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ORIGINAL ARTICLE



Desiccation of the leaf mesophyll and its implications for CO_2 diffusion and light processing

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Abstract

Leaves balance CO₂ and radiative absorption while maintaining water transport to maximise photosynthesis. Related species with contrasting leaf anatomy can provide insights into inherent and stress-induced links between structure and function for commonly measured leaf traits for important crops. We used two walnut species with contrasting mesophyll anatomy to evaluate these integrated exchange processes under non-stressed and drought conditions using a combination of light microscopy, X-ray microCT, gas exchange, hydraulic conductance, and chlorophyll distribution profiles through leaves. Juglans regia had thicker palisade mesophyll, higher fluorescence in the palisade, and greater low-mesophyll porosity that were associated with greater gas-phase diffusion (gIAS), stomatal and mesophyll (g_m) conductances and carboxylation capacity. More and highly-packed mesophyll cells and bundle sheath extensions (BSEs) in Juglans microcarpa led to higher fluorescence in the spongy and in proximity to the BSEs. Both species exhibited droughtinduced reductions in mesophyll cell volume, yet the associated increases in porosity and g_{IAS} were obscured by declines in biochemical activity that decreased g_m . Inherent differences in leaf anatomy between the species were linked to differences in gas exchange, light absorption and photosynthetic capacity, and drought-induced changes in leaf structure impacted performance via imposing species-specific limitations to light absorption, gas exchange and hydraulics.

KEYWORDS

3D leaf complexity, drought stress, leaf carbon-water exchange, leaf structure and function, light absorption profiles, mesophyll conductance, X-ray microcomputed tomography

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1 | INTRODUCTION

Photosynthesis supports plant growth, development, and reproduction, and to optimise this process, leaves must balance light absorption, carbon capture, and water loss under ever changing conditions. Anatomical and physiological leaf traits play key roles in determining the exchange of light, CO₂, and water with the environment. How the structural and physiological components of the leaf are affected by drought is of increasing importance given the increasing frequency and duration of drought globally (Brodribb et al., 2020; Choat et al., 2018). Stomata regulate the diffusion of gases across the leaf surface, where water vapour is lost in exchange for CO2 after crossing the leaf boundary layer, and respond strongly to changes in vapour pressure and soil moisture in many species to minimise water loss (Cowan & Troughton, 1971; Farguhar & Sharkey, 1982; Mott & Peak, 2013; Oren et al., 1999; Turner et al., 1984). After reaching the substomatal cavity, CO₂ molecules are subject to a series of gas and liquid phase resistances along the diffusion pathway through the intercellular airspace, cell walls, membranes, cytosol, and other cellular components to reach carboxylation sites inside chloroplasts. The inverse of the sum of these resistances is used to calculate mesophyll conductance (gm, see Table 1 for symbol definitions), (Flexas et al., 2008; Flexas et al., 2018; Tosens & Laanisto, 2018), and these resistances to the movement of CO₂ should be sensitive to changes in leaf water status. Theoretical predictions and experimental observations have found that both the physical properties of the mesophyll (e.g., cell wall thickness, palisade and spongy mesophyll cell density, mesophyll surface area exposed to the intercellular airspace -IAS) and the underlying physiology (i.e., chloroplast positioning, aquaporins, and carbonic anhydrase activity) strongly influence CO₂ diffusion within a leaf and its concentration at the sites of carboxylation (Flexas et al., 2012; Momayyezi & Guy, 2017a, 2017b, 2018; Muir et al., 2014; Théroux-Rancourt & Gilbert, 2017; Tholen & Zhu, 2011). The products of photosynthesis are then either consumed locally or exported to the vascular tissue.

Similarly, but in an opposing flow direction, water exits the vascular tissue and travels through the mesophyll, ultimately evaporating into the IAS and lost to the atmosphere via the stomata or across the epidermis. A primary role of the leaf vasculature is therefore to replace the water lost while the stomata are open to sustain the uptake of CO₂ for photosynthesis. During drought, insufficient soil moisture or declines in the hydraulic conductance of the vascular system fail to meet the evaporative demands of leaves, leading to loss of turgor in the mesophyll. Recent work has shown that turgor loss directly influences cell shape and leaf porosity even while the xylem remains functional (Scoffoni et al., 2017), and the resulting structural and physiological changes associated with leaf desiccation can significantly alter leaf hydraulic conductance (Buckley et al., 2017; Scoffoni et al., 2014), highlighting the complex sequence of events that take place inside the leaf during drought.

An additional layer of complexity can be observed in the overall structure of the leaf mesophyll and the embedded vasculature, which should not only be organised to facilitate the movement of both

Variable	Definition	Unit
A _n	Net assimilation rate	$\mu mol~CO_2~m^{-2}~s^{-1}$
A _{max}	Maximum assimilation rate at saturating CO ₂	$\mu mol \ CO_2 \ m^{-2} \ s^{-1}$
BSEs	Bundle sheath extensions	Dimensionless
Ci	Intercellular airspace CO ₂ concentration	µmol mol ^{−1}
C _i *	Intercellular CO ₂ photocompensation point	μ mol mol ⁻¹
C _c	Chloroplast CO ₂ concentration	$\mu mol mol^{-1}$
Е	Transpiration rate	mmol $m^{-2} s^{-1}$
gias	Intercellular airspace (gas phase) conductance	mol m ⁻² s ⁻¹ bar ⁻¹
g liq	Liquid phase conductance	mol m $^{-2}$ s $^{-1}$ bar $^{-1}$
g _m	Mesophyll conductance	mol $\rm CO_2~m^{-2}~s^{-1}$
gs	Stomatal conductance	mol $m^{-2} s^{-1}$
g _{smax}	Maximum stomatal conductance	mol $m^{-2} s^{-1}$
K _{leaflet}	Leaflet hydraulic conductance	$\begin{array}{c} mmol \ m^{-2} \\ s^{-1} \ MPa^{-1} \end{array}$
L _{leaf}	Leaf thickness	μm
L _{mes}	Mesophyll thickness	μm
$L_{epi-adaxial}$	Adaxial epidermis thickness	μm
L _{epi-abaxial}	Abaxial epidermis thickness	μm
PPFD	Photosynthetic photon flux density	$\mu mol\ m^{-2}\ s^{-1}$
SA_{mes}/V_{mes}	Mesophyll surface area exposed to the intercellular airspace per mesophyll volume	$\mu m^2 \ \mu m^{-3}$
V _{IAS} / V _{mes-cell}	Intercellular airspace volume to mesophyll cell volume	$m^3 m^{-3}$
WUE _i	Intrinsic water use efficiency	$\mu mol CO_2 mol^{-1} H_2O$
Ψ_{leaflet}	Leaflet water potential	MPa
Φ_{PSII}	Quantum yield of photosystem II	Dimensionless
Γ*	Chloroplast CO ₂ photocompensation point	μ mol mol ⁻¹
R _d	Dark respiration	$\mu mol\ m^{-2}\ s^{-1}$
θ_{IAS}	Mesophyll porosity	$m^3 m^{-3}$
τ_{leaf}	Tortuosity	$m^2 m^{-2}$
λ_{leaf}	Lateral path lengthening	m m ⁻¹

carbon and water, but also optimised for the opposing gradients of light and CO_2 within the leaf (Borsuk & Brodersen, 2019; Evans, 1999; Evans, 2021; Smith et al., 1997; Xiao et al., 2016). A general assumption is that the absorptive, optical, and hydraulic

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properties of leaves are optimised under well hydrated conditions with the mesophyll cells under full turgor. What then are the effects of turgor loss and the associated changes in cell shape and volume on the processing of light, CO2 and water as leaves dehydrate? As mesophyll cells lose turgor, there should be consequences for the different physiological roles that those cells contribute to, and perhaps differently in the palisade and spongy mesophyll based on cell size and shape. For example, loss of turgor in the mesophyll should lead to changes in the physical shape of the cells (Canny et al., 2012), which has implications for the surface area exposed to the IAS, the tortuosity of the diffusion pathway for both H₂O and CO₂ (i.e., decline in $g_{\rm m}$, Cano et al., 2014) by bringing the epidermis closer to the sites of evaporation within the leaf (Buckley et al., 2017). However, it should also directly affect the optical properties of those cells for light propagation and scattering, leading to sub-optimal light absorption with negative impacts on biochemical activity and light use during photosynthesis. The distribution of mesophyll cells and presence of bundle sheath extensions (BSEs) can influence light distribution with depth into a leaf (Evans & Vogelmann, 2003; Holloway-Phillips, 2019; Smith et al., 1997). Numerous studies have shown significant relationships between optical properties (e.g., absorptance and reflectance) and leaf chlorophyll concentration under water stress (e.g., Carter, 1993; Carter & Knapp, 2001; Gitelson et al., 2003), however, the functional relationship between mesophyll and light absorption with depth into a leaf and under dehydration is not known. Recent studies evaluating mesophyll anatomy at finer scales have shown links between biophysical properties of mesophyll cells and IAS conductance (g_{IAS}) . These linkages are associated with variation in airspace tortuosity (i.e., the ratio of the diffusive path length to the straight path length; τ_{leaf}), porosity (i.e., IAS volume fraction of the mesophyll; θ_{IAS}), and path lengthening as a consequence of CO₂ diffusion through each distinct stomate to IAS (λ_{leaf}) (Earles et al., 2019; Gommes et al., 2009; Harwood et al., 2021; Théroux-Rancourt et al., 2021; Tosens et al., 2016). Little is known about how water stress influences these relationships, as suggested in CO₂ and water flux models considering mesophyll and vascular geometry (Rockwell et al., 2014c, 2017).

Declines in the net assimilation rate (A_n) under water stress are well documented, and arise due to both stomatal and non-stomatal limitations. Loss of turgor in the guard cell complex creates a physical barrier for the diffusion of CO₂ into the leaf, and leads to a depletion of the internal CO₂ supply to carboxylation sites, but also negatively influences photochemistry due to increases in leaf temperature (Brodribb & Holbrook, 2003; Buckley, 2019; Buckley et al., 2017; Galle et al., 2009). Excessive leaf temperatures and desiccation can also lead to permanent damage to photosynthetic machinery (Cano et al., 2013; Chaves et al., 2009; Galmés et al., 2007; Hsiao, 1973; Nadal & Flexas, 2018; Trueba et al., 2019; Urban et al., 2017). A negative response of g_m to dehydration occurs under mild water stress, and this response is exacerbated by high light intensity (Flexas et al., 2008; Galle et al., 2009; Zhou et al., 2007), illustrating, the need to understand the coordination of multiple exchange processes since excess light

can be detrimental to the photosynthetic machinery when rates of carbon fixation decrease with water stress.

The goal of this study is to unfold the complex links between the leaf anatomical traits and functional diversity in CO2, water and light absorption. Here, we explore inherent differences in leaf structure for two walnut species with leaf anatomy contrasting in the fraction of BSEs in relation to functional responses under non-stressed condition and impacts of stress-induced changes in leaf anatomy on species performance, and tested several hypotheses based on our preliminary observations for these species. We used Juglans regia L., native to central Asia, Himalayas, China and southeastern Europe (McGranahan & Leslie, 2009) and J. microcarpa Berland. var. microcarpa, native to southwestern United States and northwestern Mexico, which are adapted to contrasting environments with different water and light availabilities (McGranahan & Leslie, 2009). Our preliminary greenhouse and field measurements indicate differences between species in gas exchange capacity and leaf anatomy, with J. microcarpa showing a higher fraction of BSEs within the leaf. We expected that inherent differences in BSEs and mesophyll cell packing will affect light absorption profiles and CO₂ diffusion in two walnut species. Juglans regia, with elongated and densely stacked palisade mesophyll and more porous lower mesophyll was hypothesised to show higher upper-mesophyll light absorption (Cui et al., 1991), and greater intercellular airspace diffusion. Previous studies reported species with more BSEs have greater structural rigidity and lower turgor loss point and show less shrinkage in leaf and mesophyll cells under dehydration (Pivovaroff et al., 2014, Scoffoni et al., 2017). Therefore, we expected J. regia leaves with less structural and functional support by BSEs (mainly known as parenchyma cells connecting veins to epidermis) to exhibit more volumetric changes through mesophyll cells, porosity, and g_{IAS} under dehydration. In contrast, J. microcarpa with higher cell packing and BSEs was expected to more reflect small changes in cell geometry through light absorption profile, as suggested in species with dense spongy mesophyll through more lower-mesophyll scattering impact (Ren et al., 2019; Smith et al., 2004). To evaluate these hypotheses, we used X-ray micro-computed tomography (microCT) imaging to observe in-depth variation in leaf and cell morphology with dehydration coupled with gas exchange measurements.

2 | MATERIALS AND METHODS

2.1 | Plant materials and growth conditions

Juglans regia cv. Chandler is the most common hybrid scion from natural populations of *J. regia* L., and *J. microcarpa*, is used in *J. microcarpa* × *J. regia* crosses to produce rootstocks with resistance to crown gall and root rot diseases (Browne et al., 2015; Hasey, 2016; McGranahan & Leslie, 2009). Juglans microcarpa is reported to be more tolerant to water deficit (Knipfer et al., 2020).

Two-year-old saplings of clonal and non-grafted *J. regia* and *J. microcarpa* were grown under consistent greenhouse light and

temperature condition, and shipped from the University of California, Davis to the Marsh Botanical Garden greenhouse at Yale University, and were allowed to acclimate under well-watered conditions (without any pre-drought hardening) for 4 weeks before use in the experiments. The gradual dry down procedure was done by reducing water application to 75% of full-irrigation during the first week and then reducing it further to 50% of full-irrigation in the second week of drying. Eight saplings for each species were randomly assigned to either a well-watered control treatment (200 ml water per day) or a water stress treatment with 50% less water than controls (100 ml water per day), equal to daily water loss from pots under each treatment. This watering regime was then maintained until the completion of the experiment. Using the method as described by Knipfer et al. (2020), water loss through transpiration and water evaporation from the soil were quantified during the experiment to calculate the required amount of water under each treatment.

During growth and experimental stages, plants were under supplemental lighting (PPFD = $500 \mu mol m^{-2} s^{-1}$) with a 16-h photoperiod, maximum temperature of 25°C during day and minimum of 18°C during night in the greenhouse, in 2.65-L pots containing a 40% pine bark, 40% sphagnum peat moss and 20% vermiculite. The two irrigation treatments were maintained for approximately 2 weeks before the measurements.

2.2 | Photosynthesis measurements

Net assimilation rate (A_n), stomatal conductance (g_s) and the intercellular airspace CO₂ concentration (C_i) were measured on the 4th or 5th leaflet of the most recent fully expanded leaf using LI-COR 6400 XT and LI-COR 6800 systems fitted with 6400-40 and 6800-01A fluorometers, respectively (see Supporting Information Method for A- C_i and A-*I* curves). All measurements were done under PPFD = 1500 (10% blue vs. 90% red) (µmol m⁻² s⁻¹), chamber temperature at 25°C, ambient chamber CO₂ concentration (C_a) at 400 (µmol mol⁻¹), flow rate at 150 (µmol air s⁻¹), and vapour pressure deficit between 1.5 and 2.0 kPa. All leaflets were dark adapted for 20 min before all other measurements to obtain the maximum quantum yield of photosystem II. The quantum yield of photosystem II (Φ_{PSII}) under actinic light was obtained by application of saturating multiphase flashes (>8000 µmol m⁻² s⁻¹) as per Genty et al. (1989).

2.3 | Stable carbon isotope discrimination method

Pre-evacuated 10 ml gas tight vials (Exetainer, Labco, UK) were used to collect air exiting the LI-COR chamber through a tube connected to the cuvette exhaust, either with $(CO_2P = plant CO_2)$ or without $(CO_2R = reference CO_2)$ leaf material inside the chamber. The air exiting the LI-COR cuvette was collected as described by Théroux-Rancourt and Gilbert (2017) and analysed for stable carbon isotope composition. A three-way valve was added to the LI-COR 6800 chamber through the exhaust tube. A ~2 m sampling tube was connected to the third port, and the valve was opened towards it. After ~5 min, the valve was returned to its primary position along the chamber exhaust tube, and 15 ml air was collected from the tube into a gas-tight glass syringe through a brass luer-lock fitting. A needle was connected to the syringe, the syringe's valve was opened, and 3 ml of air sample was flushed through the needle before purging 12 ml of the air into a vial. Sampling started with CO₂R samples, followed by CO₂P and then alternating CO₂R with one CO₂P sample. After taking the first CO₂R sample, a leaf was placed inside the chamber and light adapted for 20 min before taking the first CO₂P sample. The same protocol was followed for every plant sample, ending with a final CO₂R sample. Gas exchange and chlorophyll fluorescence measurements were recorded during each sampling for CO₂P.

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Vials were transferred to the Stable Isotope Facility, at the University of California Davis within a week for measuring carbon isotope discrimination on ThermoScientific GasBench system II interfaced to a ThermoScientific Delta V Plus isotope ratio mass spectrometer (ThermoScientific). Through a six-port rotary valve (Valco), CO_2 was sampled using a 250 µl loop programmed to switch at the maximum CO₂ concentration in the helium carrier gas. N₂O and other gases were trapped and separated from CO_2 by moving through a PoraPLOT Q column (25 m × 0.32 mm ID, 2.5 ml min⁻¹) set at 50°C at the mass spectrometer. A pure CO₂ standard tank of 400 μ mol mol⁻¹ was used to calculate provisional δ values of samples. The system was referenced against internal laboratory standards which were calibrated against NIST 8545 isotopic standards to correct provisional δ values. Final δ^{13} C values were corrected and expressed relative to the international Vienna PeeDee Belemnite standard.

2.4 | Calculation of g_m from carbon isotope discrimination

Ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco) discriminates against ¹³CO₂ relative to ¹²CO₂ during carboxylation (Guy et al., 1993). The amount of discrimination expressed in vivo depends on the diffusion gradient for CO₂ from the bulk atmosphere. By comparing the observed discrimination (Δ_o) with the predicted discrimination (Δ_i) based only on the diffusion gradient through the stomata (i.e., C_a to C_i), the gradient associated with the remaining portion of the diffusion pathway (i.e., C_i to C_c) can be estimated and used to calculate g_m (Evans et al., 1986). Smaller contributions to total discrimination, associated with respiratory (Δ_e) and photorespiratory carbon flux (Δ_f), must also be accounted for. The effect of g_m on overall isotope discrimination (Δ_{gm}) is then given by:

$$\Delta_{g_{\rm m}} = \Delta_{\rm i} - \Delta_{\rm o} - \Delta_{\rm e} - \Delta_{\rm f} \tag{1}$$

Observed discrimination was calculated according to Evans et al. (1986):

1366

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$$\Delta_{\rm o} = \frac{1000\zeta (\delta^{13}C_{\rm a} - \delta^{13}C_{\rm e})}{1000 + \delta^{13}C_{\rm a} - \zeta (\delta^{13}C_{\rm a} - \delta^{13}C_{\rm e})}$$
(2)

$$\zeta = \frac{C_{\rm e}}{(C_{\rm e} - C_{\rm a})} \tag{3}$$

where, $\delta^{13}C_e$ and $\delta^{13}C_a$ are the isotopic ratios of reference CO₂ and unconsumed CO₂, respectively. ζ is the ratio of the reference CO₂ concentration (C_e) entering the cuvette, as determined by the LI-COR 6800, and the net amount consumed in photosynthesis (i.e., $C_e - C_a$).

Predicted discrimination was calculated from gas exchange data with corrections for ternary effects as per Farquhar and Cernusak (2012):

$$\Delta_{i} = \frac{1}{(1-t)}a' + \frac{1}{(1-t)}((1+t)b - a')\frac{C_{i}}{C_{a}}$$
(4)

where *b* is the fractionation in carboxylation of ribulose bisphosphate catalysed by Rubisco (-29%; Guy et al., 1993). The ternary correction factor, *t*, is:

$$t = \frac{(1+a')E}{2g_{ac}^t}$$
(5)

where *E* is the transpiration rate and g_{ac}^t is the combination of boundary layer and stomatal conductance to CO₂. The combined factor for diffusional fractionation through stomata and the boundary layer, *a'*, is:

$$a' = \frac{a_{\rm b}(C_{\rm a} - C_{\rm s}) + a(C_{\rm s} - C_{\rm i})}{(C_{\rm a} - C_{\rm i})} \tag{6}$$

where *a* and a_b are the fractionations occurring during diffusion across the stomata (4.4‰) and through the boundary layer (2.9‰), respectively, and C_s is the CO₂ concentration at the leaf surface (Evans et al., 1986).

Discriminations associated with respiration (Δ_e) and with photorespiration (Δ_f) were calculated from Equations (9) and (10) (Farquhar & Cernusak, 2012):

$$\Delta_{\rm e} = \frac{1+t}{1-t} \left[\frac{eR_{\rm d}}{(A_n+R_{\rm d})C_{\rm a}} \right] (C_{\rm i} - \Gamma^*) \tag{7}$$

$$\Delta_{\rm f} = \frac{1+t}{1-t} \left[f \frac{\Gamma^*}{C_{\rm a}} \right] \tag{8}$$

where *e* and *f* are the fractionations associated with respiration and photorespiration, respectively. We assumed *f* to be -11.6% (Lanigan et al., 2008) and that there is no significant fractionation associated with dark respiration during the day (Wingate et al., 2007). However, because respired carbon was likely fixed during prior photosynthesis in the greenhouse, we took *e* to equal the difference between $\delta^{13}C_e$ (-32 to -37%) and the isotopic composition for atmospheric CO₂ ($\delta^{13}C_{atm}$) in the greenhouse (assumed to be -8%; Alonso-Cantabrana & von Caemmerer, 2015):

$$e = \delta^{13}C_e - \delta^{13}C_{atm} \tag{9}$$

Discrimination associated with g_m is described by Farquhar and Cernusak (2012):

$$\Delta_{gm} = \frac{1+t}{1-t} \left[b - a_i - \frac{eR_d}{A + R_d} \right] \frac{A_n}{g_m C_a}$$
(10)

where a_i is the fractionation factor associated with hydration and diffusion in water (1.8‰ at 25°C). Substituting Equation (3) into Equation (12) and rearranging, g_m was then calculated as:

$$g_{\rm m} = \frac{1+t}{1-t} \left[b - a_{\rm i} - \frac{eR_{\rm d}}{(A_{\rm n} + R_{\rm d})} \right] \frac{A_{\rm n}}{C_{\rm a}} / (\Delta_{\rm i} - \Delta_{\rm o} - \Delta_{\rm e} - \Delta_{\rm f})$$
(11)

2.5 | Calculation of C_c

Having obtained g_m by the chlorophyll fluorescence method, the CO₂ concentration in the chloroplast (C_c) was estimated:

$$C_{\rm c} = C_{\rm i} - \frac{A_{\rm n}}{g_{\rm m}} \tag{12}$$

 g_m obtained from the stable isotope discrimination method was strongly correlated with that estimated using the chlorophyll fluorescence method (see Supplementary Methods) (g_m values between 0.03 and 0.19; $R^2 = 0.8016$, p < 0.0001; Figure S1). Given the potential uncertainties with g_m estimates obtained from the variable J method, and the increased sensitivity of certain methods for leaves experiencing water stress, we chose to present g_m from data carbon isotopic discrimination technique.

2.6 | A_n - C_i and A_n -I curves

To better understand photosynthetic responses under dehydration, we constructed CO₂ (A_n - C_i) and light response (A_n -PPFD) curves for each species. A_n - C_i curves were constructed for all individuals at 1500 µmol m⁻² s⁻¹ PPFD under the following sample CO₂ concentration: 400, 50, 80, 100, 150, 200, 400, 600, 800, 1000, 1200, 1500 ppm. Leaflets from all individuals were illuminated at adaxial and abaxial surfaces, respectively at 0, 50, 100, 400, 800, 1000, 1500 µmol m⁻² s⁻¹ to measure A_n -PPFD curves at 400 µmol mol⁻¹ sample CO₂ (Figure S2).

2.7 | Leaflet water potential measurements

The two leaflets opposite the one used for gas exchange measurements were used to measure water potentials. The first leaflet was cut at petiolule base and bagged (in a clear bag) for 10 min to allow equilibration within the leaflet. Then, using a razor blade ~1 cm of

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leaflet lamina was cut from either side of the middle vein to fit the short petiolule inside the pressure chamber gasket. Chamber pressure was increased slowly until the balancing pressure was reached. The second leaflet was covered in a dark bag for 20 min before removal to obtain the water potential of the rachis for the remainder of the leaf.

2.8 | Leaflet water potential and leaflet hydraulic conductance

Leaflet water potential (Ψ_{leaflet}) was measured using a pressure chamber (PMS Instrument Company, Model 1505D) immediately after gas exchange measurements between 10 a.m. to 3 p.m. (Williams & Araujo, 2002) (see Supporting Information Method).

Leaflet hydraulic conductance (K_{leaflet}) was calculated using in situ evaporative flux method according to Brodribb and Holbrook (2003) and Simonin et al. (2015):

$$K_{\text{leaflet}} = E/\Delta \Psi_{\text{bagged leaflet-unbagged leaflet}}$$
 (13)

E is the transpiration rate (mmol $m^{-2} s^{-1}$) measured using gas exchange system, and $\Delta\Psi_{\text{bagged leaflet-unbagged leaflet}}$ is the difference between bagged leaflet and unbagged leaflet water potential (MPa). Average unbagged and bagged Ψ leaflet were -0.8 (±0.04) and -0.7 (±0.03) MPa for J. regia and -1.0 (±0.06) and -0.6 (±0.02) and J. microcarpa under well-watered, respectively, and -1.4 (±0.09) and -1.25 (±0.06) MPa for J. regia and -1.7 (±0.06) and -1.4 (±0.1) MPa for J. microcarpa under dehydration, respectively. The Ψ_{leaflet} showed 3%–6% variability between the leaflets and the average Ψ_{leaflet} was 3%–5% more negative than Ψ_{leaf} in each species. The average transpiration rate (E) within and between leaflets on the same leaf were compared for these measurements. for J. regia and J. microcarpa under well-watered (0.8 ± 0.02 and 0.7 ± 0.01 mmol m⁻² s⁻¹), and drought conditions (0.6 ± 0.03 and 0.4 ± 0.03 mmol m⁻² s⁻¹), varied by 5%-10% between leaflets. Minimal or no significant difference in E existed across individual leaflets for scaling to the total leaflet area.

To quantify the stomatal aperture under well-watered and dehydrated conditions in each species, both hypostomatous, abaxial epidermis imprints using transparent nail polish (water-based) were obtained from the same leaflets used for the gas exchange measurements. Using the imprints, stomata images were taken on a light microscope at 20x (Nikon C2⁺, Nikon Instruments Inc.) and used to measure the stomatal pore dimensions. The inner pore width (μ m) was divided by the inner pore length (μ m) to calculate the stomatal aperture ratio (Rui & Anderson, 2016). Maximum g_s (g_{smax}) was calculated using the stomata pore dimensions (Franks & Beerling, 2009) and used to interpret changes in stomata opening and g_s in the two species and further, test the precision of the stomatal aperture quantification method. To calculate stomata size, guard cells length was multiplied by total width, for closed guard cells (Franks & Beerling, 2009).

2.9 | X-ray micro computed tomography imaging and segmentation

Intact plants with their soil were sent back to the UC Davis greenhouse and potted again where water potentials and soil water content were monitored and maintained for several days until scanning them 7 days after shipping in Lawrence Berkeley National Laboratory (LBNL) Advanced Light Source (ALS). The same leaflet samples used for gas exchange at Yale were kept intact, collected, bagged and placed in a cooler at room temperature an hour before scanning in ALS. A section of the leaflet lamina from each plant was enclosed between two pieces of Kapton tape to prevent desiccation of the tissue and sample movement during the scanning. Samples were placed inside the end of a pipette tip and scanned under a continuous tomography mode at 23 keV using $10\times$ objective lens (pixel resolution of $0.65 \,\mu$ m). Raw tomographic data were reconstructed using TomoPy (Gürsoy et al., 2014) through both gridrec and phase retrieval reconstruction methods (Figure S3) (Davis et al., 1995; Dowd et al., 1999).

Five hundred consecutive slices from the grid and phase stacks were selected for segmentation. The resulting image stack was segmented using the methods presented in Théroux-Rancourt et al. (2020) (Figure S3). Six slices were labelled manually per scan and were used to train a random-forest model for automated segmentation of the whole scan image stack. The final segmented stacks had individual labels for the adaxial epidermis, abaxial epidermis, mesophyll cells, intercellular airspace, BSEs, veins, and background outside of the scanned leaf. This final stack was used to extract leaf anatomical traits, that is surface areas, volumes, and lengths.

2.10 | Mesophyll surface area and porosity

As described by Théroux-Rancourt et al. (2017), mesophyll porosity, θ_{IAS} (m³ m⁻³) was calculated as the IAS volume as a fraction of the total mesophyll volume. The IAS volume (V_{IAS}) to mesophyll cell volume ($V_{mes-cell}$) ratio and the mesophyll surface area exposed to the IAS (SA_{mes}) per mesophyll volume (V_{mes}) were calculated as V_{IAS}/V_{mes $cell}$ (m³ m⁻³) and SA_{mes}/ V_{mes} (μ m² μ m⁻³), respectively (Figure 1).

2.11 | Tortuosity and lateral path lengthening

The tortuosity factor, $\tau(m^2 m^{-2})$, was defined as the ratio of the diffusive path length within the IAS to the straightest path length in the absence of any physical obstacles to diffusion between a stomate and the cell surface:

$$\tau = \left(\frac{L_{\text{geo}}}{L_{\text{Euc}}}\right)^2 \tag{14}$$

where geodesic distance (L_{geo}) is the distance from the stoma to a cell surface, and Euclidean distance, (L_{Euc}) (Earles et al., 2018). The L_{geo} and L_{Euc} were mapped and quantified for all voxels along the mesophyll

WILEY <u><u>P</u> <u>Environment</u></u>

surface and τ was calculated for the whole 3D image array as in Earles et al. (2018). Then, leaf-level tortuosity (τ_{leaf}) was calculated as the mean of τ values at the edge of mesophyll cells. The lateral path lengthening, λ (m m⁻¹) was calculated using L_{Euc} , and a second distance map as described by Earles et al. (2018) to measure the shortest unobstructed distance in a straight line between the abaxial epidermis and all points along the mesophyll surface, L_{epi} (Legland et al., 2016):

$$\lambda = \frac{L_{\text{Euc}}}{L_{\text{epi}}} \tag{15}$$

Similarly, leaf-level lateral path lengthening (λ_{leaf}), was then calculated as the mean of λ values at the edge of mesophyll cells.

2.12 | IAS conductance

The τ_{leaf} , λ_{leaf} and θ_{lAS} were used to calculate leaf-level IAS conductance (g_{IAS}), where D_m is the diffusivity of CO₂ in air (m² s⁻¹).



FIGURE 1 Leaf three-dimensional projection for *Juglans regia* and *Juglans microcarpa* under well-watered and dehydrated conditions

Diffusion path length in gas phase was equal to half of the mesophyll thickness (L_{mes}) for hypostomatous leaves (Earles et al., 2018; Niinemets & Reichstein, 2003; Tomás et al., 2013):

$$g_{\text{IAS}} = \frac{\theta_{\text{IAS}} D_m}{0.5 L_{\text{mes}} \tau_{\text{leaf}} \lambda_{\text{leaf}}}$$
(16)

2.13 | Porosity profiles

MicroCT scans for each species under well-watered and dehydrated conditions (Figure 2) were used to determine porosity profiles from IAS distribution with leaf depth using a plot profile of grey value distribution across leaf excluding adaxial and abaxial epidermis tissue. The grey values were used to calculate air volume for a known mesophyll area (4 μ m²) and based on mesophyll thickness (μ m) per individual within each depth after converting pixel to distance (pixel resolution of 0.65 μ m).

2.14 | Palisade mesophyll cell diameter at paradermal section

The grid reconstructions of microCT images were used to compare the palisade mesophyll cell diameter through paradermal sections at three depths (20%, 40% and 60%) from the adaxial surface in wellwatered versus dehydrated leaves within 0.02 mm² of the mesophyll area.

2.15 | Relative chlorophyll distribution through the leaf profile

We used previously reported methods to obtain chlorophyll distribution (Borsuk & Brodersen, 2019; Vogelmann & Evans, 2002) and light absorption profiles (Brodersen & Vogelmann, 2010; Koizumi et al., 1998; Takahashi et al., 1994; Vogelmann & Evans, 2002; Vogelmann & Han, 2000) for each species (Figure S4). Chlorophyll distributions were obtained by calculating the relative





chlorophyll fluorescence (proportional to chlorophyll concentration) at each relative depth. Light absorption gradients, representing relative chlorophyll distribution patterns were measured using chlorophyll fluorescence imaging of leaf cross sections under direct illumination (Vogelmann & Evans, 2002; Vogelmann & Han, 2000). Fresh samples in a subset of three were cut into $\sim 1 \text{ cm}^2$ from the same leaflets and placed on top of a wet paper to protect the specimen from desiccation in a glass holder on the microscope stage (Olympus BX60, Olympus America Inc., Center Valley, PA, USA). The sample was irradiated by a broad-spectrum LED light source at cross-sectional direction (epi-illumination at 490 nm; beam radius ~1 mm) (Figure S5). For adaxial or abaxial profile imaging, leaves were irradiated with direct light in sequence with monochromatic red (660 nm), green (532 nm), or blue (488 nm) light obtained from three lasers one at a time (laser spot radius = 1 mm; red solid state laser: Model #BWN-660-10E, BandW Tek Inc.; green solid state laser: Model # DY20B, Power Technology Inc.; and blue argon gas laser: Model # Innova 300, Coherent Inc.). Using a digital Peltiercooled CCD camera (PIXIS 1024B, Princeton Instruments, Trenton, NJ, USA) with shutter times of 70-150 ms, emitted light of chlorophyll fluorescence was imaged after passing through a barrier filter (680 nm, half band width = 16 nm, S10-680F; Corion Filters). Light intensity through the leaf was measured in Image J (Rueden et al., 2017) from the adaxial edge of the mesophyll to the abaxial edge of the mesophyll using the line profile tool averaged over a width of 50–100 pixels (100-pixel width was equivalent to \sim 60 μ m at 20x magnification or ~120 µm at 10x magnification) and excluded conspicuous non-photosynthetic structures such as epidermal cells and veins. The obtained values per each profile were normalised by dividing them by the chlorophyll fluorescence depth maxima. An absolute fluorescence intensity could not be estimated, first, due to lack of flexibility in accounting variation in light exposure needed for different samples sizes with different focal points, and second, the overall decline in fluorescence intensity under a continuous supply of light over time, that is, temporal variation in detected fluorescence signal due to Kautsky decay (Borsuk & Brodersen, 2019;

2.16 Bundle sheath extensions area

Vogelmann & Han, 2000).

MicroCT scans for each species were used to calculate the ratio of the BSEs area where parenchyma cells connecting vascular tissue to both epidermises, as a percentage from the mesophyll cross sectional area (i.e., area of mesophyll cells and airspace) using Image J (Griffiths et al., 2013).

%BSEs area = BSEs area/(BSEs area + mesophyll area) (17)

Green light penetrates deeper into leaf and is absorbed more equally throughout the leaf profile (Brodersen & Vogelmann, 2010). Therefore, adaxial fluorescence images from green laser illumination, as described above, were used to determine the relative fluorescence for mesophyll tissue adjacent to BSEs and plot their distribution along the mesophyll in Image J. Fluorescence percentage near BSEs were normalised using maximum values per sample and plotted using epi- and adaxial illumination imaging data for the two species.

2.17 | Statistics

Linear regression lines were used to describe relationships between A_n and C_i and paired t-tests were used to compare differences in estimated g_m from the isotope discrimination and chlorophyll fluorescence methods using GraphPad prism 8 (GraphPad Software, Inc). Mixed linear models were used to compare treatments effects on the following physiological variables: A_n, g_m, g_s, C_i, C_c, Φ_{PSII}, Ψ_{leaflet}, τ_{leaf}, θ_{IAS} , L_{leaf} , L_{mes} , $L_{epi-adaxial}$, $L_{epi-abaxial}$, SA_{mes}/V_{mes} , $V_{IAS}/V_{mes-cell}$, λ_{leaf} and g_{IAS} in the two species under well-watered and dehydrated treatments using SAS 9.4 (SAS Institute Inc). Adjusted p-value (=0.0083) was calculated by dividing α (=0.05) by number of mean pairs per test (n = 6). Mixed linear models were used to compare absolute and percentage reductions for all the physiological variables relative to the well-watered (p = 0.05). Number of vein emboli (see results for method description), BSEs area, and palisade diameter were compared using mixed linear models (p = 0.05). Logarithm or squared transformations were performed to meet normality and equal variance assumptions where needed. Multiple *t*-tests were used for a pairwise comparison between all pairs of means (p = 0.05).

3 | RESULTS

3.1 | Mesophyll traits and IAS parameters

Total leaf thickness (L_{leaf} ; p = 0.0183), mesophyll thickness (L_{mes} ; p = 0.0203), $\theta_{\rm IAS}$ (p < 0.0001), $V_{\rm IAS}/V_{\rm mes-cell},~g_{\rm IAS}$ (p < 0.0001) and $\lambda_{\rm leaf}$ (p = 0.0023) were greater in J. regia compared to J. microcarpa under well-watered conditions (Figure 3). Lower θ_{IAS} in J. microcarpa aligned with significantly greater SA_{mes}/V_{mes} (p < 0.0001, Figure 3). Tortuosity (τ_{leaf}) (Figure 3) and adaxial and abaxial epidermis thicknesses (data not shown) were not statistically different between the species. Water stress reduced L_{leaf} (by 8% vs. 9%) and L_{mes} (by 10% vs. 13%) similarly in J. regia and J. microcarpa, respectively. Although the abaxial epidermis showed some shrinkage under dehydration, the abaxial and adaxial epidermis thickness (Lepi-abaxial, Lepi-adaxial) were not significantly reduced in either species. Dehydration increased VIAS/Vmes-cell in both species by 20% (p < 0.0001) through reducing both V_{IAS} and V_{mes-cell} but in different rates in each species (p < 0.0001, Figure 3). The reductions were in line with an increase in porosity (θ_{IAS}) in both species under dehydration, but this effect was greater for J. regia than J. microcarpa (p = 0.0065) and significantly higher SA_{mes}/V_{mes} under drought, in J. regia only (p = 0.010) (Figure 3). g_{IAS} increased equally for J. regia (by 23%) and J. microcarpa (by 21%) (p > 0.05) under

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FIGURE 3 Intercellular airspace parameters under well-watered (solid bar) and dehydrated (empty bar) treatments for Juglans regia and Juglans microcarpa. a, leaf tortuosity factor (τ_{leaf} , $m^2 m^{-2}$); b, intercellular airspace porosity (θ_{IAS} , $m^3 m^{-3}$); c, intercellular airspace to mesophyll cell volume ratio ($V_{IAS}/V_{mes-cell}$); d, leaf thickness (L_{leaf} , μ m); e, mesophyll thickness (L_{mes} , μ m); f, mesophyll surface area exposed to the intercellular airspace per mesophyll volume basis $(SA_{mes}/V_{mes}, \mu m^2 \mu m^{-3})$; g, lateral path lengthening within intercellular airspace (λ_{leaf} , m m⁻¹); h, intercellular airspace conductance (g_{IAS} , mol m⁻² s⁻¹ bar⁻¹). Data points are means of four biological replicates (ramets) per species under well-watered or dehydrated conditions (±SE)

dehydration compared to the well-watered condition. Dehydration reduced g_{IAS} contribution to g_m (calculated as described by Niinemets & Reichstein, 2003) from 22% to 9% in *J. regia*, and 23% to 8% in *J. microcarpa* (p < 0.05). Although there was a significant increase in τ_{Ieaf} (by 23%) in *J. microcarpa* under dehydration (p = 0.010) (Figure 3), path lengthening (λ_{Ieaf}) did not change in either species.

3.2 | CO₂ and light response curves

Despite species-dependent differences in photosynthetic capacity and greater A_n at ambient CO₂ (400 µmol mol⁻¹) and higher maximum carboxylation rate (V_{cmax}) and maximum electron transport rate (J_{max}) in *J. regia* as expected (Figures 4 and 5), maximum photosynthesis (A_{max} at

*C*_i greater than 750 μmol mol⁻¹) was statistically similar for the two species under well-watered conditions (Figure 4). *Juglans microcarpa* maintained its photosynthetic capacity (i.e., greater J_{max}) under dehydration to a greater extent compared to *J. regia* (17% vs. 52% decrease in A_{max} , respectively; *p* < 0.0001). Dehydration reduced A_n significantly in both species (*p* = 0.0003) with a greater percent decrease in *J. regia* (by 47%) compared to a 42% for *J. microcarpa* (*p* = 0.0023) under ambient CO₂ and saturating light (1500 µmol m⁻² s⁻¹) (Figure 4). At lower PPFD (50 to 500 µmol m⁻² s⁻¹) from adaxial illumination, the percent and absolute reductions in A_n were similar between the species (Figure S2). In general, A_n was lower with abaxial illumination, however, absolute and percent reductions in A_n were similar to those from adaxial illumination in the two species (*p* = 0.0014; Figure S2).

3.3 | Mesophyll conductance and photosynthesis at ambient CO₂

 A_n and photosystem efficiency (Φ_{PSII}) at ambient CO₂ were greater in *J. regia* than *J. microcarpa* under control conditions (Figures 5a and 4b, p < 0.0083), in agreement with higher g_s (p = 0.0080; Figure 5c), g_m (p < 0.0001, stable isotope method, Figure 5d), C_i (p = 0.0001; Figure 5e), and C_c (p < 0.0001, Figure 5f). Reduced A_n under dehydration aligned with reductions in g_s and g_m (Figure 5). On the other hand, Φ_{PSII} and C_i decreased significantly in *J. regia* only and C_c showed greater reductions with dehydration in *J. regia* compared to *J. microcarpa* (Figure 5, p < 0.0001). The stomatal aperture ratio (inner pore width/inner pore length) was greater for *J. regia* under well-watered condition (*J. regia* 0.47 ± 0.04, J. microcarpa 0.37 \pm 0.03), and dehydration induced stomatal closure and increased the ratio by 38% in J. regia versus 61% in to J. microcarpa. The relative changes in the pore dimensions were proportional to g_s reduction in J. microcarpa (by 68%), but less than reduction in J. regia (by 58%).

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Under well-watered conditions, Ψ_{leaflet} (Figure 5h) and K_{leaflet} (Figure 5g) were similar for the two species, however, they were correlated negatively in both species ($R^2 = 0.9985$, p = 0.0008). g_s responded negatively to decreasing Ψ_{leaflet} ($R^2 = 0.9091$, p = 0.0465) with a greater reduction in J. microcarpa compared to J. regia (p < 0.0001) (Figure 6), however, the reductions in K_{leaflet} induced by water stress were not linked with significant changes in the number of embolized conduits for either species (Figure 6). The percent ratio of embolized conduits (C_{emb}) per number of conduits (C) (Scoffoni et al., 2017) in representative microCT images (800 µm of each cross; n = 6) in secondary veins was similar between under well-watered and dehydrated conditions in J. regia (11.3% vs. 12.4%) and J. microcarpa (20.2% vs. 14.2%) (p > 0.05). Similarly, no significant effect on emboli formation was found in tertiary veins under well-watered versus dehydrated conditions in J. regia (7.4% vs. 10.9%) and J. microcarpa, (3.6% vs. 4.6%) (p > 0.05).

3.4 | Chlorophyll distribution, light absorption and porosity profiles

Relative chlorophyll distribution was estimated from fluorescence profiles in leaf cross sections using epi-illumination (Figures 7 and S5). The patterns showed species-specific differences; *J. regia* exhibited a

FIGURE 4 Photosynthetic CO_2 response curves, relationship between mean A_n (net assimilation rate) and C_i (intercellular airspace CO_2 concentration) at 1500 µmol m⁻² s⁻¹ photosynthetic photon flux density were constructed using FvCB model (Sharkey, 2016), averaged over four replications in *Juglans regia* (panels a and c) and *Juglans microcarpa* (panels b and d) under well-watered (solid symbols and top row) and dehydrated (empty symbols and bottom row) treatments (±SE; n = 4). Assimilation rate at saturating CO_2 (A_{max}), Rubisco and RuBP regeneration limitations are indicated for each species [Color figure can be viewed at wileyonlinelibrary.com]





MOMAYYEZI ET AL.

FIGURE 5 Photosynthetic traits under wellwatered (solid bar) and dehydrated (empty bar) treatments for Juglans regia (J. reg) and Juglans microcarpa (J. mic). a, net assimilation rate (An, μ mol CO₂ m⁻² s⁻¹); b, photosystem II efficiency (Φ_{PSII}) ; c, stomatal conductance (g_s, mol m⁻² s⁻¹); d, mesophyll conductance (g_m , mol CO₂ m⁻² s⁻¹); e, intercellular airspace CO₂ (C_i, μmol mol⁻¹); f, chloroplast CO₂ (C_c , µmol mol⁻¹); g, leaflet hydraulic conductance ($K_{leaflet}$, mmol m⁻² s⁻¹ MPa⁻¹); h, leaflet water potential (Ψ_{leaf} , MPa). Data points are means of four biological replicates (ramets) per species under wellwatered or dehydrated conditions (±SE). The measurements were taken at 400 µmol mol⁻¹ and 1500 μ mol m⁻² s⁻¹ photosynthetic photon flux density

single peak in relative fluorescence around palisade mesophyll, within 0%–20% depth from the adaxial epidermis (Figure 8a), whereas *J. microcarpa* had double peaks at 10%–40% and 80%–100% depth (Figure 8c). A rapid attenuation after 20% and leveling off at 60% of the depth in *J. regia* was different than the pattern for *J. microcarpa*, where there was a depression between 40% and 80% of leaf depth. As expected, porosity increased with depth from adaxial surface in two species, but the porosity profile complemented the fluorescence

profile better in *J. regia* with an increase in porosity after 40% of depth, around spongy mesophyll, and a maximum between 90% adaxial depth (Figure 8b). In *J. microcarpa*, porosity changed less compared to relative fluorescence suggesting that components other than cell packing are involved in fluorescence gradients across the leaf. However, the porosity increased smoothly after 20% depth and reached the maximum between 80% and 100% depth from adaxial surface (Figure 8d).



FIGURE 6 Leaflet water potential (Ψ_{leaf} , MPa) and leaflet hydraulic conductance (K_{leaflet} , mmol m⁻² s⁻¹ MPa⁻¹) (a) and stomatal conductance (g_s , mol m⁻² s⁻¹) (b) relationship under well-watered (solid) and dehydrated (empty) treatments in *Juglans regia* (circle) and *Juglans microcarpa* (square). MicroCT images on the left are representative slices comparing conduit embolism from scans of *J. regia* and *J. microcarpa* leaves under well-watered and dehydrated conditions. Six slices from each microCT scan were used to count number of emboli per secondary and tertiary veins under well-watered and dehydrated conditions. Number of emboli divided by number of veins within the red circles region did not show a significant change in either secondary or tertiary veins under dehydration from well-watered condition. Bar equals 250 µm [Color figure can be viewed at wileyonlinelibrary.com]

FIGURE 7 Top images compare microscopic cross sections for *Juglans regia* and *Juglans microcarpa* under well-watered conditions. Bottom images compare spatial chlorophyll distribution from original epi-illumination imaging and false-coloured associated pairs, representative of the two species leaves under well-watered and dehydrated conditions [Color figure can be viewed at wileyonlinelibrary.com]



Relative fluorescence +

Under dehydration, *J. microcarpa* showed an increase in relative fluorescence within the first 20% of the leaf depth and a consistent reduction between 30% and 100% depths (Figure 8c). In contrast, *J. regia* did not show a significant difference between watering conditions. Porosity increased across the leaf profile under dehydration in both species (p < 0.002) with a greater increase after 60% of adaxial depth in both species (Figure 8). Between 0% and 20% depth from the adaxial epidermis and under adaxial illumination, absorption of red light was greater in *J. microcarpa* under dehydration, but it decreased significantly after

30% depth compared to the well-watered condition in all wavelengths (Figure 9). Only within 20% depth from the adaxial surface, *J. regia* showed higher absorption under dehydration at the blue wavelength (Figure 9). Illumination direction had a significant impact on absorption depth in red and blue wavelengths; maximum absorption at adaxial irradiance occurred at first 30% of depth from adaxial surface, whereas it was at first 60% of depth from abaxial epidermis under abaxial illumination. There was no consistent difference in light absorption profiles between wellwatered and dehydrated conditions under abaxial illumination in

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FIGURE 8 Chlorophyll fluorescence profiles obtained from epi-illumination (490 nm) and porosity profiles acquired from microCT scans in *Juglans regia* (circles) and *Juglans microcarpa* (squares) under well-watered (solid) and dehydrated (empty) treatments (mean ± SE; *n* = 4–6)



FIGURE 9 Light absorption profiles in *Juglans regia* and *Juglans microcarpa* leaves irradiated with direct monochromatic light at blue (488 nm), green (532 nm) and red (650 nm) wavelength under well-watered (solid lines) and dehydrated (dashed lines) treatments averaged over four replications (n = 4). Directional lights were illuminated from adaxial (a and b) and abaxial (c and d) surfaces. Relative fluorescence (%) is presented relative to the illumination direction, adaxial (a and b) or abaxial (c and d) [Color figure can be viewed at wileyonlinelibrary.com]

J. regia, while the green wavelength showed a significantly higher absorption between 60% and 100% of depth in *J. microcarpa*.

3.5 | Paradermal cells and bundle sheath extensions

The paradermal section images at 20%, 40% and 60% depths from the adaxial epidermal surface showed a significant decrease in the palisade cell diameter (μ m) under dehydration in both species; diameters decreased by 14%, 9%, 15% in *J. microcarpa* and by 14%, 17%, 19% in J. regia (p < 0.05; see example images in Figure 10) at each increasing depth, respectively. BSEs were more prominent in J. microcarpa occupying 15% of the mesophyll volume compared to 8% in J. regia (p = 0.001). That was related to higher vein density with narrow BSEs width (Figure 11) for J. microcarpa. Using epiillumination data, J. microcarpa had higher fluorescence near the BSEs cells under well-watered than dehydrated condition, compared to J. regia. (Figure 11, p < 0.0001). Under adaxial green wavelength illumination, differences in fluorescence near BSEs between the two species were not significant (normalised data shown in Figure S6).



FIGURE 10 Paradermal sections at 20% depth from the adaxial surface in Juglans regia and Juglans microcarpa under well-watered and dehydrated treatments. Similar responses were seen at 40% and 60% depths (images not shown). Bar equals 50 µm



Plant. Cell 8



DISCUSSION 4

4.1 Inherent differences between two Juglans species linked with structure and function

The exchange of water and CO₂ and light absorption gradients are connected via mesophyll geometry and stress-related changes in leaf anatomical characteristics induce responses in gs, gm, and An. Using microCT imaging, we showed that θ_{IAS} and g_{IAS} in J. regia and J. microcarpa corresponded with species-specific differences in A_{n} , g_{s} and g_m obtained from gas exchange, chlorophyll fluorescence, and stable carbon isotope methods (Figures 3, 5 and S1). Well-watered J. regia, with thicker leaves and denser mesophyll cells in the upper palisade, had higher θ_{IAS} , $V_{IAS}/V_{mes-cell}$ and g_{IAS} aligned with greater A_n , g_s and g_m and higher chlorophyll concentration near the adaxial surface. Juglans regia mesophyll structure with higher θ_{IAS} and greater IAS distribution between mesophyll cells increase the g_{IAS} thorough an effective lateral diffusion (Figure 3). J. regia with less BSEs, forms less physical barrier to gas diffusion, thus the resistance to diffusion from gas to liquid phase decreases due to a greater lateral conductivity in this mesophyll type (e.g. homobaric) (Evans & von Caemmerer, 1996; Pieruschka et al., 2005). This inner mesophyll structure couples with a greater need for g_s to keep up with the higher demand for C_i concentration and higher g_{smax} (Figure 5). According to lateral CO₂ diffusion modelling by Pieruschka et al. (2007), larger interconnected airspace can improve CO₂ diffusivity through IAS, and enhance C_i in coordination with g_s as seen in J. regia (Figures 3 and 5). Juglans regia has larger stomata (average size, 122 vs. 72 µm²) but fewer stomata (62 per mm²) than J. microcarpa (79 per mm²), and higher λ_{leaf} in *J. regia* is linked with its lower stomatal

FIGURE 11 Fluorescence percentage normalised using maximum values per sample using epi-illumination. It indicates the spatial distribution of fluorescence in equal horizontal distance from bundle sheath extensions (BSEs) in Juglans regia (circles-left data panel) and Juglans microcarpa (squares-right data panel) under well-watered (solid) and dehydrated (empty) treatments (\pm SE; n = 4). Positive and negative values present variation in grayscale values at right or left side of BSEs. BSEs location is designated with white arrows; mean J. regia BSE width was ~4.7-5.0 µm and mean J. microcarpa BSE width was \sim 3.0–3.5 µm. Zero on the x-axis of the data plots represents the centre of the measured BSE

density, which would increase CO_2 diffusion length and higher θ_{IAS} near spongy mesophyll in hypostomatous species (similar to patterns reported by Earles et al., 2018 and Harwood et al., 2021). Therefore, J. regia benefits from a higher CO₂ diffusion capacity, and exhibits improved performance through increasing maximum carboxylation rate (V_{cmax}) and A_n under lower CO₂ concentrations (Figure 4). At ambient CO2, where RUBP- regeneration is limiting, J. regia mesophyll with lower diffusion resistance (e.g., more porous leaves) and higher enzymatic activity during CO₂ fixation and carbohydrate formation (i.e., Calvin cycle) shows greater maximum electron transport rate (J_{max}) , g_m , and A_n under well-watered condition.

Smaller mesophyll cell size potentially improves CO₂ accessibility to Rubisco by enhancing SAmes/Vmes and chloroplast surface area (Ren et al., 2019; Terashima et al., 2006; Tholen et al., 2012), and consequently increases conductance within the liquid phase (g_{lig}) , while greater θ_{IAS} is associated with higher g_{IAS} (Théroux-Rancourt et al., 2021). g_{IAS} contributed 29% and 25% to g_m in J. regia and J. microcarpa, respectively, consistent with previous findings in tree

VILEY - PC Plant, Cell 8 Environment

species; Populus tremula (23%-25%; Tosens et al., 2012), Quercus ilex L. (23%; Niinemets & Reichstein, 2003), and four Eucalyptus species (8%-21%; Harwood et al., 2021) under well-watered condition and within the estimated limitation range by g_{IAS} (3%-37%) for A_n in hypostomatous species (Parkhurst & Mott, 1990). Higher SAmes/Vmes in well-watered J. microcarpa was associated with lower θ_{IAS} and V_{IAS}/V_{mes-cell}, and less variations in airspace distribution and relative fluorescence across leaf profiles (Figures 3 and 8). Juglans microcarpa mesophyll geometry increases resistances for CO2 diffusion in the gas phase (lower g_{IAS}) through disconnecting mesophyll tissues and dividing into compartments and to the liquid phase, through potentially higher cell wall thickness (Pieruschka et al., 2008; Ren et al., 2019; Tomás et al., 2013), in line with less diffusion and carboxylation capacity, exhibited as g_m , V_{cmax} and J_{max} responses for J. microcarpa (Figures 4 and 5). However, more biomass allocations toward cell packing and extensive bundle sheaths extensions, in species like J. microcarpa, improves the structural tolerance under the environmental stresses (e.g., low water) (Hikosaka & Shigeno, 2009; Niinemets et al., 2007).

Mesophyll cell packing and porosity distribution also led to different optical properties for the two Juglans species. Juglans regia leaves maintained greater θ_{IAS} under all conditions and mesophyll cells were more densely packed in the upper palisade. This coincided with maximum relative fluorescence at 10% depth and a decrease in fluorescence with increasing depth where large airspaces occurred in the spongy mesophyll layer (Figure 8), confirming our hypothesis for light absorption in J. regia. Well-watered J. regia mesophyll had higher A_n and electron transport rate under variable light conditions (Figure S2), in agreement with our expectations for higher light absorption for a mesophyll with prominent palisade layers and high IAS volume in lower mesophyll (Cui et al., 1991; Gotoh et al., 2018), also reported for intermediate shade-tolerant species (Hanba et al., 2002). This mesophyll structure appears to be an adaptation to maximize light absorption under varying light conditions (Leegood, 2008; Terashima, 1992; Tholen et al., 2012); as described by Vogelmann et al. (1996), larger IAS acts as "hall of mirrors" and improves absorption by multiple reflections between airspace, mesophyll and epidermal cells. J. microcarpa, on the other hand, had a narrower range of airspace distribution across leaf profiles with a smaller range of relative fluorescence that was distributed more evenly throughout the mesophyll, in line with mesophyll architecture for species from a high light environment (Hanba et al., 2002). The discrepancy between porosity and relative fluorescence profiles in J. microcarpa (Figure 8) could be attributed to light scattering inside the leaf due to other cell types (e.g., more BSEs) (Vogelmann & Evans, 2002), exhibited as significant difference in relative fluorescence near BSEs in two species under epi-illumination (Figure 11). Mesophyll partitioning due to BSE presence (i.e., heterobaric leaves) is predicted to increase light penetration and overall absorption in various directions, mostly through spongy cells (Vogelmann & Martin, 1993; Xiao et al., 2016). More BSE volume in J. microcarpa compared to J. regia (15% vs. 8%, respectively) and greater fluorescence from the cells near the BSEs (Figure 11) under both well-watered and dehydrated conditions

indicate that *J. microcarpa* can utilise light deeper into the leaf (absolute data not shown). Species with BSEs that contain transparent cells with few or no chloroplasts are proposed to acclimate more effectively to drought since the light transmitted through BSEs is elevated at red and blue wavelengths and it may modify available internal light for photosynthetic tissues (Karabourniotis et al., 2000). BSE-containing species like *J. microcarpa* may rely more on the structural support by BSEs than turgor associated with water supply to sustain their leaf stiffness (Read & Stokes, 2006) as an ecological response to water shortage in their growth habitat even if it costs a reduction in number of photosynthesising cells and eventually the carbon fixation.

4.2 | Drought-induced changes in photosynthetic capacity related to structure and function changes

Dehydration had a negative impact on both species by reducing A_n , g_s and gm. Juglans regia was shown to be more susceptible to stress with a decline in Φ_{PSII} and a greater imbalance in energy distribution between PSII and PSI by more reduced distribution to PSII, also suggesting photodamage-related decreases in light use efficiency under dehydration (Figure S2). Larger IAS volume increases evaporation surface for mesophyll cells, resulting in irreversible mesophyll cell shrinkage and potential permanent damage to the photosystems (Buckley et al., 2017; Rockwell et al., 2014c; Sack & Frole, 2006), as reflected by lower Amax for J. regia under saturating CO2 (Figure 4). A further reduction in C_i at low g_s , concurrent with a decline in Φ_{PSII} suggests an increase in photorespiration (Medrano et al., 2002) as seen in J. regia. According to a sequential baseline presented by Trueba et al. (2019), g_s could decrease by 50% before TLP, whereas passing the TLP under severe stress can lead to permanent damage to the chlorophyll fluorescence as occurred in J. regia. This species exhibited a limited range of resilience to drought, as the widely cultivated walnut species in commercial nut production; this is consistent with irrigation management practices aimed at avoiding water stress in this species. In contrast, J. microcarpa with a more conservative water use strategy and a higher intrinsic water use efficiency (WUE_i, 90 μ mol CO₂ mol⁻¹ H₂O vs. 76 μ mol CO₂ mol⁻¹ H₂O for J. regia) under the well-watered condition, functions at lower water potentials under drought. The inherently lower g_s under well-watered conditions and a greater reduction in Ψ_{leaflet} under dehydration for J. microcarpa suggests that this species may maintain low g_s at lower Ψ_{leaflet} closer to its turgor loss point (TLP). Species with prominent heterobaric leaves are also expected to show more nonuniform stomatal closure in response to environment stress (Kamakura et al., 2011). Juglans microcarpa is reported to be less vulnerable than J. regia to the xylem cavitation; it shows 12% decrease in stem xylem hydraulic conductivity at lower Ψ_{xylem} (e.g., ~ -1.6 vs. -1.3 MPa in J. regia) (Jinagool et al., 2018). Tyree et al. (1993) also measured about 50% reduction in petiole hydraulic conductivity for J. regia at Ψ_{xvlem} ~ -1.4 MPa. Although TLP was not measured in our study, a greater increase in porosity and decrease in palisade cell diameter at less

negative water potential suggest that *J. regia* functioned closer to its potential TLP under dehydration. These are commensurate with previous studies reporting reductions in mesophyll cell thickness and changes in IAS thickness near TLP to be species-specific (Sancho-Knapik et al., 2011; Scoffoni et al., 2014). In leaves with a structure like *J. microcarpa*, where the epidermis is more hydraulically integrated due to the presence of BSEs, water can bypass parts of the mesophyll reaching evaporation sites near epidermis as proposed by Zwieniecki et al. (2007). In this system, stomatal function is more closely linked to changes in xylem hydraulic conductance and stomata may show delays in the closure. Therefore, the relative presence of BSEs might play a role in allowing species to operate at more negative Ψ_{xylem} .

Dehydration reduced g_s equally in both species, but decreased K_{leaflet} more severely in J. microcarpa, in relative terms (Figure 5). It has been shown that mutants lacking BSEs (e.g., Solanum lycopersicum) have lower g_s , A_n and K_{leaf} than wild-type plants (Zsögön et al., 2015). In addition, BSEs are proposed to slow down stomatal closure under stress-induced conditions by enhancing hydraulic conductance through extravascular pathway (Barbosa et al., 2019; Buckley et al., 2011). Despite the higher presence of BSEs, the lower minimum g_s (0.02 mol H₂O m⁻² s⁻¹) in J. microcarpa compared to J. regia (0.05 mol $H_2O m^{-2} s^{-1}$) may represent greater response through stomatal closure via highly reduced K_{leaflet} (driven by changes outside the xylem) at the expense of significantly lower A_n (Figures 5 and 6) and higher WUE_i at ambient CO₂ under drought. Further, changes in g_s associated with stomatal aperture were supported by epidermal imprints, where stomatal aperture ratio decreased more for J. microcarpa (by 61%) compared to J. regia (by 38%). Dehydration reduced g_s to 27% and 11% of the g_{smax} for J. regia and J. microcarpa, respectively, further supporting the inherent difference in sensitivity of g_s to dehydration. Still, finding no significant emboli formation within xylem veins in either species (Figure 6) was in agreement with recent studies suggesting that declines in leaf hydraulic conductance are mostly due to declines in outside-xylem tissue hydraulic conductance under mild to moderate dehydration and even beyond the TLP (Albuquerque et al., 2020; Scoffoni et al., 2017). Furthermore, the contradicting results for J. microcarpa may highlight the importance of aquaporins activity in changing K_{leaf} under induced conditions, such as a positive link between aquaporins abundance and Kleaf under high (vs. low) light was reported in J. regia (Cochard et al., 2007).

Drought-induced shrinkage in mesophyll cells opens up more IAS volume within mesophyll, however nonuniform changes in cell shape can increase resistance to CO_2 diffusion by reducing chloroplast surface area facing cell walls (Cano et al., 2013; Tosens et al., 2012; Xiao & Zhu, 2017). Dehydration increased porosity, leading to an increase in g_{IAS} , however, the g_{IAS} contribution to g_m was reduced to 6-8% in both species under dehydration (p < 0.05). Therefore, limitation imposed by g_{Iiq} to g_m may increase under drought via chloroplast re-positioning and activity of CAs and aquaporins (Evans et al., 2009; Miyazawa et al., 2008; Momayyezi et al., 2020; Tholen et al., 2008; Tomás et al., 2013) more so than

changes in resistance via anatomical components such as the cell wall thickness and cell wall composition (e.g., lignin deposition) (Evans, 2021; Roig-Oliver et al., 2020). We must note, however, that the porous media approach of Earles et al. (2018) to compute g_{IAS} , a step forward in the representation of the inherent 3D nature of the leaf mesophyll, does not fully account for the specificities of the diffusion within the leaf. As discussed by Harwood et al. (2021), path lengthening is a step forward to account for the discrete nature of stomata along the epidermis, but path shortening within the mesophyll would also occurs because of the gradient of carbon assimilation within the leaf profile, and would theoretically increase g_{IAS}. In the present case, as water stress decreases photosynthesis, the path shortening effect could be smaller and might cancel out the g_{IAS} increases caused by higher porosity and lower tortuosity. Thus, the present result must be seen as a potential increase in g_{IAS} caused by anatomical changes.

Dehydration-induced impacts on θ_{IAS} and mesophyll cell positioning altered the chlorophyll distribution in J. microcarpa by changing the magnitude and location of fluorescence peaks (Figure 8), in alignment with our expectations for J. microcarpa to highly reflect stress-induced changes in cell geometry through light absorption. Increases in IAS volume between mesophyll cells, as seen by significant reductions in palisade cells diameter in paradermal sections at 20%, 40% and 60% from the adaxial epidermis (by 9% to 15%) (Figure 10), combined with increased frequency of BSEs in J. microcarpa to facilitate light diffusion and increase light absorption through the leaf and more at the spongy mesophyll under drought (Figure 9b). While dehydration reduced palisade cell diameter in J. regia at 40% and 60% adaxial depth, there was no significant change in absorption profiles (Figure 8) compared to the well-watered leaves. This can further highlight the role of the spongy mesophyll arrangement (Borsuk et al., 2019) in light penetration and overall absorption efficiency through the leaf under stress.

5 | CONCLUSIONS

Mesophyll structure has a substantial role in both CO_2 diffusion and light absorption. Juglans regia mesophyll with thick palisade layers and higher IAS volume between mesophyll cells and mostly near the spongy layer, has higher g_m in line with more carboxylation capacity and greater light absorption under well-watered condition. A more porous mesophyll with less BSEs has less anatomical leverage to tolerate dehydration and maintain the gas exchange in association with hydraulic components, and increases risk of damage to photosynthetic machinery. While more mesophyll cell density with less IAS distribution and greater BSEs (e.g., heterobaric leaf) can increase resistance to CO_2 diffusion and lower overall light absorption and photosynthesis, it performs better in light absorption under drought. Greater BSEs in *J. microcarpa* leaves provide physical and hydraulic support leading to less mesophyll cell shrinkage with minimum damage to the carboxylation activity.

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CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT

Data and materials are available on request from the corresponding author.

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