients, diversity, and community structure of two phytotelm systems in a lower montane forest, Puerto Rico. *Ecol Entomol* 25: 348–356.
- Oertli B, Auderset-Joye D, Castella E, Juge R, Cambin D, et al. (2002) Does size matter? The relationship between pond area and biodiversity. *Biol Conserv* 104: 59–70.
- Vondracek B, Blann KL, Cox CB, Nerbonne JF, Nerbonne BA, et al. (2005) Land use, spatial scale, and stream systems: Lessons from an agricultural region. *Environ Manage* 36: 775–791.
- Ryberg WA, Smith KG, Chase JM (2012) Predators alter the scaling of diversity in prey metacommunities. *Oikos* 121: 1995–2000.
- Chase JM, Biro EG, Ryberg WA, Smith KG (2009) Predators temper the relative importance of stochastic processes in the assembly of prey metacommunities. *Ecol Lett* 12(11): 1210–1218.
- Valladares G, Cagnolo L, Salvo A (2012) Forest fragmentation leads to food web contraction. *Oikos* 121: 299–305.
- Bascompte J (2009) Disentangling the web of life. *Science* 325: 416–419.
- Fontaine C, Thébault E, Dajoz I (2009) Are insect pollinators more generalist than insect herbivores? *Proc R Soc B-Biol Sci* 276: 3027–3033.
- Fox JW (2006) Current food web models cannot explain the overall topological structure of observed food webs. *Oikos* 115: 97–109.
- Tylianakis JM, Laliberté E, Nielsen A, Bascompte J (2010) Conservation of species interaction networks. *Biol Conserv* 143: 2270–2279.