Legume abundance along successional and rainfall gradients in Neotropical forests


The nutrient demands of regrowing tropical forests are partly satisfied by nitrogen-fixing legume trees, but our understanding of the abundance of those species is biased towards wet tropical regions. Here we show how the abundance of Leguminosae is affected by both recovery from disturbance and large-scale rainfall gradients through a synthesis of forest inventory plots from a network of 42 Neotropical forest chronosequences. During the first three decades of natural forest regeneration, legume basal area is twice as high in dry compared with wet secondary forests. The tremendous ecological success of legumes in recently disturbed, water-limited forests is likely to be related to both their reduced leaflet size and ability to fix N₂, which together enhance legume drought tolerance and water-use efficiency. Earth system models should incorporate these large-scale successional and climatic patterns of legume dominance to provide more accurate estimates of the maximum potential for natural nitrogen fixation across tropical forests.

More than half of the tropical forest area worldwide is made up of secondary forests, which regrow after canopy removal due to natural or anthropogenic disturbances1. Secondary forests are important globally because they supply firewood and timber, regulate the hydrological cycle, benefit biodiversity, and provide carbon storage as above- and belowground biomass5, but their growth can be constrained by nitrogen (N) availability4. Symbiotic fixation is thought to provide the largest natural input of N to tropical forests5, and part of the N demand of regrowing tropical forests is satisfied by legume trees (Leguminosae) that have the capacity to fix atmospheric N₂ through interactions with rhizobia bacteria6. The abundance of N-fixing legumes is not always strictly proportional to the rates of rhizobial activity, as some legumes downregulate fixation when the costs outweigh the benefits7. Nevertheless, legume abundance as represented by total basal area may provide a good estimate of the maximum potential N fixation in an ecosystem, with the advantage that this metric can be extracted from standard forest inventory surveys. Spatially explicit estimates of legume abundance through time could help to reduce uncertainties in Earth system models that include coupled carbon and N biogeochemistry8, but assessments of legume abundance have not yet been synthesized across the successional and climatic gradients that characterize tropical forests.

The abundance of N-fixing legumes relative to non-fixing trees has been closely examined in undisturbed tropical forests5,10 and savannas1,12. However, studies of legume abundance in regenerating forests are rare and have been restricted to the wet tropics5,13–16, so are likely not representative of tropical
secondary succession globally. Due to environmental filtering\textsuperscript{15}, systematic variations in legume abundance should occur along both rainfall and successional gradients. Wet and (seasonally) dry tropical forests\textsuperscript{10,18} experience substantial differences in water and nutrient availability, which in turn may influence the competitive success of legumes in both biomes\textsuperscript{18}. N-fixing legumes should have particular advantages in drier conditions; they can access N when mineralization rates decline due to low soil moisture\textsuperscript{19}, and use their high foliar N content to maintain high growth rates and use water more efficiently relative to non-fixers\textsuperscript{21}. Because young tropical forests are often N limited\textsuperscript{1}, trees that are able to fix may be favoured during the earliest stages of forest regrowth after disturbance\textsuperscript{22,23}. Some studies indeed show that legumes are more abundant in young compared with old wet Neotropical forests\textsuperscript{8,14}, but others report successional trends in the opposite direction, with the relative abundance of these species instead increasing with stand age\textsuperscript{7,19}. Characterizing these macroecological patterns of legume abundance across climate space and through successional time, along with variations in their functional traits, is crucial to determine whether our current knowledge of legume distributions can be generalized across the tropics and to achieve a more complete understanding of the role of this exceptionally diverse plant family within secondary Neotropical forest ecosystems.

Here we evaluate how the abundance of legumes (as measured by absolute and relative basal area) varies through secondary succession using data from 42 chronosequence sites\textsuperscript{15} (Supplementary Fig. 1 and Supplementary Table 1) that span a large gradient in mean annual rainfall (from 750 to 4,000 mm yr\textsuperscript{-1}) and explain legume success based on N-fixation potential and two functional traits related to drought tolerance (leaf size and leaf type). We primarily focused our climate analysis on mean annual rainfall (‘rainfall’ hereafter), but also tested the effect of rainfall seasonality\textsuperscript{33} (the rainfall coefficient of variation from WorldClim) and climatic water deficit\textsuperscript{34} (CWD; which tracks water losses during the months where evapotranspiration exceeds rainfall) as alternative predictors. Because leaf habit or leafing phenology is a better indicator of seasonal moisture stress than total annual precipitation\textsuperscript{35}, we used this parameter to classify study sites as ‘dry’ forests if the vegetation was mostly drought deciduous (sensu ref. 10), or as ‘wet’ forests in all other instances (that is, mostly evergreen; Supplementary Table 1). As such, the terms ‘dry’ and ‘wet’ forests refer to two ecologically distinct tropical biomes with floristic compositions that differ in phylogenetic, biogeographic, functional and community ecological patterns (see refs 26–28). Therefore, although rainfall is used as the main (continuous) climatic variable to model legume abundance, we did not use this variable to classify sites as either dry or wet forests.

To understand the specific factors that enable legumes to thrive in particular tropical environments, we also investigated how the abundance of these trees related to their capacity to fix N\textsubscript{2} and a pair of crucial leaf traits that reflect drought tolerance. For each of the 398 legume species present at our sites, we assessed potential to fix N\textsubscript{2} based on positive nodulation reports and expert knowledge\textsuperscript{31} (see Methods). Both of the leaf traits we examined—leaf size and leaf type—reflect adaptations to limited water availability (Supplementary Tables 2 and 3). Smaller leaves have reduced boundary-layer resistance, which enables them to dissipate heat through conductive or convective radiation\textsuperscript{32–34}. Leaf type is considered to be associated with drought severity and seasonality because plants with compound leaves (having either pinnate or bipinnate divisions) are able to shed individual leaflets (rather than whole leaves) when faced with severe moisture stress\textsuperscript{35}. Our analysis demonstrates that the abundance of legumes indeed varies substantially and systematically across Neotropical forests, and although the ecological success (that is, high relative abundance) of these species during the very earliest stages of secondary succession is partly due to N fixation, other traits related to drought tolerance and water-use efficiency likely also offer competitive advantages.

Results

During the first three decades of forest regeneration, the total abundance of all legume trees as measured by their absolute basal area doubled in both dry and wet Neotropical forests (from 3 m\textsuperscript{2} ha\textsuperscript{-1} in 2- to 10-year-old forests to 6 m\textsuperscript{2} ha\textsuperscript{-1} in 21- to 30-year-old forests; Fig. 1a) as legume biomass gradually built up through succession. Here we define legume relative abundance (RA) as the basal area of Leguminosae trees divided by the total basal area of all trees in each plot and interpret it as a measure of ecological success that reflects legume performance relative to non-legume species. Overall, although absolute legume abundance increased with successional patterns of legume abundance were largely driven by rainfall seasonality because plants with compound leaves (having either pinnate divisions) when faced with severe moisture stress35. Our analysis demonstrates that the abundance of legumes in dry forests (compared with wet forests) may partly be a consequence of their higher initial recruitment, which is suggested by the high RA of small diameter legume trees during the first two decades of forest regeneration (Supplementary Fig. 2).

Mean annual rainfall was a strong determinant of legume RA over the entire Neotropical network (Table 1). Alternative models of legume RA that used rainfall seasonality and CWD as the main climatic predictor variable also explained a significant amount of variation in our data (Supplementary Figs. 3 and 4), but the best-supported model was based on mean annual rainfall ($R^2 = 0.62$, versus $R^2 = 0.49$ and $R^2 = 0.48$ for seasonality and CWD respectively; Supplementary Table 4). The magnitude of legume RA and its relationship with rainfall differed strongly between dry and wet secondary forests, most prominently during the first three decades of secondary succession (Fig. 2). For the 26 chronosequences from wet forests, mean legume RA was approximately 15% ($\pm 16\%$ s.d.), within the range reported previously for individual sites\textsuperscript{33–35}, and did not vary with rainfall. By contrast, legume RA in the 16 dry forest sites was much higher (41% $\pm 27\%$ s.d.) and was strongly and inversely related to annual rainfall. The transition between these two patterns occurred at approximately 1,500 mm yr\textsuperscript{-1} (Fig. 2).

The functional traits of legumes also varied across the large-scale environmental gradients in our dataset. The spatial and successional patterns of legume abundance were largely driven by N-fixing species (Fig. 1c). For nearly two-fifths of the plots in our

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network, fixers were the only type of legumes present. At the plot level, the median percentage of total legume basal area comprised by fixers was 93.5%. The proportion of N-fixer basal area to total legume basal area did not vary with rainfall or stand age, and the RA of non-fixing legumes was much lower in both dry and wet secondary forest sites (Supplementary Fig. 5). In contrast to the N fixers, the RA of non-fixing legumes remained constant throughout succession (Supplementary Table 5). When we stratified our analyses by leaf type, it was evident that the extremely high legume RA in young dry forests was largely due to the prevalence of species with bipinnate leaves (Figs. 1d and 3 and Table 1), which have significantly smaller leaflets than legumes with other leaf types (Supplementary Table 3).

Discussion

Based on our survey of secondary forests across the Neotropics, we conclude the ecological success of legume trees is markedly higher in seasonally dry forests than in wet forests, especially during the earliest stages of secondary succession. These findings agree with the expectation that Neotropical legumes are proportionately more abundant in seasonally dry forests than in wet forests, especially during the earliest stages of secondary succession. We identified a threshold in mean annual rainfall at approximately 1,500 mm yr\(^{-1}\)—nearly identical to the threshold observed in forest inventories from North America\(^6\). Below this level, legume abundance was strongly and negatively correlated to water availability. Because this relationship was driven mainly by species that are both able to fix N\(_2\) and have bipinnate leaves (Fig. 1), we suggest that the exceptional abundance of tropical legumes towards the drier end of the rainfall spectrum during secondary succession is the combined product of (1) small leaflet size, which allows for leaf temperature regulation and water conservation, and (2) N fixation, which contributes to photosynthetic acclimation, enhances water-use efficiency, and may satisfy the demand for nitrogen after the post-dry season leaf flush.

The availability of nitrogen is known to constrain biomass recovery in secondary Neotropical forests\(^{4,37}\) because it is often lost following disturbance, either through harvesting of the canopy or crops, volatilization during burning or leaching\(^7\). Our finding that Neotropical legumes are proportionately more abundant in early succession throughout the Neotropics could be due to the high demand for N in recently disturbed forests\(^{4,37}\). Under those conditions, the fixation by legume species is increased, allowing for higher abundance of N-fixing legumes in seasonal dry forests than in wet forests, especially during the earliest stages of secondary succession.

Fig. 1 | Absolute and relative basal area of legume species in Neotropical secondary forests. The RA of legume trees goes down during forest regeneration, but is markedly higher in seasonally dry forests than in wet forests, especially during the earliest stages of secondary succession. a, Plot-level total basal area of all legume species. b, Relative basal area of all legume species. c, Relative basal area of N-fixing legumes. d, Relative basal area of legumes with bipinnate leaves. Each circle represents one plot. Regression lines were drawn to highlight the successional trajectory of each of our 42 chronosequences. Dry forest sites (with dominant deciduous vegetation) are indicated in orange, and wet forests in blue. Insets show the average of all fits for absolute (a) or relative (b-d) basal area of legumes in dry and wet forests.
circumstances, fixation would allow legumes to overcome N limitation more easily than their competitors, although the degree to which initial secondary forest regrowth is limited by N availability is highly variable and influenced by local disturbance history and prior land use\cite{38,39}.

Besides providing legumes with an advantage in young secondary forests in general, N fixation could offer additional benefits to plants growing under seasonally dry conditions. Forests that experience a pronounced annual dry season are affected by recurrent seasonal declines in soil moisture and, due to the associated hiatus in transpiration, plants are not able to access nutrients in the soil solution, including N \cite{21,42}. Because many dry forest trees lose their leaves each year and are required to grow a new leaf canopy, being able to obtain N through symbiosis could allow legumes to more quickly rebuild their leaf canopy at the end of the dry season compared with non-fixers\cite{11,41}, which may need to wait until soil water is sufficiently recharged to acquire mineral N (ref. \cite{40}). Towards the high end of the Neotropical rainfall gradient, forests do not experience a strong seasonal moisture deficit, and as a result, are not usually faced with the regular nutrient scarcity that consistently affects dry forests. In addition to its role in satisfying seasonal nitrogen demands, fixation should also help legumes to further acclimate to hot, dry conditions\cite{32,43}. By investing part of their fixed N into the production of photosynthetic enzymes, plants are able to draw down their internal concentrations of carbon dioxide, thereby creating steeper diffusion gradients in their leaves\cite{41}. This adjustment allows photosynthesis to occur at lower stomatal conductance and reduced transpiration. These two advantages offered by fixation could account for the extremely high abundance of legumes early in dry forest succession, when air and soil temperatures are high, and soil water potential is at its lowest\cite{42}, as well as their continued dominance in this ecosystem over evolutionary timescales\cite{10,11}.

But N fixation is clearly not the only trait that is advantageous to legumes in Neotropical dry forests. Because this biome is dominated by legumes that have bipinnate leaves particularly during the early stages of succession, it is clear that these species also benefit from their conservative use of water. Reduced leaf size (and the accompanying thinner boundary layer) offers a significant adaptive value in hot, dry environments\cite{44} by enhancing heat dissipation and regulating leaf temperature more efficiently\cite{45}. Many legumes also have the ability to adjust the angle of their laminae to regulate diurnal incident solar radiation\cite{44}, which further reduces heat loading. Hence, having small leaflets could enable legumes at dry sites to benefit from high irradiance while preventing excessively high leaf temperatures. The bipinnate leaf type is confined exclusively to one subclade of legumes, the Caesalpinioideae\cite{45}, and half of all Caesalpinioideae species in our dataset (mainly those that fix N)\cite{46} have bipinnate leaves. By contrast, only one-quarter of all N fixers have this trait, which means a relatively small subset of taxa is extremely well represented

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Across the Neotropics, the abundance of legumes in secondary forests can be predicted by rainfall, stand age and their interaction. Linear mixed-effects models were run for absolute and relative basal area of all legume species (\(1\) and \(2\)), and for relative basal area of N-fixing and bipinnate legumes separately (\(3\) and \(4\)). Stand age, mean annual rainfall (‘rainfall’) and their interaction were included as fixed effects. Random site intercepts account for between-site variation in initial legume basal area, and random slopes for the variation of the effect of stand age on legume basal area among sites (\(\ast P \leq 0.05; \ast\ast P \leq 0.01; \ast\ast\ast P \leq 0.001\)). The standardized regression coefficients compare the effect of the independent variables on the dependent variable. Values of marginal ($R^2$ (m)) and conditional ($R^2$ (c)) $R^2$ indicate the proportion of the variance explained by the fixed effects of the model, and the fit of the whole model with fixed and random effects, respectively. For all models, $N=42$ chronosequence sites.
in the secondary forests of the dry Neotropics. This strong phylogenetic signal highlights the importance of drought tolerance traits as an adaptation to seasonal dry forests\textsuperscript{1,47}, and demonstrates that leaf traits enhancing moisture conservation are equally as important as the potential to fix N\textsubscript{2} when explaining patterns of legume abundance. N-fixing legumes that have small bipinnate leaves thus hold a double advantage because those traits combine to minimize water loss during C acquisition and lead to increased water-use efficiency.

Across the Neotropics, a substantial amount of the variation in the relative abundance of legumes in secondary forests (17\%) can be predicted by rainfall, stand age and their interaction. The strong negative relationship between legume RA and water availability was not sensitive to our choice of climate variable (either mean annual rainfall, rainfall seasonality or CWD; Fig. 2 and Supplementary Figs. 3 and 4), which suggests that legumes have a competitive advantage in dry climates that are characterized by recurrent seasonal droughts and strong water deficits during the growing season. The climate transition near 1,500 mm yr\textsuperscript{-1}, where the relationship between legume RA and rainfall switched from strongly inverse to flat, coincides with a known cross-over point in woody regeneration\textsuperscript{48}. In regenerating dry forests, the canopy of established plants serves to moderate the harsh microclimatic conditions, thus facilitating the establishment of seedlings\textsuperscript{49}, while in some wet forests, N-fixing legumes act to inhibit the growth of neighbouring trees

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**Fig. 2 | Legume relative basal area across a rainfall gradient in the Neotropics.** For forests in the wet Neotropics, the RA of legume trees is not influenced by variations in mean annual rainfall. However, below a threshold at approximately 1,500 mm yr\textsuperscript{-1}, legume RA increases as total rainfall diminishes. Results are stratified by stand age: a, 2- to 10-year-old forests; b, 11- to 20-year-old forests; c, 21- to 30-year-old forests; d, 31- to 40-year-old forests; e, 41- to 50-year-old forests; f, 61- to 100-year-old forests. Results of a segmented linear fit are shown in each panel (**P < 0.01; ****P < 0.0001); N indicates the number of chronosequence sites with plots in each age category. Grey lines represent fitted values obtained using segmented regression models. Each circle represents site-level legume relative basal area averaged for each stand age category. Dry forest sites (with dominant deciduous vegetation) are indicated in orange, and wet forests in blue. Some dry forest sites receive higher average annual rainfall than some wet forest sites, but these sites are classified as ‘dry’ or ‘wet’ because of their functional composition (evergreen or deciduousness), not their mean climatology.

**Fig. 3 | Relative basal area of legumes for 5- and 20-year-old forests as a function of mean annual rainfall.** a-c. The exceptional ecological success of legumes in recently disturbed, water-limited forests is mainly due to species that (1) are able to fix nitrogen and (2) have bipinnate leaves. Fitted values were obtained using a linear mixed-effects model, with stand age and rainfall as fixed effects and site as random effect (see (3)-(5) in Table 1 for full model results). Relative basal area and stand age were arcsin and log-transformed, respectively, before analysis. Models were computed separately for N-fixing legume species (a), legumes with bipinate leaves (b) and legumes with pinnate leaves (c).
during secondary succession\(^4\). A large fraction (45%) of the variance in legume abundance was associated with site identity (represented in our model as a random factor), which could be related to site-specific factors such as resource limitation (phosphorus, molybdenum, iron\(^{16-23}\)) or biotic interactions (dispersal, herbivory) that influence fixation in individual forest stands. More research is needed to determine how and to what extent these factors influence legume dominance at finer spatial scales.

N-fixing legume species attain their greatest levels of ecological success in Neotropical dry forests, where the benefits of fixation and co-occurring traits related to water conservation outweigh their costs throughout decades of succession. It is thus clear that insights about legumes derived from studies conducted exclusively in wet forests (for example, ref. \(^3\)) cannot be extrapolated across all Neotropical forests, and in particular are not valid for dry forests. As an alternative, future efforts to model the tropical N cycle must account for the strong heterogeneity exhibited by this hyperdiverse family of plants. Incorporating these large-scale abundance trends into Earth system models should allow for more accurate estimates of the potential for symbiotic N fixation across tropical forests. Our study also demonstrates that, even though the potential to fix N\(^2\) through symbiosis is a crucial element of their success, it is not the only trait that accounts for the exceptionally high abundance of legumes in Neotropical forests. Leaf traits related to drought tolerance and water-use efficiency are also key adaptive features of dry forest legumes. In conclusion, our results provide a deeper mechanistic explanation for the abundance of Neotropical Leguminosae trees, which should be increasingly relevant to natural forest regeneration and ecosystem functioning as global temperatures warm and dry conditions become more widespread in the tropics\(^15,16\).

Methods

Chronosequence database. We extracted plot-scale legume abundance (m\(^2\) ha\(^{-1}\)) from 42 previously published chronosequence studies\(^2\) (2ndFOR network; Supplementary Fig. 1 and Supplementary Table 1). Our dataset includes lowland forests between 2 and 100 years old. Plot size varied from 0.008 to 1.3 ha, network, mean annual rainfall varied from 750 to 4,000 mm yr\(^{-1}\) for which a minimum diameter at breast height of 10 cm was used. Across the per site was 14, ranging from 2 to 272. Prior land use in our sites included cattle grazing, shifting cultivation or a combination of the two. In each plot, all woody trees, shrubs and palms \(\geq 5\) cm in diameter were measured and identified, with the exception of sites in Costa Rica (Santa Rosa and Palo Verde) and Puerto Rico (Cayey) for which a minimum diameter at breast height of 10 cm was used. Across the network, mean annual rainfall varied from 750 to 4,000 mm yr\(^{-1}\). Based on local site knowledge, study sites were classified as ‘dry’ forests when the vegetation is mostly drought deciduous, or ‘wet’ forests (mostly evergreen) in all other instances (Supplementary Table 1). Because some tropical wet forests experience annual rainfall averages that overlap with the range exhibited by dry forests (particularly around 1,500 mm yr\(^{-1}\)) \(1,\) we used seasonality in leaf habit (drought deciduous or evergreen) to distinguish between the two main biome types. We also repeated our analysis using two additional measures of water availability in the dry season, when water availability is a stronger limiting factor for plant growth and functioning: the rainfall coefficient of variation from WorldClim, which is an index of seasonality\(^2\), and the climatic water deficit (CWD, in mm yr\(^{-1}\)) defined as months where evapotranspiration exceeds rainfall\(^3\), http://chave.ups-tlse.fr/pantropical_allometry.htm) which reflects the relative intensity of water loss during dry months.

Functional traits. Across all sites, 398 legume species were present (Supplementary Table 2). We classified the potential of each species to fix N\(^2\) based on positive nodulation reports for each species\(^(\text{J.I.S.})\) (I.S. determined the fixation potential of all legume species within our dataset. To accurately represent the morphological variation of leaflets, for each leaf we averaged measurements made on leaflets from the bottom, middle and top of the axis. Leaflet size was measured on three leaves of each species (N = 3 individuals per species) using Neotropical specimens from the University of Minnesota Herbarium (133 of 398 species) or from online databases\(^14,16\). Leaflet length and width were highly correlated \((R^2 = 0.82, P < 0.0001)\). Across our dataset, intraspecific variation in leaflet size was small compared with the proportion of variance explained by species (80% and 81% for leaf length and width, respectively), consistent with refs \(^14,16\). All N-fixation potential and leaf size data for each species are provided in Supplementary Table 3.

Legume abundance. We calculated (1) total legume basal area (m\(^2\) ha\(^{-1}\)) to serve as an approximate estimate of aboveground legume biomass, and (2) relative basal area (%) which summarizes the abundance of legumes compared with all species within each plot and is an ecologically meaningful indication of community composition relative to basal area as a measure of legume ecological success because it reflects legume performance relative to other species. To determine whether N fixation and leaf type influence legume success, we also calculated relative basal area (%) for legume trees grouped by fixation potential and by leaf type. Because the basal area of small diameter trees during early stages of forest regeneration (an approximation of recruitment, we stratified legume basal area by tree diameter classes. We focused our analysis on legume relative basal area as a measurement of legume relative abundance (RA) because it reflects biomass accumulation, but across our dataset, this variable was positively and significantly correlated to relative stem density \((R^2 = 0.38, P < 0.0001)\).

Statistical analyses. To evaluate how legume abundance changed over successional time and along the rainfall gradient, we modelled legume RA as a function of stand age and mean annual rainfall with a linear mixed model using the lmer package \((v. 1.11.11)\) in R. We applied an arcsin and natural-logarithm transformation to improve the normality of RA and stand age, respectively. We included random site intercepts and we expected between-site variation in initial legume abundance and random slopes to account for the variation of the effect of stand age on legume abundance among sites (Fig. 1). P values for the effect of stand age were calculated from the lmerTest package \((v. 2.0.30)\), and random effect P values were estimated via the likelihood ratio test. We obtained \(R^2\) values for the effect of stand age (marginal \(R^2\)) using the entire model \((R^2\text{marg})\) and the null model \((R^2\text{null})\) function in the MuMIn package \((v. 1.15.6)\). To determine whether other climatic variables also predicted legume abundance, we repeated models for legume RA using rainfall seasonality or CWD as the main climatic predictor variable. We compared the three models based on Akaike’s information criterion adjusted for small sample sizes \((\text{AICc})\) and selected the best-supported model with the lowest \(\text{AICc}\). To determine the effect of rainfall on legume abundance at different stand age categories \((2 \leq 10\), 10 \(\leq 20\), 20 \(\leq 30\), 30 \(\leq 40\), 40 \(\leq 60\) and 60 \(\leq 100\) of forest age), we performed a piecewise linear regression using the Segmented package \((v. 0.5.1.4)\). Lastly, to understand the successional change in legume basal area of trees of different size classes \((\leq 10\) cm, \(\leq 20\) cm, \(\leq 30\) cm and \(> 30\) cm), we performed a multiple regression on arsin transformed RA with stand age and forest type \((\text{dry} or \text{wet})\) as covariates. All analyses were performed in R version 3.2.2 (ref. \(^3\)).

Data availability. Plot-level legume basal area data from the 42 Neotropical forest sites are available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.3plk3d2.

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References

Author contributions
M.G. and J.S.P. conceived the idea, all co-authors coordinated the data compilations, M.G. and M.D.G. collected leaf traits data, M.G. analysed the data, D.M.A.R. contributed to the analytical approach, M.G. and J.S.P. wrote the paper, and all co-authors collected field data, discussed the results, gave suggestions for further analyses and commented on the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41559-018-0559-6.
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Correspondence and requests for materials should be addressed to J.S.P.
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Experimental design

1. Sample size
   - Describe how sample size was determined.
   - Sample sizes were determined by the number of plots that individual research teams contributed to the pooled dataset.

2. Data exclusions
   - Describe any data exclusions.
   - The only data exclusions were sites for which the minimum threshold tree diameters were insufficient.

3. Replication
   - Describe whether the experimental findings were reliably reproduced.
   - Our study relied on pooling data from many individual chronosequence studies. As such, we can consider each chronosequence a replicated site.

4. Randomization
   - Describe how samples/organisms/participants were allocated into experimental groups.
   - This does not apply to our dataset.

5. Blinding
   - Describe whether the investigators were blinded to group allocation during data collection and/or analysis.
   - Not applicable.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.

6. Statistical parameters
   - For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

n/a Confirmed

- [x] The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)
- [ ] A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- [ ] A statement indicating how many times each experiment was replicated
- [x] The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section)
- [ ] A description of any assumptions or corrections, such as an adjustment for multiple comparisons
- [x] The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted
- [ ] A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)
- [x] Clearly defined error bars

See the web collection on statistics for biologists for further resources and guidance.
Software

7. Software

Describe the software used to analyze the data in this study.

All statistical analyses were performed in R.

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). Nature Methods guidance for providing algorithms and software for publication provides further information on this topic.

Materials and reagents

8. Materials availability

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

Not applicable.

9. Antibodies

Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

Not applicable.

10. Eukaryotic cell lines

a. State the source of each eukaryotic cell line used.

Not applicable.

b. Describe the method of cell line authentication used.

Describe the authentication procedures for each cell line used OR declare that none of the cell lines used have been authenticated OR state that no eukaryotic cell lines were used.

c. Report whether the cell lines were tested for mycoplasma contamination.

Confirm that all cell lines tested negative for mycoplasma contamination OR describe the results of the testing for mycoplasma contamination OR declare that the cell lines were not tested for mycoplasma contamination OR state that no eukaryotic cell lines were used.

d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use.

Provide a rationale for the use of commonly misidentified cell lines OR state that no commonly misidentified cell lines were used.

Animals and human research participants

11. Description of research animals

Provide details on animals and/or animal-derived materials used in the study.

None used.

Policy information about studies involving human research participants

12. Description of human research participants

Describe the covariate-relevant population characteristics of the human research participants.

None used.