Xylem traits mediate a trade-off between resistance to freeze–thaw-induced embolism and photosynthetic capacity in overwintering evergreens

Brendan Choat1, Danielle E. Medek1, Stephanie A. Stuart1, Jessica Pasquet-Kok2, John J. G. Egerton1, Hooman Salari1, Lawren Sack2 and Marilyn C. Ball1

1Plant Science Division, Research School of Biology, The Australian National University, Canberra, ACT 0200, Australia; 2Department of Ecology and Evolutionary Biology, University of California, Los Angeles, 621 Charles E. Young Drive South, Los Angeles, CA 90095-1606, USA

Summary

• Hydraulic traits were studied in temperate, woody evergreens in a high-elevation heath community to test for trade-offs between the delivery of water to canopies at rates sufficient to sustain photosynthesis and protection against disruption to vascular transport caused by freeze–thaw-induced embolism.

• Freeze–thaw-induced loss in hydraulic conductivity was studied in relation to xylem anatomy, leaf- and sapwood-specific hydraulic conductivity and gas exchange characteristics of leaves.

• We found evidence that a trade-off between xylem transport capacity and safety from freeze–thaw-induced embolism affects photosynthetic activity in overwintering evergreens. The mean hydraulically weighted xylem vessel diameter and sapwood-specific conductivity correlated with susceptibility to freeze–thaw-induced embolism. There was also a strong correlation of hydraulic supply and demand across species; interspecific differences in stomatal conductance and CO2 assimilation rates were correlated linearly with sapwood- and leaf-specific hydraulic conductivity.

• Xylem vessel anatomy mediated an apparent trade-off between resistance to freeze–thaw-induced embolism and hydraulic and photosynthetic capacity during the winter. These results point to a new role for xylem functional traits in determining the degree to which species can maintain photosynthetic carbon gain despite freezing events and cold winter temperatures.

Introduction

Low temperatures are important determinants of species’ distributions, with freezing presenting major challenges to evergreen species in frost-prone climates (Sakai & Larcher, 1987). Woody plants that tolerate freezing undergo seasonal changes in acclimation which enable living cells to tolerate the dehydration that accompanies extracellular ice formation (Xin & Browse, 2000). In addition, water transport to leaves is threatened when xylem conduits become embolized following freeze–thaw events (Hacke & Sperry, 2001). Such a loss of water transport can reduce the capacity of a plant to take advantage of warm, sunny conditions, particularly when growth recommences in spring. Hence, among woody evergreens, interspecific differences in vulnerability to freeze–thaw-induced embolism could influence growth, competitive outcomes and distribution along gradients in minimum temperature. Although freeze–thaw-induced embolism has been studied in a variety of species (Sperry et al., 1994; Langan et al., 1997; Cavender-Bares & Holbrook, 2001; Feild & Brodribb, 2001; Ball et al., 2006), its influence on carbon balance is not well understood.

The mechanism of freeze–thaw-induced embolism differs from that of drought-induced tension (Hacke & Sperry, 2001). Ice is crystalline and the solubilities of gases in ice are much lower than in liquid water. Consequently, when water freezes, gases dissolved in water can be forced out of solution, forming bubbles surrounded by ice. When frozen xylem sap thaws, bubbles formed during freezing can either...
dissolve (in water released by melting ice) or expand until the xylem conduit is filled with gas, a process known as cavitation. According to the Young–Laplace equation, whether a bubble dissolves or nucleates a cavitation depends on its internal pressure ($P_b$), which is a function of its radius ($r$), the surface tension of xylem sap ($\gamma$) and the pressure of xylem sap ($P_s$), where $P_b = (2\gamma/r) + P_s$. Small bubbles or those subject to large positive external pressure tend to collapse or to dissolve, and so the probability that a bubble will nucleate a cavitation increases with an increase in the radius of curvature of the bubble and the tension in the xylem stream. These two components are highly variable. Values of $r$ are thought to depend on the size of xylem conduits; larger conduits contain more water and hence more air is forced out of solution during freezing, thereby increasing the probability that a bubble with a critical radius for nucleation will form on thawing (Sperry & Sullivan, 1992). Indeed, many studies have reported that narrow conduits have an advantage in resisting freeze–thaw-induced embolism (Sperry & Sullivan, 1992; Logullo & Salleo, 1993; Davis et al., 1999; Pittermann & Sperry, 2003; Stuart et al., 2007). Such an advantage for narrow conduits contrasts with anatomical adaptations to maintaining xylem function during drought, which relate more to the structure of intervessel pits (Choat et al., 2008). Although narrow conduits have lower probabilities of freeze–thaw-induced embolism (Davis et al., 1999), they may impose a limit on the carbon and water economy of plants when conditions are favourable for photosynthesis and growth. A small reduction in conduit diameter considerably decreases vessel conductivity, because, as the Hagen–Poiseuille equation states, conductivity is proportional to vessel radius raised to the fourth power (Tyree & Zimmermann, 2002). Thus, narrow conduits that benefit resistance to freeze–thaw-induced embolism involve a trade-off in maximum hydraulic conductivity (Wang et al., 1992; Sperry et al., 1994; Feild & Brodribb, 2001).

Such a trade-off would mean that narrow conduits may also limit growth (Castro-Diez et al., 1998). Although the role of stomata in regulating water loss and photosynthetic rates is well established (Givnish, 1986), the hydraulic architecture and hydraulic conductivity of stem xylem ($K_s$) may also constrain the productivity of trees (Zhang & Cao, 2009). Indeed, there is increasing evidence of functional coordination between the hydraulic capacity of stems to supply water to leaves and the gas exchange characteristics of leaves (Meinzer et al., 1995; Franks & Farquhar, 1999; Hubbard et al., 2001; Sack & Holbrook, 2006). These relationships are complicated by allocation patterns that affect leaf-specific conductivity $K_s$ [$K$ multiplied by the sapwood : leaf area ratio, i.e. the Huber value (HV)], the most direct metric for the influence of stem conductivity on leaf gas exchange (Brodribb et al., 2003; Santiago et al., 2004). For example, even a stem with low $K_s$ can support a high stomatal conductance and favourable rates of CO$_2$ assimilation without causing tension-induced cavitation if $K_s$ is sufficiently high (i.e. if the stem supports less leaf area for a given $K_s$). Thus, if hydraulic traits associated with the minimization of freeze–thaw-induced embolism constrain the rate of delivery of water to leaves, we predict that morphological adaptations that alter the scaling between photosynthetic and nonphotosynthetic tissues will have significant impacts on productivity.

Although the coordination of hydraulic and photosynthetic traits has been demonstrated in a number of studies, these have typically focused on optimal conditions during the primary growing season (Brodribb & Feild, 2000; Santiago et al., 2004; Chen et al., 2009), and there have been no such studies to our knowledge for woody plants facing winter. In the boreal zone, evergreen plants have low and often negligible photosynthetic rates during the winter (Mooney & Ehleringer, 1997). However, in the temperate zone, winter photosynthetic activity may contribute significantly to annual growth in broadleaf evergreens (Miyazawa & Kikuzawa, 2005). In Australia, winter photosynthesis supports root growth (Blennow et al., 1998; Egerton et al., 2000) and contributes substantially to the rapid growth of eucalypts during early spring when soil resources are relatively abundant (Ball et al., 1997). Hence, the ability to resist freeze–thaw-induced embolism may be a strong benefit for overwintering evergreens, although few data are available.

We examined the interrelationships between resistance to freeze–thaw-induced embolism, hydraulic capacity and leaf gas exchange in a high-elevation heath community dominated by evergreen angiosperms regularly exposed to winter freezing. The wide range of growth forms and diverse xylem structure present in the heath flora provided an ideal opportunity to study the functional diversity of adaptation in hydraulic traits to low-temperature stress. Measurements of xylem vessel diameter and hydraulic conductivity were made in conjunction with experimental manipulations to determine the loss of hydraulic capacity associated with a severe freeze–thaw event, and were related to field measurements of stomatal conductance and CO$_2$ assimilation rates. We tested three key hypotheses of hydraulic limitation in this system: that stem hydraulic conductivity is negatively related to resistance to freeze–thaw-induced embolism; that stem hydraulic conductivity correlates across species with leaf gas exchange; and that this variation in xylem vessel diameter underlies both relationships.

**Materials and Methods**

**Study site**

Field measurements were made in July (mid-winter) 2009 in Kanangra Boyd National Park (33°59′9″S,
15°6′37″E), within the Greater Blue Mountains World Heritage Area in south-eastern Australia. The site was located on an exposed plateau at 1070 m above sea level adjacent to the Kanangra Walls. No weather data are available for this remote site. The closest weather station is located c. 20 km from the site at Oberon (elevation 1190 m above sea level) in very different topography, and so gives an indication of regional conditions. Oberon receives a mean annual rainfall of 725 mm yr\(^{-1}\)) gives an indication of regional conditions. Oberon receives a mean annual rainfall of 725 mm yr\(^{-1}\)) and high levels of hydration during winter.

Eleven shrub and tree species were selected from the heath vegetation on Boyd Plateau, as well as *Acacia* and *Allocasuarina* (Table 1). Three individuals of each species were selected for measurements of anatomy, hydraulic traits and leaf gas exchange.

### Xylem anatomy

Branch samples were collected for each species before dawn and returned to the field laboratory. For each of the 11 species, we measured the maximum vessel length using the air injection technique (Zimmermann and Jeje 1981). Branch samples of 5–10 mm in diameter were stored in 50% ethanol for anatomical analysis. Transverse sections were then taken on a sliding microtome and observed under a compound microscope (Nikon, Tokyo, Japan). Images were collected using a digital camera (Canon Powershot S5; Canon, Tokyo, Japan) and analysed using imageJ software (US National Institutes of Health, Bethesda, MD, USA). A region of sapwood from pith to cambium was examined, and the xylem diameter was estimated by measuring the areas of the vessel lumens in transverse section and calculating the diameter of a circle with the same area. Conduits with diameters < 5 \(\mu\)m were excluded from the analysis. The hydraulically weighted vessel diameter (\(D_h\)) was calculated as \(D_h = (\Sigma D^4/N)^{1/4}\), where \(N\) is the number of vessels in cross-section (Tyree & Zimmermann, 2002).

### Hydraulic conductivity and freeze–thaw-induced embolism

Branch sections longer than the longest vessel were collected before dawn, placed into plastic bags with moist paper towels and returned to the field laboratory, a 30 min drive away. For each replicate individual, two branches were collected, one for the freeze–thaw treatment and one as an untreated control. In the laboratory, branches were separated into two groups. Treatment branches were placed into an insulated freezing chamber (Stuart et al., 2007) and control branches were left in plastic bags for the duration of the treatment. The temperature of the freezing chamber was controlled by a programmable water bath (Julabo, Seelbach, Germany). The air temperature of the chamber and the temperature of stems and leaves were monitored throughout the experiment by copper-constantan thermocouples referenced against a platinum resistance thermometer and connected to a DT800 Datatker (Datatker, Scoresby, Australia). The chamber was cooled from 2°C at a rate of 2°C per hour, with stems reaching average nadir temperatures of −7.5°C at the conclusion of the freezing cycle (Table 2). Freezing exotherms were recorded first in the leaves in almost all cases at an average temperature of −4.8°C. Samples were held at the nadir temperature for 30 min, and then removed from the freezing chamber and allowed to thaw to room temperature over 30 min. Samples were placed back into humidified plastic

Table 1 | Species and families selected from heath vegetation with average species’ height at site
---|---|---|---
Species | Family | Growth form | Height (m)
---|---|---|---
*Acacia obtusifolia* (Ao) | Fabaceae | Tree | 3
*Allocasuarina littoralis* (Al) | Casuarinaceae | Tree | 6
*Baeckia brevifolia* (Bb) | Myrtaceae | Shrub | 2
*Banksia ericifolia* (Be) | Proteaceae | Tree | 2
*Banksia spinulosa* (Bs) | Proteaceae | Tree | 2
*Eucalyptus burgessiana* (Eb) | Myrtaceae | Tree | 10
*Eucalyptus sieberi* (Es) | Myrtaceae | Tree | 15
*Hakea dactyloides* (Hd) | Proteaceae | Tree | 3
*Isopogon anemonifolius* (Ia) | Proteaceae | Shrub | 1
*Leptospermum polygalifolium* (Lp) | Myrtaceae | Tree | 2
*Petrophile pulchella* (Pp) | Proteaceae | Shrub | 2

Table 2 | Leaf nucleation temperature and nadir stem temperature for branches in freezing chamber (averages ± SE)
---|---|---
Species | Leaf nucleation temperature (°C) | Nadir stem temperature (°C)
---|---|---
*Allocasuarina littoralis* | −5.14 ±0.51 | −8.32
*Acacia obtusifolia* | −7.04 ±0.64 | −7.65
*Baeckia brevifolia* | −5.54 | −7.00
*Banksia ericifolia* | −4.29 ±0.31 | −6.62
*Banksia spinulosa* | −5.45 ±0.32 | −8.24
*Eucalyptus burgessiana* | −3.33 ±0.74 | −7.81
*Eucalyptus sieberi* | −3.46 ±0.43 | −7.64
*Hakea dactyloides* | −4.79 ±0.33 | −7.46
*Isopogon anemonifolius* | −4.46 ±0.39 | −7.79
*Leptospermum polygalifolium* | −6.10 ±1.10 | −7.43
*Petrophile pulchella* | −3.72 ±0.22 | −7.08

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Hydraulic conductivity was measured on branch sections, 5 cm in length and 5–10 mm in diameter, cut from treatment and control branches under water and trimmed with a fresh razor blade. Segments were connected to water-filled tubing running to a reservoir raised above the stems to deliver 5 kPa of hydraulic pressure. The perfusate used was 10 mM KCl, filtered to 0.2 μm with Millipore syringe filters (Millex GS Filter; Millipore Corrigtwohill Co, Cork, Ireland). The rate of water flow through the stems was measured by tracking the movement of the meniscus in a calibrated pipette connected by tubing to the distal end of the stem. The position of the meniscus was recorded every 30 s by a digital camera (Canon Powershot G10) connected to a laptop computer using Canon Utilities RemoteCapture DC version 3.1.0.5; Canon). The flow rate was measured for 10–20 min, with the final flow rate value averaged over a 3 min period once the flow had stabilized. After an initial flow rate measurement, the samples were flushed at a pressure of 125 kPa for 20 min to remove emboli. The flow rate was then measured again to determine the maximum conductivity and the loss of conductivity caused by embolism. The hydraulic conductivity ($K_h$) was calculated as $K_h = FL/ΔP$, where $F$ is the flow rate, $L$ is the segment length and $ΔP$ is the pressure difference across the sample. The leaf area distal to the segment ($A_s$) and the conductive sapwood area ($A_i$) were also measured to calculate the sapwood-specific hydraulic conductivity ($K_s$) and leaf-specific hydraulic conductivity ($K_l$) as $K_s = K_h/A_i$ and $K_l = K_h/A_s$, respectively. The percentage loss of hydraulic conductivity (PLC) was calculated as $PLC = (K_{max} - K_{initial})/K_{max} \times 100$, where $K_{initial}$ and $K_{max}$ are the initial and maximum hydraulic conductivities, respectively. The PLC caused by a freeze–thaw cycle was calculated as the PLC of treated branches minus the PLC of control branches. For each of the 11 species, a treatment branch and a control branch from each of three individuals were measured to determine the hydraulic parameters.

The segment length of 5 cm was chosen to allow for easy flushing of the segments and standardization across a broad range of branch morphology. It is recognized that open vessels occur in these short segments and that this will lead to higher values of $K_s$ and $K_l$ than in longer segments with no open vessels, because endwall resistance has been removed. Therefore, caution must be used when comparing these measurements directly with those made on longer segments with no open vessels. However, because lumen and pit resistances are coordinated across angiosperm species (Sperry et al., 2005), it is expected that conductivity values obtained from short segments will hold their species’ ranks for comparison with leaf gas exchange parameters.

Leaf gas exchange

Measurements of stomatal conductance and CO$_2$ assimilation rate were made during the morning from 09:00 to 11:00 h over a 3 wk period in July 2009 with an Li-Cor portable gas exchange system (Li-Cor 6400; Lincoln, NE, USA). Measurements were made on one block per week, with each block containing one individual of each of the 11 species. Measurements were made at ambient temperature and relative humidity with a photosynthetic photon flux density of 1200 μmol m$^{-2}$ s$^{-1}$ and a chamber flow rate of 400 μmol s$^{-1}$. There was no significant difference ($P = 0.48$) between species in leaf temperature, which ranged from average values of 9.6 ± 0.1°C for Eucalyptus sieberi to 11.4 ± 1.2°C for Acacia obtusifolia. The reference CO$_2$ concentration was maintained at 380 μmol mol$^{-1}$ by the LI-6400 CO$_2$ injector using a high-pressure liquefied CO$_2$ cartridge.

Statistics

For initial and maximum values of $K_h$ and $K_l$, and for xylem anatomical traits, differences among species were established using a general linear model with trait data averaged for each replicate tree ($n = 3$; SPSS Statistics 18; IBM Corporation, New York, NY, USA). To test the effect of the freeze–thaw treatments, we conducted a repeated-measures ANOVA, comparing the impact on each individual of each species, determining the effect of species, treatment and interaction (Minitab Pty. Ltd., Minitab Release 14; State College, PA, USA). Bivariate relationships were assessed using parametric correlation coefficients ($r$), and lines were fitted by linear regression or nonlinear curve fitting using the simplex procedure (Origin 8.1; OriginLab Corporation, Northampton, MA, USA).

Results

Xylem anatomy

Xylem anatomy varied significantly among the 11 species (Table 3). The mean hydraulically weighted vessel diameter ($D_h$) ranged from 14.1 μm in Hakea dactyloides to 33.3 μm in E. sieberi. The hydraulically weighted vessel diameter was negatively related to vessel density and positively related to maximum vessel length ($L_{v,max}$) ($r = -0.79$, $P = 0.004$ and $r = 0.86$, $P < 0.001$, respectively) (Fig. 1).

Hydraulic conductivity and freeze–thaw-induced embolism

The sapwood-specific hydraulic conductivity ($K_s$) varied widely among the 11 species ($P < 0.001$; ANOVAs on initial and flushed stems), with two shrub species, Isopogon...
anemonifolius and Petrophile pulchella, having the lowest values and the two Eucalyptus species having the highest values (Fig. 2a). For $K_s$, ranks were exchanged between the species because of variation in HV (Fig. 2b). HV increased with decreasing $K_s$ following a power function ($r^2 = 0.66$, $P < 0.0001$), indicating that the species with less efficient sapwood tend to support their leaf area with greater cross-sectional area of stem (Fig. 3). Thus, HV tended to partially compensate for $K_s$ differences and, as a result, $K_s$ was less variable than $K_L$ across species, and did not correlate significantly with $K_L$ across species (Figs 2a, b, 3).

The loss of hydraulic conductivity induced by the experimental freeze–thaw treatment was significant across the species (repeated-measures ANOVA testing the percentage loss in conductivity between the control and treated shoots; species effect, $P = 0.006$; treatment, $P = 0.001$; species × treatment, $P = 0.045$). Freeze–thaw-induced loss of conductivity was negligible for Baeckia brevifolia and Leptospermum polygalifolium and varied from 4% for I. anemonifolius to 56% for E. sieberi (Fig. 4). Species with narrower vessels and lower conductivities were less vulnerable to freeze–thaw-induced embolism than species with wider vessels (Fig. 5); the magnitude of PLC caused by a freeze–thaw cycle was significantly related to $D_h$ ($r = 0.65$, $P = 0.03$) and $K_s$ ($r = 0.71$, $P = 0.01$). No relationship was observed across species between stem water potential before freezing and freeze–thaw-induced PLC; however, stem water potentials were relatively high ($0.15$ to $0.98$ MPa) and were above $-0.5$ MPa in all species except B. brevifolia (overall mean = $-0.32$ MPa; Table 3).

Relationships between leaf gas exchange and hydraulic traits

Stomatal conductance and CO$_2$ assimilation rates were significantly related to $K_{Lmax}$ ($r = 0.87$, $P < 0.001$ and $r = 0.73$, $P = 0.01$, respectively) for untreated control branches (Fig. 6), demonstrating the association of leaf level gas exchange and the capacity of stem tissue to supply water to the leaves. Stomatal conductance and $A$ were related to $K_{smax}$ ($r = 0.67$, $P = 0.02$ and $r = 0.71$, $P = 0.01$, respectively).

### Table 3

<table>
<thead>
<tr>
<th>Species</th>
<th>$D_h$ (μm)</th>
<th>VD</th>
<th>$L_{Vmax}$ (cm)</th>
<th>$\Psi_s$ (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acacia obtusifolia</td>
<td>25.2 (2.0) bcd</td>
<td>115 (11) a</td>
<td>70</td>
<td>$-0.46$ (0.17)</td>
</tr>
<tr>
<td>Allocasuarina littoralis</td>
<td>19.8 (1.1) abc</td>
<td>537 (7) bcd</td>
<td>68</td>
<td>$-0.24$ (0.05)</td>
</tr>
<tr>
<td>Baeckia brevifolia</td>
<td>15.7 (2.3) a</td>
<td>398 (4) abc</td>
<td>29</td>
<td>$-0.98$ (0.42)</td>
</tr>
<tr>
<td>Banksia ericifolia</td>
<td>18.1 (2.7) ab</td>
<td>367 (17) abc</td>
<td>7</td>
<td>$-0.15$ (0.08)</td>
</tr>
<tr>
<td>Banksia spinulosa</td>
<td>19.3 (1.8) abc</td>
<td>460 (23) abc</td>
<td>14</td>
<td>$-0.19$ (0.09)</td>
</tr>
<tr>
<td>Eucalyptus burgessiana</td>
<td>27.8 (1.6) cd</td>
<td>185 (16) ab</td>
<td>64</td>
<td>$-0.23$ (0.09)</td>
</tr>
<tr>
<td>Eucalyptus sieberi</td>
<td>33.3 (0.4) d</td>
<td>152 (1) ab</td>
<td>120</td>
<td>$-0.17$ (0.09)</td>
</tr>
<tr>
<td>Hakea dactyloides</td>
<td>14.1 (1.4) a</td>
<td>762 (9) cd</td>
<td>10</td>
<td>$-0.36$ (0.13)</td>
</tr>
<tr>
<td>Isopogon anemonifolius</td>
<td>16.2 (0.5) a</td>
<td>367 (6) abc</td>
<td>14</td>
<td>$-0.23$ (0.03)</td>
</tr>
<tr>
<td>Leptospermum polygalifolium</td>
<td>19.3 (2.2) abc</td>
<td>436 (6) abc</td>
<td>52</td>
<td>$-0.15$ (0.08)</td>
</tr>
<tr>
<td>Petrophile pulchella</td>
<td>15.6 (1.0) a</td>
<td>872 (16) d</td>
<td>15</td>
<td>$-0.33$ (0.21)</td>
</tr>
</tbody>
</table>

Values are means from three individuals ($n = 3$) with SE in parentheses. Different lower case letters represent significant differences ($P < 0.05$) within each column.
tively), although the linear regression explained less of the variation than the relationships between gas exchange parameters and $K_{L\text{max}}$ (Fig. 6). Correlations were also found between $A$ and $D_h$ ($r = 0.65$, $P = 0.03$) and PLC ($r = 0.73$, $P = 0.01$) (Fig. 7).

**Discussion**

We observed coordination in a suite of traits governing resistance to freeze–thaw-induced embolism, hydraulic efficiency of the xylem and photosynthetic capacity of leaves in 11 heathland species. Across these species, significant positive relationships between leaf gas exchange ($A$ and $g_s$) and $K_L$ indicated that photosynthetic and hydraulic traits were highly interdependent. Vulnerability to freeze–thaw-induced embolism increased with $K_s$, a relationship driven by variations in xylem vessel characteristics; species more susceptible to freeze–thaw-induced embolism had wider and longer xylem vessels with lower vessel density per square millimetre. These results show that the tremendous variation in anatomy and physiology underlies a continuum of traits along which tolerance to freezing conditions is bal-

![Fig. 2] Hydraulic conductivity of stems for 11 woody heath species. Initial (open bars) and maximum (closed bars) hydraulic conductivity is shown normalized to leaf area distal to the stem ($K_s$) (a) and xylem cross-sectional area ($K_L$) (b). Bars show mean values from three individuals ($n = 3$) with error bars showing standard error. Abbreviations of the species' names are given in Table 1.

![Fig. 3] Relationship between maximum sapwood-specific conductivity ($K_{s\text{max}}$) and Huber value (HV) (a), and $K_{s\text{max}}$ and maximum leaf-specific conductivity ($K_{L\text{max}}$) (b). Each point is the mean from three individuals ($n = 3$) with error bar showing standard error.

![Fig. 4] Percentage loss of hydraulic conductivity (PLC) caused by freeze–thaw-induced embolism in stems of woody heath species. Bars show mean values from three individuals ($n = 3$) with error bar showing standard error. Abbreviations of the species' names are given in Table 1.
advanced by hydraulic efficiency and photosynthetic capacity, even during the winter. Notably, the heath vegetation at this high-elevation site maintained a significant degree of photosynthetic activity during a winter characterized by cool, above zero, daytime temperatures and high levels of hydration. Therefore, a significant potential for growth exists during the winter despite the regular imposition of freezing temperatures and the consequent possibility of freeze–thaw-induced cavitation.

Xylem anatomy, hydraulic efficiency and resistance to freeze–thaw-induced embolism

Xylem vessel diameter and length emerged as central to the coordination of traits linked to stress tolerance and productivity. In our dataset, there was a significant positive relationship between vessel diameter and maximum vessel length (Fig. 1). Although the full frequency distribution of vessel lengths was not measured, published distributions suggest that median and maximum vessel lengths tend to be well correlated (Zimmermann & Jeje, 1981; Sperry et al., 2005). This xylem vessel allometry suggests that species with wider vessels should also have fewer vessel endings per unit length, and therefore the hydraulic resistance caused by intervessel pit crossings would be reduced as vessels become wider (Choat et al., 2006). The number of vessels in cross-section per unit area decreased with increasing vessel diameter, a well-established trend following the geometric packing limit (Baas, 1986; McCulloh et al., 2010; Zanne et al., 2010). However, $K_s$ increased with $D_h$ and $V_L$ despite lower vessel density, because of the exponential increase in conductivity with conduit diameter.

Wider conduit diameter was also associated with greater vulnerability to freeze–thaw-induced embolism, consistent with previous studies of both coniferous and angiosperm species (Sperry et al., 1994; Davis et al., 1999; Feild & Brodribb, 2001; Pittermann & Sperry, 2003; Cobb et al., 2007; Stuart et al., 2007). Here, we found a correlation within one community representing a continuum of freeze–thaw-induced PLC and vessel diameter, similar to that observed in Tasmanian heath species (Feild & Brodribb, 2001). Notably, none of the heath species in the present study exhibited the extreme vulnerability of ring porous species and vines in the Northern Hemisphere (Sperry et al., 1994; Cobb et al., 2007). Indeed, the heath species were relatively resistant to freeze–thaw-induced embolism, with the majority suffering PLCs of 20% or less in response to a $-10^\circ$C freeze.

However, care must be taken when comparing results of different studies, given that the magnitude of tension and time of freezing can also be important factors in freeze–thaw-induced embolism (Davis et al., 1999; Mayr et al., 2007; Stuart et al., 2007). For example, although, in our study, freeze–thaw-induced PLC values ranged from 20 to 60%, a previous study reported freeze–thaw-induced PLCs ranging from 40 to 80% for a range of angiosperm tree species with vessel diameters similar to those observed here (15–30 $\mu$m), including some congeners with our dataset, such as Eucalyptus and Leptospermum (Feild & Brodribb, 2001). However, in that study, branches were dehydrated to c. $-0.5$ MPa before the freeze–thaw treatment, whereas branches in the present study were frozen at the naturally occurring stem water potentials, which were high (i.e. mostly above $-0.5$ MPa) as a result of winter rainfall, and we saw no relationship between stem water potential before freezing and PLC. Thus, high water status may have contributed to the apparent high level of resistance to freeze–thaw-induced embolism in the heath species.

Coordination of stem hydraulic traits and leaf gas exchange

A strong positive relationship was observed between $K_s$ and leaf gas exchange parameters ($g_s$ and $A$) across the 11 species (Fig. 6). Such direct scaling between light-saturated photosynthetic rates and the maximum hydraulic conductivity of

![Fig. 5 Relationship between percentage loss of hydraulic conductivity (PLC) caused by freeze–thaw-induced embolism and maximum sapwood-specific hydraulic conductivity ($K_{smax}$) (a) and mean hydraulically weighted vessel diameter ($D_h$) (b). Each point is the mean from three individuals ($n = 3$) with error bar showing standard error.](image-url)
stems (Santiago et al., 2004; Lovelock et al., 2006; Chen et al., 2009) or leaves (Brodribb et al., 2007) has been reported for a range of temperate and tropical species. We found that $g_s$ and $A$ were more closely related to $K_L$ than $K_s$. Most previous studies have also found that gas exchange parameters are more closely related to $K_L$ than to $K_s$, suggesting that the coordination of hydraulic supply and demand incorporates the allometry of the sapwood : leaf area ratio, rather than simply conforming to the hydraulic capacity of the stem xylem. In addition, HV decreased in a log-linear fashion with increasing $K_s$, indicating that a greater amount of leaf area was supported per cross-sectional area of stem as the efficiency of sapwood increased. Because changes in $K_s$ are largely controlled by changes in vessel diameter and length, increasing vessel size was correlated with greater investment in leaf area for a given investment in sapwood tissue. This association is consistent with the idea that hydraulic traits related to whole-plant relative growth rates are part of a suite of coordinated traits, i.e. faster growing species have more allocation to leaf area relative to stem area for competitive light capture, as well as higher hydraulic and photosynthetic efficiency. Thus, the relationship between $A$ and $K_s$ is probably the result of a long-term coordination of hydraulic and photosynthetic traits, with far-reaching consequences for all aspects of plant morphology and function. The fact that the plants were in well-watered soil indicates that the correlation was not a result of differential stomatal closure across species to maintain hydraulic function. Rather, the relationship appears to reflect species optimizing maximum gas exchange relative to maximum hydraulic capacity, and this coordination is maintained when gas exchange and hydraulic conductance decline in the winter.

Conclusions

This study revealed strong integration between resistance to freezing stress and hydraulic and photosynthetic capacity across a functionally diverse range of overwintering broadleaf evergreens. Trade-offs were mediated by xylem anatomy, with narrow xylem vessel diameter correlated with resistance of stems to freeze–thaw-induced embolism, and higher rates of leaf gas exchange being related to greater hydraulic capacity ($K_L$ and $K_s$), which was derived principally from wider xylem vessels. Trade-offs in hydraulic and photosynthetic capacity strongly imply that many of the shrub species would be limited in productivity by xylem traits that afford better resistance against freeze–thaw-induced embolism induced by episodic severe freezing events. By contrast, the taller stature Eucalyptus species exhibited higher photosynthetic rates that came at the expense of more significant losses of hydraulic capacity after freezing. Given the high water status of material examined in this study, the potential for more catastrophic loss of hydraulic capacity may accompany
unusually dry periods during the winter. This is of particular importance given predictions of greater variation in regional precipitation patterns.

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