Deterministic tropical tree community turnover: evidence from patterns of functional beta diversity along an elevational gradient

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Explaining the mechanisms that produce the enormous diversity within and between tropical tree communities is a pressing challenge for plant community ecologists. Mechanistic hypotheses range from niche-based deterministic to dispersal-based stochastic models. Strong tests of these hypotheses require detailed information regarding the functional strategies of species. A few tropical studies to date have examined trait dispersion within individual forest plots using species trait means in order to ask whether coexisting species tend to be more or less functionally similar than expected given a null model. The present work takes an alternative approach by: (i) explicitly incorporating population-level trait variability; and (ii) quantifying the functional beta diversity in a series of 15 tropical forest plots arrayed along an elevational gradient. The results show a strong pattern of decay in community functional similarity with elevation. These observed patterns of functional beta diversity are shown to be highly non-random and support a deterministic model of tropical tree community assembly and turnover.

Keywords: community ecology; functional traits; leaf area; Puerto Rico; specific leaf area; wood density

1. INTRODUCTION

Tropical communities are renowned for their biodiversity. Not only is there an elevated level of species diversity within communities (i.e. alpha diversity) with respect to less diverse temperate communities, but species turnover between tropical communities (i.e. beta diversity) is also elevated. Perhaps, nowhere are patterns of tropical diversity more often documented than in tree communities (e.g. [1–5]), yet a substantial debate still exists on the extent to which these patterns are the result of deterministic or stochastic processes (e.g. [6–8]). Understanding the degree to which patterns of diversity are the result of deterministic or stochastic processes is particularly critical for those trying to conserve and predict the fate of tropical biodiversity.

The majority of research into patterns of tropical tree diversity and turnover has focused on one type of biodiversity: species diversity. The functional component of biodiversity in these communities has been less well quantified (but see [8–10]). This is unfortunate because documenting the functional strategies of species in communities may provide refined or novel mechanistic insights into their structure and diversity [11].

One principle concern for a community ecologist is that traditional analyses of species beta diversity are unable to determine whether or not the communities are functionally similar or dissimilar. In figure 1, we present one example of this where species may be dispersal limited causing complete species turnover between two communities, but the environment also plays a key role in determining the community assembly, thereby generating functionally analogous communities (figure 1). This would result in a decoupling of species and functional beta diversity where species beta diversity is high and functional beta diversity is low. We would expect this result to be particularly important when comparing assemblages between regions that share similar environments. Despite the potential insights one can gain from analysing the functional similarity of communities, the concept of ‘functional beta diversity’ is rather new and it still remains to be deeply explored (but see [12,13]). To date, the majority of the work on this topic has regarded the development of functional beta diversity metrics similar to those designed to analyse species beta diversity (e.g. [14]) or phylogenetic beta diversity [15,16]. To our knowledge, there have been no analyses of functional beta diversity of communities arrayed along broad environmental gradients and no analyses of diverse ecosystems such as tropical rainforests. Thus, tests using such datasets and multiple metrics are needed.

When comparing communities along an environmental gradient it is probable that species beta diversity and functional beta diversity will be, on average, positively related. In such instances, the question becomes whether the functional turnover between communities is higher or lower than that expected given the species turnover. For example, a higher than expected functional turnover between communities given the species turnover could occur if the species in the two communities are more
functionally divergent than expected given the species pool. This would indicate that the species turnover along the gradient is not simply owing to dispersal limitation and that the underlying environmental gradient plays a key role in influencing species turnover and community assembly. Thus the given importance of quantifying the functional strategies of species within and among communities, ecologists are increasingly promoting a functional trait-based approach to community ecology [11].

Plant community ecologists are increasingly integrating functional ecology into their research programmes by quantifying functional traits on the individuals and species in their study systems (e.g. [17–21]). Functional traits are morphological and physiological traits that are indicative of the ecological strategies of species. An advantage of functional traits is that they allow for a rapid and standardized inventory of several major axes of plant functional differentiation across all species in diverse communities [22–24]. Indeed this advantage has led to the pioneering work into the relative influence of deterministic and stochastic processes on patterns of species coexistence in tropical tree communities [8].

Despite the promise of a functional trait-based paradigm in community ecology there are several outstanding challenges. First, the appropriateness of the functional traits used in community ecology investigations is routinely questioned and determining which are the ‘right’ traits to use is important. In recent years, there has been a great deal of focus on several plant functional traits that have been proposed to be important indicators of plant ecological strategies [22–24]. These ‘soft’ functional traits are proxies of actual physiological processes and they are relatively easy to measure compared with ‘hard’ functional traits that are often more closely linked to the physiological processes of interest, but much more difficult to quantify across diverse communities [25]. While the debate of which are the best traits or the right traits for community ecology investigations continues, an initial step is to determine when and where those often used soft functional traits can provide novel insights into the mechanisms determining community assembly and diversity.

A second challenge has been that of intraspecific functional trait variability [20,26]. Many functional trait investigations of communities have assigned species-level trait means to all individuals or populations (e.g. [8,10]). In some cases, these mean values are calculated from a few individuals from the study system, and in others these values come from global databases where...
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2. MATERIAL AND METHODS

(a) Study area and forest inventory plots

The present study took place in the Luquillo Experimental Forest (LEF) located in eastern Puerto Rico (18°19.6′N, 65°49.4′W). The LEF is approximately 11 330 ha in area with elevations ranging from 100 to 1075 m above sea level (a.s.l.) [27]. At 100 m a.s.l. the temperature averages 24.5°C during the warmest month of September and annual rainfall totals 2300 mm on average. At 1075 m a.s.l. the temperature averages 20°C during September and the annual rainfall totals 3600 mm on average. The elevational gradient spans from premontane rainforest to cloud forest. Permanent forest inventory plots were established from 2001 to 2002 along the Sonadora River watershed within the LEF [28]. The forest plots were arrayed every 50 m in elevation from 300 to 1000 m a.s.l. The sites selected for each forest plot had little to no previous human disturbance and did not include ravines [28]. Each forest plot is 0.1 ha in area (50 × 20 m) with all free-standing woody stems 1 cm in diameter at breast height (130 cm above the ground) mapped, tagged and identified. As the flora of the LEF is well-known, over 99 per cent of the stems could be identified to species and voucher specimens were deposited in the El Verde Field Station Herbarium. There are a total of 6562 stems and 102 species found in the 15 forest plots. In each plot, we calculated the species alpha and beta diversity. The species beta diversity was calculated using a Bray–Curtis Distance in the R package ‘vegan’.

(b) Functional traits and dendrogram

A total of six traits were chosen for this study to quantitatively represent several axes of plant functional strategy [24]. Specific leaf area (SLA), leaf nitrogen content (%N) and leaf phosphorus content (%P) were used to represent the well-documented leaf economics spectrum [29]. Wood-specific gravity was used to represent a wood economics spectrum [30]. Leaf area was used to represent a trade-off between area deployed for light capture and leaf temperatures [31], and maximum height was used to represent the adult light niche of the species [32].

Each trait was measured on multiple individuals in each forest inventory plot following the protocols in Cornelissen et al. [33]. When possible, 15 individuals were sampled for leaf and height trait measurements and 10 individuals were used for wood-specific gravity measurements. A recent power analysis of leaf functional traits in a tropical forest community has suggested that a similar sampling intensity is necessary for reliable estimations of population mean trait values [26]. Sampling was fewer than 15 and 10 individuals respectively, only when that many individuals could not be located at the given elevation.

For each trait, a distance matrix was constructed to represent the trait similarity between all populations using population-level mean trait values for each species. This distance matrix and UPGMA clustering were used to construct a dendrogram for each trait [34]. This dendrogram represented the trait similarity between populations. Two of the functional dissimilarity metrics shown below, $D_{\text{na}}$ and $D_{\text{pat}}$, can also be implemented without using trait dendrograms, whereas the other, $F_{\text{ext}}$, cannot. Thus, for consistency we have chosen to use the dendrograms for each metric. Further studies are needed to address the degree to which using dendrograms versus raw trait distances influence functional beta diversity metrics. As this study was also interested in a composite measure of functional similarity between species, we used a principle coordinates analysis (PCA). To perform the PCA, we first logged the trait data and converted it to standard normal deviates to identify the major axes of multivariate functional similarity. The first three PCA axes explained over 96 per cent of the variation (see the electronic supplementary material, table S1) and the Euclidean distance between populations in this three-dimensional space and UPGMA clustering were used to construct a composite functional dendrogram representing the overall functional similarity between populations.
(c) Functional beta diversity

The present study implemented three metrics of functional beta diversity. The concept and measurement of functional beta diversity are still in their infancy and further work will be needed to compare and contrast the relative benefits of various metrics. Here, we have chosen to use three potential metrics of functional similarity between communities as a starting point without making any particular judgement with respect to their relative merits. The first metric used is the Functional Sorensen’s Index, $F_{sor}$. This metric is presence–absence weighted and defined as:

$$F_{sor} = \frac{\text{BL}_{h;k_2}}{(\text{BL}_{h;k_1} + \text{BL}_{h;k_2})^{1/2}},$$

where $\text{BL}_{h;k_1}$ is the total dendrogram branch length common to all species in communities $k_1$ and $k_2$, and $\text{BL}_{h;k_2}$ is the total dendrogram branch length common to the species within communities $k_1$ and $k_2$, respectively. This metric is a functional trait analogue of the phylogenetic metric, PhyloSor, introduced by Bryant et al. [35]. It is also analogous to a traditional Sorensen’s Index. Thus, the $F_{sor}$ metric provides an overall indicator of the shared function driven by communities. Thus we may not expect it to correlate with the species within communities.

The next two metrics we employed use abundance information. First, we employed an abundance weighted nearest neighbour metric, $D_{nn}$, between two communities, $k_1$ and $k_2$. This metric was originally published by Ricotta & Burasciano [14] and is defined as:

$$D_{nn} = \sum_{i=1}^{n_k} f_i \min \delta_{ki} + \sum_{j=1}^{n_j} f_j \min \delta_{jk},$$

for species $i \neq$ species $j$

where $n_k$ is the number of species in community $k_1$, $f_i$ the relative abundance of species $i$ in community $k_1$, and $\delta_{ki}$ the dendrogram branch lengths between species $i$ in community $k_1$ and its nearest functional neighbour in community $k_2$ that is not the same species. Thus if two communities share functionally analogous or nearly analogous species, the average nearest neighbour distances will be much lower than two communities that have very dissimilar community trait distributions.

The third metric was an abundance weighted pairwise dissimilarity, $D_{pw}$, between communities $k_1$ and $k_2$ modified from the $D_{nn}$ metric described above. This metric is defined as:

$$D_{pw} = \frac{\sum_{i=1}^{n_k} f_i \delta_{ki} + \sum_{j=1}^{n_j} f_j \delta_{jk}}{2},$$

for species $i \neq$ species $j$

where $\delta_{ki}$ is the mean pairwise distance on the dendrogram between all species in community $k_1$ and all species in community $k_2$ excluding conspecific species. The $D_{pw}$ is somewhat different than most beta diversity metrics because it compares all pairwise distances between species in the two communities. Thus we may not expect it to correlate with patterns of species beta diversity as well as the previous two metrics.

(d) Null model and standardized effect sizes

As discussed above, it is reasonable to expect species and functional beta diversity to be correlated. While we were interested in documenting patterns of functional beta diversity in the study system, we were also interested in determining whether the functional beta diversity was higher or lower than expected given the species beta diversity. In order to accomplish this we performed null modelling analyses. A null distribution of functional beta diversity values was generated for each trait by randomizing the names of the populations across the tips of the trait dendrograms 9999 times. During each iteration, the functional beta diversity was calculated across the entire elevational gradient. These values were used to generate the null distribution. Therefore, the randomization procedure only randomized the functional similarity of populations while maintaining the observed species occupancy rates, community species richness levels, community abundance distributions and species beta diversity. Further, as the null model maintains the observed distributional patterns of species and their populations, the observed dispersal limitation of species is conserved in the null model.

A standardized effect size (SES; [36]) was calculated for functional beta diversity using the mean and standard deviation of the null distribution as follows:

$$\text{SES} = \frac{X_{obs} - X_{null}}{\text{s.d.}(X_{null})},$$

where $X_{obs}$ is the observed dissimilarity value (i.e. $D_{pw}$) between two communities, $X_{null}$ the mean of the null distribution and s.d.($X_{null}$) the standard deviation of the null distribution. Values greater than 1.96 indicate a higher than expected functional dissimilarity between the communities and values below $-1.96$ indicate a lower than expected functional dissimilarity between the two communities. Non-random SES values could be generated by functionally non-random replacement of species along the gradient and/or population-level trait means shifting directionally along the gradient.

3. RESULTS

(a) Species alpha and beta diversity

There was a general decline in species alpha diversity with increasing elevation (see the electronic supplementary material, figure S1) although the pattern may have become unimodal if lower elevation forests could have been censused. The species alpha diversity in the plots ranged from 18 species at 950 m a.s.l. to a maximum of 52 species at 300 m a.s.l. The species beta diversity was also calculated and the results show a general increase in community dissimilarity with increasing elevation using a Bray–Curtis Distance (see the electronic supplementary material, figures S1 and S2).

(b) Functional beta diversity and standardized effect sizes

The functional beta diversity of the forest plots was determined using three different metrics. In the main text, we only present the functional dissimilarity between the plot at the lowest elevation and the other 14 forest plots along the gradient, but all pairwise comparisons between plots are presented in the electronic supplementary material. The results show a general decay in functional similarity with elevation across all traits using both the presence–absence weighted $F_{sor}$ metric and the abundance weighted nearest neighbour metric, $D_{nn}$ (figure 2 and electronic supplementary material, figure S3). The results were not as consistent using the abundance weighted pairwise metric, $D_{pw}$. Specifically, dissimilarity
increased with elevation for maximum height, leaf area, leaf nitrogen content, leaf phosphorus content and PCA, but the wood-specific gravity results were unimodal (figure 2 and electronic supplementary material, figure S3).

In an additional analysis we compared the trait dissimilarity values between plots with the Bray–Curtis Distance between plots. The Bray–Curtis Distance was positively correlated with the \( F_{\text{ora}} \) and \( D_{\text{nn}} \) trait dissimilarity metrics (see the electronic supplementary material, figure S4). The \( D_{\text{pw}} \) metric on the other hand was generally not correlated with the Bray–Curtis Distance (electronic supplementary material, figure S4). Given the strong relationship between some of the functional beta diversity metrics and species beta diversity and our desire to know whether the functional beta diversity was higher or lower than expected given the species beta diversity, we conducted null model analyses. The SESs for the \( F_{\text{ora}} \), \( D_{\text{nn}} \) and \( D_{\text{pw}} \) metrics were generally positive across all traits and consistent across elevation with effect sizes often greater than 1.96 (figure 3). The constancy in the positive values greater than 1.96 is indicative of higher than expected functional trait dissimilarity or turnover with elevation. A notable exception to this general pattern is the SLA results for the \( D_{\text{pw}} \) metric where there was lower than expected turnover initially along the gradient followed by a larger than expected turnover when comparing the lowest plot to the high elevation plots (figure 3).

4. DISCUSSION

The present study asked whether the turnover of tropical tree communities along an elevational gradient is deterministic or stochastic with respect to species function. To date, tropical tree community turnover has often been analysed using lists and abundances of species that are treated as functionally equivalent or binned into broad categorically defined functional groupings. Recently,

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**Figure 2.** The beta functional diversity along the elevational gradient in the Luquillo Experimental Forest presented as the dissimilarity between the plot at the lowest elevation (300 m a.s.l.) and the other 14 forest plots spaced every 50 m in elevation. (a) Calculates beta functional diversity using the presence–absence weighted \( F_{\text{ora}} \) metric, (b,c) calculate beta functional diversity using the abundance weighted nearest neighbour metric, \( D_{\text{nn}} \) and the pairwise metric, \( D_{\text{pw}} \), respectively.
species mean functional trait values have been used in two tropical forest plots to quantify whether the functional alpha diversity in forest subplots is higher or lower than expected given the species richness [8, 10]. To date, this work has not addressed the turnover of tree species in these diverse systems and it has not explicitly dealt with the potentially high level of intraspecific variation in functional trait values. In this study, to our knowledge, we have presented the first analysis of functional beta diversity in tropical tree communities, and we have provided the first functional trait analysis of tropical tree communities that explicitly incorporates intraspecific trait variation. The functional beta diversity results showed a general non-random compositional turnover of function along the elevational gradient, suggesting a large role for deterministic processes in controlling community turnover along the gradient. Below, we discuss these results and their implications in more detail.

The primary goal of the present study was to ask whether the turnover of tropical tree communities is relatively deterministic or stochastic with respect to species function. To address this we quantified the functional beta diversity or dissimilarity between the forest plot at the lowest elevation to the other 14 forest plots at higher elevations using three different metrics (figure 2 and electronic supplementary material, figure S3). In general, the functional beta diversity increased with elevation using all three metrics (figure 2 and electronic

Figure 3. The beta functional dispersion along the elevational gradient in the Luquillo Experimental Forest. The data are presented using the standardized effect size (SES). SES values above 1.96 (the upper dashed grey line) indicate significantly higher than expected beta functional diversity, or dissimilarity, given the observed beta species diversity. SES values below −1.96 (the lower dashed grey line) indicate significantly lower than the expected beta functional diversity, or dissimilarity, given the observed beta species diversity.
supplementary material, figure S3). In other words, there was a clear distance decay in functional similarity from the lowest to the highest forest plot. Thus, the functional composition of the tree communities along the elevational gradient changes directionally for all of the traits studied.

Further analyses revealed that two of the metrics used were strongly correlated with species beta diversity, thereby making it difficult to determine whether functional turnover along the gradient was any different from that expected given the underlying species turnover. Consequently, it is tenuous to reject a stochastic model of community turnover and assembly from functional distance decay results alone and null model analyses are necessary.

The results from the null model analyses for all of the metrics were generally non-random with SESs often greater than 1.96 (figure 3). Thus, the functional composition of the communities along the elevational gradient turns over faster than expected given the species compositional turnover and changes in species abundance. The faster than expected turnover in function between communities could occur for two reasons. First, those species that replace each other from one community to the next are functionally more divergent than expected given the species pool. For example, say that there was complete turnover in the species composition when transitioning from 300 to 900 m elevation. If the species at 900 m are a non-random subset of the species pool that is functionally more distant than expected at random from those found at 300 m, then this would indicate that the species turnover is not simply stochastic. Second, if there are substantial directional trends in population-level trait means along the environmental gradient, the functional turnover would be faster than that expected given the species turnover. While there is a high level of species turnover along the gradient studied (see the electronic supplementary material, figures S1 and S2), nowhere is there a complete turnover in species composition between forest plots, and a few species span the entire elevational gradient. We therefore suggest that both of the above processes probably generated the faster than expected functional turnover reported in this study. Further, both a functionally non-random replacement of species along a gradient and directional trends in population-level trait values along a gradient are evidence for deterministic species turnover and community assembly along the gradient.

The primary goal of the current study was to document patterns of functional turnover in tree communities along a substantial environmental gradient in the tropics. To achieve this goal, we implemented three metrics of functional beta diversity. In conducting the work we found that the first two metrics, $F_{diss}$ and $D_{diss}$ were strongly related to a Bray–Curtis Distance while the third metric, $D_{pat}$ was not (see the electronic supplementary material, figure S4). Thus, the first two metrics may be considered more closely aligned with what is traditionally considered to be ‘beta diversity’ while the third metric may just serve as an alternative measure of similarity that cannot be easily fitted into a diversity-partitioning framework. As the topic of functional beta diversity becomes more broadly studied, detailed investigations into when and why these particular metrics and other metrics are preferred will be needed.

In summary, here we have provided, to our knowledge, the first functional beta diversity analyses of a series of tropical tree communities arrayed along a substantial ecological gradient, to address whether the compositional turnover between these communities is relatively stochastic or deterministic. We also present, to our knowledge, the first analysis of functional beta diversity that explicitly incorporates intraspecific variation in traits along a broad gradient. The results presented are largely non-random and the turnover of the functional composition of communities is generally faster than that expected along the gradient. The results are also consistent across all of the traits studied and each of the metrics used, which indicates that the compositional change along the gradient involves several axes of plant functional differentiation and is not just a shift in any one trait or axis. These results suggest the predominate role of deterministic factors promoting the compositional turnover of the tropical tree communities along the gradient and a lesser role for stochasticity.

5. CONCLUSIONS AND FUTURE DIRECTIONS

Here, we have presented an initial test of whether or not the compositional similarity of tropical tree communities is non-random with respect to species function. The results show that tropical tree community similarity in our study system is indeed functionally non-random and supports a deterministic model of tropical tree community assembly and turnover. Future work should aim to repeat a similar approach across several gradients within and outside the tropics to determine whether or not similar distance decay relationships occur in other systems.

The work has also successfully integrated intraspecific functional trait variation into the analyses. Accounting for this variation will become increasingly important as the scale of functional trait investigations into community structure broadens and even more so as functional beta diversity becomes increasingly measured across broad ecological gradients. Additional studies will be needed to quantify to what degree intraspecific variability will bias functional beta diversity analyses that use species-level mean trait values and how this bias may be enhanced or mitigated in systems with more or less variable species.

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