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Eleinis Ávila-Lovera

Gregory R. Goldsmith

Kathleen M. Kay

Jennifer L. Funk

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Comments

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STUDIES

Above- and below-ground functional trait coordination in the Neotropical understory genus *Costus*

Eleinis Ávila-Lovera^{1,3,*}, Gregory R. Goldsmith¹, Kathleen M. Kay² and Jennifer L. Funk^{1,4}

¹Schmid College of Science and Technology, Chapman University, Orange, CA 92866, USA,

²Department of Ecology and Evolutionary Biology, University of California, Santa Cruz, CA 95060, USA

³Present address: Smithsonian Tropical Research Institute, PO Box 0843-03092, Balboa, Ancon, Republic of Panama

⁴Present address: Department of Plant Sciences, University of California, Davis, CA 95616, USA

*Corresponding author's e-mail address: eleinis.avila@gmail.com

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Abstract

The study of plant functional traits and variation among and within species can help illuminate functional coordination and trade-offs in key processes that allow plants to grow, reproduce and survive. We studied 20 leaf, above-ground stem, below-ground stem and fine-root traits of 17 *Costus* species from forests in Costa Rica and Panama to answer the following questions: (i) Do congeneric species show above-ground and below-ground trait coordination and trade-offs consistent with theory of resource acquisition and conservation? (ii) Is there correlated evolution among traits? (iii) Given the diversity of habitats over which *Costus* occurs, what is the relative contribution of site and species to trait variation? We performed a principal components analysis (PCA) to assess for the existence of a spectrum of trait variation and found that the first two PCs accounted for 21.4 % and 17.8 % of the total trait variation, respectively, with the first axis of variation being consistent with a continuum of resource-acquisitive and resource-conservative traits in water acquisition and use, and the second axis of variation being related to the leaf economics spectrum. Stomatal conductance was negatively related to both above-ground stem and rhizome specific density, and these relationships became stronger after accounting for evolutionary relatedness, indicating correlated evolution. Despite elevation and climatic differences among sites, high trait variation was ascribed to individuals rather than to sites. We conclude that *Costus* species present trait coordination and trade-offs that allow species to be categorized as having a resource-acquisitive or resource-conservative functional strategy, consistent with a whole-plant functional strategy with evident coordination and trade-offs between above-ground and below-ground function. Our results also show that herbaceous species and species with rhizomes tend to agree with trade-offs found in more species-rich comparisons.

Keywords: Ecophysiology; functional strategies; rhizome traits; specific root length; stem specific density; tropics; variance component analysis.

Introduction

Functional traits are defined as any morphological, physiological or phenological characteristic that indirectly influences fitness through their effects on growth, reproduction and survival

(*Violle et al. 2007*). However, traits do not usually work in isolation. Indeed, ecologists often use suites of correlated functional traits, i.e. plant strategies, to provide insights into

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the way that plants acquire, use and conserve resources (Reich et al. 2003; Wright et al. 2004; Zanne et al. 2010; Reich 2014). Resource-acquisitive strategies involve traits that allow for fast acquisition and use of resources (high turnover), which in turn results in fast growth rates and low capacity to tolerate stress. On the other hand, resource-conservative strategies involve traits that allow for slow use of resources and slow growth, but with increased capacity to tolerate stress. For instance, leaf traits related to carbon and nutrient economy have been found to covary among plant species worldwide resulting in a 'leaf economics spectrum' (LES) (Reich et al. 1997; Westoby et al. 2002; Wright et al. 2004). In this spectrum, leaves with high specific leaf area (SLA) and high nutrient concentration also have high photosynthetic and respiration rates, but a short lifespan (resource-acquisitive strategy), whereas leaves with low SLA and nutrient concentration also have low photosynthetic and respiration rates, but a long lifespan (resource-conservative strategy) (Reich et al. 1997; Wright et al. 2004). This economic spectrum has been extended to stems (Chave et al. 2009) and, with mixed support, to roots (Kong et al. 2015, 2019; Roumet et al. 2016). The existence of spectra of variation within organs suggests that coordination (positive covariance) and trade-offs (negative covariance) among traits limit organ function, and likely whole-plant function.

Despite the importance of below-ground processes on plant physiology and performance (Laliberté 2017), below-ground traits have only recently been incorporated into plant functional strategy frameworks (McCormack et al. 2012; Mommer and Weemstra 2012; Weemstra et al. 2016). Root traits that allow plants to acquire large amounts of water and nutrients, such as high root length density (cm root cm⁻³ soil) or specific root length (SRL; m g⁻¹), should be beneficial in resource-rich environments, whereas traits that allow plants to avoid water stress by accessing more stable sources of water and restricting resource loss (or encouraging conservation), such as high root diameter or root tissue density (RTD; g cm⁻³), should be beneficial in resource-poor environments (Bowsher et al. 2016). Specific root length is suggested to be the below-ground analogue to SLA, but SRL is often orthogonal to the main axis of root variation usually formed by root diameter and RTD (Freschet et al. 2010; Liu et al. 2010; Bowsher et al. 2016; Kramer-Walter et al. 2016; Weemstra et al. 2016), complicating efforts to identify a single axis characterizing below-ground function that aligns with above-ground function (Reich 2014). More recently, the inclusion of mycorrhiza (e.g. mycorrhizal colonization) in root functional trait frameworks has shown the existence of a fungal collaboration gradient that dominates the root economic spectrum in a large data set of species (McCormack and Iversen 2019; Bergmann et al. 2020; Weigelt et al. 2021). However, the question remains as to what extent plants align their above-ground and below-ground traits, i.e. are traits coordinated across organs reflecting a single unified whole-plant functional strategy?

Other below-ground traits have been completely left out from recent plant functional strategy frameworks. For example, some plant species use below-ground stems (i.e. rhizomes) as important anchoring structures, for carbohydrate and water storage, and for vegetative reproduction. Traits such as rhizome water content (RhWC; %) and rhizome specific density (RhSD; g cm⁻³) provide information on water storage capacity or investment in structural carbon in rhizomes. Evaluating rhizome traits of perennial herbs can give us insights into the growth strategy of these plants. We expect that species with below-ground resource-conservative traits would also have low RhWC and high RhSD, indicating

a greater investment in structural carbon rather than in water storage, with opposite values of traits corresponding to resource-acquisitive. These rhizome traits would also align with fine-root traits, and above-ground traits, if a whole-plant economic spectrum does exist (Reich 2014).

We studied closely related species from the genus *Costus* because (i) species co-occur at multiple sites, (ii) a well-resolved phylogeny exists (Vargas et al. 2021) and (iii) they have speciated rapidly in recent history, giving opportunities to study traits that have recently evolved. By using congeneric species, we reduced the effect of large divergence patterns (e.g. across families or life forms) on observed trait values.

Given the potential importance for above-ground and below-ground relationships in determining whole-plant functional strategies, we studied a group of closely related tropical species in the genus *Costus* living in contrasting habitats to answer the following questions: (i) Do congeneric species show above-ground and below-ground trait coordination and trade-offs consistent with theory of resource acquisition and conservation? (ii) Is there correlated evolution among traits? (iii) Given the diversity of habitats over which *Costus* occurs, what is the relative contribution of site and species to trait variation? Given the habitat variability across sites, and the morphological differences among species living at the same site, we hypothesized that *Costus* will show a diversity of strategies (i.e. combination of traits), ranging from resource-acquisitive to resource-conservative strategies that matches their habitats. For example, species in wet habitats are expected to have traits that allow for greater water use, such as low rhizome and stem specific density and high stomatal conductance. We also hypothesized that species identity plays a significant role in explaining trait variation. Research to date on tropical plant functional traits has largely been confined to woody species and to the context of community assembly (Kraft et al. 2008), with little work on how functional traits and trade-offs can help understand the physiological mechanisms by which herbaceous species respond to environmental variation.

Methods

Study sites

We measured leaf, above-ground stem, below-ground stem (rhizome from now on) and fine-root traits on individual plants of 17 species of *Costus* in six sites in Costa Rica and two sites in Panama (6 of these 17 species were present at more than one site; Table 1) during the rainy season. Field sites varied in elevation (Fig. 1), which affects mean annual temperature (MAT) and precipitation (MAP) as well as precipitation seasonality. We used the latitude and longitude of the sampled individuals to download bioclimatic variables from WorldClim 2.0 (Fick and Hijmans 2017) and then averaged by species and sites. Lowland wet forests have high MAT and MAP and low precipitation seasonality, whereas highland wet montane and pre-montane forests have relatively lower MAT and MAP and high precipitation seasonality (E. Ávila-Lovera et al., submitted for publication). Lowland seasonal forests have a more pronounced dry season than lowland wet forests (see Supporting Information—Table S1 for more bioclimatic data of field sites). Permit information can be found in Supporting Information—Notes S1.

Study species

The genus *Costus* (Costaceae) comprises approximately 60 species in the Neotropics, and it occupies habitats that range from lowland to montane forests, from deep shade understory to

Table 1. List of species studied, abbreviation use in figures, sites where they are present, elevation and habitat type. BDT: Bocas del Toro (Panama), LA: Las Alturas (Costa Rica), LC: Las Cruces (Costa Rica), LG: La Gamba (Costa Rica), LS: La Selva (Costa Rica), MV: Monteverde (Costa Rica), PLR: Pipeline Road (Panama), TG: Tortuguero (Costa Rica).

Species	Abbreviation	Site	Elevation (m asl)	Habitat type
<i>C. aff. wilsonii</i>	aff.wils	MV	1519.3	Montane forest, streams
<i>C. alleni</i>	alle	PLR	113.5	Wet forest, deep shade
<i>C. bracteatus</i>	brac	LS	77.5	Wet forest
		TG	12.3	Wet forest
<i>C. guanaiensis</i> var. <i>macrostobilus</i>	guan	PLR	69.2	Seasonal forest
<i>C. laevis</i>	laev	LC	1216.8	Pre-montane forest, streams
		LG	113.2	Wet forest
		LS	61.5	Wet forest
		PLR	92.0	Seasonal forest
		TG	20.0	Wet forest
<i>C. lima</i>	lima	LG	82.0	Wet forest, riverine
<i>C. malortieanus</i>	malo	LS	56.0	Wet forest
<i>C. montanus</i>	mont	MV	1569.3	Montane forest
<i>C. osae</i>	osae	LG	122.2	Wet forest, streams
<i>C. plicatus</i>	plic	LG	112.2	Wet forest, riverine
<i>C. pulverulentus</i>	pulv	LG	130.5	Wet forest
		LS	68.8	Wet forest, treefall gaps
		PLR	73.0	Seasonal forest
		TG	16.4	Wet forest
<i>C. ricus</i>	ricu	LG	211.25	Wet forest
<i>C. scaber</i>	scab	LG	227.7	Wet forest
		LS	78.4	Wet forest, streams
		PLR	74.8	Seasonal forest
		TG	26.3	Wet forest
<i>C. stenophyllus</i>	sten	LG	180.2	Wet forest
<i>C. villosissimus</i>	vill	PLR	70.0	Seasonal forest, forest edges
<i>C. wilsonii</i>	wils	LA	1559.0	Montane forest
		LC	1216.8	Pre-montane forest
<i>C. woodsonii</i>	wood	BDT	0	Beach
		TG	3.8	Beach

high light gaps and from dry forest edges to ravines and swamps (Kay et al. 2005). Therefore, it is an interesting group to study plant trait coordination and trade-offs. Furthermore, species in the genus have the ability to reproduce vegetatively via rhizomes, which also store water, carbohydrates and nutrients (Klimešová et al. 2018).

We sampled adult individuals of species of the genus *Costus* during the wet seasons of 2018 (July, Costa Rica) and 2019 (June, Panama). For all traits measured, we aimed to sample six individuals per species per site (Table 1; see Supporting Information—Table S1), but could not do it for the following species: *C. bracteatus* and *C. laevis* in Tortuguero ($n = 3$ each), and *C. lima* ($n = 2$) and *C. ricus* ($n = 4$) in La Gamba.

Plant functional traits

We chose plant functional traits related to the carbon and water economy of plants, that vary in response to environmental conditions, including precipitation and temperature regime (E. Ávila-Lovera et al., submitted for publication), and that are commonly measured to allow for comparison with other studies. We also included less common traits, such as RhWC and RhSD, as species in the *Costus* genus have perennating rhizomes that are important for the life of the plants. We studied a total of 20 traits among leaves, above-ground stems, rhizomes and fine roots [see Supporting Information—Table S1; Fig. S3–S22].

Above-ground traits. All leaf traits were measured in one fully expanded leaf per individual, usually from the fourth–sixth

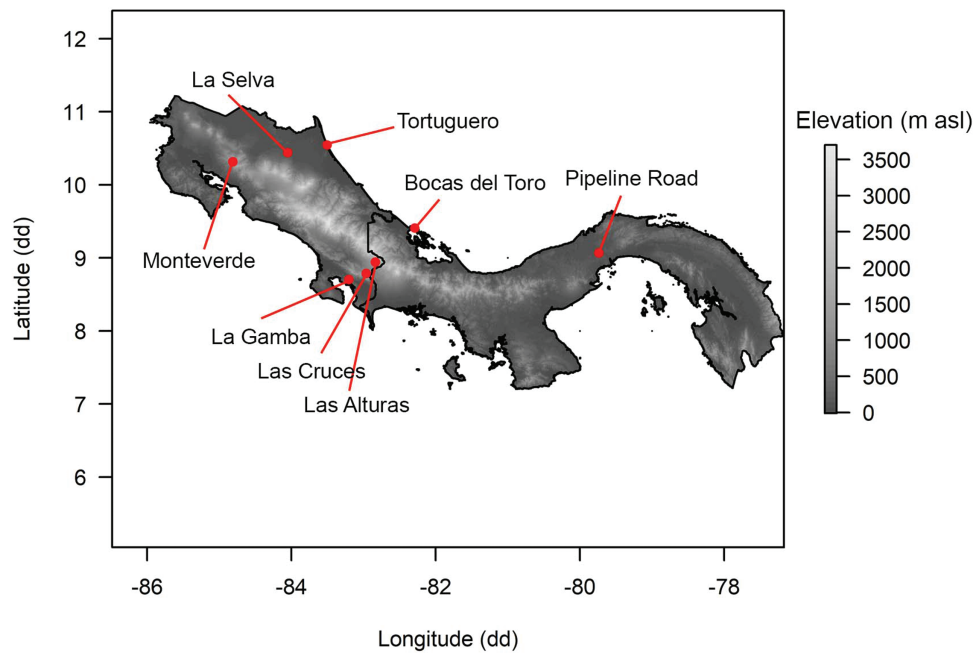


Figure 1. Map of Costa Rica and Panama showing the geographic location of the eight field sites. Shading corresponds to elevation (m asl).

node of the plant to standardize leaf age across individuals. We chose to include traits that could vary with light environment and water and nutrient availability, the so-called economic traits, and followed standard protocols (Cornelissen *et al.* 2003; Pérez-Harguindeguy *et al.* 2013).

Leaf relative chlorophyll content (Chl; SPAD units) and maximum stomatal conductance (g_s ; $\text{mmol m}^{-2} \text{s}^{-1}$) were measured *in situ* between 0730 and 1200 h preferably on rainless days. Chlorophyll content was measured using a digital chlorophyll meter (SPAD 502, Konica Minolta Sensing Inc., Japan), whereas g_s was measured with a steady-state leaf porometer (SC-1, Meter Environment, USA). After these measurements were taken, the leaf, including the petiole, was collected, placed in a zip lock bag and transported to the lab for further processing.

In the lab, we measured leaf thickness (LT; mm) in the middle portion of the leaf (avoiding major veins) using a digital micrometer (Mitutoyo IP65, Global Industrial, Port Washington, NY, USA); then, the whole leaf (lamina + petiole) was photographed against a white background including a ruler. We determined leaf size as leaf area (LA; cm^2) using ImageJ software (Rasband 1997). The leaf was then weighed whole to obtain leaf fresh mass and dried at 60 °C for 72 h to obtain leaf dry mass. From these variables we calculated multiple leaf traits: leaf dry matter content (LDMC; mg g^{-1}), calculated as leaf dry mass divided by leaf fresh mass, i.e. what proportion of the whole leaf is not water; SLA ($\text{cm}^2 \text{g}^{-1}$), calculated as LA divided by leaf dry mass; and finally, we calculated two traits, lamina dry mass to petiole dry mass ratio (LM:PM ratio; g g^{-1}) and leaf area to petiole dry mass ratio (LA:PM ratio; $\text{cm}^2 \text{g}^{-1}$), that have been previously studied in palms and heliconias as a measure of the costs of leaf mass support (Chazdon 1986; Rundel *et al.* 1998).

Dry leaf samples were ground to a fine powder using a mill (Wiley mini-mill, Thomas Scientific, Swedesboro, NJ, USA) and analysed for phosphorus (P; %) and potassium (K; %) concentrations at the Analytical Laboratory of the University of California, Davis (UC Davis). Ground leaf samples were also sent to the UC Davis Stable Isotope Facility for determination of carbon isotopic composition ($\delta^{13}\text{C}$; ‰), carbon concentration (C; %), nitrogen isotopic composition

($\delta^{15}\text{N}$; ‰) and nitrogen concentration (N; %). Values of $\delta^{13}\text{C}$ were standardized against Vienna Pee Dee Belemnite.

From the same plant sampled for leaf traits, we collected a c. 10-cm-long stem sample subtending the leaf previously sampled, which was placed in a zip lock bag and transported to the lab for further processing. From the collected stem, we sectioned a 2-cm-long piece and measured its fresh volume using the water mass displacement method (De Guzman *et al.* 2017). Then, the stem sample was dried at 60 °C for 72 h to obtain stem dry mass. We calculated stem specific density (SSD; g cm^{-3}) as stem dry mass divided by stem fresh volume.

Below-ground traits. Rhizomes with attached roots were dug up, placed in a zip lock bag and transported to the lab for further processing. Rhizomes were washed, and a portion was sectioned, blotted dry and its fresh mass measured. Fresh volume was determined as in stems, and the rhizome portion was dried at 60 °C for 72 h to obtain rhizome dry mass. We calculated RhWC (%) as rhizome water mass divided by rhizome fresh mass and multiplied by 100, and RhSD (g cm^{-3}) was calculated as rhizome dry mass divided by rhizome fresh volume.

Fine roots (<2 mm thick) that were attached to the rhizomes via coarse roots were collected from 0 to 10 cm soil depth. These fine roots were measured for length and dried at 60 °C for 72 h to obtain fine-root dry mass. Specific root length (m g^{-1}) was calculated as fine-root length divided by fine-root dry mass. Photos of the fresh roots were taken and fine-root diameter (FRD; mm) was measured using ImageJ. Length and diameter of fine roots were then used to calculate fine-root volume, and dry mass and volume were used to calculate RTD (g cm^{-3}). We could not obtain fine-root traits for samples in Panama for logistic reasons.

Statistical analyses

We performed one principal components analysis (PCA) for above-ground and below-ground traits to extract the main axes of variation using both individual data points and species means across sites and the 'prcomp' function in R v.3.6.6 (R Core Team 2020).

Traits were standardized to mean 0 and standard deviation 1 before running the analysis. We tested for phylogenetic signal in traits using species means across sites (H_0 of Blomberg's $K = 0$, significance at $P < 0.05$) and using the 'phylosig' function of the 'phytools' package in R (Revell 2012, 2019), and found no phylogenetic signal for any trait [see Supporting Information—Table S2]; thus, we did not include phylogenetic information in the PCAs. These results contrast with what has been found for some root traits in a large data set (Valverde-Barrantes et al. 2017).

To test for pairwise correlations among traits, we performed an analysis on the species means across sites using the 'rcorr' function of the 'Hmisc' package in R, and the plots were performed with the function 'corrplot' from the 'corrplot' package in R (Wei et al. 2021). To infer correlated evolution of traits, we estimated phylogenetic independent contrasts (PICs) of species means across sites using the function 'pic' of the 'ape' package in R (Paradis et al. 2004, 2018) and a phylogenetic tree [see Supporting Information—Fig. S1] constructed from a larger *Costus* phylogeny publicly available (Vargas et al. 2021). Finally, we ran a correlation analysis using the PICs. We decided not to correct for multiple testing, as recommended by Moran (2003).

To determine the contribution of site, species and individuals to the observed trait variation, we performed a variance component analysis. We ran a general linear mixed-effects model using the 'lmer' function from the 'lme4' package (Bates et al. 2015, 2020), to determine the proportion of variance explained by the three factors: site, species and individuals. The model used the raw trait data, not mean values.

Results

PCAs performed on species means across sites and on individual plant values were qualitatively similar; hence, we only present and discuss the former here (the latter can be found in Supporting Information—Fig. S2). The first two PC axes together explained 39.2 % of the total variation, with PC1 explaining 21.4 % and PC2 explaining 17.8 % (Fig. 2; Table 2). PC1 captured a trade-off between resource acquisition and conservation of water: species with a resource-acquisitive strategy had high LA, g_s , LT, P and RhWC mostly indicating high water use; whereas species with a resource-conservative strategy had high LA:PM, SSD and RhSD mostly indicating low water use and transport (Fig. 2). PC2 explained a similar amount of variation as PC1, with a resource-conservation versus resource-acquisition trade-off that was in accordance with the leaf economic spectrum: Chl, LDMC, $\delta^{13}C$ and C loaded positively with PC2 indicating high investment in structure, and SLA and K loaded negatively with PC2 indicating high capacity for leaf photosynthesis (Fig. 2). Interestingly, most below-ground traits, especially fine-root traits, strongly loaded with PC3, being orthogonal to the rest of the traits (Table 2).

The pairwise correlations supported the relationships found among traits in the PCA (see all results in Supporting Information—Table S3), for example, LA and LT were positively related with each other (Fig. 3A), whereas g_s was negatively related to SSD (Figs 3A and 4A). Similarly, g_s was negatively related to RhSD (Figs 3A and 4B) and positively related to RhWC (Fig. 3A). Furthermore, when correlations were performed on the PICs, the number of significant correlations increased from 24 to 27 and the strength of most correlations (r -value) increased as well (Fig. 3). Some interesting correlations using both species means and PICs were between: (i) leaf size and

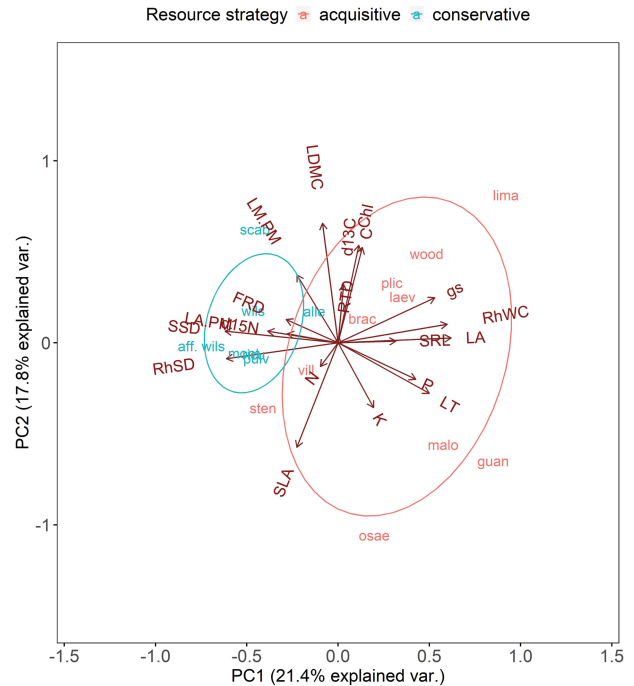


Figure 2. Principal components analysis (PCA) biplot of the studied functional traits. Groupings denote species with resource-acquisition or resource-conservation strategies. Species are abbreviated as shown in Table 1. Chl: chlorophyll concentration; g_s : stomatal conductance; LT: leaf thickness; LA: leaf area; LDMC: leaf dry matter content; SLA: specific leaf area; LM:PM: lamina dry mass to petiole dry mass ratio; LA:PM: leaf area to petiole dry mass ratio; P: leaf phosphorus concentration; K: leaf potassium concentration; $\delta^{13}C$: leaf carbon isotopic composition; C: leaf carbon concentration; $\delta^{15}N$: leaf nitrogen isotopic composition; N: leaf nitrogen concentration; SSD: stem specific density; RhWC: rhizome water content; RhSD: rhizome specific density; SRL: specific root length; FRD: fine-root diameter; RTD: root tissue density.

thickness (Fig. 3); (ii) g_s and structural above-ground stem and rhizome traits: SSD, RhWC and RhSD (Figs 3 and 4); (iii) SRL and leaf N and $\delta^{15}N$ (Fig. 3); and (iv) SRL and RTD (Fig. 3).

The variance component analysis showed that there was substantial trait variation explained by individuals: 32–87 % of total trait variation compared to 5–58 % explained by species, and 0.4–42 % explained by sites (Fig. 5). Individual variation was particularly high in FRD and g_s (Fig. 5). Remarkably, site only explained a relatively high proportion of variance in RTD (26 %), leaf $\delta^{15}N$ (35 %) and SSD (42 %) (Fig. 5). Specific leaf area was the trait with the lowest variation ascribed to site (Fig. 5), despite its important role describing the leaf economic spectrum.

Discussion

We studied 20 leaf, above-ground stem, rhizome and fine-root traits of 17 *Costus* species in eight sites that span lowland seasonal and wet forests to pre-montane and montane wet forests. We found evidence for trait coordination and trade-offs among functional traits, as well as correlated evolution. Furthermore, trait variation ascribed to individuals was high across all traits measured, indicating a high contribution of individual variation to total within-genus variation for *Costus*.

We found coordination and trade-offs among traits that are consistent with two distinct axes of resource acquisition and conservation related to different functions. The first two axes of the PCA roughly explained a similar amount of the total trait

Table 2. Results from the PCA analysis on species means, including the eigenvalue of the first three PCs, the percent of total variance explained by the first three PCs and the cumulative variance explained. Trait loadings are also included, where bolded values indicate the highest loading of the trait among the three first PC axes.

	PC1	PC2	PC3
Eigenvalue	4.28	3.56	3.14
Percent of total variance explained (%)	21.4	17.8	15.7
Cumulative variance explained (%)	21.4	39.2	54.9
Trait loadings			
Above-ground traits			
Chlorophyll concentration	0.14	0.73	0.32
Stomatal conductance	0.66	0.34	0.14
Leaf thickness	0.62	-0.38	-0.25
Leaf area	0.77	0.04	-0.24
Leaf dry matter content	-0.11	0.89	-0.13
Specific leaf area	-0.28	-0.78	0.31
Leaf mass to petiole mass ratio	-0.28	0.51	-0.12
Leaf area to petiole mass ratio	-0.48	0.09	0.13
Phosphorus concentration	0.53	-0.27	0.43
Potassium concentration	0.24	-0.48	-0.19
Carbon isotopic composition	0.04	0.44	-0.11
Carbon concentration	0.16	0.71	-0.03
Nitrogen isotopic composition	-0.34	0.07	0.32
Nitrogen concentration	-0.12	-0.18	0.74
Stem specific density	-0.77	0.09	-0.11
Below-ground traits			
Rhizome water content	0.74	0.14	0.53
Rhizome specific density	-0.76	-0.12	-0.53
Specific root length	0.39	0.01	-0.78
Fine-root diameter	-0.35	0.17	0.69
Root tissue density	0.01	0.04	0.45

variation, precluding the existence of a single major axis of variation as observed in some studies (Freschet et al. 2010; Liu et al. 2010). For example, PC1 showed coordination and trade-offs among traits, both above-ground and below-ground, related to water acquisition, use and movement. On the other hand, traits loading on PC2 were consistent with the leaf economic spectrum, with some species having high leaf C and LDMC, aligning with a resource-conservation strategy, whereas others had low LDMC and high nutrient concentrations, corresponding with a resource-acquisition strategy. Taken together, these results are consistent with the global plant economic spectrum (Reich 2014; Díaz et al. 2016) and support the idea of a unified whole-plant functional strategy. Interestingly, fine-root traits loaded strongly with the third PC, not being related to the first two axes of variation.

Few studies have examined below-ground traits and how they relate to above-ground traits and results from these studies are often contradictory. For example, some studies found no coordination between leaf, stem and root traits (Fortunel et al.

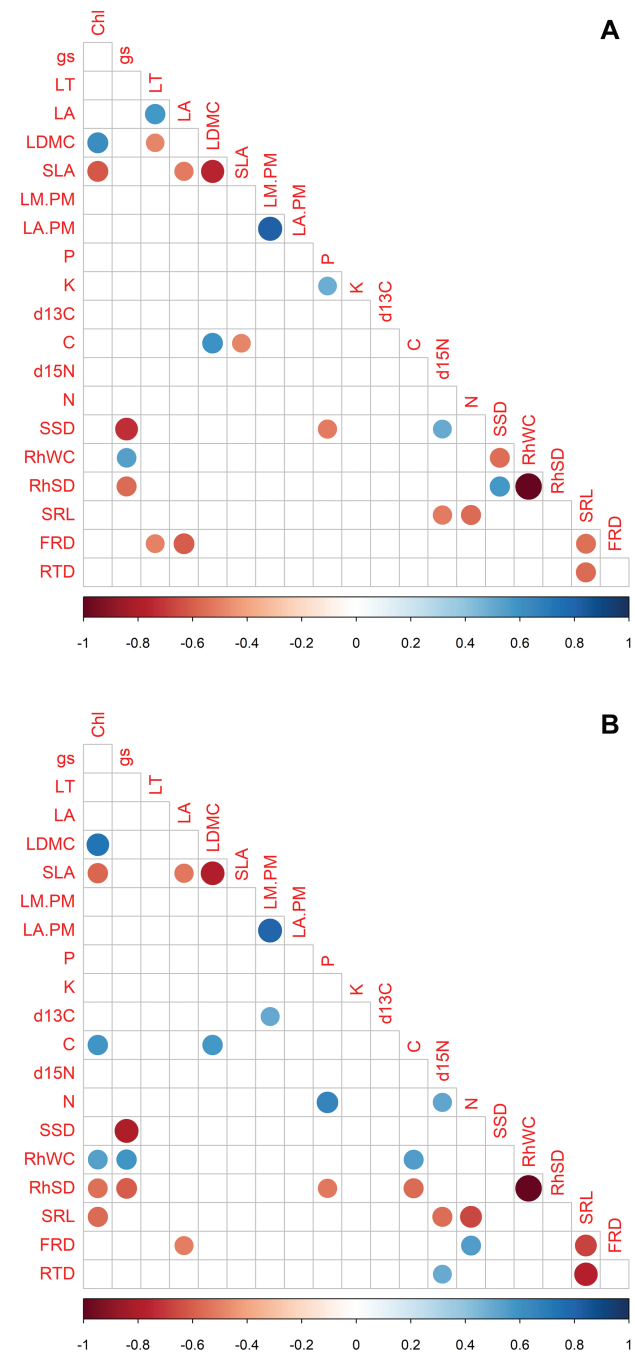


Figure 3. Correlation plots showing significant correlations only ($P < 0.05$). (A) Cross-species correlations. (B) Correlations using phylogenetic contrasts. Chl: chlorophyll concentration; g_s : stomatal conductance; LT: leaf thickness; LA: leaf area; LDMC: leaf dry matter content; SLA: specific leaf area; LM:PM: lamina dry mass to petiole dry mass ratio; LA:PM: leaf area to petiole dry mass ratio; P: leaf phosphorus concentration; K: leaf potassium concentration; $\delta^{13}C$: leaf carbon isotopic composition; C: leaf carbon concentration; $\delta^{15}N$: leaf nitrogen isotopic composition; N: leaf nitrogen concentration; SSD: stem specific density; RhWC: rhizome water content; RhSD: rhizome specific density; SRL: specific root length; FRD: fine-root diameter; RTD: root tissue density.

2012; Bowsher et al. 2016; Silva et al. 2017), whereas others found leaf-root functional coordination (Freschet et al. 2010; Liu et al. 2010). In our system, rhizomes provide structural support and water transport to above-ground organs (Maas 1972) and thus may be critical for herbaceous plants to achieve tall heights

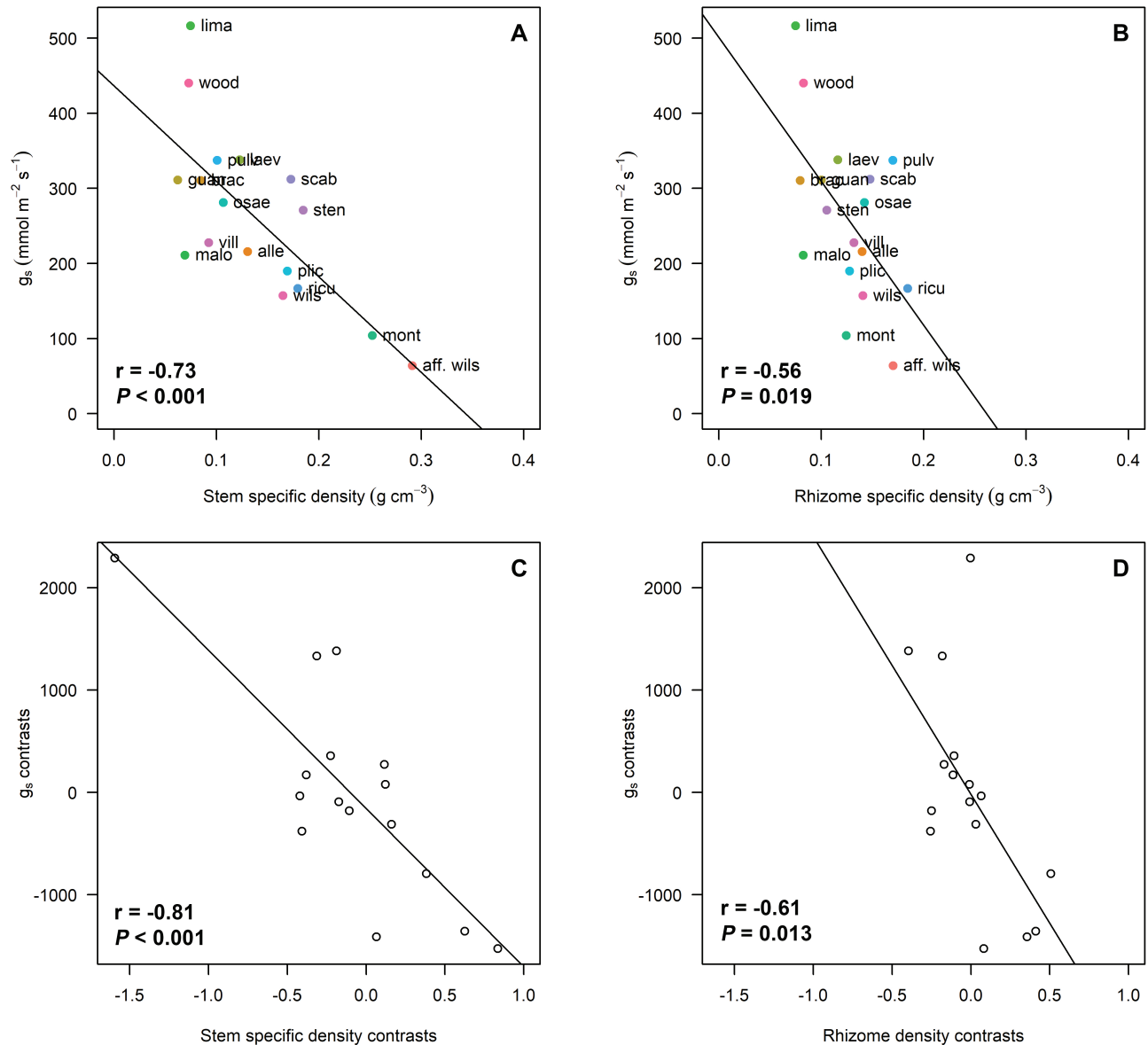


Figure 4. (A) Cross-species correlation between stomatal conductance and stem specific density and (B) between stomatal conductance and rhizome specific density. (C) Correlation between stomatal conductance contrasts and stem specific density contrasts, and (D) stomatal conductance contrasts and rhizome specific density contrasts. Contrasts were calculated as the difference between trait values of sister species divided by branch length. Trend line is included when correlations were significant. Species are abbreviated as shown in Table 1.

(e.g. 2 m in *C. montanus*) given their lack of woody tissue; hence, they play an important role in plant functioning. We found that rhizome traits related to above-ground traits loading on PC1, but were orthogonal to SRL, indicating that below-ground function in these species may be multidimensional, as has been previously found in a review of tree species (Weemstra et al. 2016), seedlings of temperate tree species (Kramer-Walter et al. 2016) and temperate herbaceous plants (Zhou et al. 2018). Being perennial organs, rhizomes perform multiple functions: they provide support for aerial shoots, serve as carbohydrate storage organs and are a mean for vegetative reproduction (clonality); as anatomical stems, they hydraulically connect roots and aerial shoots. The importance of rhizomes for the life of *Costus* plants is evidence in their coordination with above-ground function.

One reason for the no coordination between above-ground and fine-root traits is that multiple combinations of these traits can be favoured in the Neotropical forests sampled.

A previous study on trait–environment relationships using the same species studied here suggested that few functional traits respond to environmental variation (E. Ávila-Lovera et al., submitted for publication). In the current study, for example, closely related *C. osae* and *C. lima* are found in La Gamba, Costa Rica, and experience similar macroclimate conditions (MAT and MAP). However, these species have different suites of above-ground traits: *C. osae* has low LDMC and high nutrients (K and P), whereas *C. lima* has high leaf C, low N and g_s , albeit having similar values of RhWC, RhSD and SRL. These results may indicate adaptation to different light microhabitats: *C. osae* occurs in shady ravines, where there is high water availability (low LDMC is favoured) but low light availability (where leaf nutrients can enhance photosynthetic activity), and *C. lima* is found in sun-exposed habitats characterized by both high water and light availability (high g_s is favoured).

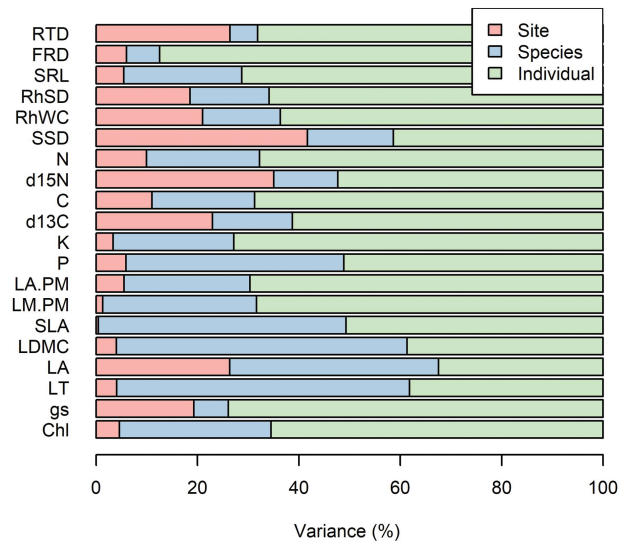


Figure 5. Variance component analysis of the traits studied. We partitioned the total trait variation into three levels: sites, species and individuals. Chl: chlorophyll concentration; g_s : stomatal conductance; LT: leaf thickness; LA: leaf area; LDMC: leaf dry matter content; SLA: specific leaf area; LM:PM: lamina dry mass to petiole dry mass ratio; LA:PM: leaf area to petiole dry mass ratio; P: leaf phosphorus concentration; K: leaf potassium concentration; $\delta^{13}C$: leaf carbon isotopic composition; C: leaf carbon concentration; $\delta^{15}N$: leaf nitrogen isotopic composition; N: leaf nitrogen concentration; SSD: stem specific density; RhWC: rhizome water content; RhSD: rhizome specific density; SRL: specific root length; FRD: fine-root diameter; RTD: root tissue density.

The relatively weak coordination between above-ground and below-ground traits in these herbaceous plants contrasts with those from woody congeneric species at higher latitudes, such as aspen (Hajek et al. 2013) and oak (Cavender-Bares et al. 2004, 2020), and community-level studies (Withington et al. 2006; Liu et al. 2010), where a clear coordination between below-ground and above-ground function exists. The only study performed in the tropics that we are aware of in which above-ground and below-ground function was studied, also found little coupling between above- and below-ground functional traits in dry forest seedlings (Arrieta-González et al. 2021), and the authors suggested the existence of multiple strategies to cope with water deficit. When comparing our results to those found at higher latitudes, seasonality experienced by the woody species rather than difference in growth form may explain such differences, such that only certain combinations of traits may be successful in highly seasonal environments at high latitudes (but see Pivovarov et al. 2016). The limited coordination among below-ground traits, but also among below-ground and above-ground traits may be due to the multiple functions that below-ground organs perform, and hence a single main axis of trait variation may be precluded altogether. Variation in plant form and function within and among species creates the basis for species co-existence, plasticity and evolvability (Silva et al. 2017). This way, multiple combinations of traits that lead to different strategies among congeneric species may facilitate their co-existence within highly diverse plant communities (Bruehlheide et al. 2018), especially in the Neotropics.

Correlations can sometimes better evidence the co-variation nature of traits among organs. Even though fine-root traits were strongly related to each other, they were rarely related to other traits, below- or above-ground. One of the few notable relationships was between SRL and both leaf $\delta^{15}N$ and N, where species with high SRL also had low leaf $\delta^{15}N$ and N. This is an interesting combination of traits given that high SRL indicates fast acquisition of water and nutrients, which can be beneficial if

paired with high rates of carbon acquisition mediated by high N concentration. However, in our data set, species with low SRL had high leaf N and leaf $\delta^{15}N$. Low SRL indicates low ability to explore soil for water and nutrient sources; however, these species have leaves with high N concentration, characteristics of the fast-return end of the LES. One aspect of below-ground function that we did not explore and could explain these trait-trait associations is the capacity of plants to form associations with mycorrhizal fungi. It has been recently reported that the fungal collaboration gradient dominates the root economic spectrum in a large data set of species (Bergmann et al. 2020). More work is needed to unravel this mystery; it is possible that additional traits not measured here (e.g. relative growth rate, whole-plant biomass allocation patterns, rates of nutrient uptake) may shed light on these seemingly contradictory relationships.

Correlated evolution among traits within groups of closely related species is common (Santiago and Kim 2009; Kembel and Cahill 2011; Liu et al. 2012; Savage and Cavender-Bares 2012; Sedio et al. 2012; Zhang et al. 2014; Bruy et al. 2018; Gallaher et al. 2019). We found correlated evolution among LA and LT. Leaf area determines the capacity to intercept light (Díaz et al. 2016) and has known impacts on leaf energy and water balance (Cornelissen et al. 2003). In *Costus*, species with large leaves, and likely high competitive ability, also have thick leaves of high succulence. High succulence allows for greater metabolite storage and has implications for structure and defence (Guterman and Chausser-Volfson 2000; Mason et al. 2016). We also found correlated evolution among some above-ground and below-ground traits, which supports the hypothesis of the existence of the plant economic spectrum (Freschet et al. 2010; Reich 2014). For example, g_s was negatively correlated with SSD and RhSD, and positively related to RhWC, indicating that these traits evolved together and that species with high stomatal opening and profligate water use have low structural investment in above-ground stems and rhizomes. This low structural investment may indicate short lifespan but may promote high hydraulic efficiency (anatomical work to test this relationship is underway). Finally, SRL, our fine-root trait analogous to SLA of the LES, only evolved in a correlated fashion with chlorophyll concentration and leaf nutrients, but in the opposite direction to what it is expected: species with high SRL had low leaf chlorophyll concentration, leaf N and $\delta^{15}N$. This unexpected relationship between SRL and leaf N requires further study in environments where water and nutrient availability can be controlled and independently modified.

Plant functional traits usually vary as a function of climate (Cavender-Bares et al. 2004; Wright et al. 2004; Violle et al. 2007; Moles et al. 2014; Mitchell et al. 2015; Blonder et al. 2017). In our study however, some traits, such as leaf (LT, LA, LDMC and SLA) and rhizome structural traits (RhSD), had low percent of variation ascribed to site. Even within a single genus, we expected trait variation ascribed to sites given the wide macroclimatic conditions experienced by species in sites that differ in elevation [see Supporting Information—Table S1], and the fact that some traits do respond to climate variation in these *Costus* species (E. Ávila-Lovera et al. submitted for publication). However, our results align with other studies that have found low variation due to site in SLA, LA and LT in tropical riparian plant communities (Liu et al. 2018). That there was no variation due to site indicates that (i) those structural traits in the genus *Costus* are less labile than other traits (i.e. physiological and nutrient traits), indicating that they have lower capacity to be adjusted during the course of the plant lifetime (Scheiner 1993), or (ii) that there are phylogenetic constraints in those traits, which is further supported by slightly higher Blomberg's K values for rhizome structural traits (RhWC and RhSD) than for other traits [see Supporting Information—Table S2].

Even though we found little trait variation that can be ascribed to site, it is important to note that there is habitat variation within sites, and species may be responding to this variation, rather than macroclimatic conditions. Future studies in which habitat variation can be better characterized will add to our understanding of trait variation in this genus.

Across all traits, a high proportion of total trait variation was ascribed to individual variation. This is consistent with results from a recent meta-analysis that showed a high contribution of intraspecific trait variation to total plant trait variation (Siefert et al. 2015), and from other studies that found that half of the variation in the LES is within-species variation (Fajardo and Siefert 2018). Intraspecific trait variation can also be substantial at regional scales in tropical and subtropical forests (Choat et al. 2007; Umaña and Swenson 2019) and temperate forests (Fajardo and Piper 2011; Hajek et al. 2013; Fajardo and Siefert 2018). However, our results contrast with those that have found that functional traits, especially root structural traits, are highly associated with phylogeny at levels above family (Valverde-Barrantes et al. 2017). The fact that we used a single genus may explain the discrepancy. Nevertheless, the implications of our results are profound, as high intraspecific trait variation can drive variation in whole-plant performance (Westerband et al. 2021), and this may help explain the patterns we observed in *Costus* species. High individual variation may result from species responses to environmental conditions (trait plasticity) or ecotypic differentiation of populations within species, as several species in our study are found at more than one site. A recent study of *Costus* across environmental gradients suggested that plasticity is one of the strongest drivers of trait–environment relationships (E. Ávila-Lovera et al., submitted for publication). Regardless of the mechanism, high individual variation highlights the ability of *Costus* species to adjust leaf, above-ground stem, rhizome and fine-root traits to match the local environmental conditions, which can help mediate responses to changes in climate. However, further studies are necessary to evaluate if high individual variation in physiology has fitness advantages (Nolting et al. 2020).

Conclusions

We conclude that *Costus* species show two apparent trade-offs between resource acquisition and conservation, one relating to water use and one to the LES. Taken together, these axes determine a unified whole-plant functional strategy for each *Costus* species. There was correlated evolution among multiple traits, especially those related to water movement and use. Finally, there was little variation in traits ascribed to site, but high individual variation in most traits, indicating high within-site and within-species variation.

Supporting Information

The following additional information is available in the online version of this article—

Figure S1. Phylogenetic tree used to perform phylogenetic independent contrasts analysis and phylogenetic signal tests. Numbers following species names are as in Table 1 and Fig. 2. Species names match those in Vargas et al. 2021).

Figure S2. Principal components analysis (PCA) biplots using individual data points grouped by (A) species, and (B) site. Species and site abbreviations are as shown in Table 1.

Figure S3. Individual values of chlorophyll concentration by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S4. Individual values of stomatal conductance by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S5. Individual values of leaf thickness by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S6. Individual values of leaf area by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S7. Individual values of leaf dry matter content by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S8. Individual values of specific leaf area by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S9. Individual values of lamina dry mass to petiole dry mass ratio by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S10. Individual values of leaf area mass to petiole dry mass ratio by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S11. Individual values of leaf phosphorus concentration by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S12. Individual values of leaf potassium concentration by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S13. Individual values of leaf carbon isotope composition by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S14. Individual values of leaf carbon concentration by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S15. Individual values of leaf nitrogen isotope composition by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S16. Individual values of leaf nitrogen isotope composition by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S17. Individual values of stem specific density by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S18. Individual values of rhizome water content by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S19. Individual values of rhizome specific density by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S20. Individual values of specific root length by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S21. Individual values of fine-root diameter by species and site. Abbreviations are as in Table 1 and Fig. 2.

Figure S22. Individual values of root tissue density by species and site. Abbreviations are as in Table 1 and Fig. 2.

Table S1. Data set used for the analyses of this article (separate excel file).

Table S2. Table showing values of Blomberg's K and P -values associated with the test of $H_0: K = 0$. P -values lower than 0.1 are in italics.

Table S3. Correlation coefficients and P -values for the cross-species correlations and the correlations using phylogenetic independent contrasts (PICs) (separate excel file).

Notes S1. Research permit information.

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Conflict of Interest

None declared.

Contributions by the Authors

J.L.F., G.R.G. and K.M.K. conceived the ideas and designed methodology; E.A.-L., G.R.G. and J.L.F. conducted field work; E.A.-L. analysed the data and wrote the manuscript; other authors provided editorial advice.

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Data Availability

All the trait data can be found in [Supporting Information—Table S1](#). Additionally, data and R codes can be found in 10.5281/zenodo.5639572.

Literature Cited

- Arrieta-González R, Paez J, Dominguez-Haydar Y, Salgado-Negret B. 2021. Limited evidence of coupling between above and belowground functional traits in tropical dry forest seedlings. *Revista de Biología Tropical* 69:9.
- Bates D, Mächler M, Bolker B, Walker S. 2015. Fitting linear mixed-effects models using lme4. *Journal of Statistical Software* 67. doi:10.18637/jss.v067.i01.
- Bates D, Maechler M, Bolker B, Walker S, Christensen R, Singmann H, Dai B, Scheipl F, Grothendieck G, Green P, Fox J. 2020. *lme4: Linear Mixed-Effects Models using 'Eigen' and S4*. R Package version 1.1-27.1.
- Bergmann J, Weigelt A, Plas F van der, Laughlin DC, Kuyper TW, Guerrero-Ramirez N, Valverde-Barrantes OJ, Bruelheide H, Freschet GT, Iversen CM, Kattge J, McCormack ML, Meier IC, Rillig MC, Roumet C, Semchenko M, Sweeney CJ, Ruijven J van, York LM, Mommer L. 2020. The fungal collaboration gradient dominates the root economics space in plants. *Science Advances* 6:eaba3756.
- Blonder B, Salinas N, Patrick Bentley L, Shenkin A, Chambi Porroa PO, Valdez Teixeira Y, Violle C, Fyllas NM, Goldsmith GR, Martin RE, Asner GP, Díaz S, Enquist BJ, Malhi Y. 2017. Predicting trait–environment relationships for venation networks along an Andes–Amazon elevation gradient. *Ecology* 98:1239–1255.
- Bowsher AW, Mason CM, Goolsby EW, Donovan LA. 2016. Fine root tradeoffs between nitrogen concentration and xylem vessel traits preclude unified whole-plant resource strategies in *Helianthus*. *Ecology and Evolution* 6:1016–1031.
- Bruelheide H, Dengler J, Purschke O, Lenoir J, Jiménez-Alfaro B, Hennekens SM, Botta-Dukát Z, Chytrý M, Field R, Jansen F, Kattge J, Pillar VD, Schrodt F, Mahecha MD, Peet RK, Sandel B, van Bodegom P, Altman J, Alvarez-Dávila E, Arfin Khan MAS, Attorre F, Aubin I, Baraloto C, Barroso JG, Bauters M, Bergmeier E, Biurrun I, Bjorkman AD, Blonder B, Čarni A, Cayuela L, Černý T, Cornelissen JHC, Craven D, Dainese M, Derroire G, De Sanctis M, Díaz S, Doležal J, Farfan-Rios W, Feldpausch TR, Fenton NJ, Garnier E, Guerin GR, Gutiérrez AG, Haider S, Hattab T, Henry G, Hérault B, Higuachi P, Hölzel N, Homeier J, Jentsch A, Jürgens N, Kačák Z, Karger DN, Kessler M, Kleyer M, Knollová I, Korolyuk AY, Kühn I, Laughlin DC, Lens F, Loos J, Louault F, Lyubanova MI, Malhi Y, Marcenò C, Mencuccini M, Müller JV, Munzinger J, Myers-Smith IH, Neill DA, Niinemets Ü, Orwin KH, Ozinga WA, Penuelas J, Pérez-Haase A, Petřík P, Phillips OL, Pärtel M, Reich PB, Römermann C, Rodrigues AV, Sabatini FM, Sardans J, Schmidt M, Seidler G, Silva Espejo JE, Silveira M, Smyth A, Sporbert M, Svenning JC, Tang Z, Thomas R, Tsiropidis I, Vassilev K, Violle C, Virtanen R, Weiher E, Welk E, Wesche K, Winter M, Wirth C, Jandt U. 2018. Global trait–environment relationships of plant communities. *Nature Ecology & Evolution* 2:1906–1917.
- Bruy D, Hattermann T, Barrabé L, Mouly A, Barthélémy D, Isnard S. 2018. Evolution of plant architecture, functional diversification and divergent evolution in the genus *Atractocarpus* (Rubiaceae) for New Caledonia. *Frontiers in Plant Science* 9:1775.
- Cavender-Bares J, Fontes CG, Pinto-Ledezma J. 2020. Open questions in understanding the adaptive significance of plant functional trait variation within a single lineage. *The New Phytologist* 227:659–663.
- Cavender-Bares J, Kitajima K, Bazzaz FA. 2004. Multiple trait associations in relation to habitat differentiation among 17 Floridian oak species. *Ecological Monographs* 74:635–662.
- Chave J, Coomes D, Jansen S, Lewis SL, Swenson NG, Zanne AE. 2009. Towards a worldwide wood economics spectrum. *Ecology Letters* 12:351–366.
- Chazdon RL. 1986. The costs of leaf support in understory palms: economy versus safety. *The American Naturalist* 127:9–30.
- Choat B, Sack L, Holbrook NM. 2007. Diversity of hydraulic traits in nine *Cordia* species growing in tropical forests with contrasting precipitation. *The New Phytologist* 175:686–698.
- Cornelissen JHC, Lavorel S, Garnier E, Díaz S, Buchmann N, Gurvich DE, Reich PB, Steege H ter, Morgan HD, Heijden MGA van der, Pausas JG, Poorter H. 2003. A handbook of protocols for standardised and easy measurement of plant functional traits worldwide. *Australian Journal of Botany* 51:335–380.
- De Guzman ME, Santiago LS, Schnitzer SA, Álvarez-Cansino L. 2017. Trade-offs between water transport capacity and drought resistance in neotropical canopy liana and tree species. *Tree Physiology* 37:1404–1414.
- Díaz S, Kattge J, Cornelissen JH, Wright IJ, Lavorel S, Dray S, Reu B, Kleyer M, Wirth C, Prentice IC, Garnier E, Bönsch G, Westoby M, Poorter H, Reich PB, Moles AT, Dickie J, Gillison AN, Zanne AE, Chave J, Wright SJ, Sheremet'ev SN, Jactel H, Baraloto C, Cerabolini B, Pierce S, Shipley B, Kirkup D, Casanoves F, Joswig JS, Günther A, Falczuk V, Rüger N, Mahecha MD, Gorné LD. 2016. The global spectrum of plant form and function. *Nature* 529:167–171.
- Fajardo A, Piper FI. 2011. Intraspecific trait variation and covariation in a widespread tree species (*Nothofagus pumilio*) in southern Chile. *The New Phytologist* 189:259–271.
- Fajardo A, Siefert A. 2018. Intraspecific trait variation and the leaf economics spectrum across resource gradients and levels of organization. *Ecology* 99:1024–1030.
- Fick SE, Hijmans RJ. 2017. WorldClim 2: new 1-km spatial resolution climate surfaces for global land areas. *International Journal of Climatology* 37:4302–4315.
- Fortunel C, Fine PVA, Baraloto C, Dalling J. 2012. Leaf, stem and root tissue strategies across 758 Neotropical tree species. *Functional Ecology* 26:1153–1161.
- Freschet GT, Cornelissen JHC, van Logtestijn RSP, Aerts R. 2010. Evidence of the 'plant economics spectrum' in a subarctic flora. *Journal of Ecology* 98:362–373.
- Gallaher TJ, Adams DC, Attigala L, Burke SV, Craine JM, Duvall MR, Klahs PC, Sherratt E, Wysocki WP, Clark LG. 2019. Leaf shape and size track habitat transitions across forest–grassland boundaries in the grass family (Poaceae). *Evolution* 73:927–946.
- Gutterman Y, Chauser-Volfson E. 2000. The distribution of the phenolic metabolites barbaloin, aloeresin and aloenin as a peripheral defense strategy in the succulent leaf parts of *Aloe arborescens*. *Biochemical Systematics and Ecology* 28:825–838.
- Hajek P, Hertel D, Leuschner C. 2013. Intraspecific variation in root and leaf traits and leaf–root trait linkages in eight aspen demes (*Populus tremula* and *P. tremuloides*). *Frontiers in Plant Science* 4:415.
- Kay KM, Reeves PA, Olmstead RG, Schemske DW. 2005. Rapid speciation and the evolution of hummingbird pollination in neotropical *Costus* subgenus *Costus* (Costaceae): evidence from nrDNA ITS and ETS sequences. *American Journal of Botany* 92:1899–1910.
- Kemmel SW, Cahill JF. 2011. Independent evolution of leaf and root traits within and among temperate grassland plant communities. *PLoS One* 6:e19992.
- Klímešová J, Martinková J, Ottaviani G. 2018. Belowground plant functional ecology: towards an integrated perspective. *Functional Ecology* 32:2115–2126.
- Kong D, Wang J, Kardol P, Wu H, Zeng H, Deng X, Deng Y. 2015. The root economics spectrum: divergence of absorptive root strategies with root diameter. *Biogeosciences Discussions* 12:13041–13067.
- Kong D, Wang J, Wu H, Valverde-Barrantes OJ, Wang R, Zeng H, Kardol P, Zhang H, Feng Y. 2019. Nonlinearity of root trait relationships and the root economics spectrum. *Nature Communications* 10:2203.
- Kraft NJ, Valencia R, Ackerly DD. 2008. Functional traits and niche-based tree community assembly in an Amazonian forest. *Science* 322:580–582.

- Kramer-Walter KR, Bellingham PJ, Millar TR, Smissen RD, Richardson SJ, Laughlin DC. 2016. Root traits are multidimensional: specific root length is independent from root tissue density and the plant economic spectrum. *Journal of Ecology* **104**:1299–1310.
- Laliberté E. 2017. Below-ground frontiers in trait-based plant ecology. *New Phytologist* **213**:1597–1603.
- Liu G, Freschet GT, Pan X, Cornelissen JH, Li Y, Dong M. 2010. Coordinated variation in leaf and root traits across multiple spatial scales in Chinese semi-arid and arid ecosystems. *The New Phytologist* **188**:543–553.
- Liu H, Edwards EJ, Freckleton RP, Osborne CP. 2012. Phylogenetic niche conservatism in C₄ grasses. *Oecologia* **170**:835–845.
- Liu R, Liang S, Long W, Jiang Y. 2018. Variations in leaf functional traits across ecological scales in riparian plant communities of the Lijiang river, Guilin, Southwest China. *Tropical Conservation Science* **11**:194008291880468.
- Maas PJM. 1972. *Costoideae (Zingiberaceae)*. *Flora Neotropica*. New York: Haner, **8**:1–139.
- Mason CM, Bowsheer AW, Crowell BL, Celoy RM, Tsai CJ, Donovan LA. 2016. Macroevolution of leaf defenses and secondary metabolites across the genus *Helianthus*. *The New Phytologist* **209**:1720–1733.
- McCormack ML, Adams TS, Smithwick EAH, Eissenstat DM. 2012. Predicting fine root lifespan from plant functional traits in temperate trees. *The New Phytologist* **195**:823–831.
- McCormack ML, Iversen CM. 2019. Physical and functional constraints on viable belowground acquisition strategies. *Frontiers in Plant Science* **10**:1215.
- Mitchell N, Moore TE, Mollmann HK, Carlson JE, Mocko K, Martinez-Cabrera H, Adams C, Silander JA Jr, Jones CS, Schlichting CD, Holsinger KE. 2015. Functional traits in parallel evolutionary radiations and trait–environment associations in the cape floristic region of South Africa. *The American Naturalist* **185**:525–537.
- Moles AT, Perins SE, Laffan SW, Flores-Moreno H, Awasthy M, Tindall ML, Sack L, Pitman A, Kattge J, Aarssen LW, Anand M, Bahn M, Blonder B, Cavender-Bares J, Cornelissen JHC, Cornwell WK, Díaz S, Dickie JB, Freschet GT, Griffiths JG, Gutierrez AG, Hemmings FA, Hickler T, Hitchcock TD, Keighery M, Kleyer M, Kurokawa H, Leishman MR, Liu K, Niinemets Ü, Onipchenko V, Onoda Y, Penuelas J, Pillar VD, Reich PB, Shiodera S, Siefert A, Sosinski EE, Soudzilovskaia NA, Swaine EK, Swenson NG, van Bodegom PM, Warman L, Weiher E, Wright IJ, Zhang H, Zobel M, Bonser SP. 2014. Which is a better predictor of plant traits: temperature or precipitation? *Journal of Vegetation Science* **25**:1167–1180.
- Mommer L, Weemstra M. 2012. The role of roots in the resource economics spectrum. *The New Phytologist* **195**:725–727.
- Moran MD. 2003. Arguments for rejecting the sequential Bonferroni in ecological studies. *Oikos* **100**:403–405.
- Nolting KM, Prunier R, Midgley GF, Holsinger KE. 2020. Intraspecific trait variation influences physiological performance and fitness in the South Africa shrub genus *Protea* (Proteaceae). *Annals of Botany* **1–13**. doi:10.1093/aob/mcaa060.
- Paradis E, Blomberg S, Bolker B, Claude J, Cuong HS, Desper R, Didier G, Durand B, Duthéil J, Gascuel O, Guillaume T, Heibl C, Ives A, Jones B, Krahe F, Lawson D, Lefort V, Legendre P, Lemon J, McCloskey R, Nylander J, Opgen-Rhein R, Popescu A-A, Royer-Carenzi M, Schliep K, Strimmer K, de Vienne D. 2018. *ape: Analyses of Phylogenetics and Evolution*. R Package version 5.5.
- Paradis E, Claude J, Strimmer K. 2004. APE: analyses of phylogenetics and evolution in R language. *Bioinformatics* **20**:289–290.
- Pérez-Harguindeguy N, Díaz S, Garnier E, Lavorel S, Poorter H, Jaureguiberry P, Bret-Harte MS, Cornwell WK, Craine JM, Gurvich DE, Urcelay C, Veneklaas EJ, Reich PB, Poorter L, Wright IJ, Ray P, Enrico L, Pausas JG, de Vos AC, Buchmann N, Funes G, Quétier F, Hodgson JG, Thompson K, Morgan HD, ter Steege H, van der Heijden MGA, Sack L, Blonder B, Poschlod P, Vaieretti MV, Conti G, Staver AC, Aquino S, Cornelissen JHC. 2013. New handbook for standardised measurement of plant functional traits worldwide. *Australian Journal of Botany* **61**:167.
- Pivovarov AL, Pasquini SC, De Guzman ME, Alstad KP, Stemke JS, Santiago LS. 2016. Multiple strategies for drought survival among woody plant species. *Functional Ecology* **30**:517–526.
- R Core Team. 2020. *R: a language and environment for statistical computing*. Vienna, Austria: R Foundation for Statistical Computing.
- Rasband WS. 1997. *ImageJ*. Bethesda, MD: U.S. National Institutes of Health.
- Reich PB. 2014. The world-wide ‘fast–slow’ plant economics spectrum: a traits manifesto. *Journal of Ecology* **102**:275–301.
- Reich PB, Walters MB, Ellsworth DS. 1997. From tropics to tundra: global convergence in plant functioning. *Proceedings of the National Academy of Sciences of the United States of America* **94**:13730–13734.
- Reich PB, Wright IJ, Cavender-Bares J, Craine JM, Oleksyn J, Westoby M, Walters MB. 2003. The evolution of plant functional variation: traits, spectra, and strategies. *International Journal of Plant Sciences* **164**:S143–S164.
- Revell LJ. 2012. phytools: an R package for phylogenetic comparative biology (and other things). *Methods in Ecology and Evolution* **3**:217–223.
- Revell LJ. 2019. *phytools: Phylogenetic Tools for Comparative Biology (and Other Things)*. R Package version 0.7-80.
- Roumet C, Birouste M, Picon-Cochard C, Ghestem M, Osman N, Vrignon-Brenas S, Cao KF, Stokes A. 2016. Root structure-function relationships in 74 species: evidence of a root economics spectrum related to carbon economy. *The New Phytologist* **210**:815–826.
- Rundel PW, Sharifi MR, Gibson AC, Esler KJ. 1998. Structural and physiological adaptation to light environments in neotropical *Heliconia* (Heliconiaceae). *Journal of Tropical Ecology* **14**:789–801.
- Santiago LS, Kim S. 2009. Correlated evolution of leaf shape and physiology in the woody *Sonchus* Alliance (Asteraceae: Sonchinae) in Macaronesia. *International Journal of Plant Sciences* **170**:83–92.
- Savage JA, Cavender-Bares J. 2012. Habitat specialization and the role of trait lability in structuring diverse willow (genus *Salix*) communities. *Ecology* **93**:S138–S150.
- Scheiner SM. 1993. Genetics and evolution of phenotypic plasticity. *Annual Review of Ecology and Systematics* **24**:35–68.
- Sedio BE, Wright SJ, Dick CW. 2012. Trait evolution and the coexistence of a species swarm in the tropical forest understorey. *Journal of Ecology* **100**:1183–1193.
- Siefert A, Violle C, Chalmandrier L, Albert CH, Taudiere A, Fajardo A, Aarssen LW, Baraloto C, Carlucci MB, Cianciaruso MV, de L Dantas V, de Bello F, Duarte LD, Fonseca CR, Freschet GT, Gaucherand S, Gross N, Hikokasa K, Jackson B, Jung V, Kamiyama C, Katabuchi M, Kembel SW, Kichenin E, Kraft NJ, Lagerström A, Bagousse-Pinguet YL, Li Y, Mason N, Messier J, Nakashizuka T, Overton JM, Peltzer DA, Pérez-Ramos IM, Pillar VD, Prentice HC, Richardson S, Sasaki T, Schamp BS, Schöb C, Shipley B, Sundqvist M, Sykes MT, Vandewalle M, Wardle DA. 2015. A global meta-analysis of the relative extent of intraspecific trait variation in plant communities. *Ecology Letters* **18**:1406–1419.
- Silva JLA, Souza AF, Caliman A, Voigt EL, Lichstein JE. 2017. Weak whole-plant trait coordination in a seasonally dry South American stressful environment. *Ecology and Evolution* **8**:4–12.
- Umaña MN, Swenson NG. 2019. Intraspecific variation in traits and tree growth along an elevational gradient in a subtropical forest. *Oecologia* **191**:153–164.
- Valverde-Barrantes OJ, Freschet GT, Roumet C, Blackwood CB. 2017. A worldwide of root traits: the influence of ancestry, growth form, climate and mycorrhizal association on the functional trait variation of fine-root tissues in seed plants. *The New Phytologist* **215**:1562–1573.
- Vargas OM, Goldston B, Grossenbacher DL, Kay KM. 2021. Patterns of speciation are similar across mountainous and lowland regions for a Neotropical plant radiation (Costaceae: *Costus*). *Evolution*. doi:10.1111/evo.14108.
- Violle C, Navas M-L, Vile D, Kazakou E, Fortunel C, Hummel E, Garnier E. 2007. Let the concept of trait be functional! *Oikos* **116**:882–892.
- Weemstra M, Mommer L, Visser EJ, van Ruijven J, Kuyper TW, Mohren GM, Sterck FJ. 2016. Towards a multidimensional root trait framework: a tree root review. *The New Phytologist* **211**:1159–1169.
- Wei T, Simko V, Levy M, Xie Y, Zemla J, Freidank M, Cai J, Protivinsky T. 2021. *corrplot: Visualization of a Correlation Matrix*. R Package version 0.90.
- Weigelt A, Mommer L, Andraczek K, Iversen CM, Bergmann J, Bruehlheide H, Fan Y, Freschet GT, Guerrero-Ramírez NR, Kattge J, Kuyper TW, Laughlin DC, Meier IC, van der Plas F, Poorter H, Roumet C, van Ruijven J, Sabatini FM, Semchenko M, Sweeney CJ, Valverde-Barrantes OJ, York LM, McCormack ML. 2021. An integrated framework of plant form and function: the belowground perspective. *The New Phytologist* **232**:42–59.

- Westerband AC, Funk JL, Barton KE. 2021. Intraspecific trait variation in plants: a renewed focus on its role in ecological processes. *Annals of Botany* **127**:397–410.
- Westoby M, Falster DS, Moles AT, Vesk PA, Wright IJ. 2002. Plant ecological strategies: some leading dimensions of variation between species. *Annual Review of Ecology and Systematics* **33**:125–159.
- Withington JM, Reich PB, Oleksyn J, Eissenstat DM. 2006. Comparisons of structure and life span in roots and leaves among temperate trees. *Ecological Monographs* **76**:381–397.
- Wright IJ, Reich PB, Westoby M, Ackerly DD, Baruch Z, Bongers F, Cavender-Bares J, Chapin T, Cornelissen JH, Diemer M, Flexas J, Garnier E, Groom PK, Gulias J, Hikosaka K, Lamont BB, Lee T, Lee W, Lusk C, Midgley JJ, Navas ML, Niinemets U, Oleksyn J, Osada N, Poorter H, Poot P, Prior L, Pyankov VI, Roumet C, Thomas SC, Tjoelker MG, Veneklaas EJ, Villar R. 2004. The worldwide leaf economics spectrum. *Nature* **428**:821–827.
- Zanne AE, Westoby M, Falster DS, Ackerly DD, Loarie SR, Arnold SE, Coomes DA. 2010. Angiosperm wood structure: global patterns in vessel anatomy and their relation to wood density and potential conductivity. *American Journal of Botany* **97**:207–215.
- Zhang SB, Sun M, Cao KF, Hu H, Zhang JL. 2014. Leaf photosynthetic rate of tropical ferns is evolutionarily linked to water transport capacity. *PLoS One* **9**:e84682.
- Zhou M, Bai W, Zhang Y, Zhang W-H. 2018. Multi-dimensional patterns of variation in root traits among coexisting herbaceous species in temperate steppes. *Journal of Ecology* **106**:2320–2331.