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Aquatic invertebrate communities in tank bromeliads: how well do classic ecological patterns apply?

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- 18

19 Summary

Tank bromeliads (Bromeliaceae) often occur in high densities in the Neotropics and represent a key freshwater habitat in montane forests, housing quite complex invertebrate communities. We tested the extent to which there are species richness–altitude, richness–environment, richness– size, richness–habitat complexity and richness–isolation relationships for the aquatic invertebrate communities from 157 bromeliads in Cusuco National Park, Honduras.

25 We found that invertebrate species richness and abundance correlated most strongly, and 26 positively, with habitat size, which accounted for about a third of the variance in both. Apart from 27 bromeliad size (equivalent of the species-area relationship), we found remarkably little evidence 28 of classic biogeographic and ecological relationships with species richness in this system. 29 Community composition correlated with altitude, bromeliad size and position, though less than 30 20% of the variation was accounted for by the tested variables. The turnover component of 31 dissimilarity between the communities correlated with altitude, while the nestedness-resultant 32 component was related to bromeliad size. The unexplained variance could reflect a large 33 stochastic component in the system, associated with the ephemerality of the habitat patches (both 34 the plants themselves and the fluctuations in their water content) and stochasticity due to the 35 dispersal dynamics in the system.

We conclude that there is a small contribution of classic biogeographic factors to the diversity and community composition of aquatic invertebrates communities in bromeliads. This may be due to the highly dynamic nature of this system, with small patch sizes and high emigration rates. The patterns may mostly be driven by factors affecting colonization success.

40 41 Introduc

41 Introduction

42 Bromeliads (Bromeliaceae) are a characteristic component of Neotropical forests. Found from 43 ground level to high in the canopy, they contribute significantly to the habitat complexity 44 (Benzing, 2000), in particular for invertebrates. Bromeliads in a large subset of the family, called 45 tank bromeliads, are capable of holding considerable quantities of water in their leaf axils, 46 creating aquatic habitats that are inhabited by aquatic invertebrate communities (Fish, 1976; 47 Greeney, 2001; Frank and Lounibos, 2009). Tank bromeliads can occur in high densities, and, 48 based on their three-dimensional distribution in forests, may be the phytotelm (plant-held water 49 body) habitat occurring in the highest densities anywhere. For example, Sugden and Robins 50 (1979) recorded a mean density of 17.5 plants per square metre of ground area in a cloud forest in 51 Colombia. If the volume of water retained per plant is on average of the order of 100 cm³ (the 52 average for the bromeliads in our data), then such densities translate into tens of thousands of 53 litres of water available for colonization by aquatic animals, per hectare. In the absence of other 54 lentic water bodies, as is often the case in mountainous tropical forest areas, phytotelm habitats 55 provide an important freshwater habitat. The profusion of bromeliads, and their use as breeding 56 habitats by vectors for human diseases such as malaria and dengue, render bromeliads important 57 from a range of perspectives. In addition, bromeliads represent self-contained aquatic 58 communities for the aquatic stages of invertebrates, present naturally and at high replication, 59 making them potentially valuable as a study system for tackling prominent ecological and evolutionary questions (Srivastava et al., 2004). Well-defined aquatic communities occurring in 60 61 clusters are highly suitable for studying metacommunity dynamics (Leibold et al., 2004). Further, 62 bromeliads can be effectively imitated by artificial containers (Srivastava, 2006). These features allow easy manipulation and great flexibility in research design. 63

64

65 Despite the great advantages conferred by tank bromeliads, knowledge of their aquatic 66 invertebrate communities, and what structures them, remains limited—even though research on 67 aquatic invertebrates in phytotelmata dates back at least to 1915 (Picado, 1915; see also Laessle, 68 1961; Maguire, 1971; Frank and Lounibos, 1983; Kitching, 2000). Most studies on aquatic 69 invertebrates in bromeliads to date have focused on cataloguing species not previously known in 70 phytotelmata (e.g. Mendes et al., 2011). Recently, however, ecological studies have started to 71 contribute to the understanding of this habitat (e.g., Armbruster et al., 2002, Jabiol et al., 2009; 72 Brouard et al., 2011).

73

74 The highly dynamic nature of the system (the plants have limited life spans and there can be 75 considerable drying and wetting), and the wide environmental range in which bromeliads are 76 found, mean a lot of variability. This variability offers considerable opportunities but also 77 complicates the study of (invertebrate) community-structuring mechanisms. Additionally, the 78 mixing of terrestrial and (semi-)aquatic components of invertebrate communities hampers 79 straightforward interpretation of results. Earlier ecological bromeliad invertebrate studies included 80 both terrestrial and aquatic species, analyzing them as single communities (e.g., Cotgreave et al., 81 1993). Although terrestrial-aquatic links are present (Cereghino et al., 2011), the two components 82 are structured differently and should ideally be analysed separately if both groups are included in 83 the study. Tank bromeliads represent discrete habitat units for aquatic invertebrates, but less so for 84 most terrestrial invertebrates. A large proportion of the terrestrial invertebrates found in 85 bromeliads comprises occasional vagrants, increasing noise in analyses. Also, the very high 86 diversity of terrestrial invertebrates in tropical forests presents formidable challenges in terms of 87 identification, typically pushing the taxonomic resolution to ecologically less interesting levels 88 such as that of the family. Ecological studies aiming to unravel community structuring should take 89 these differences into consideration, and for all of these reasons, we restrict our analyses herein to 90 aquatic invertebrates.

91

92 Bromeliads can be seen as islands of aquatic habitat in a forest matrix and results from recent

93 studies indicate that these communities fit with at least one well-established biogeographic pattern

94 for islands: the species (richness)–area relationship. Jabiol et al. (2009) found that aquatic insect

95 richness and abundance were positively associated with water volume, a proxy for island/habitat

96 size. In studies considering both terrestrial and aquatic invertebrate communities, positive

97 correlations between invertebrate species richness and bromeliad size have been observed

- 98 (Armbruster et al., 2002; Montero et al., 2010). Apart from this, little is known about the extent to 99 which aquatic invertebrates in bromeliads follow classic island biogeographic patterns, such as the
- species-isolation and species-altitude relationships. Altitude, for example, is known to affect
- species-isolation and species-antitude relationships. Antitude, for example, is known to affect species richness in a wide range of taxa and habitats globally (Rahbek, 1995; McCain, 2007).
- Further, the extent to which these aquatic bromeliad communities are structured according to
- 103 classic ecological and biogeographic rules remains fragmentarily evaluated: environment (often
- 104 measured as productivity; Field et al., 2009) and habitat complexity (Hortal et al., 2009) are also

- 105 factors affecting species richness in many taxa around the world. For bromeliad invertebrate
- 106 communities, some studies point towards the importance of light and organic material (a proxy for
- productivity; Srivastava et al., 2008) in influencing community assembly (e.g. Dezerald et al.,
 2013). Habitat complexity, measured as the number of leaves, may affect the invertebrate system,
- as judged by results combining terrestrial and aquatic components (Armbruster et al., 2002).
- 110

111 Much island biogeography theory, including the classic 'equilibrium theory of island 112 biogeography' (ETIB, MacArthur and Wilson, 1967), is based on the idea that species richness on 113 an island (whether a true island or a habitat island) is the result of a dynamic equilibrium between 114 influx and local loss of organisms. Influx includes both colonization from an external source pool 115 and local addition through speciation. Loss of species may result from both emigration of 116 individuals and the deaths of individuals culminating in local extinction. The ETIB focuses on 117 immigration and local extinction, and not speciation (which we do not consider relevant for our 118 dataset, and do not consider further) or emigration. It also assumes some dispersal limitation from 119 the source pool to the island. Bromeliads, although easily recognised as insular habitat patches, 120 may have a differing relative importance of processes to those that underlie the ETIB and related 121 theories. During the aquatic phase of the invertebrates' life-cycles, dispersal limitation is likely to 122 be very strong, especially for the active dispersers, many of which actively avoid leaving the 123 bromeliads at this stage. However, once they have emerged as flying insects, the distances 124 between bromeliads may present almost no barrier to dispersal; instead, limitation may be mainly 125 due to their ability to locate suitable habitat, and this may favour colonization of large bromeliads 126 and those in clusters. In terms of species loss, although competition may play a role, we consider 127 predation (particularly for passive dispersers) and emigration (particularly for active dispersers) to 128 be far more important. It is of interest to ask whether both the different colonization dynamics and 129 the substitution of emigration and predation for demographic extinction are associated with 130 similar biogeographic patterns to those associated with islands more closely matching the 131 assumptions of the ETIB.

132

133 A long-term, time-series dataset would be ideal for analysing the processes associated with influx 134 and loss of species in bromeliads. However, investigating patterns of community composition 135 should yield interesting results that are informative about community assembly. For example, the 136 separation of species replacement (turnover) and species loss without replacement (nestedness) 137 when comparing communities, gives insight into community structuring factors (Baselga, 2010). 138 Nestedness of species assemblages—when the lists of species in species-poor communities are 139 subsets of those in species-rich communities-reflects a non-random limitation of species in a 140 community, which could be driven by habitat size or colonization limitation. Spatial turnover-141 the replacement of some species by others through space—could result from environmental 142 sorting or dispersal constraints. More generally, examining altitudinal and environmental 143 relationships addresses key aspects of biodiversity patterning (Rohde, 1992), reflecting the fact 144 that bromeliads represent independent replicates of aquatic invertebrate communities.

145

146 We investigate classic ecological and biogeographic relationships for a large sample of naturally 147 occurring bromeliad aquatic invertebrate communities in montane tropical forest, including cloud 148 forest, in Honduras. . We test the following specific predictions. (1) Larger bromeliads are 149 occupied by more species (equivalent to the species-area relationship). (2) More isolated 150 bromeliads (from other bromeliads) contain fewer species (species-isolation relationship). (3) 151 Bromeliads with more leaves contain more species (habitat heterogeneity hypothesis). (4) There is 152 a positive relationship between detritus content (productivity hypothesis) and species richness. (5) 153 There is a negative relationship between altitude and species richness. In addition, we analyze 154 ecological community structuring parameters in a metacommunity setting, again in relation to 155 size, isolation, altitude, habitat complexity and environmental variables. To gain a better insight 156 into the processes underlying tank bromeliad meta-community patterns and test how well these

- 157 communities follow island biogeographic patterns, we include an analysis of the dissimilarity
- 158 between the component communities (beta diversity), partitioning it into turnover and nestedness-
- 159 resultant components (Baselga, 2010).
- 160

161 **Material and Methods**

- 162 Field site
- 163 The bromeliad sampling took place from June to August 2006 and 2007 in Cusuco National Park
- (CNP), situated in the Merendon Mountain range in north-western Honduras. The core zone of the 164
- 165 park consists of lower montane tropical rain forest (a mix of primary and secondary), with patches
- of primary cloud forest and upper montane rain forest characterized by high densities of 166
- 167 bromeliads. We collected 157 bromeliads from five main sampling areas (centred on the field
- 168 camps 'Base Camp', 'El Cortecito', 'Guanales', 'El Danto' and 'Cantiles'). For a detailed
- 169 description of the area and permanent sampling lines, see Field and Long (2007).
- 170
- 171 Sampling protocol
- 172 In order to minimize the influences of physical structure and possible biochemical differences
- 173 between species, we only sampled individuals of *Tillandsia guatemalensis* Smith. This is one of
- 174 the more abundant bromeliad species in CNP, with enough water to accommodate aquatic
- 175 invertebrate communities. We sampled invertebrate communities completely, by dismantling each
- 176 bromeliad, leaf by leaf. For a fuller description of the sampling protocol, including the
- 177 randomization procedure, see Jocque et al. (2010a). We only sampled bromeliads large enough to
- 178 contain water, which translated into minimum leaf spread of 18 cm, all but four being 20 cm or
- 179 larger.
- 180

181 Before sampling each bromeliad, we recorded a range of environmental variables: altitude

- 182 ("altitude", in metres above sea level), attachment height on the tree ("attach"), the width
- ("width") and height ("height") of the bromeliad, the number of other bromeliads within a two-183
- 184 metre radius ("R-Brom"), the amount of light ("light", openness of the canopy, ten-point scale) 185 and the openness to receive water from precipitation ("rain", effectively an inverse shelter
- 186 measure, ten-point scale). The attachment height on the tree was measured as the shortest distance
- 187 in cm between the forest floor and the underside of the bromeliad core. The bromeliad core is the
- 188 central axis where all leaf bases join, and is also the origin of the roots (or 'holdfasts' because 189 they are only used for attachment). For plant width and height we measured from the point of
- 190 water catchment on the leaves: leaves are angled upwards from the core of the plant until they
- 191 (particularly outer leaves) bend downwards from the weight of the leaf. Up to this point of
- 192 bending downwards, the water intercepted by the leaves runs into the leaf axils; beyond that point
- 193 most water does not run into the tank of the bromeliad. We measured the width of the plant as the
- 194 largest horizontal distance between the water catchments points of two opposing leaves (cm). We
- 195 measured the height of the plant from the base of the bromeliad core to the highest water 196 catchment point (cm). We counted the number of other bromeliads on the same tree as the
- 197 sampled bromeliad, and also on the trees within a two-metre radius. In 2007 we additionally
- 198 stratified sampling of bromeliads, into two types: individuals that were the only bromeliad
- 199 attached to the tree (SOLO) and individuals that were one of at least two bromeliads attached to
- 200 the same tree (MULTI). In MULTI, we targeted bromeliads that were underneath others on the
- 201 tree, thus allowing dispersal into them via water flow from other bromeliads. This was mostly geared towards the dispersal of the passive dispersers (Ostracoda and Anomopoda).
- 202
- 203
- 204 After the measurement of these variables in the field, we placed each sampled bromeliad in a
- 205 bucket and transported it to the nearest camp for immediate processing. We collected the water 206 contained and then took the plant apart, leaf by leaf, rinsing every leaf with 64 µm-filtered river
- 207 water. During this process, we measured additional variables: amount of water held by the plant
- 208 (ml), the total number of leaves, fresh weight of the cleaned plant (g) and circumference of the

- 209 core (mm). The total number of leaves comprised all the green leaves and the leaves with at least
- 210 the base still green. All washed parts of the plant were weighed with a 500 g Pesola spring meter,
- 211 once excess water had been removed. We measured the circumference (mm) of the bromeliad
- 212 core after removing the leaves. Animals were picked out alive from the rinsing water, and fixated 212 is 70% of -100 km s⁻¹ km
- in 70% ethanol. After removing all the invertebrates, we manually removed the larger organic
 debris and then filtered the rest using a 22 µm sieve, to determine detritus content (g). We
- debris and then filtered the rest using a 22 µm sieve, to determine detritus content (g). We
 processed the invertebrates in a laboratory using an OLYMPUS SZX-12 stereomicroscope and
- 216 identified all to morphospecies; full species identification (and description, in some cases, e.g.
- 217 Mendes et al., 2011) is ongoing and gives us confidence in the matching of our morphospecies to
- 218 known species.
- 219
- 220 Statistical analyses

221 To test predictions 1–5 we first examined linear correlations between all variables. When needed 222 to remove skew and normalize the errors associated with best-fit lines, we either square-root or 223 log transformed variables for further analysis. We used regression to determine the individual 224 contributions of the recorded variables relevant to predictions 1–5, in accounting for the variation 225 in both richness and total abundance. To assess whether any improvement could be made on the 226 simple model for species richness resulting from this exploratory analysis, we used multi-model 227 inference. This ran 16,383 regression models, comparing all against each other using Akaike's 228 Information Criterion (AICc). Many of the potential explanatory variables were highly collinear, 229 particularly measures of bromeliad size and measures of bromeliad position. We therefore ran 230 principal components analyses (correlation method) of these two groups of variables, to create 231 two orthogonal principal components of each phenomenon; in doing so, we square root-

- transformed most of the variables, to reduce or remove skew.
- 233

Complementary to the correlations with richness and abundance to test predictions 1-5 we
analysed community composition using both ordination and analysis of beta diversity. In
ordination, the choice of linear or unimodal analysis methods is traditionally based on the amount
of variation present in the dataset, reflected as the length of the environmental gradient. Because

- the environmental gradient in our dataset was less than four, we opted for the linear response (ter
- Braak and Smilauer, 2002). We used the linear direct analysis (RDA) with forward selection
- based on 999 Monte Carlo permutations to build a model. We square root-transformed theabundance data to reduce the impact of high abundances. We standardized species abundances
- abundance data to reduce the impact of high abundances. We standardized species abundances(dividing them by the standard deviation of values) to focus on community composition. We also
- removed rare species (defined as only 1 or 2 individuals in the total dataset) from the analysis;
- these were three beetle species, a chironomid, a culicid, two Diptera and a copepod species. We
- included all the measured variables in the initial analyses and, using a forward selection
- 246 procedure, isolated the factors accounting for the most variance in the dataset.
- 247

248 When examining for possible effects of altitude (prediction 5), as well as correlating diversity 249 with the continuous altitude data, we looked for patterns in diversity in altitude categories. The 250 altitudinal range was from 1347 m to 2084 m, but samples were not equally spread over all altitudes. We used four categories: <1500; 1500-1600; 1600-1900; >1900. Based on the lowest 251 252 number of bromeliads sampled in a category (18 below 1500 m), we reduced all the other groups 253 to 20 bromeliads, selecting bromeliads randomly. We then used Kruskal–Wallis tests to test for 254 any differences between the altitudinal categories, for Shannon, Simpson and Margalef diversity, 255 average species richness, dominance Index, evenness and total richness.

- 256
- 257 To gain insight into the underlying metacommunity structuring processes, we partitioned beta
- diversity following the method of Baselga (2010). Splitting overall beta diversity into its (spatial)
- turnover and nestedness components allows the identification of species replacement or species
- loss, respectively, as driving factors in community assembly (Baselga, 2010). We used the

- 261 'betapart' package in R. This calculates the pair-wise Sorensen dissimilarities between all the
- bromeliad communities (overall dissimilarity or 'beta diversity'), and partitions that into its
 turnover (Simpson dissimilarity) and nestedness-resultant components. The part of the Sorensen
- dissimilarity that is due to nestedness rather than turnover is simply the difference between the
- 265 Sorensen and Simpson dissimilarity measures. We repeated this analysis using Jaccard
- 266 dissimilarity measures, but because both gave qualitatively identical results in all cases, we focus
- 267 mainly on the Sorensen–Simpson method. We thus obtained a series of six distance matrices (of
- 268 pairwise dissimilarities: Sorensen, Simpson, nestedness (Sorensen minus Simpson), Jaccard,
- 269 Jaccard turnover, Jaccard nestedness), each of which we then correlated with a matrix of
- 270 geographic distances between the bromeliads, using Mantel tests in the R package 'vegan', with
- 999 permutations to determine significance. We repeated this for other types of environmental
- distance, focusing on correlating the six sets of community dissimilarities with pairwisedifferences in altitude, bromeliad size and bromeliad position.
- 273

For statistical analyses we used R (Rstudio, Inc. 2012), STATISTICA (StatSoft, Inc. 2012) and SAM (Spatial Analysis in Macroecology, Rangel et al., 2006).

277278 Results

For this study we recognized 42 (morpho)species (Table 1). Ongoing determinations have resulted in a more conservative identification of the recognized morphospecies, with reductions of the numbers of species in the Chironomidae (2), Tipulidae (2), Culicidae (5), Syrphidae (4) and the additions of a Psychodidae species and a copepod, compared to a previous study on the same bromeliads (Jocque et al., 2010a).

284

285 The correlation matrix of the measured variables (Table 2) suggests no correlation between species richness or total invertebrate abundance and detritus content (refuting prediction 4), 286 287 altitude (refuting prediction 5), attachment height or the number of nearby bromeliads 288 (inconsistent with prediction 2). It also indicates two blocks of correlated explanatory variables. 289 The first is all factors associated with phytotelm size and complexity: weight, width, height, core 290 diameter, water content, detritus content and number of leaves. Weight was by far the strongest 291 correlate (r = 0.95) of the first axis of the principal components analysis of these variables, this 292 axis accounting for 62% of the variation in the data. The second block of correlated variables 293 includes factors associated with the positioning of the bromeliad in the environment: the openness 294 to light and rainfall and the number of other bromeliads nearby (Table 2). From the positional 295 variables in the second group, the estimated exposure to light and rainfall were the only variables 296 correlating significantly with species richness, but each only accounted for 4% of the variance in 297 species richness. 298

299 Of all the putative explanatory variables, the total fresh weight of the leaves ('weight') correlated 300 most strongly with both species richness and the total number of individuals (abundance) in the 301 bromeliads (Table 2). This supports prediction 1. Log-transformed weight accounted for slightly more of the variation in richness (r = 0.58, $r^2 = 0.33$) than the untransformed (Figure 1). In a 302 303 partial regression using log(weight) and the number of leaves as explanatory variables, while 11% 304 of the variation in species richness was accounted for uniquely by bromeliad weight, only 0.4% 305 was uniquely accounted for by the number of leaves and this contribution was not significant; 306 shared explained variance was 22%. Very similar results were obtained when analysing total 307 abundance of invertebrates, rather than species richness. This is inconsistent with prediction 3. 308 The correlation between species richness and log(weight) was also stronger than that between 309 species richness and the first axis from the principal components analysis on all the size variables. 310 Further, no combination of explanatory variables improved on log(weight) alone, in accounting 311 for variation in species richness, as judged by Akaike's Information Criterion (AICc) in multi-312 model inference. This supports prediction 1 and is inconsistent with predictions 2–5.

313

314 A large number of recorded environmental variables together accounted for a relatively small 315 proportion of the community composition in our dataset. The forward selection of the linear 316 redundancy analyses (RDA) isolated altitude, water, number of leaves, total weight of the bromeliad, exposure to precipitation, the total number of bromeliads, detritus content and 317 318 attachment height of the bromeliads as the strongest explanatory variables (Figure 2), in total 319 accounting for 20% variation of the dataset (sum of all canonical eigenvalues = 0.200, F = 4.563, 320 *p-value* =0.001). In the biplot of species and environmental variables, three groups of variables 321 can be distinguished, with particular species associated with them (Figure 2). One comprises 322 variables measuring the size and complexity of the bromeliad habitat (number of leaves, detritus 323 content, plant weight and the water content). Most of the species associated with variation in these 324 variables are Diptera. The second group contains two position variables: attachment height of the 325 bromeliad on the tree and the exposure to rainfall. Most strongly associated with these variables 326 are passive dispersers (the two ostracod species and the two water fleas). The final group is only 327 altitude, which is most associated with several species of Diptera.

328

329 The results of the beta diversity partitioning are shown in Table 3 and nicely complement the 330 ordinations. The dissimilarity in species composition between bromeliads was positively 331 correlated with both the difference in altitude and the difference in size between bromeliads. It 332 was the turnover component that correlated with altitude and the nestedness-resultant component 333 that correlated with bromeliad size. Thus, invertebrate species tended to replace each other along 334 the altitudinal gradient, while dissimilarity related to bromeliad size was due to smaller 335 bromeliads tending to contain a subset of the invertebrate species found in larger ones. These 336 relationships were quite weak, but strongly significant (Table 3). The correlations with geographic 337 distance were qualitatively identical to those with altitudinal distance, but were quantitatively 338 much weaker and less significant, suggesting that the trend with geographic distance was an 339 indirect result of the altitudinal relationship. Indeed, using partial Mantel tests, when controlling 340 for altitudinal differences no significant correlations between dissimilarity and geographic 341 distance remained, while the correlations with altitude remained almost unchanged when 342 controlling for geographic distance. This suggests no effect of geographic distance in our study 343 system. 344

- Based on the selection of altitude in the multivariate analysis, its importance in the beta diversity analysis, but its lack of significance in the species richness analysis, we examined possible altitudinal patterns in various measures of species diversity, using our four altitude categories (see Methods). Again refuting prediction 5, we found no significant differences between the categories in any of species richness (H = 1.345, P = 0.718), total abundance (H = 0.273, P = 0.435), Shannon–Weiner (H = 2.85, P = 0.415), Margalef (H = 1.13, P = 0.770) or evenness (H = 4.76, P= 0.190).
- 352

359

353 Finally, species richness did not differ (F = 0.005, p = 0.941) between isolated phytotelmata

- (SOLO) and phytotelmata occurring in clusters (MULTI) (prediction 2). Nor did community
 composition differ between the two, as judged by RDA analysis, either with SOLO versus
- composition differ between the two, as judged by RDA analysis, either with SOLO versus MULTI as the single explaining variable (total sum of the eigenvalues = 0.015, F = 0.774, P =
- 0.687) or additionally with the different sampling locations as a covariable (total sum of the
- 358 eigenvalues = 0.015, F = 0.888, P = 0.554).

360 Discussion

361 With the exception of the influence of habitat patch size, there is remarkably little evidence of

- 362 classic biogeographic and environmental relationships affecting the diversity in the aquatic
- 363 invertebrate bromeliad system. These observations support the notion that bromeliads, although
- 364 easily recognised as an insular habitat system (eg Richardson 1999), do not follow the classic

365 island biogeography theory in a strict sense. After the publication of the ETIB in 1967, the 366 concept was applied to a wide diversity of habitats with insular characteristics. This was further 367 reinforced by application of the metacommunity concept (see Leibold et al. 2004), a conceptual 368 framework based on well delineated communities. The small community size, temporal instability 369 and highly dynamic colonisation and emigration of most of its inhabitants position the bromeliad 370 phytotelm rather near one extreme of a habitat continuum to which the theory could be applied. In 371 particular the high emigration rate (associated with the emergence of the insect larvae) sets this 372 habitat apart from most other island habitats to which island biogeographic theories (generally) 373 are applied. The dynamics of the insect-dominated invertebrate communities leave little room for 374 extinction or speciation to play significant roles in individual bromeliad communities, and their 375 richness is probably due largely to factors determining the colonisation of bromeliads. The 376 dynamics of the passively dispersing members in these communities are expected to be quite 377 different. Previous research on Ostracoda in Jamaican bromeliads (Little and Hebert 1996) 378 recorded a high diversification rate in bromeliads, most probably due to the limited dispersal 379 between bromeliad clusters of these organisms. This subset of the bromeliad invertebrate 380 communities could lean more towards the classic ETIB. In our system this was difficult to test

- 381 because of the limited occurrence of microcrustaceans in the sampled bromeliads.
- 382

383 The species-area relationship is the most pervasive of the classic relationships assessed in this 384 study, and it was by far the strongest physical or geographic determinant of either species richness 385 or abundance in the invertebrate communities we analyzed. The size of the bromeliad (600 g 386 range in fresh weight) was positively related to the number of species in the community (Figure 387 1), a semi-log relationship typical of a species-area curve. Size accounted for about one third of 388 the variance in species richness. This is consistent with prediction 1 from the biogeographic 389 theory. The strong inter-correlation of the variables measuring bromeliad size suggests that most 390 of those could be used reliably to quantify habitat size. About a quarter of the invertebrate species 391 in our data tended to be found more in larger bromeliads; in most cases this remained true after

- 392 accounting for the number of species in the community (by regressing abundance on bromeliad
- 393 size with species richness as a covariate, on a species-by-species basis). This is also reflected in
- the relationship between bromeliad size and the nestedness component of communitydissimilarity: smaller bromeliads tend to lack some species found in larger ones.
- 396

397 The positive relationship between phytotelm size and community size or structure supports 398 previous findings, both for bromeliad communities specifically (Richardson 1999) and for aquatic 399 communities more generally (e.g. Srivastava and Lawton, 1998; Kitching, 2000; Armbruster et 400 al., 2002; Frank et al., 2004). Mechanistic interpretation of this association, however, is not 401 straightforward because habitat size is strongly collinear with various likely influences, as is 402 usually the case. Mechanistic elements associated with habitat size that are often thought to affect 403 community size and structure include a larger target for dispersing individuals or any component 404 of size that positively influences colonisation, larger populations and thus smaller extinction 405 probability, and greater habitat complexity or diversity (e.g. Hortal et al., 2009).

406

407 For invertebrate communities inhabiting bromeliads, the number of leaves is often used as an 408 indicator for the complexity or diversity of the bromeliad habitat (Srivastava et al., 2006). The 409 aquatic habitat in the phytotelm is composed of many small compartments, associated with the 410 individual leaves, arranged in a spiral, and one relatively large central compartment at the centre. 411 These compartments are isolated in the sense that they collect their own water and organic debris 412 but are all connected in that most aquatic organisms can move from one leaf-axil compartment to 413 the next. Aquatic invertebrate species in bromeliads such as mosquito larvae partition space in 414 bromeliads to co-exist (Gilbert et al. 2008) and a more complex habitat is expected to be able to 415 house more diverse communities. In our data, while the number of leaves did correlate positively

416 with species richness (Table 2), in partial regression this variable did not significantly add to the

- 417 variation accounted for by weight, while weight added a lot to the variation accounted for by the
- 418 number of leaves. Very similar results were obtained when analyzing total abundance of
- 419 invertebrates, rather than species richness. These results suggest that habitat size, rather than
- 420 habitat complexity, is what matters for the size and structure of the aquatic invertebrates living in
- 421 the bromeliads—consistent with prediction 1 but not prediction 3.
- 422

423 While the diversity in larger habitats can be a direct result of better survival of populations, with the 424 larger habitat size allowing larger populations, which tend to persist longer, it is more plausible that 425 the driving factors behind community structure in bromeliads is associated with the factors 426 affecting the colonisation of the habitat patch. Larger aquatic habitats may have higher immigration 427 because they represent a larger target (Dodson, 1992) or there might be active selection, whereby 428 individuals select the larger habitat patches because this increases survival chances—a behaviour 429 observed in other aquatic invertebrates (Binckley and Resetarits, 2005) and also for Odonata in 430 bromeliads (Srivastava et al. 2008). Also the exposure time to colonisation could play a role. Larger 431 bromeliads are typically older ones, available for colonisation longer. However, the highly dynamic 432 nature of aquatic invertebrate communities in bromeliads, with most larval stages emerging and 433 emigrating, and thus communities being reassembled frequently, may decrease the effect of longer 434 exposure to colonisation. Older bromeliads are also usually the ones with more leaves, and this did

- 435 not add to size in accounting for species richness or abundance.
- 436

437 The strong colonization–emigration dynamics also suggest a strong rescue effect (Brown and 438 Kodric-Brown, 1977). However, the strongest correlate of species richness in our dataset was total 439 abundance (r = 0.71 using log(abundance)), which was related primarily to bromeliad size. The 440 population size mechanism may operate partly through a sampling effect, with more colonizing 441 individuals representing more species by neutral or random assembly. We suggest that this might 442 operate in combination with a preference for larger bromeliads, disproportionally decreasing the 443 immigration to smaller bromeliads.

444

445 Inconsistent with prediction 2 (species richness-isolation relationship), we found no significant 446 spatial structure in our species richness data. On a very local scale, positioning of the phytotelm 447 (relative to water and resource inputs and other bromeliads) had little or no relationship with the 448 overall invertebrate community, but was relevant for the passive dispersers (Crustacea). The 449 number of bromeliads in the cluster (within 2 m) affected the community composition but there 450 was no effect of presence/absence of other bromeliads on the same tree. On a larger spatial scale, 451 geographic distance (9 km range) appeared to have no effect on species richness or community 452 composition.

453

454 Altitude was not correlated with species richness or abundance, inconsistent with prediction 5. This 455 was despite the 800-m altitudinal range sampled, which in ecological terms is very significant: from 456 the lowest occurrences of bromeliads (in numbers greater than the odd scattered individual) to the 457 upper montane dwarf forest at the highest altitude in the study area. Other overall diversity-related 458 parameters (species diversity, evenness) did not vary significantly with altitude, nor could we detect 459 the mid-altitude diversity bulge that occurs commonly in altitudinal studies (Rahbek, 1995, 2005). 460 Currently, consensus on the mechanisms driving this pattern remains elusive, but climatic variables 461 and an area effect are dominant elements in the discussion (Grytnes & McCain, 2007; McCain, 462 2007; Karger et al., 2011). In our study area, both climate and area change markedly with altitude area because the mountains are approximately conical in shape. Temperature and evaporation 463 464 decrease with altitude, while humidity increases; the tops of the mountains in Cusuco National Park 465 are typically in cloud. The north-west side of the mountain range, facing the Altantic Ocean, 466 receives a particularly large amount of rainfall and has higher air humidity. Yet we found no 467 significant relationship between altitude and species richness.

468

469 Altitude was, however, associated with beta diversity and community composition, with the

470 turnover component of community dissimilarity tending to increase with greater differences in

471 altitude between host bromeliads. This corresponded with a tendency (usually weak) for some of

the invertebrate species to occur primarily at either relatively low (some Dipteran larvae and aColeopteran) or relatively high altitudes (some Chironomid and Culicid larvae). Thus, although

474 the assemblage-level patterns seem invariant to altitude, some species replacement is evident.

475

476 Some studies on aquatic invertebrates suggest that bromeliad-specific local environment affects 477 insect communities (Ngai et al. 2008), in particular the availability of resources (Srivastava et al. 478 2008) (prediction 4). Bromeliads obtain their nutrients from the decomposition of organic material 479 that falls in the bromeliad (Richardson, 1999). The decomposing detritus is the main source of 480 nitrogen for epiphytic bromeliads (Reich et al., 2003), at least for shaded ones. Recent studies 481 show that food webs in exposed phytotelmata are driven by primary production; Srivastava et al. 482 (2008) found detrital mass to be a strong predictor of species richness in bromeliads. In our data, 483 although detritus content correlated strongly with bromeliad size (r = 0.87), its correlation with 484 total invertebrate abundance was much weaker (r = 0.44). Indeed, detritus content added nothing 485 to bromeliad size in accounting for variation in total abundance in a partial regression, and the 486 same was true when modelling species richness. The same was also true when adding any variable 487 related to bromeliad position (e.g. openness to light or rainfall input) to bromeliad size, in 488 regressions to account for variation in total abundance or species richness. We thus found no 489 evidence to support the idea that resource input is causing variation in community size or

490 structure, though other measures of resource input might provide such evidence.

491

492 Overall, surprisingly few variables had significant explanatory power and the overall variance in 493 community composition accounted for, using the measured physical and geographic variables in 494 this study, was relatively low (around 20%). This may be partly due to environmental factors not 495 recorded in this study. Possible candidates include primary production by bacteria and algae 496 within the bromeliads, though we would expect such an influence to be reflected in our detritus 497 variable. More meaningful representation of the positioning of the bromeliad plants in relation to 498 the forest canopy may provide some explanatory power. Even so, there is probably a large 499 stochastic component present in the system. Bromeliad-held aquatic communities may be highly 500 dynamic because of the ephemerality of the habitat patches, both the plants themselves and their 501 water content (which fluctuates, including a seasonal component). Most of the invertebrates 502 remain for a relatively short time, after which they emerge and emigrate from the phytotelm. 503 When dispersing, most inhabitants are therefore highly mobile, either flying or using dispersal 504 vectors that are highly mobile (Lopez et al., 2002). The importance of stochastic elements and the 505 absence of strong short-distance dispersal limitation are in keeping the lack of any differences in 506 diversity or community composition between bromeliads on trees host to no other bromeliads, and 507 those that are found in clusters—even for the passively dispersing species. Chance and stochastic 508 effects may therefore play a large role in these systems.

509

510 The low explanatory power of altitude and other measured environmental variables may also reflect 511 a lack of meaningful differences in habitat within the bromeliads, for the invertebrates inhabiting 512 them. Water temperature in bromeliads does vary with altitude, but also fluctuates considerably 513 both seasonally and on a daily basis (Jocque and Kolby, 2012). Such an environmental regime may 514 favour generalist species (Jocque et al., 2010b), rendering the environmental differences observed 515 in the study rather small for the study organisms. These considerations are important because the 516 system of bromeliad-held aquatic invertebrate communities appears to be an exception to the 517 dominant pattern of strong diversity and community-composition changes with altitude (prediction 518 5). Further, given the strongly increasing density of tank bromeliads with altitude (in the study area, 519 at least; Jocque et al., 2010a), the usual negative relationship between altitude and habitat area is 520 broken. That is, while mountains are typically conical, giving lower area of higher altitudinal bands,

- 521 the counteracting effect of increasing bromeliad density with altitude may cancel out, or perhaps
- 522 even reverse, the usual trend.
- 523
- 524 In conclusion, most classic biogeographic and ecological relationships appear not to apply to
- 525 aquatic invertebrate communities inhabiting bromeliads, perhaps because they are such a highly
- 526 dynamic habitat system characterized by small patch sizes. Only the species-area relationship was
- 527 strongly supported, and even this may have been mostly driven by factors affecting colonization.
- 528 Aquatic metacommunities in bromeliads (and other phytotelmata) may be highly suited to
- 529 research on the effects of colonization sequence and immigration rate on the stability of
- 530 community composition. Particularly interesting here would be to investigate priority effects (De
- 531 Meester et al., 2002), habitat selection and fixed colonization sequences.
- 532

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- 541

542 **References**

- Armbruster, P., R. A. Hutchinson & P. Cotgreave, 2002. Factors influencing community structure in a South
 American tank bromeliad fauna. Oikos 96: 225–234.
- Baselga, A., 2010. Partitioning the turnover and nestedness components of beta diversity. Global Ecology and
 Biogeography 19: 134–143.
- 547 Binckley, C. A. & W. J. Resetarits, 2005. Habitat selection determines abundance, richness and species
 548 composition of beetles in aquatic communities. Biology Letters 1: 370–374.
- 549 Brouard, O., A.H. Le Jeune, C. Leroy, R. Cereghino, O. Roux, L. Pelozuelo, A. Dejean, B. Corbara & J.F.
 550 Carrias, 2011. Are algae relevant to the detritus-based food web in tank-bromeliads? PLoS ONE 6, e20129. doi: 10.1371/journal.pone.0020129.
- Brown, J.H. & A. Kodric-Brown, 1977. Turnover rates in insular biogeography effect of immigration on
 extinction. Ecology 58: 445–449.
- Cereghino, R., C. Leroy, J. F. Carrias, L. Pelozuelo, C. Segura, C. Bosc, A. Dejean & B. Corbara, 2011. Ant–
 plant mutualisms promote functional diversity in phytotelm communities. Functional Ecology 25: 954–
 963.
- Cotgreave, P., M. J. Hill & D. A. J. Middleton, 1993. The Relationship between body-size and population-size in
 bromeliad tank faunas. Biological Journal of the Linnean Society 49: 367–380.
- De Meester, L., A. Gomez, B. Okamura & K. Schwenk, 2002. The Monopolization Hypothesis and the
 dispersal-gene flow paradox in aquatic organisms. Acta Oecologica-International Journal of Ecology 23:
 121–135.
- 562 Dézerald O., T. Stanislas, C. Leroy, J. Carrias, B. Corbara, A. Dejean & R. Céréghino, 2013. Environmental
 563 determinants of macroinvertebrate diversity in small water bodies: insights from tank-bromeliads.
 564 Hydrobiologia 1-10.
- 565 Dodson, S., 1992. Predicting Crustacean Zooplankton Species Richness. Limnology and Oceanography 37: 848–
 566 856.
- Field, R., B.A. Hawkins, H.V. Cornell, D. J. Currie, J. A. F. Diniz-Filho, J.-F. Guégan, D. M. Kaufman, J. T.
 Kerr, G. G. Mittelbach, T. Oberdorff, E. M. O'Brien & J. R. G. Turner, 2009. Spatial species-richness gradients across scales: a meta-analysis. Journal of Biogeography 36: 132–147.
- 570 Field, R. & P. R. Long, 2007. *Cusuco National Park, Honduras: ecology of a Meso-American cloud forest.*571 Operation Wallacea, Ltd, Old Bolingbroke, UK.
- 572 Fish, D., 1976. Structure and composition of the aquatic invertebrate community inhabiting epiphytic
 573 bromeliads in South Florida and the discovery of an insectivorous bromeliad. PhD dissertation.
 574 University of Florida.
- Frank, J. H. & L. P. Lounibos, 1983. *Phytotelmata: terrestrial plants as hosts for aquatic insect communities.* Medford, New Jersey; Plexus.

- Frank, J. H., S. Sreenivasan, P. J. Benshoff, M. A. Deyrup, G. B. Edwards, S. E. Halbert, A. B. Hamon, M. D.
 Lowman, E. L. Mockford, R. H. Scheffrahn, G. J. Steck, M. C. Thomas, T. J. Walker & W. C.
 Welbourne, 2004. Invertebrate animals extracted from native Tillandsia (Bromeliales: Bromeliaceae) in
 Sarasota County, Florida. Florida Entomologist 87: 176–185.
- Frank, J. H. & L. P. Lounibos, 2009. Insects and allies associated with bromeliads: a review. Terrestrial
 Arthropod Reviews 1: 125–153.
- Grytnes, J. A. & C. M. McCain, 2007. Elevational patterns in species richness. *Encyclopedia of Biodiversity* (Ed
 S. Levin), Elsevier, Inc.
- Greeney, H. F., 2001. The insects of plant-held waters: a review and bibliography. Journal of Tropical Ecology
 17: 241–260.
- Hortal, J., K. A. Triantis, S. Meiri, E. Thebault & S. Sfenthourakis, 2009. Island species richness increases with
 habitat diversity. American Naturalist 174: E205–E217.
- Jabiol, J., B. Corbara, A. Dejean & R. Cereghino, 2009. Structure of aquatic insect communities in tank bromeliads in an East-Amazonian rainforest in French Guiana. Forest Ecology and Management 257:
 351–360.
- Jocque, M., A. Kernahan, A. Nobes, C. Willians & R. Field, 2010a. How effective are non-destructive sampling
 methods to assess aquatic invertebrate diversity in bromeliads? Hydrobiologia 649: 293–300.
- Jocque, M., R. Field, L. Brendonck & L. De Meester, 2010b. Climatic control of dispersal–ecological
 specialization trade-offs: a metacommunity process at the heart of the latitudinal diversity gradient?
 Global Ecology and Biogeography 19: 244–252.
- Jocque, M. & J. Kolby, 2012. Acidity of tank bromeliad water in a cloud forest, Cusuco National Park,
 Honduras. International Journal of Plant Physiology and Biochemistry 4: 59–70.
- Karger, D. N., J. Kluge, T. Kromer, A. Hemp, M. Lehnert & M. Kessler, 2011. The effect of area on local and
 regional elevational patterns of species richness. Journal of Biogeography 38: 1177–1185.
- Kitching, R. L., 2000. Food webs and container habitats: the natural history and ecology of phytotelmata.
 Cambridge University Press, Cambridge.
- 603 Laessle, A. M., 1961. A micro-limnological study of Jamaican bromeliads. Ecology 42: 499–517.
- Leibold, M. A., M. Holyoak, N. Mouquet, P. Amarasekare, J. M. Chase, M. F. Hoopes, R. D. Holt, J. B. Shurin,
 R. Law, D. Tilman, M. Loreau & A. Gonzalez, 2004. The meta-community concept: a framework for
 multi-scale community ecology. Ecology Letters 7: 601–613.
- Little, T and P. D. N. Hebert. 1996. Endemism and ecological islands: the ostracods from Jamaican bromeliads.
 Freshwater Biology 36: 327–338.
- MacArthur, R. H. & E. O. Wilson, 1967. The theory of island biogeography. Princeton University Press, New Jersey.
- Maguire, B. 1971. Phytotelmata: Biota and community structure determination in plant-held waters. Annual
 Review of Ecology and Systematics 2: 439–464.
- 613 McCain, C. M., 2007. Area and mammalian elevational diversity. Ecology 88: 76–86.
- Mendes, H. F., T. Andersen & M. Jocque, 2011 A new species of *Polypedilum* Kieffer from bromeliads in
 Parque Nacional Cusuco, Honduras (Chironomidae: Chironominae). Zootaxa 3062: 46–54.
- Montero G., C. Feruglio & I. M. Barberis, 2010. The phytotelmata and foliage macrofauna assemblages of a
 bromeliad species in different habitats and seasons. Insect Conservation and Diversity 3, 92–201.
- 618 Picado, C., 1913. Les Bromeliacees Epiphytes. Bulletin Scientifique Tome XLVII: 216–360.
- 619 Rahbek, C., 1995. The elevational gradient of species richness a uniform pattern. Ecography 18: 200–205.
- Rahbek, C., 2005. The role of spatial scale and the perception of large-scale species-richness patterns. Ecology
 Letters 8: 224–239.
- Rangel, T., J. A. F. Diniz-Filho & L. M. Bini, 2006. Towards an integrated computational tool for spatial
 analysis in macroecology and biogeography. Global Ecology and Biogeography 15: 321–327.
- Reich, A., J. J. Ewel, N. M. Nadkarni, T. Dawson & R. D. Evans, 2003. Nitrogen isotope ratios shift with plant
 size in tropical bromeliads. Oecologia 137: 587–590.
- Richardson, B. A., 1999. The bromeliad microcosm and the assessment of faunal diversity in a Neotropical
 forest. Biotropica 31: 321–336.
- R Studio, 2012. RStudio: Integrated development environment for R (Version 0.96.122) [Computer software].
 Boston, MA. Retrieved May 20, 2012.
- 630 Srivastava, D. S. & J. H. Lawton 1998. Why more productive sites have more species: An experimental test of
 631 theory using tree-hole communities. American Naturalist 152: 510–529.
- 632 Srivastava, D. S., 2006. Habitat structure, trophic structure and ecosystem function: interactive effects in a
 633 bromeliad-insect community. Oecologia 149: 493–504.
- 634 Srivastava, D. S., J. Kolasa, J. Bengtsson, A. Gonzalez, S. P. Lawler, T. E. Miller, P. Munguia, T. Romanuk, D.
 635 C. Schneider & M. K. Trzcinski, 2004. Are natural microcosms useful model systems for ecology? Trends
 636 in Ecology & Evolution 19: 379–384.

Srivastava, D. S., M. K. Trzcinski, B. A. Richardson & B. Gilbert, 2008. Why are predators more sensitive to habitat size than their prey? Insights from bromeliad insect food webs. American Naturalist 172:761–771.

- 639 StatSoft, Inc., 2012. STATISTICA (data analysis software system), version 11. www.statsoft.com.
- Sugden, A. M. & R. J. Robins, 1979. Aspects of the ecology of vascular epiphytes in Colombian cloud forests .1. Distribution of the epiphytic flora. Biotropica 11: 173–188.
- ter Braak, C. J. F. & P. Šmilauer, 2002. CANOCO Reference Manual and CanoDraw for Windows User's Guide: Software for Canonical Community Ordination (version 4.5). Microcomputer Power (Ithaca NY, USA).

Table 1. Number of unidentified morphospecies recorded in the sampled bromeliads.

Class	Family	Genus	Species	No. of spp.
Diptera				6
1	Chironomidae			5
	Ceratopogonida	e <i>Bezzia</i>		3
	Tipulidae	Trentepohlia		1
	Culicidae	Culex		1
	Culicidae	Aedes		2
	Culicidae	Toxorhychites		1
	Culicidae	Wyeomia		1
	Syrphidae	Ocyptamus		1
	Syrphidae	Copestylum		4
	Syrphidae	Meromacrus		1
	Psychodidae	Telmatoscopus		2
Coleoptera				5
	Scirtidae	Scirtes		1
Hemiptera	Mesoveliidae			1
Ostracoda	Limnocytherida	e Elpidium		1
	Candonidae			1
Branchiopoda	Daphniidae	Ceriodaphnia	laticaudata	1
	Chydoridae	Alona	bromelicola	1
Copepoda				1
Oligochaeta	Enchytraeidae	Hemienchytraeu	5	1
Turbellaria				1
Total				42

Table 2. Correlation matrix of the explanatory variables, the species richness (S) and total Abundance (Abund) of invertebrates recorded in the sampled bromeliads. Values shown are Pearson's *r* for untransformed variables. Significance is indicated as * 0.05 > P > 0.01, ** 0.01 > P > 0.001, *** P < 0.001. Width, height and core (diameter of) all measure physical bromeliad size. Water content, weight (fresh weight of leaves), number of leaves and detritus content are also related to bromeliad size. Attachment height, access to light and rainfall, and the total number of bromeliads within a radius of 2 m (R-Brom) all quantify position in the forest canopy. The number of invertebrate species (S) and the total number of invertebrates (Abund) per bromeliad are response variables. Mean values with standard deviation (Stdev) and the maximum and minimum recorded values of each variable are presented at the lower part of the table.

	Width	Height	Core	Water	Weight	#leaves	Detritus	Attach	Light	Rain	R-Brom	S	Abund
Altitude	-0.26**	-0.26**	0.15	6 0.11	-0.11	-0.14	-0.07	-0.02	-0.26**	-0.15	-0.07	-0.1	-0.13
Width		0.55***	0.55***	0.45***	0.80***	0.63***	0.70***	0.11	0.18*	0.20*	0.15	0.48***	0.45***
Height			0.31***	0.18*	0.40***	0.30***	0.46***	0.01	0.37***	0.42***	0.28***	0.26**	0.33***
Core				0.39***	0.69***	0.47***	0.52***	0.16*	0.1	0.18*	0.11	0.42***	0.47***
Water					0.61***	0.52***	0.25**	0.05	-0.04	0.05	0.03	0.35***	0.32***
Weight						0.74***	0.74***	0.15	0.14	0.15	0.09	0.54***	0.49***
#leaves							0.54***	0.11	0.14	0.13	0.08	0.47***	0.32***
Detritus								0.09	0.08	0.14	0.11	0.45***	0.34***
Attach									-0.01	-0.07	-0.07	-0.01	-0.02
Light										0.83***	0.51***	0.19*	0.25**
Rain											0.62***	0.20*	0.31***
R-Brom												0.06	0.08
S													0.62***
	Width	Height	Core	Water	Weight	#leaves	Detritus	Attach	Light	Rain	R-Brom	S	Abund
	(cm)	(cm)	(mm)	(ml)	(g)		(g)	(mm)					
Mean	37.3	3 12.8	8 8.8	8 83.3	174.0	26	28.2	140.1	4.1	3.2	13.0	8.0	55.8
Stdev	12.8	8 8.3	3 3.0	85.1	132.5	8	24.5	60.0	2.0	2.7	18.0	3.2	46.6
Max	84	52	2 27.5	5 410	613	49	130	369	8	9	114	17	227
Min	17	/ () 3.5	5 1	9	11	1	30	1	0	0	1	4

Table 3. Beta diversity partitioning: results of Mantel tests correlating pairwise dissimilarity of aquatic invertebrate communities inhabiting bromeliads with pairwise distance or differences in environmental variables. Size was measured as the first principal component of the size variables. Values given are Mantel's *r*, with significance indicated as * 0.05 > P > 0.01, ** 0.01 > P > 0.001, *** P = 0.001. P-values were derived from 999 permutations using the 'vegan' package in R, which returns the number of permutations in which the observed *r* is exceeded plus one, then divided by 1000; thus *** represents cases where none of the permutations exceeded the observed correlation. Because the 'P'-value returned is one-tailed, we have doubled it before binning into significance categories, to approximate two-tailed testing (note: this assumes a symmetric distribution of *r* in permutations; no cases were marginal). For negative correlations, we first subtracted the returned '*P*' from 1.

Dissimilarity measure	Geographic distance	Altitudinal distance	Size difference	
Sorensen	0.05**	0.19***	0.16***	
Simpson (turnover)	0.05*	0.16***	0.02	
Nestedness-resultant	-0.01	-0.02	0.17***	
Jaccard	0.06**	0.19***	0.16***	
Turnover (Jaccard)	0.05*	0.14***	0.01	
Nestedness-resultant (Jaccard)	-0.02	-0.03	0.12***	



FIGURE 1. Scatter plot of species richness and total wet weight of the washed bromeliad leaves in grams. The correlation is significant and a semi-logarithmic fit is displayed (species richness = $-1.66 + 4.65 \times \log(\text{weight})$), which accounts for 33% of the variance in species richness.



FIGURE 2. RDA biplot of the species and environmental variables. See Table 1 for a list of the species and Table 2 for explanation of the variable names.