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Limited acclimation in leaf anatomy to experimental drought in tropical rainforest trees

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Dry periods are predicted to become more frequent and severe in the future in some parts of the tropics, including Amazonia, potentially causing reduced productivity, higher tree mortality and increased emissions of stored carbon. Using a long-term (12 year) through-fall exclusion (TFE) experiment in the tropics, we test the hypothesis that trees produce leaves adapted to cope with higher levels of water stress, by examining the following leaf characteristics: area, thickness, leaf mass per area, vein density, stomatal density, the thickness of palisade mesophyll, spongy mesophyll and both of the epidermal layers, internal cavity volume and the average cell sizes of the palisade and spongy mesophyll. We also test whether differences in leaf anatomy are consistent with observed differential drought-induced mortality responses among taxa, and look for relationships between leaf anatomy, and leaf water relations and gas exchange parameters. Our data show that trees do not produce leaves that are more xeromorphic in response to 12 years of soil moisture deficit. However, the drought treatment did result in increases in the thickness of the adaxial epidermis (TFE: 20.5 ± 1.5 µm, control: 16.7 ± 1.0 µm) and the internal cavity volume (TFE: 2.43 ± 0.50 mm\textsuperscript{3} cm\textsuperscript{-2}, control: 1.77 ± 0.30 mm\textsuperscript{3} cm\textsuperscript{-2}). No consistent differences were detected between drought-resistant and drought-sensitive taxa, although interactions occurred between drought-sensitivity status and drought treatment for the palisade mesophyll thickness (P = 0.034) and the cavity volume of the leaves (P = 0.025). The limited response to water deficit probably reflects a tight coordination between leaf morphology, water relations and photosynthetic properties. This suggests that there is little plasticity in these aspects of plant anatomy in these taxa, and that phenotypic plasticity in leaf traits may not facilitate the acclimation of Amazonian trees to the predicted future reductions in dry season water availability.

**Keywords**: Amazon, anatomical plasticity, gas exchange, leaf physiology, through-fall exclusion, water relations, water stress.

Introduction

A key issue in the prediction of future climate change is understanding how forests, significant stores of carbon (Pan et al. 2011, Grace et al. 2014), will respond to current and future changes in temperature and water availability (Bonan 2008). Tree mortality has reportedly increased in response to episodic severe drought (Breshears et al. 2005, Allen et al. 2010), including in the tropics (Nakagawa et al. 2000, Meir and Grace 2005, Phillips et al. 2009, Brienen et al. 2015), and understanding the physiology underlying drought-induced mortality is essential for estimating forest sensitivity to drought (Christensen et al. 2013, Allen et al. 2015, Meir et al. 2015b). Although spatial variation in predicted rainfall patterns is substantial, current consensus suggests that precipitation extremes in the tropics, and especially in Amazonia, are likely to become more frequent, with extended dry seasons of particular note (Christensen et al.
These changed conditions will exert a selection pressure affecting the next generation of trees, but the persistence of the current generation depends on their capacity for acclimation or resilience in the face of climate change. Investigating the capacity of trees to cope with drought in tropical forests is consequently of paramount importance in estimating the magnitude of biosphere–atmosphere feedbacks.

Species differ in their ability to cope with water stress (da Costa et al. 2010, Bartlett et al. 2012, Chaot et al. 2012, Meir et al. 2015a, Rowland et al. 2015b) and establishing exactly what traits account for this differential susceptibility is complex, particularly in species-diverse communities. Globally, there have been reports of drought-induced tree mortality with the implicated cause being either carbon starvation or hydraulic failure, or a mixture of the two (McDowell 2011, Anderegg et al. 2012, Galiano et al. 2012, Anderegg et al. 2013, Hartmann et al. 2013, McDowell et al. 2013, Brödribb and McAdam 2015, Gleason et al. 2015). For the tropics, however, there have been few studies, with initial suggestions hinting at a role for progressive carbon starvation (Metcalfe et al. 2010) superseded by more recent evidence pointing towards hydraulic deterioration as a principal trigger for drought-induced mortality (Rowland et al. 2015). Water stress in plants is commonly represented by the percent loss of hydraulic conductivity (PLC) of the xylem, whereby P50, the leaf or branch tissue water potential at 50% PLC, is used as a metric of the drought resistance of a plant or species, with higher (less negative) P50 indicating less resistance to the loss of conductivity through embolism. Attempts have been made to map P50 onto xylem anatomy, where wood density, conduit diameter and conduit wall thickness have been found to be weakly predictive of cavitation resistance (Hacke et al. 2001, Hajek et al. 2014, Gleason et al. 2015). Certain characteristics of leaves have also been shown to be associated with drought resistance in plants, e.g., turgor loss point (Bartlett et al. 2012) and the elastic modulus (Bowman and Roberts 1985), but the mechanistic relationships between leaf anatomy and drought resistance remain poorly understood.

Distinct morphological characteristics of leaves occur in environments of especially low or high water availability, and are termed xeromorphic and hygromorphic, respectively. Xeromorphic leaves tend to be small in area, with a multi-layered epidermis, thick cuticle, compactly arranged mesophyll with little air space, high stomatal density and high vein density (Maximov 1929, Cutler et al. 1977). By contrast, hygromorphic leaves tend to show the opposite features (Schimper 1903, Roth 1985). In certain species spanning gradients of rainfall, traits such as leaf area, thickness, specific leaf area, density, and stomatal morphology have been shown to vary with water availability (Geeske et al. 1994, Cunningham et al. 1999, Warren et al. 2005, McLean et al. 2014). Drought experiments have also shown reduced cell size in the mesophyll and epidermis, and increases in cell wall thickness, stomatal density, vein density and cuticle thickness in droughted vs non-droughted plants (Maximov 1929, Morton and Watson 1948, Shields 1950, Cutler et al. 1977).

Most of the experimental research on the adaptations of leaf anatomy to water stress was conducted over half a century ago on mesophytic crop plants (Maximov 1929, Morton and Watson 1948, Shields 1950, Cutler et al. 1977), but its relevance is current in the context of predicted changes in the hydrological regimes of tropical rainforests. Moreover, recent work has attempted to improve the mechanistic understanding of water movement from veins to stomata in leaves, by modelling the hydraulic pathway through cells, cell walls and as vapour through the internal airspaces (Rockwell et al. 2014a, 2014b, Buckley 2015, Buckley et al. 2015). Therefore, understanding the plastic response of leaves at the tissue level, may well be informative of the influence of cell structure on the hydraulic pathway, and how this affects water use at the leaf level. Thus, the question arises: can trees respond to long-term reductions in water availability by producing leaves that exhibit more xeromorphic characteristics? Such a capacity for acclimation could confer a significant advantage for long-lived canopy tree species (Nicotra et al. 2010), and could be important for determining the sensitivity of a forest to drought and the difference in drought sensitivity among species. Existing studies addressing this question in natural communities are observational (Geeske et al. 1994, Cunningham et al. 1999, McLean et al. 2014), thus, here we test the plasticity of leaf morphology experimentally, in tropical rainforest trees.

This study uses a long-term (>12 years) through-fall exclusion (TFE) experiment in the lowland Amazon Rainforest (da Costa et al. 2010, Meir et al. 2015) to address the following questions: (i) do trees respond to long-term imposed soil moisture deficit through changes in leaf structure or anatomy? and (ii) do differences in anatomy, or anatomical plasticity, explain contrasts in drought sensitivity among taxa? We also examine any further associations between leaf anatomy, water relations and gas exchange traits and drought using multivariate analyses. The expectation was that xeromorphic traits should be found particularly under conditions where leaves have to cope with drought stress. In particular, higher stomatal and vein density, smaller cell size in the spongy and palisade mesophyll, and lower cavity volume were expected to occur in the experimentally droughted forest and/or in individuals from drought-resistant genera. Additionally, to highlight potential links between anatomical and physiological properties of leaves, multivariate analyses were carried out combining leaf tissue properties with plant water relation traits and gas exchange parameters.

**Methods**

**Study site**

The field work was conducted in the Caxiuana National Forest Reserve in the eastern Amazon Rainforest (1°43’S, 51°27’W).
The field site is situated in lowland *terra firme* rainforest approximately 10–15 m above river level, has a mean temperature of ca. 25 °C, receives 2000–2500 mm of rainfall annually and has a dry season in which rainfall is <100 mm per month between June and November.

**Large-scale TFE experiment**

The TFE experiment consists of a 1 ha plot from which approximately 50% of canopy through-fall has been excluded using plastic panels located 1–2 m above the ground since 2002. A 1 ha control plot, <50 m from the TFE, which has received normal rainfall, was also studied. The plots are divided into 10 m × 10 m subplots, of which the outer-most subplots were excluded from the study to account for possible edge effects on tree growth. Further details on experimental setup and results can be found in Fisher et al. (2007), Meir et al. (2009), Metcalfe et al. (2010), da Costa et al. (2010) and Rowland et al. (2015b).

**Study specimens and drought vulnerability status**

All measurements (Table 1) were taken from six genera common to both the TFE and the control plot of which *Manilkara*, *Eschweilera* and *Pouteria* have been classified ‘drought sensitive’ and *Protium*, *Swartzia* and *Licania* as ‘drought resistant’, based on analysis of rates of drought-induced mortality (da Costa et al. 2010, Rowland et al. 2015b). These will be subsequently referred to as sensitive and resistant species. The sensitivity status of a genus is based on mortality in response to the imposed drought conditions. Where possible, a single species was used to represent a genus (*Pouteria anomala* (Pires) T.D. Penn., *Manilkara bidentata* (A.DC.) A. Chev., *Swartzia racemosa* (Benth.)), but more than one species was used where there were too few individuals in a species per plot. So *Eschweilera* is represented by the species *E. coriacea* (DC.) S.A. Mori, *E. grandiflora* (Aubl.) Sandwith, and *E. pedicellata* (Rich) S.A. Mori, *Licania* by *L. membranacea* (Sagot ex Laness) and *L. octandra* (Kuntze) and *Protium* by *P. tenuifolium* Engl. and *P. paniculatum* Engl. This approach was necessary to obtain sufficient numbers of each genus per plot to enable a comparison of drought sensitivity groups, i.e., in order that drought-sensitive and -resistant taxa were represented by three genera in both plots, and was used in two previous studies undertaken at the same site (Rowland et al. 2015b, Binks et al. 2016). In Binks et al., variance was consistently greater among genera than amongst individuals within genera. Similarly, in this study variance was also greater among, than within, genera in 13 out of 17 traits, the exceptions being leaf area, leaf mass per area and the proportional tissue thicknesses of the spongy mesophyll and abaxial epidermis (Table 2).

For each of the variables, an attempt was made to measure at least two leaves per individual tree, and three individual trees per genus per plot which would have resulted in 18 leaves from nine individuals per drought sensitivity status, per plot. Unfortunately, it was not always possible to achieve this number of samples, due to difficulties obtaining suitable leaves and performing the analysis under field conditions. Moreover, peculiarities in some of the specimens made particular analyses difficult or impossible; e.g., leaves from the genus *Manilkara* were densely packed with sclereids, which obscured the vasculature and made accurate analysis of vein density impossible under the conditions available. Therefore, the minimum and mean number of leaves per genus, genus per plot and drought sensitivity status

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**Table 1.** Data transformations and final model structures used in analysis for the effect of treatment (T, control plot vs TFE) and drought-sensitivity status (S, sensitive or resistant) on tissue parameters, and gas exchange parameters used in PCA. The random effects were tree individual nested inside genera for all models with the exception of CVprop, for which tree individual was not used because of sample size limitations (see Table S1 available as Supplementary Data at Tree Physiology Online).

| Leaf properties | Response variable | Symbol | Transformation | Model structure |
|-----------------|-------------------|--------|---------------|----------------|
| Structural properties | Leaf area | A | log | T S |
| | Leaf thickness | T | log | T S |
| | Leaf mass per area | LMA | log | T S |
| | Vein density | VD | y^2 | T S |
| | Stomatal density | SD | – | T S |
| Tissue properties | Spongy mesophyll thickness | SM | – | T S |
| | Adaxial epidermis thickness | Ad | log | T S |
| | Abaxial epidermis thickness | Ab | log | T S |
| | Internal cavity volume | CV | – | T S |
| | Proportional SM thickness | SMprop | y^2 | T-1 S |
| | Proportional Ab thickness | Abprop | log | T S |
| | Proportional Ad thickness | Adprop | log | T S |
| | Proportional CV thickness | CVprop | y^2 | T S |
| Cellular properties | SM cell volume | SM_{cell_volume} | y^2 | T S |
| | Pal cell volume | Pal_{cell_volume} | log | T S |
| Gas exchange parameters | Rubisco carboxylation | V_{cmax} | – | PCA |
| | Electron transport | J_{max} | – | PCA |
| | Dark respiration | R_{dark} | – | PCA |
Table 2. Variance accounted for by separate components in the mixed models and the conditional and marginal $r^2$ of each model.

| Variable | Variance (%) | $r^2$ Conditional | $r^2$ Marginal |
|----------|--------------|-------------------|----------------|
|          | Fixed | Random | ID  |         |         |        |        |
| T        | 5.8   | 58.3   | 29.9 | 6.0    | 0.94   | 0.06   |
| A        | 2.0   | 32.8   | 33.9 | 31.3   | 0.69   | 0.02   |
| Ad       | 1.8   | 71.5   | <0.1 | 26.7   | 0.73   | 0.02   |
| SM       | 8.0   | 45.5   | 27.8 | 18.8   | 0.81   | 0.08   |
| Pal      | 7.2   | 39.8   | 35.9 | 17.0   | 0.83   | 0.07   |
| CV       | 9.0   | 55.2   | 29.9 | 5.9    | 0.94   | 0.09   |
| Ab       | 4.2   | 45.1   | 24.5 | 26.1   | 0.74   | 0.04   |
| Adprop   | 16.7  | 63.1   | 7.2  | 13.0   | 0.87   | 0.17   |
| Smprop   | 19.0  | 26.2   | 33.0 | 29.8   | 0.70   | 0.14   |
| Palprop  | 11.8  | 33.1   | 28.5 | 26.6   | 0.73   | 0.12   |
| CVPprop  | 18.1  | 47.4   | 34.5 | 35.9   | 0.76   | 0.06   |
| Abprop   | 14.3  | 15.7   | 40.2 | 29.8   | 0.70   | 0.14   |
| VD       | 1.9   | 69.3   | 6.7  | 22.0   | 0.78   | 0.02   |
| SD       | 3.8   | 39.9   | 38.6 | 17.8   | 0.82   | 0.04   |
| LMA      | 4.1   | 13.1   | 44.8 | 38.0   | 0.62   | 0.04   |
| SMcell_volume | 23.8 | 20.5  | 14.7 | 40.9   | 0.59   | 0.24   |
| Palcell_volume | 20.3 | 49.0  | 10.1 | 20.6   | 0.79   | 0.20   |

Table 3. Mean value by genus of each of the leaf tissue parameters ± 1 standard error.

|          | Drought sensitive | Drought resistant |
|----------|-------------------|-------------------|
|          | Eschweilera       | Manilkara         | Pouteria         | Licania         | Protium        | Swartzia        |
| T (µm)   | 164.7 ± 4.7       | 247.2 ± 15.9      | 168.9 ± 6.7      | 165.5 ± 27.7    | 123.5 ± 9.9    | 248.4 ± 8.4    |
| A (cm²)  | 69.5 ± 4.4        | 29.9 ± 1.5        | 43.4 ± 1.7       | 40.8 ± 1.8      | 47.1 ± 2.1     | 39.4 ± 2       |
| Pal (µm) | 35.9 ± 1.3        | 66.7 ± 8.1        | 60.6 ± 6.5       | 56.8 ± 6.5      | 42.8 ± 3.2     | 66.8 ± 3.8     |
| SM (µm)  | 100.6 ± 4.7       | 150.5 ± 9.1       | 82.9 ± 7.7       | 71.9 ± 17.7     | 59.2 ± 5.4     | 139.9 ± 5.5    |
| Ad (µm)  | 15.9 ± 0.8        | 13.1 ± 1.5        | 15.6 ± 0.8       | 34.3 ± 1.8      | 12.4 ± 1.5     | 19.9 ± 1.1     |
| Ab (µm)  | 12.3 ± 0.6        | 15.4 ± 1.9        | 13.1 ± 0.7       | 15.6 ± 2.3      | 9.2 ± 0.9      | 22.5 ± 2       |
| CV (mm³·cm⁻²) | 2.1 ± 0.3     | 3.3 ± 0.2         | 0.1 ± 0          | 3.1 ± 0.7       | 0.3 ± 0        | 2.5 ± 0.4      |
| VD (mm²) | 8.4 ± 0.3         | –                 | 9.6 ± 0.2        | 11.4 ± 0.6      | 11.5 ± 0.3     | 6.3 ± 0.2      |
| SD (mm⁻¹) | 409.7 ± 12.2     | 317.3 ± 20.6      | 350.1 ± 28.8     | 306.1 ± 32.3    | 563.5 ± 12.7   | 301 ± 10.5     |
| LMA (m)  | 91.1 ± 1.7        | 125.8 ± 5.8       | 103.3 ± 3.6      | 116.6 ± 4.8     | 90.6 ± 1.6     | 115.4 ± 2.7    |
| Palprop (%) | 22.2 ± 1.1     | 26.3 ± 2.1        | 35.9 ± 3.7       | 37.1 ± 3.4      | 34.9 ± 1.1     | 26.4 ± 1.3     |
| Smprop (%) | 60.6 ± 1.6      | 63.7 ± 2.2        | 46.8 ± 3.4       | 41.6 ± 4.3      | 47.7 ± 1.1     | 55.9 ± 2.1     |
| Abprop (%) | 7.6 ± 0.4       | 5.2 ± 0.5         | 7.9 ± 0.5        | 9.8 ± 0.8       | 7.5 ± 0.5      | 9.5 ± 1.1      |
| Adprop (%) | 9.6 ± 0.4       | 4.9 ± 0.6         | 9.4 ± 0.6        | 24.1 ± 2.8      | 10.9 ± 0.9     | 8.3 ± 0.5      |
| SMcell_volume (µm³) | 3456 ± 421      | 2614 ± 224        | 2641 ± 295       | 884 ± 67        | 946 ± 193      | 2656 ± 348     |
| Palcell_volume (µm³) | 1220 ± 106      | 3695 ± 263        | 4371 ± 440       | 823 ± 109       | 1797 ± 204     | 2046 ± 683     |
at a pressure of 18 kPa. The cavity volume (CV) was determined by subtracting the fresh leaf mass from the perfused leaf mass expressed as mm$^3$ air space cm$^{-2}$ leaf surface.

**Vein density**

Small squares of leaf approximately 1 cm$^2$ in size taken from midway between the tip and base, and midrib and margin of the leaf were cleared using 5% NaOH and briefly 10% NaClO when necessary to remove the last of the colour (Scoffoni et al. 2010). The cleared leaf sections were then placed in a 1% solution of toluidine blue for several seconds before being rinsed in water; this process was repeated until sufficient dye was judged to have infiltrated the sample and the veins were clearly visible. The samples were photographed using a Moticam 2 digital camera on a Motic B3 microscope (Motic, Canada). An objective lens of ×10 magnification was used for most images, but ×4 magnification was used where Student’s t-test revealed no significant difference (P > 0.05) in the vein densities calculated from either magnification. Vein density (VD, mm mm$^{-2}$) was derived by tracing and measuring vein length in a known area using ImageJ software (Schneider et al. 2012). Where there was a clear distinction in size between second and third order veins, only those in the third order and above were included in the analysis (where the midrib counts as the first order).

**Stomatal density**

Dental impression gel was used to cover a minimum area of 2 cm$^2$, situated midway between the tip and base, of the abaxial surface of four leaves per individual and the adaxial surface of one leaf per individual. Clear nail varnish was applied to the surface of four leaves per individual and the adaxial surface of leaves were scanned to enable the determination of area using ImageJ software (Schneider et al. 2012) and then dried at 70 $^\circ$C to a constant dry mass. Clear nail varnish was applied to the surface of four leaves per individual and the adaxial surface of one leaf per individual. Clear nail varnish was applied to the surface of four leaves per individual and the adaxial surface of leaves were scanned to enable the determination of area using ImageJ software (Schneider et al. 2012) and then dried at 70 $^\circ$C to a constant dry mass.

**Pressure volume analysis**

Pressure–volume (PV) curves were carried out as per the bench-drying protocol described in Tyree and Hammel (1972). Briefly, leaves were taken from branches that had been rehydrated by allowing to stand overnight in a bucket of water filtered to 0.2 µm. The leaves were allowed to dehydrate over a period of 3–8 h, during which time water potential and mass were measured repeatedly using a Scholander pressure bomb (PMS Instruments Co., Corvalis, OR, USA) and mass balance accurate to 0.1 mg, respectively. After the final set of measurements, leaves were scanned to enable the determination of area using ImageJ software (Schneider et al. 2012) and then dried at 70 $^\circ$C in an oven for >48 h to find dry mass. The parameters osmotic potential at full turgor ($\Psi_s$, MPa), turgor loss point ($\Psi_{tlp}$, MPa), saturated water content (the ratio of water mass to leaf dry mass in a fully saturated leaf, SWC, g g$^{-1}$), relative water content at x$_{tip}$ (RWC$_{tip}$, %), modulus of elasticity ($\epsilon$, MPa) and hydraulic capacitance (C, mol MPa$^{-1}$ m$^{-2}$) were calculated as per Sack and Pasquet-Kok (2011). Differences in PV parameters across drought sensitivity status and plots are reported in a separate paper (Binks et al. 2016). Here, we present a correlation analysis including PV, leaf anatomy and gas exchange parameters in the Supplementary Data. The PV parameters were averaged across the wet and dry season for the correlation analysis as the magnitude of seasonal variation is less significant than the differences between species (Bartlett et al. 2014), and thus the benefit of doubling the sample size outweighed the cost of the slight increase in variance. Moreover, seasonal variation was only significant in SWC, RWC$_{tip}$ and C, but not in $\Psi_s$, $\Psi_{tlp}$ or $\epsilon$ (Binks et al. 2016).

**Photosynthesis**

For a detailed description of how the gas exchange parameters were measured, refer to Rowland et al. (2015b), in which these parameters were presented in the context of the experimental drought. Photosynthesis was measured on canopy top branches using LICOR 6400 portable photosynthesis systems (LI-COR, Lincoln, NE, USA). The parameters $V_{cmax}$ (the maximum rate of Rubisco carboxylation) and $J_{max}$ (the maximum rate of electron transport) were derived from A–C, curves performed under saturating photosynthetically active radiation (PAR), and data were temperature corrected to 25 $^\circ$C following Sharkey et al. (2007). To measure dark respiration ($R_{dark}$) leaves were covered in tinfoil for 30 min prior to, and during, gas exchange measurements, and these data were also temperature corrected to 25 $^\circ$C according to Atkin and Tjoelker (2003).

**Predawn water potentials**

Water potential was measured in three leaves per individual tree, between 5.30 and 7.00 in the morning at the end of the dry season (October) in 2013. Values were averaged for each individual tree.

**Analysis**

The data were analysed using linear mixed effects models in the packages lmer and lmerTest in R (R Core Team 2015). Because the study was focused on finding treatment and drought sensitivity effects, and not the responses of individual genera, genus was designated as a random effect, and tree individual was nested inside genus, where there were >2 individuals per genus per plot. This statistical design removes the variance attributable to individuals within genera, and between genera, in order to selectively find the influence of the fixed effects, e.g., treatment and drought sensitivity status. Models (Table 1) were simplified by comparing their respective AIC. The distributions of all of the variables were checked for normality using the Shapiro–Wilk test and either log or power transformed depending on the starting...
distribution. The powers employed for transformation were determined using a Box–Cox transformation function in the MASS package in R (Venables 2002). The marginal and conditional $r^2$ were calculated according to Nakagawa and Schielzeth (2013).

All of the anatomy variables were tested for correlations with the parameters derived from the PV analysis, predawn water potential ($\Psi_{PD}$) and gas exchange (correlation matrix, see Table S2 available as Supplementary Data at Tree Physiology Online). The analysis was carried out using Pearson correlation analyses in the R package ’psych’ (Revelle 2015) and all variables were transformed as per the mixed effects models (Table 1).

A principal component analysis (PCA) was performed on the thicknesses of leaf tissues, vein and stomatal density and the gas exchange-derived parameters $V_{cmax}$, $J_{max}$ and $R_{dark}$ to highlight possible relationships between anatomy and photosynthesis. Absolute values of tissue thickness were used because of the relevance of distance, i.e., tissue thickness, to the molecular diffusion processes.

**Results**

Leaf traits by genus are presented in Table 3. Leaf thickness varied from 78 to 370 $\mu$m with a combined mean and standard error of all genera and treatments of 187.6 ± 7.2 $\mu$m and a mean relative thickness with standard error of palisade, spongy mesophyll, abaxial and adaxial epidermis of 29.5 ± 1.3, 52.9 ± 1.49, 8.1 ± 0.3 and 11.5 ± 0.9%, respectively (Figure 1).

Significant differences in response to treatment ($P < 0.05$) amongst all taxa combined were found for the absolute measure of cavity volume (TFE 2.43 ± 0.50 mm$^3$ cm$^{-2}$, control $1.77 ± 0.30$ mm$^3$ cm$^{-2}$, Table 4) and the thickness of the adaxial epidermis (TFE 20.5 ± 1.5 $\mu$m, control 16.7 ± 1.0 $\mu$m, Table 4, Figure 2). Total leaf thickness ($P = 0.070$), palisade thickness ($P = 0.086$), the proportional cavity volume ($P = 0.069$) and LMA (leaf mass per area, $P = 0.098$) were found to be marginally significant (0.05 < $P < 0.1$). None of the measured variables showed significant differences between the two drought sensitivity classes (Table 4). However, CV and palisade thickness showed significant interactions between treatment and sensitivity status ($P = 0.025$ and 0.034, respectively), whereby the treatment effect was stronger (i.e., values were lower in the control plot) amongst the resistant compared to the sensitive genera (Figure 3).

The variance accounted for by the fixed effects (see Table 1 for model structures) varied from 1.8% for the adaxial epidermis to 23.8% for the spongy mesophyll cell volume with a mean of 10.1% over all of the variables (Table 2). In all but 4 out of 17 traits variance was higher among genera than within genera, averaging 42.7 and 27.1%, respectively, suggesting that the analysis of the effects was robust to pooling. Of the four traits where more variance occurs within a genus, leaf area, SM$_{prop}$, Ab$_{prop}$ and LMA, none were found to have significant treatment or drought sensitivity effects.

The correlation matrix of all the anatomical parameters, the PV parameters, the gas exchange parameters and predawn water potentials is given in Table S2 available as Supplementary Data at Tree Physiology Online. The thickness of the spongy mesophyll and adaxial epidermis correlated with more parameters than the other tissue layers, suggesting that they were tightly associated with other leaf traits, although they appeared to operate antagonistically, i.e., Ad$_{prop}$ and SM$_{prop}$.
were negatively correlated with each other, and had opposite associations with the other leaf traits. Both Ad and Adprop were positively correlated with $\varepsilon$ and $\Psi_{PD}$. The photosynthesis parameters correlated positively with both palisade and spongy mesophyll thickness and negatively with vein and stomatal density, but did not correlate with the thickness of the epidermal layers.

In the PCA, combining anatomical and gas exchange traits, the first and second axes explained 57.6 and 16.2% of the variance, respectively (Figure 4). Palisade thickness was grouped with $I_{\text{max}}$ and $V_{\text{max}}$ at high values of both axis 1 and 2, in the opposite quadrant to $R_{\text{dark}}$. For the other parameters, there was a gradient of high vein and stomatal density, at low values of axis 1 and high values of axis 2, to high thickness of spongy mesophyll and epidermis in the opposite quadrant, which was orthogonal to the photosynthesis traits.

### Discussion

The results of this study reveal that little anatomical change occurred in response to the soil moisture deficit following 12 years of imposed through-fall exclusion. In addition, the drought sensitivity classification in these tropical forest trees, as determined by their increased mortality risk during drought stress, is not linked to specific leaf anatomy traits. We hypothesized that leaves would become more xeromorphic in character in response to the treatment, i.e., thicker, with smaller area, having lower internal cavity volume and higher stomatal and vein density. In fact, the only traits that did vary significantly in response to the treatment were the thickness of the adaxial epidermis (Figure 2) and the cavity volume, the second of which, contrary to expectation, increased in response to the imposed drought (Figure 3a).

### Drought sensitivity status

There were no significant differences in leaf anatomy based on sensitivity status, implying that other aspects of plant physiology determine sensitivity to water stress. However, the interaction between sensitivity status and treatment for cavity volume and palisade thickness indicates a possible link between drought sensitivity and plasticity in these traits. Values of cavity volume were similar among sensitive and resistant genera in the TFE, whereas the values of the resistant genera in the control plot were lower than the overall mean (Figure 3a). In other words, the acclimation response to the drought stress brought the value for CV amongst the resistant taxa in the TFE in-line with the values of the sensitive taxa in the TFE, bringing into question any drought-related benefit of plasticity in this trait. Therefore, there is no strong evidence in our dataset to suggest that the leaves of drought-resistant species are consistently different from those of drought-sensitive species.

Despite the strong general relationship between palisade thickness and photosynthesis (Chabot and Chabot 1977, Smith et al. 1998, Hanba et al. 2002, Catoni et al. 2015), and also observed in this data (see Table S2 available as Supplementary Data at Tree Physiology Online), earlier work by Rowland et al. (2015b) on the same experiment demonstrated no effect of plot or drought sensitivity on photosynthetic capacity or leaf nitrogen content. Therefore, the palisade is significantly thicker for the resistant species on the TFE (with a marginally significant difference between plots, Figure 3b, Table 4), but this has not resulted in higher photosynthetic capacity as determined by maximum rates of Rubisco carboxylation ($V_{\text{max}}$) or electron transport ($I_{\text{max}}$) (Rowland et al. 2015b). Moreover, the lack of significant change in N content suggests that Rubisco content (or Rubisco activation) has not changed considerably between treatments. These findings demonstrate that, at least in this case, the change measured in palisade thickness is not related to the maximum photosynthetic capacity. However, at least one other

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**Table 4.** Probability values of the fixed effects included in the mixed models listed in Table 1, values in bold indicate a significant effect at $P < 0.05$. Factors with a dash were not included in the final model.

| Variable | Treatment | Drought sensitivity | Interaction treat. × sens. |
|----------|-----------|---------------------|---------------------------|
| T        | 0.070     | 0.421               | 0.181                     |
| A        | 0.666     | 0.826               | 0.834                     |
| Ad       | 0.038     | –                   | –                         |
| SM       | 0.543     | 0.488               | 0.757                     |
| Pal      | 0.086     | 0.618               | 0.034                     |
| CV       | 0.009     | 0.565               | 0.025                     |
| Ab       | 0.195     | 0.893               | 0.611                     |
| Adprop   | 0.634     | 0.266               | 0.750                     |
| Smprop   | 0.786     | 0.417               | 0.147                     |
| Palprop  | 0.848     | 0.632               | 0.212                     |
| CVprop   | 0.069     | 0.832               | 0.054                     |
| APprop   | 0.972     | 0.173               | 0.832                     |
| VD       | 0.756     | 0.540               | 0.379                     |
| SD       | 0.797     | 0.638               | 0.470                     |
| LMA      | 0.098     | 0.591               | 0.278                     |
| SMcell_volume | 0.914 | 0.083 | 0.896 |
| Palc_volume | 0.193 | 0.131 | 0.098 |

**Figure 2.** Treatment effect on the thickness of the adaxial epidermis of all studied taxa combined ($P = 0.038$).
experiment has shown that the palisade mesophyll thickness increased in response to water deficit (Boughalleb and Hajlaoui 2011), suggesting that this trait may also influence water use within the leaf, or be influenced by water status during ontogeny.

**The drought effect**

The cause of the higher CV in the drought plot (Figure 3a) is unknown as this is a feature of leaf morphology that has been explored little in the context of water stress, and the response of CV to long-term conditions of low VPD is not consistent among studies (Leuschner 2002, Aliniaeifard et al. 2014). We speculate that a higher internal cavity volume may reduce internal vapour pressure, both because of the effect of the larger cavity (greater distance between adjacent cell walls) and/or because of longer apoplastic path lengths resulting in lower local water potentials, with the consequence of potentially increasing photosynthetic water use efficiency (Mediavilla et al. 2001).

Several previous studies have linked thicker adaxial epidermis with drought resistance (Bacelar et al. 2004, Boughalleb and Hajlaoui 2011), although the causes for this remain uncertain. The results of this study show that the adaxial epidermis is thicker in the TFE and correlates positively with predawn water potential (see Table S2 available as Supplementary Data at *Tree Physiology* Online), while the results of a previous study, on the same taxa and individuals, showed that the thickness of the adaxial epidermis correlates negatively with osmotic potential at full turgor \((\Psi \pi)\) and turgor loss point \((\Psi \pi)\) and positively with the elastic modulus \((\varepsilon)\) (Binks et al. 2016, see Table S2 available as Supplementary Data at *Tree Physiology* Online). If it is assumed that these correlations arose because turgor and/or osmotic properties in the Ad differ from those in other parts of the leaf, then the Ad would appear to be a particularly drought resistant tissue, e.g., low \(\Psi \pi)\) and \(\Psi \pi)\), and high \(\varepsilon\). Thus, a thicker Ad would be linked to higher drought resistance which may explain why it is significantly thicker in the drought plot (Table 4, Figure 2).

**Water deficit and leaf expansion during ontogamy**

The Lockhart equation explains the mechanical relationship between turgor pressure and the rate of cell expansion \((E)\) in which \(E = m(\Psi \pi) - \Psi \pi)\) where \(\Psi \pi\) is turgor pressure, \(\Psi \pi)\) is the threshold turgor pressure below which growth does not occur, and \(m\) is the cell wall extensibility (Lockhart 1965). Leaves will, therefore, be smaller if their expansion phase occurs during periods of water stress (Shields 1950). Because vascular tissue and stomata are differentiated prior to expansion, reductions in leaf size effected during growth can be associated with increases in vein and stomatal density (Hsiao 1973, Schoch et al. 1980, Carins Murphy et al. 2014). However, no differences were detected in leaf area, cell size, vein or stomatal density in response to the drought treatment indicating that turgor pressure must not have dropped below the threshold minimum for long enough during the leaf expansion phase to influence these parameters in the mature leaves. All of the study species were evergreen but a partial leaf flush occurs at the beginning of the wet season (unpublished data) during which most leaves complete their growth. Thus, the wet season water supply on the TFE appears to be non-limiting to growth, which may be facilitated by reduced stomatal conductance to maintain adequately high cell
turgor, and the reduction in leaf area index on the TFE (Metcalfe et al. 2010).

One factor that is interesting to consider in relation to the effects of experimental drought, is the potentially challenging aspect of separating mechanical effects of reduced turgor on growth, from the active expression of plastic traits that facilitate drought resistance, i.e., passive versus active plasticity (Valladares et al. 2007). In the context of leaves, this leads to the question of whether the traits that emerge as a consequence of expansion under sub-optimal turgor pressure, e.g., smaller area, higher vein and stomatal density and smaller mesophyll cells, actually provide an adaptive advantage for drier conditions, or are simply the product of drought stress during ontogeny. The similarity between leaf traits that emerge in response to experimental drought (Maximov 1929, Cutler et al. 1977), and those that differ along precipitation gradients (Geeske et al. 1994, Cunningham et al. 1999, Warren et al. 2005, McLean et al. 2014) presumably suggests the former: that these traits offer an adaptive advantage to coping with water stress.

Correlations between leaf anatomy, gas exchange, predawn water potential and pressure volume traits

The correlation matrix revealed that several leaf tissues appear to be associated with many water relation and gas exchange traits (see Table S2 available as Supplementary Data at Tree Physiology Online), suggesting that these tissues are particularly representative of overall leaf physiology despite the limited treatment effects. These results must be interpreted with caution as the traits in this analysis are not independent; for example, leaf thickness is the sum of the thicknesses of all tissues layers, and similarly $\Psi_{tlp}$ is a function of $\Psi_p$ and $\epsilon$, so these traits inevitably correlate. Having said that, the proportional thickness of the adaxial epidermis and the absolute thickness of the spongy mesophyll correlated with a larger number of traits than the other tissues, and always in opposite directions (see Table S2 available as Supplementary Data at Tree Physiology Online). For example, SM correlates positively with $\Psi_{tlp}$, SWC and RWC$^{lp}$ and negatively with SD, VD and $\Psi_{PD}$, while $Ad_{prop}$ shows the opposite relationships. Thus, leaves with thicker adaxial epidermis and thinner spongy mesophyll and, therefore, low $\Psi_{tlp}$ and high $\Psi_{PD}$, should be associated with greater drought resistance.

A high value of the elastic modulus is generally associated with drought resilience because it results in a greater change in $\Psi$ for a given amount of water loss (Bowman and Roberts 1985), thus increasing the potential gradient and the capacity for rehydration. The correlation matrix reveals a positive relationship between $\epsilon$ and $\Psi_{PD}$ supporting this theory, suggesting that high elastic modulus could be advantageous for nocturnal drought stress recovery in these taxa. In a previous study (Binks et al. 2016), $\epsilon$ was not found to vary significantly with drought sensitivity status but was significantly higher across all groups in the TFE than the control plot.

Principal component analysis

Given the association of the palisade mesophyll with photosynthesis, it is not surprising that it is clustered with $J_{max}$ and $V_{cmax}$ in the PCA (Figure 4). However, it is surprising that stomatal and vein density are on a vector orthogonal to the photosynthesis traits, as in other studies they have been shown to correlate positively with photosynthesis (Brodribb et al. 2010, Walls 2011, ...
Muller et al. 2014, Zhang et al. 2014), while, in this dataset, they correlate negatively (see Table S2 available as Supplementary Data at Tree Physiology Online). Acting in the opposite direction to VD and SD are the other leaf traits, SM, Ab and Ad, which may suggest that increases in the thickness of these tissues can compensate for the functions of otherwise higher vein and stomatal density. Past research analysing the movement of dye from leaf veins into the surrounding tissue indicates that the epidermal layers play a role in lateral water transport (Wylie 1943), which might explain the negative correlation between vein density and the epidermal layers (Figure 4, see Table S2 available as Supplementary Data at Tree Physiology Online).

**Acclimation to drought**

The level of acclimation detected in this study was lower than expected suggesting a limitation to levels of plasticity in the measured traits in response to the experimental drought. The conditions that are thought to favour the selection for phenotypic plasticity are predictable variations in the environment within certain limits. If, in a given environment, a particular abiotic factor fluctuates very little, unpredictably, or to too extreme an extent for plastic responses to incur a significant increase in fitness, then phenotypic plasticity is not selected for in a population (Valladares et al. 2007). Thus, it is possible that the taxa in this study have limited capacity to acclimate to drought because of the historical stability or unpredictability in water availability. Other factors that may contribute to the limited response include the concept of ‘integrated phenotype’ where traits are so tightly interdependent that changes in one aspect of physiology impact, perhaps negatively, on other aspects (Gianoli 2001, Valladares et al. 2007), or the effects of resource limitation inhibiting a plastic response (Van Kleunen and Fischer 2005). Moreover, plants are rarely specialised to cope with more than one kind of abiotic stress (Niinemets and Valladares 2006), so perhaps the same is true of plasticity in certain traits, and the studied taxa may show higher levels of plasticity in response to, e.g., different levels of irradiance, which may be more advantageous to rainforest species.

Although the changes that were expected did not occur, e.g., higher stomatal and vein density, smaller cell size in the spongy and palisade mesophyll and lower cavity volume, other traits that have been found to arise in droughted plants, such as cell wall and cuticle thickness (Schimper 1903, Maximov 1929, Shields 1950), were not measured in this study. So it is possible that such changes did occur but were not detected. However, leaf thickness and LMA (or its inverse measure, specific leaf area) have been found to vary with water availability, and had marginally significant treatment effects (Table 4). Therefore, larger sample sizes might have revealed significant plot effects for these parameters.

**Summary**

Changing climate is likely to exert selection pressure on the next generation of forest trees, but as the climate change is so rapid, the persistence and vigour of the current generation will be dependent to some extent on their ability to resist or acclimate to the new conditions (Nicotra et al. 2010). The taxa in this study responded to the imposed drought via changes in aspects of leaf anatomy that were not known to influence drought resistance, whilst exhibiting none of the expected changes. This might indicate that the experimental drought is not severe enough to influence leaf anatomy in the way expected, or that the traits typically associated with drought resistance are tightly constrained by other aspects of plant physiology. In the latter case, the restricted capacity for acclimation is suggestive of high sensitivity of this forest to climate change. The extent to which leaf anatomy determines the capacity of plants to cope with changes in water availability could have wide-reaching implications in understanding the drought sensitivity of plants; yet it is the subject of little research (Maximov 1929, Morton and Watson 1948, Shields 1950, Cutler et al. 1977) and has not been explored in an ecological context. Given that rainfall regimes are predicted to continue changing (Stocker 2014), and that the tropics in particular are likely to undergo more frequent and severe dry seasons (Christensen et al. 2013, Fu et al. 2013, Reichstein et al. 2013, Boisier et al. 2015), an improved understanding of these subjects could be invaluable to future estimates of forest vulnerability.

**Supplementary data**

Supplementary data for this article are available at Tree Physiology Online.

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