Species Divergence and Phylogenetic Variation of Ecophysiological Traits in Lianas and Trees

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Abstract

The climbing habit is an evolutionary key innovation in plants because it is associated with enhanced clade diversification. We tested whether patterns of species divergence and variation of three ecophysiological traits that are fundamental for plant adaptation to light environments (maximum photosynthetic rate $A_{\text{max}}$, dark respiration rate $R_d$, and specific leaf area SLA) are consistent with this key innovation. Using data reported from four tropical forests and three temperate forests, we compared phylogenetic distance among species as well as the evolutionary rate, phylogenetic distance and phylogenetic signal of those traits in lianas and trees. Estimates of evolutionary rates showed that $R_d$ evolved faster in lianas, while SLA evolved faster in trees. The mean phylogenetic distance was 1.2 times greater among liana species than among tree species. Likewise, estimates of phylogenetic distance indicated that lianas were less related than by chance alone (phylogenetic evenness across 63 species), and trees were more related than expected by chance (phylogenetic clustering across 71 species). Lianas showed evenness for $R_d$, while trees showed phylogenetic clustering for this trait. In contrast, for SLA, lianas exhibited phylogenetic clustering and trees showed phylogenetic evenness. Lianas and trees showed patterns of ecophysiological trait variation among species that were independent of phylogenetic relatedness. We found support for the expected pattern of greater species divergence in lianas, but did not find consistent patterns regarding ecophysiological trait evolution and divergence. $R_d$ followed the species-level pattern, i.e., greater divergence/evolution in lianas compared to trees, while the opposite occurred for SLA and no pattern was detected for $A_{\text{max}}$. $R_d$ may have driven lianas’ divergence across forest environments, and might contribute to diversification in climber clades.

Introduction

Climbing plants, in particular woody vines (lianas), are a distinctive component of mature forests in both tropical and temperate regions [1–3]. Data from long-term plots indicate that the dominance of lianas relative to trees is increasing in tropical forests [4,5]. Moreover, liana abundance is negatively associated with tree carbon storage in tropical forests [6,7]. The climbing habit has independently arisen numerous times throughout plant evolution [1,8], and it seems to be a key innovation in angiosperms: climbing plant lineages have greater species richness than their non-climbing sister groups [9]. Thus, evidence from both ecological and macroevolutionary patterns suggests a performance advantage of lianas over trees.

Explanatory factors for the increased abundance and biomass of lianas in tropical forests include increasing forest disturbance, which increases local resource availability, and rising levels of atmospheric CO₂ [8]. Moreover, increased abundance of lianas in seasonal forests during the dry season, as compared to trees, has been related to their increased efficiency in water uptake and transport, and higher photosynthetic rates [10–12]; but see [13]). Thus, data suggest that lianas are better than trees at exploiting resource pulses. When providing functional arguments for the key innovation of the climbing habit (sensu [14]), Gianoli [9] suggested that ecological specialization may arise as a consequence of an hypothetically expanded light niche of lianas in the forest, which would result from the co-occurrence of unsupported (creeping) and supported (climbing) individuals that go up and down the forest canopy. This would maximize interactions with a wide array of antagonistic and mutualistic species [15,16] that, in turn, might promote diversification [17]. It is increasingly recognized that purported evolutionary key innovations may be tested at an ecological time scale [14,18–20].

Ecophysiological traits are fundamental components of plant adaptation to the environment [21,22]. Specifically, $A_{\text{max}}$ (maximum photosynthetic rate), $R_d$ (dark respiration rate) and SLA (specific leaf area) play a key role in the phenotypic adjustment to heterogeneous light environments in both lianas and trees [23–26]. Thus, they reflect the balance between carbon gain ($A_{\text{max}}$) and carbon use ($R_d$), and the allocation of leaf biomass to light interception (SLA), which together determine plant growth and performance across light environments [27,28]. Importantly, variation in plant functional traits observed at the population level is likely to be paralleled by evolutionary divergences under contrasting environments [29]. Moreover, the analysis of the phylogenetic structure of communities can provide insights to our understanding of trait evolution [30]. Recent studies have addressed phylogenetic variation in ecophysiological traits in climbing plants and trees [31–34], but their approach has been either exploratory (aiming to report global patterns) or method-
ological (testing new analytical tools); to our knowledge, a hypothesis-driven analysis is wanting.

Using data reported for several liana and tree species coexisting in tropical and temperate forests, and focusing on three key ecophysiological traits involved in plant adaptation across forest light gradients: \( A_{\text{max}} \), \( R_{\text{d}} \) and SLA, we hereinafter compare lianas and trees in terms of trait evolutionary rates, phylogenetic diversity, phylogenetic trait diversity, and the phylogenetic signal. Thus, we compared the rate at which variance in the traits is accumulated among species per unit time at the tips of the phylogenetic tree [35,36]. We also evaluated how similar is the average pair of species of lianas and trees both in terms of mean phylogenetic distance and trait variation [37]. We finally evaluated in lianas and trees the tendency for phylogenetically related species to resemble each other, i.e., the phylogenetic signal [38]. We tested the hypotheses that if the climbing habitat enhances clade diversification [9], and ecological divergence is the process underlying this pattern, then lianas should show higher trait evolutionary rates and greater species and trait divergence than trees under common environmental scenarios.

Materials and Methods

Data collection

We searched the literature for field studies in forest ecosystems where lianas and trees were analyzed for at least one of three ecophysiological traits: \( A_{\text{max}} \) on an area basis, \( R_{\text{d}} \) and SLA. We only chose those studies carried out in mature forests, where light heterogeneity across microsites is the greatest [3]. We only included native species because they have a long history of adaptation to the environment. We focused on angiosperms because of the availability of tools to reconstruct their phylogenetic history and estimate trait evolution (see below). The final data set included a pool of 63 liana species and 71 tree species belonging to four tropical forests (Gamboa, Panama; San Lorenzo, Panama; Riberalta, Bolivia; Xishuangbanna, China) and three temperate forests (Yakushidake, Japan; Beltsville, USA; Puyehue, Chile). We pooled species from all sites into growth forms, thus we had one liana “super-community” and one tree “super-community”. Phylogenetic analyses were conducted on these super-communities (see below). Detailed information, including study species, traits, sites, and data sources, is available in Supporting Information S1.

Phylogeny reconstruction

We produced a phylogeny of all species using a backbone tree based on the angiosperm megatree provided by the PhyloDiversity Network in cooperation with the Angiosperm Phylogeny Group (APG; http://www.mobot.org/MOBOT/research/APweb/). Our tree was generated using Phylomatic (http://www.phylodiversity.net/phylomatic/phylomatic.html), a program that returns a working phylogenetic tree after matching the genus and family names of study species to those contained in the angiosperm phylogeny [39]. Comparative inferences require branch lengths for the tree, which were calculated based on the branch length adjustment algorithm (BLADJ) implemented in Phylocom v. 4.2 (www.phylodiversity.net/phylocom) [40]. This algorithm fixes a subset of nodes in the tree to specified ages and evenly distributes the ages to the remaining nodes. Age estimates for major nodes in our tree were taken from [41]. To avoid inaccuracies in tree calibration and to have an updated version of our tree, we corrected the ages file with age estimates in [41] included in Phylocom. Corrections followed procedures suggested recently [42]. We also checked and updated age estimates of internal order-level clades according to a net diversification rate estimate of angiosperms [43]. The few polytomies in the working tree were resolved randomly using the multi2phy function in R. Values of functional traits of closely related species resulting from such random resolutions were very similar, so results of the final comparative tests were highly robust to topological uncertainty.

These and all subsequent analyses were conducted using the R statistical environment version 3.0.2 [44]. Reconstructed phylogenetic trees with associated trait variation are shown in Figures 1-3.

Rate of trait evolution

To assess differences in ecophysiological trait evolution between liana and tree species, we compared estimates of evolutionary rate for \( A_{\text{max}} \), \( R_{\text{d}} \) and SLA. To this end, we first used stochastic character mapping, a Bayesian method that uses Monte Carlo simulations to sample the posterior probability distribution of ancestral states and timings of transitions on phylogenetic branches under a Markov process of evolution [45,46]. We built stochastic character-mapped reconstructions for each trait/growth form combination using the make_simmap function in the phytools package of R [36]. We thus simulated character history evolution of all three traits in relation to growth form as an initial step, as suggested recently [35,47]. To test the hypothesis that a discrete character state had influenced the rate of a continuous character, one should first stochastically map the discrete trait (e.g., climbing habit), and then test if one state of the discrete character has a different evolutionary rate for the continuous trait of interest (e.g., ecophysiological trait) than the other discrete state [35,47].

The resulting reconstructions of trait states and phylogeny represented a set of phylogenetic topologies, branch lengths and growth forms sampled in proportion to their posterior probabilities. Reconstructions were then used in subsequent analyses as a way of integrating over uncertainty in phylogeny and ancestral states. Finally, we fitted the evolutionary models of character history on the trees to trait data using a likelihood method [35]. This is a maximum likelihood approach that estimates rates of evolution (\( \sigma^2 \)). The parameter \( \sigma^2 \) was calculated using the function brownie.lite in the phytools package [36]. \( \sigma^2 \) is interpreted as the Brownian motion process most likely to have produced the data at the tips of the tree, i.e., the rate at which variance is accumulated among species over time. 95% confidence intervals were calculated for each \( \sigma^2 \) to infer differences between lianas and trees in ecophysiological trait evolution.

Trait and phylogenetic diversity

To compare the phylogenetic relatedness among liana species against the phylogenetic relatedness among tree species we used measures of phylogenetic structure. Specifically, we calculated, based on a phylogenetic distance matrix, the mean phylogenetic distance (MPD) and the standardized effect size of the mean phylogenetic distance (SES_MPD) [48] between pairs of species for each group. Interspecific phylogenetic distance matrices were obtained from the reconstructed tree of phylogenetic relationships among taxa using the cophenet function in R and unweighted pair-group average (UPGMA) as the clustering method.

Standardized effect sizes describe the difference between average phylogenetic distances in the observed super communities or groups (lianas and trees) compared to null distributions generated for each group with randomization procedures, standardized by the standard deviation of phylogenetic distances in the null data [48]. We compared observed mean distances (branch length) against a null model generated by calculating 999 times the mean phylogenetic distance between 8911 random pairs of species (without replacement) drawn from the matrix of
phylogenetic distances between all liana and tree species. In all cases 999 iterations were found to be suitable for our randomization procedures as they were sufficient to attain convergence. The null model was constructed by reshuffling the distance of species labels across the phylogenetic tree using the `ses.mpd` function and the `taxa.labels` algorithm of the `picante` package of R. Positive values of SES MPD (mpd.obs.z) and high quantiles (p-values, 0.95) indicate significant phylogenetic evenness, while negative values of SES MPD and low quantiles (p-values, 0.05) indicate significant phylogenetic clustering [48]; these outcomes correspond to scenarios where species are more distantly or more closely related than expected by chance, respectively [48,49]. Authors often refer to (weak) evenness or clustering when p-values are slightly lower than 0.95 or slightly higher than 0.05, respectively (e.g., [50,51]). Finally, to assess how similar are the average pair of species within each group in terms of ecophysiologial traits; we used the SES MPD as a trait diversity measure. This was done by replacing the phylogenetic distance matrix in the analysis with a trait distance matrix, and proceeding accordingly to calculate standardized values of mean phylogenetic trait distance (SESMTD). These results are interpreted in the same way as those of SES MPD with regard to phylogenetic evenness or clustering [48].

**Phylogenetic signal**

To quantify the degree to which phylogenetic relatedness predicts the similarity of species in functional traits for both trees and lianas, we calculated separately phylogenetic signal for \( \lambda_{\text{max}} \), \( R_d \), and SLA. Phylogenetic signal indicates to what extent phenotypic expression is explained by the lineage to which a species belongs. Figure 1. Phylogenetic relationships among tree and liana species and species values of dark respiration rate (\( R_d \)). Grey circle size represents the proportional magnitude of the trait across species. Square tip symbols represent climbing habit (grey squares = liana, black squares = tree). Timescale is in millions of years before present. doi:10.1371/journal.pone.0099871.g001

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**Figure 1. Phylogenetic relationships among tree and liana species and species values of dark respiration rate (\( R_d \)).** Grey circle size represents the proportional magnitude of the trait across species. Square tip symbols represent climbing habit (grey squares = liana, black squares = tree). Timescale is in millions of years before present. doi:10.1371/journal.pone.0099871.g001
species belongs, and it can be compared among clades and among traits [52]. We quantified phylogenetic signal using both Blomberg’s K [50] and Pagel’s λ [51] statistics for quantitative traits. To calculate these parameters, we first pruned two separate phylogenies, one for the group of lianas and one for the group of trees, using the original tree as a base phylogeny. Then we pruned a tree for each group-trait combination independently, removing taxa for which trait information was not available. The number of species included in each trait/plant growth habit analysis ranged from 26 (Rd/lianas) to 67 (A max/trees), thus meeting the N>20 threshold to achieve good statistical power [38].

Values of $K = 1$ imply that a trait shows exactly the amount of phylogenetic signal expected under a null, stochastic branch of character evolution (Brownian motion evolution) [38]. $K$-values $> 1$ and $<1$ imply that close relatives are more similar and less similar, respectively, than expected under a Brownian motion model of trait evolution [38]. If $K$ does not differ from zero it is concluded that the trait has no phylogenetic signal. Statistical significance of $K$ [38] was assessed via permutation tests with 1000 randomizations. The significance of the phylogenetic signal was based on the variance of phylogenetically independent contrasts relative to tip shuffling randomization implemented by the {phylignal} function of the {picante} package in R [48]. $P$-values were determined by comparing the variance of standardized independent contrasts for the tip values against variances for randomized data.

The parameter $\lambda$ scales tree structure in terms of expected variances and covariances in trait change [54]. Thus, $\lambda$ is a phylogenetic transformation that maximizes the likelihood of the data given a Brownian motion model [54]. When $\lambda = 1$, the trait is consistent with a Brownian motion evolution based on branch lengths represented by the variance-covariance in trait change. Values between 0 and 1 indicate less phylogenetic signal than expected under a Brownian motion model, while values $>1$ indicate more signal than expected, although $\lambda$ is not always defined for values greater than one [54]. Values of $\lambda$ were estimated using the fitContinuous function of the {geiger} package. To determine the significance of $\lambda$, as an indicator of phylogenetic signal, we compared the maximum likelihood estimate of $\lambda$ against the maximum likelihood of models when $\lambda = 1$ using likelihood ratio tests (LRT).

### Results and Discussion

#### Rate of trait evolution

In general, lianas and trees presented homogenous evolution of ecophysiological traits. In all cases evolutionary rates, as estimated by $\sigma^2$, were not significantly different from a single-rate Brownian motion process of evolution (Table 1). Parameter estimate values of $\sigma^2$, however, did differ between lianas and trees in two of the three ecophysiological traits considered (Table 1). The evolutionary rate for dark respiration rate (Rd) in lianas was 1.8 times greater than in trees. In the case of the biomass allocation trait (specific leaf area, SLA), the evolutionary rate was 1.2 times greater in trees than in lianas. Evolutionary rates for maximum photosynthetic rate ($A_{\text{max}}$) did not differ between lianas and trees; overall, this trait showed the lowest evolutionary rate among the traits considered (Table 1).

The patterns observed suggest that for all ecophysiological traits a change along any given branch in the phylogeny is independent of both previous changes and changes in other branches of the reconstructed tree. Evolutionary rates in both lianas and trees did not differ from a single-rate Brownian motion model of evolution, which assumes that variance among species in the phylogenetic tree accumulates as function of their time of independent evolution [55]. Thus, it cannot be ruled out that ecophysiological traits evolve at a constant rate over time. A Brownian motion process, however, is not equal to a neutral model of evolution. Brownian motion simply describes the distribution of observed trait changes and may be consistent with adaptive models of evolution [53,56]. Therefore, natural selection could be a plausible force behind the alteration in rate change of traits in relation to growth form (climbers vs. non-climbers).

Evolutionary rates ($\sigma^2$) differed between lianas and trees in two of the three ecophysiological traits considered, but in opposite trends. Thus, Rd evolved at a higher rate in lianas, while SLA evolution occurred at a higher rate in trees. This suggests that the outcome of modifications in the selective regime related to the climbing habit depends on the particular plant traits that are under selection (gas-exchange traits vs. biomass allocation traits). Gas-exchange traits have been shown to be of selective value for the exploitation of light availability in mature forests for trees [57], vines [58] and ferns [59]. Our findings suggest that climbers are more evolutionary responsive with regard to Rd than trees. Assuming that (adaptive) ecological speciation is the process behind species divergence in this trait [20], the next step would be to address whether this results from a greater magnitude of selection on Rd or from greater trait heritability [21,60]. Conversely, SLA showed a higher greater evolutionary rate across tree species. This somewhat supports the view of SLA as an essential attribute for tree performance and carbon gain [57,61].

#### Trait and phylogenetic diversity

We found that mean phylogenetic distance (MPD, non-standardized values) was greater among liana species (259.9 Myr) than among tree species (229.6 Myr). Moreover, there was a clear-cut difference between lianas and trees in the standardized mean phylogenetic distance among species (SESMPD). Whereas lianas showed greater distances between species relative to the null model (SESMPD = 2.271; $p$-value = 0.99), i.e., phylogenetic evenness, trees showed a pattern of phylogenetic clustering (SESMPD = −3.622; $p$-value = 0.006).

Lianas and trees differed in their patterns of trait diversity. For one of the gas-exchange traits (Rd), lianas showed phylogenetic evenness (SESMTD = 1.266; $p$-value = 0.993), which means that trait dissimilarity among liana species was higher than expected by chance, while trees showed phylogenetic clustering (SESMTD = −1.863, $p$-value = 0.039), indicating that tree species were more phenotypically similar than expected by chance (Figure 1). In contrast, for the biomass allocation trait (SLA), lianas exhibited phylogenetic clustering (SESMTD = −1.194, $p$-value = 0.122) and trees showed phylogenetic evenness (SESMTD = 1.193, $p$-value = 0.877) (Figure 2). Finally, the other gas-exchange trait, $A_{\text{max}}$, did not show phylogenetic structure in both lianas (SESMTD = 0.996, $p$-value = 0.536) and trees (SESMTD = −0.598, $p$-value = 0.277) (Figure 3).

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**Figure 2. Phylogenetic relationships among tree and liana species and species values of maximum photosynthetic rate ($A_{\text{max}}$).** Grey circle size represents the proportional magnitude of the trait across species. Square tip symbols represent climbing habit (grey squares = liana, black squares = tree). Timescale is in millions of years before present.

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First, in agreement with the study hypotheses, we found greater phylogenetic distance among species within the group of lianas (63 species) than within the group of trees (71 species). This agrees with a recent study in Australian rainforests, where standardized values of mean phylogenetic distance indicated that climbers show weak evenness or no phylogenetic structure, while trees/shrubs show weak to significant phylogenetic clustering [30]. Second, the average phenotypic distance among species for one gas-exchange trait (but not for the other two ecophysiological traits) was greater in the phylogenetic tree of lianas than in that of trees. The environmental gradient experienced by the study species was roughly the same for lianas and trees because data were obtained from sites where trees and lianas coexisted. Therefore, these patterns of (partial) increased phenotypic distance and greater phylogenetic divergence in lianas are consistent with the notion that lianas have a greater differentiation potential than trees [9]. Several plant attributes have been associated with evolutionary rates in angiosperms. For instance, it has been shown that trees and shrubs have lower rates of molecular evolution than herbaceous plants [62,63], and that taller plants have lower rates of molecular evolution [64]; in both cases the outcome is thought to be linked to differences in generation time, which in turn is related to mutation rate. In our study all climbers were woody species (lianas) so herbaceousness should not be a confounded factor. However, there is no available information to reject the possibility that there were longer generation times in the group of trees (see General Conclusions). As to the plant height factor, it is a rather problematic issue, because trees are usually taller than lianas in terms of freestanding height, but if total length is considered, then canopy lianas may be taller. Both issues deserve further scrutiny.

In the realm of community phylogenetics, patterns of phylogenetic evenness in resource-use traits are often interpreted to reflect niche differentiation processes [49]. If trait-based niche differentiation facilitates evolutionary responses to divergent selection, which in turn may lead to incipient speciation [65], then our results of phylogenetic evenness in a liana ecophysiological trait might be linked to the ecological/evolutionary processes that underlie the key innovation of the climbing habit in plants [9]. Whereas results of trait phylogenetic distance suggest that $R_d$ may have played a role as driver of lianas’ adaptive divergence, SLA showed greater phenotypic divergence among tree species, as was shown for rates of trait evolution (see above). However, this trait distribution pattern across the phylogenetic tree was not accompanied by an overall greater phylogenetic distance among tree species. This might be interpreted as SLA contributing to tree adaptation to environmental challenges at local scales but do not driving taxonomic divergence across clades.

### Phylogenetic signal

Overall, lianas and trees presented mixed but comparable patterns of phylogenetic signal (or lack thereof) in ecophysiological traits (Table 2). In lianas, $\Lambda_{\text{max}}$ showed no significant phylogenetic signal, but values were lower than expected under a Brownian model of evolution (with $K<1$ and $\lambda<1$). $R_d$ showed mixed results, with significant phylogenetic signal indicated by $K$, and lower than expected under a Brownian model of evolution, but no significant signal as indicated by $\lambda$ ($=1$). There was no phylogenetic signal detected for specific leaf area (SLA) using $K$ but strong signal using $\lambda$, and lower than expected under a Brownian model of evolution (Table 2). In trees, whereas no phylogenetic signal was found for both $\Lambda_{\text{max}}$ and $R_d$ as indicated by $K$ and $\lambda$, a significant phylogenetic signal was found for SLA when $K$ is considered (with $K<1$) but no signal was detected by $\lambda$, with values lower than expected under a Brownian model of trait evolution ($\lambda=0$) (Table 2). Summarizing, in all cases both lianas and trees tended to show patterns of ecophysiological trait variation among species that were independent of phylogenetic relatedness.

Our results are consistent with the general pattern that physiological traits tend to show low values of phylogenetic signal [38]. A global analysis of trait variation in climbing plants reported that SLA showed no phylogenetic signal [34], as found in the present study. Likewise, in agreement with our results, a global-scale study in Angiosperms reported that $\Lambda_{\text{max}}$ (on an area basis) showed no consistent phylogenetic signal [32]. Conversely, a genus-level study in trees [31] found that $\Lambda_{\text{max}}$ exhibited significant phylogenetic signal, which seemingly opposes our findings. However, this study used a metric other than Blomberg’s $K$ and Pagel’s $\lambda$, and given that different indices of phylogenetic signal often lead to contrasting outcomes ([68]; and Table 2), these results are not necessarily contrary to those reported here. Another group of ecophysiological traits that could have been studied to seek phylogenetic and evolutionary differences between trees and lianas is that of hydraulic characters. Regarding hydraulic traits, lianas have wider and longer vessels compared to trees, features that enable them to supply a large leaf area with a relatively small allocation to xylem tissue [33,66,67]. However, xylem vessel length did not show significant phylogenetic signal in a recent global analysis including lianas, shrubs and trees [33].

Results indicate that, in both lianas and trees, ecophysiological traits related to light use and carbon economy have undergone evolutionary trajectories different to those expected after phylogenetic relationships, assuming a Brownian motion model of trait evolution [68]. These phylogenetic signal results do not match the patterns of trait divergence and trait evolutionary rates found here. Although under some circumstances (e.g., fluctuating selection in related lineages) a negative association between $K$ and evolutionary rate may be found [32,55], it is generally considered that changes in trait evolutionary rates—and ensuing phenotypic divergence—do not influence phylogenetic signal for continuous characters [32,55].

### General conclusions

Lianas and trees differ in a number of anatomical, physiological, morphological and life history traits [69–71]. Among the main differences, trees show a greater allocation of biomass (and carbon) to stems and lianas have lower costs of height gain and larger total leaf area potential. Moreover, compared to shrubs and trees, lianas have lower leaf mass per area (LMA, the inverse of SLA), higher foliar N and higher mass-based photosynthetic rate, which is consistent with the characterization of lianas as fast metabolism/rapid turnover species [71]. This could be related to hypothetical differences in generation time between lianas and trees that could explain their differential evolutionary rates, as shown here. Nonetheless, when it comes to explain species distribution across the light gradient in forests [72], the life history trade-off between juvenile growth and survival is observed alike in trees and lianas [73].
### Table 1. Parameter estimates and 95% confidence intervals (CI) of the evolutionary rate ($\sigma^2$) for ecophysiological traits in lianas and trees.

| Trait | Lianas | | | | Trees | | | |
|-------|--------|---|---|---|--------|---|---|---|
|       | $\sigma^2$ | 95% CI | $\chi^2$ | $p$-value | $\sigma^2$ | 95% CI | $\chi^2$ | $p$-value |
| $A_{\text{max}}$ | 0.075 | 0.072–0.079 | -6.8 | 0.948 | 0.075 | 0.072–0.078 | -10.6 | 0.995 |
| $R_d$ | 0.148 | 0.132–0.165 | -4.1 | 0.999 | 0.083 | 0.076–0.091 | -13.7 | 0.991 |
| SLA | 0.093 | 0.087–0.097 | -11.8 | 0.999 | 0.103 | 0.099–0.108 | -11.2 | 0.999 |

$A_{\text{max}}$ = maximum photosynthetic rate; $R_d$ = dark respiration rate; SLA = specific leaf area. Values of $\sigma^2$ represent rates at which variance accumulates among species per unit time through a phylogeny with branch lengths in units of millions of years. $p$-values are for likelihood ratio tests against the chi-square distribution between a single-rate (homogenous) and a heterogeneous Brownian motion process. Bold cells indicate $\sigma^2$ values for which 95% CI do not overlap between lianas and trees.

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### Table 2. Phylogenetic signal, quantified as Blomberg’s $K$ and Pagel’s $\lambda$, for three ecophysiological traits in lianas and trees.

| Trait | Lianas | | | | Trees | | | |
|-------|--------|---|---|---|--------|---|---|---|
|       | $K$ | $p$-value | $\lambda$ | $p$-value | $K$ | $p$-value | $\lambda$ | $p$-value |
| $A_{\text{max}}$ | 0.518 | 0.157 | 0.338 | 0.281 | 0.308 | 0.459 | 0.223 | 0.503 |
| $R_d$ | **0.874** | 0.015 | 1.000 | 0.096 | 0.284 | 0.573 | 0.125 | 0.713 |
| SLA | 0.511 | 0.166 | <0.001 | <0.001 | **0.638** | 0.001 | 0.096 | 0.282 |

$A_{\text{max}}$ = maximum photosynthetic rate; $R_d$ = dark respiration rate; SLA = specific leaf area. Significant phylogenetic signals are shown in bold.

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Phylogenetic information is increasingly used to test macroevolutionary hypotheses of trait evolution [74–76]. The study hypotheses, arising from the macroevolutionary pattern of increased taxonomic diversification in lianas [9], received mixed support. Overall, mean phylogenetic distance among liana species was larger than that of trees. Lianas showed a high evolutionary rate for a gas-exchange trait (Rd), but the biomass allocation trait (SLA) evolved at a higher rate in trees. Likewise, average trait divergence across the phylogenetic tree was greater in lianas for Rd, but it was greater in trees for SLA. Therefore, although we have found support for the expected pattern of increased species divergence in lianas compared to trees, we did not find consistent patterns regarding ecophysiological trait evolution and divergence. Rd followed the species-level patterns, i.e., greater divergence/evolution in lianas compared to trees, while the opposite was found for SLA. R may have driven lianas’ divergence across forest environments and, furthermore, might contribute to the pattern of increased diversification in climber clades.

Supporting Information

Acknowledgments

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Author Contributions

Conceived and designed the experiments: EG CS-L RSR. Performed the experiments: EG CS-L RSR. Analyzed the data: EG CS-L RSR. Contributed reagents/materials/analysis tools: EG CS-L RSR. Wrote the paper: EG CS-L RSR.

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