Community Structure of Skipper Butterflies (Lepidoptera, Hesperiidae) along Elevation Gradients in Brazilian Atlantic Forest Reflects Vegetation Type Rather than Altitude

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Abstract

Species turnover across elevational gradients has matured into an important paradigm of community ecology. Here, we tested whether ecological and phylogenetic structure of skipper butterfly assemblages is more strongly structured according to altitude or vegetation type along three elevation gradients of moderate extent in Serra do Mar, Southern Brazil. Skippers were surveyed along three different mountain transects, and data on altitude and vegetation type of every collection site were recorded. NMDS ordination plots were used to assess community turnover and the influence of phylogenetic distance between species on apparent community patterns. Ordinations based on ecological similarity (Bray-Curtis index) were compared to those based on phylogenetic distance measures (MPD and MNTD) derived from a supertree. In the absence of a well-resolved phylogeny, various branch length transformation methods were applied together with four different null models, aiming to assess if results were confounded by low-resolution trees. Species composition as well as phylogenetic community structure of skipper butterflies were more prominently related to vegetation type instead of altitude per se. Phylogenetic distances reflected spatial community patterns less clearly than species composition, but revealed a more distinct fauna of monocot feeders associated with grassland habitats, implying that historical factors have played a fundamental role in shaping species composition across elevation gradients. Phylogenetic structure of community turned out to be a relevant additional tool which was even superior to identify faunal contrasts between forest and grassland habitats related to deep evolutionary splits. Since endemic skippers tend to occur in grassland habitats in the Serra do Mar, inclusion of phylogenetic diversity may also be important for conservation decisions.

Introduction

Since the early observations by Forster [1] and von Humboldt [2], species turnover across altitudinal gradients has matured into an important paradigm of community ecology. High mountain chains are usually selected for analyses, mainly because their wide elevational ranges increase the power of detecting patterns in community structure [see [3,4]]. Geologically older mountain chains whose peaks are considerably lower as a consequence of long exposition to erosion forces have received less attention, since effects of elevation on species diversity or species composition might not be as evident. In such places, historical events may play an additional fundamental role in assemblage turnover in addition to altitude. More recently, the integration of phylogenetic structure into community ecology has yielded important insights as to how historical factors influence community structure [3].

The Serra do Mar is located near the south-eastern coast of Brazil and represents a good example of an old mountain chain in the southern hemisphere, where animal and plant communities might have been structured through both ecological and historical processes. The start of its rising processes has been dated at about 90 mya, soon after the splitting of Gondwana into its two biggest daughter continents, viz. South America and Africa [6]. Despite its moderate altitudinal amplitude today (peaks generally between 800–1500 m), distinct climatic and pedological shifts occur along these rather short elevational gradients, together with a high endemism rate in certain plant taxa such as Bromeliaceae, Orchidaceae and Gesneriaceae [6,7].

Environmental dimensions change in a predictable manner with increasing altitude in the Serra do Mar. Mean annual temperature decreases by an average of 0.5–0.6 K per 100 m altitude, while soil depth decreases and wind intensity increases [8]. Correlated to these abiotic factors and modulated through superimposed effects of anthropogenic land-use, four different vegetation types can be observed on these mountains: Atlantic rain forest, early succession flora, cloud forest, and grasslands [7]. While Atlantic rain forest prevails in the lowlands (up to 1100 or 1400 m, depending on the mountain), grasslands are generally restricted to altitudes above
These grasslands are relict vegetation ecosystems which had been far more widespread in south and southeast Brazil during dry cold periods of the late Quaternary [9]. Since then, this vegetation has become increasingly restricted to mountain tops, due to forest expansion, with many animal species related to this kind of ecosystem being included in red lists [10,11]. Early successional flora is not only found after human interference, but also due to natural disturbance events such as forest fires or landslides [7,8].

Communities of host-specific herbivores, whose life histories are intrinsically related to the vegetation of their habitats, may either respond directly to abiotic environmental gradients, or alternatively their species turnover may rather track biotic changes in vegetation types [12,13]. The challenge remains to disentangle whether abiotic (climate, area size) or biotic conditions (vegetation) are more important as drivers of their community composition and species turnover along elevational gradients [14,15], and how phylogenetic history contributes to understanding contrasting patterns between taxa [5,16].

Various groups of nocturnal Lepidoptera have frequently served as targets to study elevational diversity patterns in the tropics [17–19], yet surprisingly few quantitative butterfly studies do exist from elevational gradients on tropical mountains [20,21]. Especially, the few studies addressing phylogenetic structure in butterfly communities along elevational gradients are extra-tropical [22,23], where faunas are distinctly less diverse (but see [24] for a recent example on a rich Neotropical moth assemblage). Skipper butterflies (Lepidoptera, Hesperioidea) might also have the potential to address elevational patterns and underlying processes, but thus far no specific case study on this somehow elusive family exists from mountain ranges anywhere in the tropics.

Recent studies revealed contrasting patterns of phylogenetic community structure along environmental gradients, i.e. different assemblages respond to environmental or biotic factors depending on their specific requirements, evolutionary history and biogeography [25–28]. Skippers might have particular potential to reveal how vegetation types and altitude influence phylogenetic composition along altitudinal gradients, since major skipper lineages are conservative and contrasting in relation to their larval food plant affiliations [29]. While Hesperinae larvae feed exclusively on monocotyledonous plants, Pyrginae larvae are bound to various families of dicotyledonous plants, mainly Fabaceae, Malvaceae and Rutaceae [29–31]. In addition, skippers are the only group of butterflies commonly observed throughout all sections of altitudinal gradients in south Brazil (pers. observ.), and they comprise far

Table 1. Two different sets of sample unit delimitations used to analyze skipper assemblages in the Serra do Mar (Brazil).

| Samples 1 (m) | Mountain | Veg. Type | Samples 2 (m) | Mountain | Veg. Type | Altitude Class |
|--------------|----------|-----------|---------------|----------|-----------|----------------|
| 1000–1100    | Anhangava| FOR       | 998–1060      | Anhangava| FOR       | low            |
| 1100–1200    | Anhangava| FOR+ESV   | 1061–1122     | Anhangava| FOR       | low            |
| 1200–1300    | Anhangava| ESV+GRA   | 1123–1206     | Anhangava| ESV       | medium         |
| 1300–1400    | Anhangava| GRA       | 1207–1289     | Anhangava| ESV       | medium         |
| 1400–1500    | Anhangava| GRA       | 1290–1364     | Anhangava| GRA       | medium*        |
| 900–1000     | Aracáutuba| FOR+ESV  | 1365–1440     | Aracáutuba| GRA       | high           |
| 1000–1100    | Aracáutuba| ESV       | 912–938       | Aracáutuba| FOR       | low            |
| 1100–1200    | Aracáutuba| GRA       | 939–1019      | Aracáutuba| ESV       | low            |
| 1200–1300    | Aracáutuba| GRA       | 1020–1099     | Aracáutuba| ESV       | low            |
| 1300–1400    | Aracáutuba| GRA       | 1100–1175     | Aracáutuba| GRA       | low            |
| 1400–1500    | Aracáutuba| GRA       | 1176–1250     | Aracáutuba| GRA       | medium         |
| 1500–1600    | Aracáutuba| GRA       | 1251–1325     | Aracáutuba| GRA       | medium         |
| 1600–1700    | Aracáutuba| GRA       | 1326–1400     | Aracáutuba| GRA       | medium         |
| 900–1000     | Caratuva  | FOR       | 1401–1475     | Aracáutuba| GRA       | high           |
| 1000–1100    | Caratuva  | FOR+ESV   | 1476–1550     | Aracáutuba| GRA       | high*          |
| 1100–1200    | Caratuva  | ESV       | 1551–1625     | Aracáutuba| GRA       | high           |
| 1200–1300    | Caratuva  | ESV       | 1625–1682     | Aracáutuba| GRA       | high           |
| 1300–1400    | Caratuva  | FOR       | 980–1031      | Caratuva  | FOR       | low            |
| 1400–1500    | Caratuva  | FOR+GRA   | 1032–1083     | Caratuva  | FOR       | low            |
| 1800–1900    | Caratuva  | GRA       | 1084–1158     | Caratuva  | ESV       | low            |
| 1159–1233    | Caratuva  | ESV       | medium        |
| 1234–1306    | Caratuva  | ESV       | medium        |
| 1307–1362    | Caratuva  | FOR       | medium        |
| 1363–1418    | Caratuva  | FOR       | medium        |
| 1419–1488    | Caratuva  | GRA       | high          |
| 1800–1860    | Caratuva  | GRA       | high*         |

1 Samples 1: delimited only by altitude; Samples 2: delimited by vegetation type and altitude. Each location is assigned to mountains, elevational belts and vegetation types. Note that the delimitation by altitude plus vegetation increases the number of sample units. Abbreviations: FOR: forest; ESV: early successional vegetation; GRA: grassland.
larger numbers of grassland endemics when compared to Papilionoidea families in the region [32].

The present study aimed to elucidate the structure of Hesperiidae assemblages along altitudinal gradients of moderate extension, using ecological and phylogenetic measures of commu-
nity similarity, on three different mountains, in relation to different vegetation types. Specifically, the following hypotheses were tested: 1. Assemblages of skippers are structured according to vegetation type as well as altitude; 2. Phylogenetic similarity is more informative for community responses to altitude and vegetation than ecological similarity (e.g. species composition); 3. Vegetation has a stronger influence on skipper assemblages than altitude; 4. Subfamilies of skippers show discordant community patterns because they differ in life-history traits governing their habitat preferences.

Methods
Study location and sampling methods
Three mountains located in the Serra do Mar of Paraná state, Brazil were sampled in this study: Anhangava (25°23′30″S; 49°00′15″W), Araçatuba (25°54′07″S; 48°59′37″W) and Caratuva mountain (25°13′30″S; 48°51′40″W). The three locations sampled are embedded in protected areas regulated by the state environmental agency (IAP/PR). For location details see [33]. Capture of specimens and their transport to the laboratory for subsequent identification were permitted by licences n° 59.08 (IAP/PR) and 14.595-1 (IBAMA/Sisbio). Hesperiidae specimens were captured with insect nets during up- and down-walking of transects. Each transect was walked 11 times from 2009 to 2011, between 9:00 and 16:00 h. All recorded specimens were immediately labeled according to the elevation (measured to the nearest 10 m using a Garmin 60Cx GPS device) and vegetation type of their sampling locality. In all, 1578 records of 155 species make up the data on which all analyses are based. No endangered species were recorded in this study.

On the mountains in Serra do Mar up to four different vegetation types are present above 900 m: montane forest, cloud forest, early successional vegetation, and grassland. Montane forest, early successional vegetation, and grassland stands have a taller canopy stratum. Succession vegetation is dominated by bracken fern Pteridium aquilinum (Dennstaedtiaceae), and it is often located where fire or human impact have occurred recently [8]. Grassland sites present a varied floral composition, dominated by several species of Poaceae and Cyperaceae, but occasionally with scattered low trees and shrubs [7].

Although these vegetation types are in part related to altitude (e.g. grasslands are located on mountain tops), their ranges also vary according to a mountain’s relief, soil depth, or biogeographical history [7]. This peculiarity allowed us to examine whether vegetation or altitude per se plays the more important role on community differentiation. Therefore, skipper samples were delimited and analyzed in two different ways: first, the whole altitudinal range on each mountain was divided into belts of 100 m elevational extent, without considering changes of vegetation types within; second, samples were again delimited by altitude, but in addition the prevalent vegetation type was superimposed to delimit sample sites (Table 1). As a consequence, these latter operational units were not equally sized according to elevational bands, but varied from 50 to 75 m extension each. Because only two specimens of skippers were collected in cloud forest, this vegetation type was excluded from all analyses.

Ecological and phylogenetic community structure analyses
NMDS ordinations were used to search for both ecological and phylogenetic structure in skipper assemblages along elevational gradients. This methodology enables the recognition of spatial gradients across communities through comparisons of pairwise similarities, or distances, between all samples [34]. Brehm & Fiedler [35] evaluated different ordination methods for identifying elevational gradients with incompletely sampled communities and concluded that different techniques performed quite similarly. Furthermore, NMDS has the advantage of fixing a priori the number of dimensions to be considered for analysis, and to be grounded on rank statistics which renders this ordination method very robust [36,37].

In a first series of NMDS explorations, based on Bray-Curtis matrix similarities, it was assessed whether inclusion, or exclusion, of hilltopping species (i.e. where adult butterflies aggregate at mountain tops for mating location [38]), or the segregation of samples only by altitude (100 m belts), or by altitude plus vegetation type (50–75 m belts), would affect ordination patterns, as already shown for species richness patterns [33]. Since both these factors indeed influenced the ordination of assemblages (Fig. S1), in the subsequent main series of analyses all hilltopping species were omitted and sample sites were classified according to elevation plus vegetation type. Species that exhibit hilltopping behavior are listed in Carneiro et al. [33]. Because hilltopping species were quite numerous, two samples, one from the 1800–1900 m band on Caratuva and the 1476–1550 m band on Araçatuba, became too small and therefore had to be excluded.

Table 2. Spearman rank correlation coefficients r (plus associated p-values) between altitude of sample sites and the site scores along the two ordination axes extracted from NMDS ordinations.

|                  | NMDS Axis 1 | NMDS Axis 2 |
|------------------|-------------|-------------|
|                  | r     | p     | r     | p     |
| Bray-Curtis      | 0.67  | 0.001 | 0.31  | 0.148 |
| MPD All1         | 0.54  | 0.008 | 0.07  | 0.758 |
| MPD Grafen’s Rho | 0.53  | 0.009 | 0.06  | 0.792 |
| MNTD All1        | 0.54  | 0.008 | 0.60  | 0.002 |
| MNTD Grafen’s Rho| 0.49  | 0.019 | 0.49  | 0.017 |

1 Different sets of Hesperiidae assemblages were considered with different measures of species or phylogenetic composition, sampled along altitudinal gradients in Serra do Mar, Paraná, Brazil. ‘All 1’ refers to equal branch lengths assigned to the tree topology while ‘Grafen’s Rho’ refers to Grafen’s branch length transformation method [45]. Correlations that remain significant after applying a table-wide false discovery rate approach are printed in bold face.

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altogether. As shown before [33], neither vegetation type nor altitude influenced the efficiency of sampling skippers. Therefore, differences in sample coverage are not expected to affect our results.

The Bray-Curtis similarity index was used to measure ecological similarities of assemblages. Complementarily, the incidence-based Chao-Soerensen index [39] was also calculated, but this did not yield any deviant patterns (data not shown). Additionally, two indexes were calculated to assess phylogenetic distances between assemblage samples: the Mean Pairwise Distance (MPD) and the Mean Nearest Neighbor Distance (MNTD) [5]. Both indices were compared because they provide different perspectives of phylogenetic similarities, as an overall pattern of relatedness (MPD) or as how closely related species can be (MNTD) [40]. Additionally, MNTD is more affected by changes at the terminal branches of a phylogeny, whereas MPD is more sensitive to changes at the basis of a phylogeny [40].

To calculate measurements of phylogenetic structure, a phylogenetic hypothesis of the sampled taxa is obviously required. Skipper phylogeny is still very imperfectly resolved [29,41]. Based on the latest phylogenetic approach [29], a tentative community supertree was constructed. Topology of high rank taxa was recovered after [29]. Groups (G) and subgroups (SG) stated by Evans [42] were maintained only when they did not conflict with the current tree topology [29]. Species were clumped according to their respective genera (Fig. S2). When genera or species sampled in our study had not been included in the phylogenetic analyses, they were replaced by their closest tribal affiliates, using expert taxonomical arrangement as best possible surrogate of phylogenetic support [40,43]. Because branch length estimates were not available, and even node ages and fossil calibrations are still largely questioned in butterfly systematics [44], equal branch lengths were arbitrarily assigned to the tree topology (which is the same approach as the nodal distance method; [40]). We then compared these results with branch lengths using Grafen’s Rho transformation \( r = 0.5 \) [45] (Fig. S2). Additional branch length transformation methods were also calculated (e.g. Pagel’s and Nee’s methods: [46,47]), but results did not systematically differ from those presented below (data not shown).

To ascertain that phylogenetic structure of communities obtained from the supertree and branch transformation techniques were different from what would be expected at random, we correlated the empirical phylogenetic distance matrices with matrices produced by four distinct null models (means from 1000 randomizations) through Spearman matrix rank correlation tests (procedure RELATE in PRIMER: [48]). These null models included the shuffling of species labels across the phylogeny (null 0), randomization of species from the sample pool (null 1), randomization of species from the phylogeny pool (null 2), or by swapping versions of sample/species matrix (see [49,50] for details). The choice of an adequate null model is relevant, since their different assumptions might lead to contrasting results [51]. Because our phylogenetic trees do not contain branch length information based on genetic differences, we opted for challenging the results obtained through all these null models and then interpret results.

To test whether taxonomic scale may be informative to community similarities between different altitude and vegetation types, the analyses was first performed for the whole family Hesperiidae, and then separately for communities composed only of Pyrginae+Pyrrhopyginae species (i.e. dicot feeders, hereafter called Pyrginae); or only of Hesperini (i.e. monocot feeders). *Urbanus teleus* (Hübner, 1821) was included in the Hesperini analyses, since this species has confirmed host-plant records only in
### Table 4. Spearman matrix rank correlation coefficients $r$ (999 permutations) between phylogenetic distance matrices MPD (mean pairwise distance) and MNTD (mean nearest neighbor distance) and four types of null models: 0–3 [49,50].

|        | Model 0 |        | Model 1 |        | Model 2 |        | Model 3 |        |
|--------|---------|--------|---------|--------|---------|--------|---------|--------|
|        | $r$     | $p$    | $r$     | $p$    | $r$     | $p$    | $r$     | $p$    |
| Hesperidae |         |        |         |        |         |        |         |        |
| MPD    |         |        |         |        |         |        |         |        |
| All 1  | 0.41    | 0.001  | 0.05    | 0.725  | 0.02    | 0.580  | 0.22    | 0.030  |
| Grafen Rho | 0.06    | 0.239  | 0.05    | 0.303  | 0.05    | 0.300  | 0.04    | 0.648  |
| MNTD   |         |        |         |        |         |        |         |        |
| All 1  | 0.83    | 0.001  | 0.48    | 0.001  | 0.48    | 0.001  | 0.50    | 0.001  |
| Grafen Rho | 0.17    | 0.013  | 0.27    | 0.003  | 0.27    | 0.001  | 0.37    | 0.001  |
| Hesperiinae |         |        |         |        |         |        |         |        |
| MPD    |         |        |         |        |         |        |         |        |
| All 1  | 0.60    | 0.001  | 0.00    | 0.524  | 0.06    | 0.782  | −0.23   | 0.996  |
| Grafen Rho | 0.02    | 0.376  | 0.05    | 0.248  | −0.02   | 0.671  | −0.09   | 0.935  |
| MNTD   |         |        |         |        |         |        |         |        |
| All 1  | 0.69    | 0.001  | 0.33    | 0.001  | 0.33    | 0.001  | 0.33    | 0.001  |
| Grafen Rho | 0.28    | 0.001  | 0.18    | 0.003  | 0.28    | 0.001  | 0.20    | 0.002  |
| Pyrginae |         |        |         |        |         |        |         |        |
| MPD    |         |        |         |        |         |        |         |        |
| All 1  | 0.28    | 0.013  | 0.11    | 0.209  | −0.08   | 0.753  | 0.36    | 0.023  |
| Grafen Rho | 0.18    | 0.081  | −0.32   | 0.973  | −0.05   | 0.658  | −0.07   | 0.665  |
| MNTD   |         |        |         |        |         |        |         |        |
| All 1  | 0.44    | 0.001  | 0.24    | 0.018  | 0.18    | 0.097  | 0.31    | 0.001  |
| Grafen Rho | 0.19    | 0.091  | 0.18    | 0.092  | 0.14    | 0.153  | 0.17    | 0.097  |

1 Two different branch length transformation methods were applied with the mean of 999 randomly generated matrices, according to four null models assumptions (see Methods section). ‘All 1’: equal branch lengths assigned to the tree topology; ‘Grafen’s Rho’: Grafen’s branch length transformation method [45]. Significant correlations here indicate that observed sample matrices are not substantially different from random expectations. Null model tests were applied to the three taxa Hesperidae, Hesperiinae and Pyrginae. Correlations that remain significant after applying a table-wide false discovery rate approach are printed in bold face. doi:10.1371/journal.pone.0108207.t004
Poaceae [29], disregarding a probably misleading record from Fabaceae [31]. Pyrginae and Hesperiinae similarity matrices were thereafter compared across the same common sampling sites by Spearman matrix rank correlation, aiming to test whether taxa with such contrasting hostplant associations differ in their ecological and phylogenetic similarity patterns in response to
elevation and vegetation change. Vegetation and altitude were classified as factors and tested in two-way ANOSIM analyses, where R statistics values served as a measure of effect size. Vegetation was divided into forest, early successional vegetation, and grassland, while the elevational categories included in this analyses were ‘low’ (900 m–1150 m), ‘medium’ (1150 m–1400 m), and ‘high’ (1400 m–1650 m; see Table 1).

Analyses were performed at three different taxonomic levels (entire family Hesperiidae, and two major subfamilies), and for two different branch length transformation methods: ‘All 1’: equal branch lengths assigned to the tree topology; ‘Grafen’s Rho’: Grafen’s branch length transformation method [45]. Results printed in bold remained significant after applying a false discovery rate approach.

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Table 5. Two-way ANOSIM results (R statistics and associated p-values), evaluating the effects of vegetation type and altitude on ecological similarities (Bray-Curtis) and phylogenetic distances (MPD) of skipper assemblages.

| Vegetation          | Altitude          | Hesperiidae |          |          | Hesperiidae |          |          |
|---------------------|-------------------|-------------|-----------------|-----------------|-------------|-----------------|-----------------|
| Bray-Curtis         | R^2               | p            | Bray-Curtis     | R^2               | p            | Bray-Curtis     | R^2               | p            |
| All                 | 0.47              | 0.004        | 0.10            | 0.001            | 0.10            | 0.001            |
| Grafen Rho          | 0.35              | 0.006        | 0.01            | 0.5              |

Pyrginae

| Bray-Curtis         | R^2               | p            | Bray-Curtis     | R^2               | p            | Bray-Curtis     | R^2               | p            |
|---------------------|-------------------|-------------|-----------------|-------------------|-------------|-----------------|-------------------|-------------|
| All                 | 0.64              | 0.003        | 0.76            | 0.03              | 0.76            | 0.03             |
| Grafen Rho          | 0.06              | 0.507        | 0.40            | 0.856             |

Results

Ecological and phylogenetic community structure at family level

The total skipper fauna collected along all transects comprised of 1578 specimens representing 155 species. Spatial resolution revealed by NMDS ordinations uncovered a nearly even gradient-like pattern mainly represented by the first ordination axis, along which communities were ordered from high to low altitudes (Fig. 1a, ‘Bray-Curtis’ in Table 2). However, assemblages from grassland sites (from elevations between 1100–1650 m elevation) became segregated from all others, whereas skipper assemblages from forest and early successional vegetation tended to be spatially unordered along both axes. Therefore, an elevational gradient was not evident when looking exclusively to points within each vegetation type, indicating that elevation is secondary to vegetation type in shaping species composition.

When skipper assemblages were ordinated by their phylogenetic distances in the MPD model, spatial patterns were remarkably similar, and NMDS representations achieved almost equal goodness-of-fit (Figs. 1b–c). Application of various branch length transformations did not alter the general pattern observed in NMDS ordinations, but two-dimensional representations had poorer fit (i.e. higher stress values and lower correlations of ordination axis scores with altitude) than if assuming equal branch lengths set to unity. Ordinations based on MPD and MNTD measures, respectively, showed similar patterns with regard to group clustering. Axis 2 significantly correlated with altitude in ordinations based on MNTD, but not for those using MPD (Table 2). Spearman rank correlations between the ecological Bray-Curtis similarity matrix and the four phylogeny-based distance matrices (MPD and MNTD with two branch-length options each) were always highly significant (p<0.001), but MNTD index showed higher coefficient values with Bray-Curtis ecological index (0.84 to 0.87), compared to MPD (0.50 to 0.60) (Table 3). Strikingly, phylogenetic indexes showed the lowest coefficient values of correlation between each other (0.35 to 0.52).

Differences between MPD and MNTD distance measures were particularly obvious with regard to null-model tests (Table 4). The MNTD matrix correlated strongly with random matrices created under any of the four null models compared. In contrast, correlations between the MPD matrix and null models were only significant (and much weaker so) when all branch lengths were assigned to one, i.e. in null models 0 and 3 (shuffling of phylogenetic terminal labels and swapping species between samples, respectively). Therefore, MNTD results cannot be reliably interpreted as relating to ecological conditions and we restricted further considerations to MPD measures.

Effects of vegetation, altitude and taxon scale: Pyrginae vs. Hesperiinae

When analyzed separately, the two subfamilies Hesperiinae and Pyrginae revealed important differences in their patterns of assemblage similarities (Fig. 2). The NMDS ordination of Hesperiinae rendered a similar configuration as the entire family (Fig. 2a). Grassland skipper assemblages were distinctly set apart, with no clear differentiation between communities associated with
Effects of vegetation, altitude and taxon scale: Pyrginae vs. Hesperiinae

While numerous studies have elucidated how elevational gradients influence community similarities, only a few specifically addressed the relevance of vegetation types in relation to merely abiotic gradients on animal assemblages [12, 60–62]. This integration, however, is essential, since high altitude habitats often represent different vegetation types which not necessarily are concordant to climatic change with elevation alone [14]. In cases where vegetational and abiotic dimensions were related to butterfly or moth assemblages, altitude (and climate) usually emerged as the single best predictor of faunal composition or richness instead of habitat type [12, 60, 62]. Accordingly, vegetation type at most emerged as modulating patterns of elevational faunal change [61, 63, 64].

In contrast, vegetation type played the major role with regard to community differentiation in skipper butterflies of Serra do Mar, instead of altitude per se. Along elevational transects of moderate extent, factors beyond mere lapse rate may gain higher relative importance, as temperature differences are less prominent than on high mountain ranges like the Andes or Himalaya [4]. Accordingly, the presence of distinct vegetation types should attain higher weight in elevation gradients studies, addressing not only a novel ecological constraint (e.g. humidity and solar incidence are clearly contrasting between grasslands and forests), but also evolutionary dimensions, such as insect-plant interactions.

Life history traits are particularly related to the diversification of Hesperioidea. The two major subfamilies are rather conservative in larval food plant affiliations with either monocot or dicot plant families [29]. Hesperioidea (exclusive monocot feeders) are a more recent lineage than all other dicot feeders, with almost twice the global species richness of Pyrginae [29]. Hence diversification rate in this group must be distinctly higher, in contrast to the relatively low diversification rate estimated for the entire family [65]. Another unrelated group of monocot feeding butterflies (viz. Satyrinae) displays a similar pattern of speciation, and the expansion of grassland habitats around the world has been linked to this “explosive diversification” [66]. The high number of Hesperioidea species endemic to grasslands ecosystems in south Brazil reinforces this statement.

Hesperioidea assemblages of grassland sites were more distinctly clustered when analyses were based on phylogenetic distances rather than abundance-weighted species lists. This may hint towards a historical relationship with grassland ecosystems through evolutionary time. In other words, skipper species in this highly distinct vegetation type represent lineages with unique evolutionary history [16]. Consequently, environmental filtering caused by altitudinal climate shifts seems to play only a secondary role in structuring those assemblages.

Implications for Conservation

Conservation of natural grasslands ecosystems is of transcontinental concern [67]. Atlantic forests are biodiversity hotspots not only because of high species richness of organisms, but also because of high numbers of endemics [68]. Because mountain grassland patches in the Serra do Mar are inserted amongst this forest landscape, endemic organisms of grasslands are routinely counted to this ecosystem, especially because fine scale distribution of grassland patches is hardly represented in distribution maps. Grassland ecosystems embedded in Atlantic Forest are phylogenetically linked to unique skipper assemblages, in analogy to high-altitude grassland butterflies in the European Alps [23]. The reduction of grassland ecosystems in Atlantic Forest is historically related to climate fluctuations, but nowadays these ecosystems are
also threatened by occasional anthropogenic fires, invasive exotic grasses and extensive tourism [69]. The present findings highlight the need of conserving these high altitude habitats also from the perspective of a unique skipper butterfly fauna. Although difficult to identify, Neotropical skippers comprise rich assemblages (compared for example to frugivorous butterflies), and are suitable sensitive biological indicators in Atlantic Forest [68]. Hence, further taxonomic and ecological studies into this family are desired, since for most skipper species we still lack information on hostplant associations, phylogenetic relationships, and geographical distribution records. Even, quite a number of unknown species remain to be described.

Supporting Information

Figure S1 Explorative NMDS ordination plots of Hesperiidae assemblages along elevational gradients in Serra do Mar, Brazil. Ordination patterns were first assessed based on Bray-Curtis similarities, with samples partitioned into 100 m altitudinal bands (a), into 100 m altitudinal bands, but excluding hilltopping species (b), and partitioned according to altitude and vegetation types (excluding hilltopping species) c). Arrows indicate mountain summit samples.

Figure S2 Phylogenetic relationships of Hesperiidae (Insecta, Lepidoptera) species recorded in Serra do Mar, Paraná, Brazil. Topology of high rank taxa was recovered after Warren et al. [29]. Groups (G) and subgroups (SG) stated by Evans [42] were maintained only when not conflicting with the topology published in Warren et al. [29]. Species were chumped according to its respectively genera. As no branch lengths are still available for skipper phylogeny, equal branch lengths (above) and Grafen’s Rho transformation (below) were arbitrarily assigned to quantify phylogenetic differences between species.

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Author Contributions

Conceived and designed the experiments: EC OHHM MMC KF. Performed the experiments: EC. Analyzed the data: EC KF. Contributed reagents/materials/analysis tools: EC OHHM MMC KF. Wrote the paper: EC OHHM MMC KF.

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