Breakthrough Technologies

Optical Measurement of Stem Xylem Vulnerability

Timothy J. Brodribb, Marc Carriqui, Sylvain Delzon and Christopher Lucani

Abstract: The vulnerability of plant water transport tissues to a loss of function by cavitation during water stress is a key indicator of the survival capabilities of plant species during drought. Quantifying this important metric has been greatly advanced by non-invasive techniques that allow embolisms to be directly viewed in the vascular system. Here we present a new method for evaluating the spatial and temporal propagation of embolising bubbles in the stem xylem during imposed water stress. We demonstrate how the “optical method”, previously used in leaves, can be adapted to measure the xylem vulnerability of stems. Validation of the technique is carried out by measuring the xylem vulnerability of 13 conifers and two short vesselled angiosperms and comparing results with measurements made using the “cavitron” centrifuge method. Very close agreement between the two methods confirms the reliability of the new optical technique, and opens the way to simple, efficient and reliable assessment of stem vulnerability using standard flatbed scanners, cameras or microscopes.

INTRODUCTION

In modern tracheophytes xylem cavitation constitutes a fundamental limitation to the functionality of water transport systems. As a consequence, the ability of species to resist or avoid cavitation forms a primary axis of adaptation and ecological variation among land plants (Xu et al., 2016). However, despite the tremendous ecological and physiological insights that await a detailed understanding of the limits and spread of xylem cavitation in plant species, rapid progress has been limited by technical difficulties. These difficulties are largely associated with replicating, under experimental conditions, the metastable hydraulic environment that characterizes water flowing in the xylem when exposed to the large tensions that exist during rapid transpiration or soil water deficit (Cochard et al., 2013).

Most traditional methods of quantifying the degree of xylem embolism require excision of plant parts (stems, roots or leaves), causing air or exogenous water to be rapidly sucked into the vasculature, thereby substantially perturbing the vascular system prior to measurement (Ennajeh et al., 2011; Rockwell et al., 2014). A substantial advance in recent years has been the use of imaging technology that allows water to be viewed inside intact plants, revealing the location and formation of embolisms inside stems (Brodersen et al., 2013), roots (Cuneo et al., 2016), leaves (Bouche et al., 2015; Brodribb et al., 2016; Scoffoni et al., 2017), and flowers (Zhang and Brodribb, 2017). These studies have substantially changed our view of xylem cavitation and repair, indicating that cavitation can propagate quickly between plant organs (Skelton et al., 2017), and that air blockages (embolisms) are not rapidly repaired in trees after re-watering (Choat et al., 2015; Charrier et al., 2016).

Cavitation is now widely viewed as a long-term damage to the water transport system of trees, that...
occurs under significant water stress, and that is repaired by regrowth of new xylem tissue (Brodribb et al., 2010; Cochard and Delzon, 2013).

Imaging with x-ray provides unrivalled spatial information about where cavitation occurs in stems and can be used to determine the vulnerability of xylem to cavitation in plant species (Choat et al., 2015; Nolf et al., 2017). However the damaging nature of the x-ray beam means that high frequency imaging during the hours and days required to dehydrate plants to water stresses sufficient to cause cavitation is not possible. Magnetic resonance imaging on the other hand can provide spatial and temporal information about cavitation, but low image resolution (pixel sizes larger than the vessels of most species) means that MRI can only be used to resolve embolisms in species with very large vessels. Both techniques require large and expensive hardware and are not currently usable in the field, thus having limited application for measuring large sample sizes. An alternative to these hardware-intensive methods was recently developed using an optical technique measures changes in visual light transmission caused by cavitation in leaf veins (Brodribb et al., 2016). This technique was developed following observations of cavitation bubbles in excised conifer tracheids (Ponomarenko et al., 2014), and provides detailed information about the spatial and temporal evolution of cavitation in the venation network of leaves exposed to water stress. The calculated “vulnerability” of leaf xylem to cavitation (expressed as a $P_{50}$, or potential required to deactivate 50% of xylem function) using this Optical Vulnerability (OV) method agrees closely with hydraulically measured $P_{50}$ in leaves (Brodribb et al., 2016), indicating the utility of the method for quantifying hydraulic failure. Importantly, the OV method requires only a flatbed scanner or camera to collect vulnerability information, thus providing a cost effective and portable means of assessing leaf xylem vulnerability.

Although the OV method has a demonstrated capacity to reveal leaf vulnerability to water stress, one of the primary applications of xylem cavitation physiology is in the prediction of tree mortality (Anderegg et al., 2015) and species distribution (Larter et al.), and in these applications stem vulnerability may provide a more definitive mortality threshold than leaves. Studies of potted plants have shown that failure of the stem xylem corresponds closely to the point of tree mortality during acute drought stress (Brodribb and Cochard, 2009; Urli et al., 2013), as might be expected considering the fact that embolism of the stem effectively isolates the leaves from soil water. A vulnerability gradient from stems to leaves is evident in some species (Tyree et al., 1993) (but probably not in herbs (Skelton et al., 2017)), and is hypothesized to be a way that woody plants protect their more energy expensive stem investment by sacrificing leaves during extreme drought (Zimmermann, 1983; Hochberg et al., 2017). Given the importance of understanding stem vulnerability in woody plants we sought here to extend the highly efficient OV method in leaves, to stems. We postulated that the same principle used to identify cavitation in leaves, recording changes in light transmission caused by a transition from liquid to air filled xylem conduits during cavitation, could be used in stems. Indeed it has been known for a long time that air bubbles can be visualized in stems by light microscopy (Vesque, 1883), and the same principle was used 80 years ago as a way of identifying the presence of water or air in branches by the evolution of light coloured streaks in the wood after it had been pricked with a sharp scalpel (Haines, 1935). Here we utilize the principal that a transition from a water-containing, to an air-filled conduit during cavitation will cause a distinct colour change in visible conduits from translucent (typically dark) to reflective (white) tissue. Thus we quantify spatially discrete changes in the refractive index of the stem. Continuous
observation of drying stems should thus allow the timing and pattern of cavitation to be recorded and quantified in relation to concurrent measurements of stem water potential.

In order to cross-validate the new stem optical method here we use a traditional hydraulic centrifuge method as a standard reference for comparison. The centrifuge method has long been considered an accurate method for assessing xylem vulnerability, except in cases where maximum vessel lengths are similar to the diameter of the centrifuge rotor (Cochard et al., 2013). For this reason we focussed on a diverse group of conifers which lack long xylem conduits, and two short vessel angiosperms were also included to maximize the breadth of the species sample.

RESULTS

Cavitation was easily resolved visually, and could be readily quantified by applying image difference to distinguish fast changes in light reflection due to xylem cavitation from slow movements associated with branch deformation during drying. The onset of cavitation was recorded on average 1308 minutes after branch excision, but ranged from 420 to 2230 minutes. In all species, the cumulative total of cavitations recorded followed an approximately sigmoidal function, although this was never a completely smooth function, typically being punctuated by blocks of major cavitation (Fig.1, Fig. 2). These blocks of cavitation often involved hundreds of tracheids in the conifers, and typically became larger as water potential approached $P_{50}$, before diminishing in size towards the end of the drying process. Typically, many cavitation events were recorded in the same part of the stem due to the multiple layers of overlaying xylem that were represented in the 2D image differences. The total cavitated area was typically 150-200% of the 2D area of the exposed stem (due to multiple layers of conduits).

Cavitation in the two species of angiosperms also appeared to involve groups of conduits, particularly during the period of maximum intensity of cavitation around $P_{50}$ (Fig. 2). But smaller events, presumably representing individual conduits, were often observed as early events, or as a tail towards the end of the cavitation process (Fig. 2).

Large differences in $P_{50}$ were recorded between species using the optical method, with means ranging from -1.2MPa in *Retrophyllum comptonii* to – 9.1MPa in *Disemla archeri*. Within species variation was also significant in many species, reaching a maximum in *Diselma archeri* where $P_{50}$ ranged between -6.7 and -11.2 MPa between individuals. On average the coefficient of variation in $P_{50}$ among replicate branches was 16.2% using the optical method and 9.2% using the cavitron. Mean slopes of the vulnerability curves for each species (between 12% and 88% loss of function) were correlated between the two methods, but the optical method produced steeper slopes on average.

Among the conifer species there was strong agreement between $P_{50}$ determined with the optical technique and centrifuge techniques. A regression slope of 0.997 ($r^2=0.93$) was found between optical and centrifuge $P_{50}$ in the 12 conifer species, and the ranking of $P_{50}$ was very similar using both methods.
One of the two angiosperms sampled showed a significant difference between the optical and centrifuge $P_{50}$. Although both techniques found *Rosmarinus* samples to be highly cavitation resistant, $P_{50}$ on the centrifuge (-12MPa) was 32% more negative than the optical method (-8.1MPa).

DISCUSSION

A new optical method for visualizing the process of xylem cavitation in plants is shown here to quantify the vulnerability of stem xylem to cavitation-induced reductions in hydraulic function of the stem xylem. The process of cavitation damage to the stem vascular system during water stress could be tracked in time and space on the stems of a diversity of species including woody conifers and angiosperms. This novel technique represents a very easy and cheap new method for assessing stem vulnerability in woody species using excised branches. In principle, the method can also be used on attached branches, although this was not tested here.

The optical technique allows direct visualization of the process of cavitation in stems under realistic conditions of plant desiccation (as opposed to centrifugation or stem pressurization). Apart from its simplicity, the advantage of this technique is that it provides a complete view of the spatial and temporal progression of cavitation in stems during increasing water stress. This new perspective of stem cavitation means that continuous monitoring of stem cavitation is possible as bubbles propagate axially in the stem during the development of increasing water deficit. Although cross-validation of the technique was performed using woody stems, the technique also works well in herbaceous species, where more translucent stems often do not require phloem removal.

The precisely resolved temporal dynamics of stem cavitation in both conifers and angiosperms studied here all yielded vulnerability curves that were highly sigmoidal in shape, characterized by an initial, extended period of stem desiccation before any stem cavitation events were recorded. This sigmoidal form of xylem vulnerability measured by the OV technique closely matches the form of cavitron (Lamy et al., 2011) and x-ray CT (Choat et al., 2015) vulnerability curves. The majority of data collected using traditional bench drying methods of measuring xylem vulnerability also produce sigmoidal vulnerability curves, but more linear curves are often reported in species with highly stress-resistant xylem (Markesteijn et al., 2011; Vinya et al., 2013). One important benefit of the OV and CT methods of assessing vulnerability is that they report the responses of functioning xylem without reference to a “flushed” condition. The flushing procedure is required by other hydraulic techniques, whereby samples are subjected to high water pressure to fill all airspaces in the sample and provide a theoretical maximum conductance. Flushing has the potential to activate (refil) xylem that was non-functional xylem in the intact plant, as well as introducing bubble nuclei, both of which can produce erroneous vulnerability curves (Rockwell et al., 2014).

Among the range of alternative methods for measuring xylem vulnerability, the cavitron was selected here as a standard for comparison because it is considered to be highly reliable when used to measure species with short conduits such as conifers (Cochard et al., 2013). For this reason most of our sample set was taken from the conifer clade, using the same individuals for both optical (sampled in 2016) and cavitron (sampled in 2012) techniques. The accuracy of centrifuges for measuring angiosperm xylem vulnerability is the subject of considerable debate due to probable artefacts associated with long vessels (Torres - Ruiz et al., 2014; Hacke et al., 2015). For this reason
we only measured two species of angiosperms, selected to cover a range of sensitivity to water
stress, but both of which had maximum xylem vessel lengths that were approximately half that of
the rotor diameter. Despite the huge difference in vulnerability between the two angiosperms
measured here, both were found to produce a sigmoidal form in their vulnerability curves using both
optical and cavitron methods. Our predawn sampling of well watered trees ensured that sampled
branches started drying from water potentials close to zero, thus ensuring a minimum of native
embolism in the measured samples.

The optical method assesses the loss in xylem function in terms of a cumulative area of stem
cavitated in each frame of image sequences. This area-based calculation does not account for the
profound influence of xylem conduit radius, in the order of \( r^4 \), that should determine the flow
penalty incurred by cavitation of any particular conduit in the stem (Sperry et al., 2006). Despite this
apparent limitation there was very strong agreement in \( P_{50} \) between the optical method (reporting
area of cavitated conduits) and the centrifuge method (quantifying losses in hydraulic conductance).
The explanation for the strong agreement between techniques despite different metrics of
cavitation is clearly evident from the spatiotemporal distribution of cavitation in stems observed
here. Most significant is the evolution of cavitation in large blocks of connected conduits as opposed
to discrete conduits, particularly as stems approached the \( P_{50} \) water potential. These large
interconnected cavitation events are also seen in x-ray images of stems (Choat et al., 2015b), and
produce a steep slope in the vulnerability curve around \( P_{50} \). Assuming that cavitation in stems on the
centrifuge also proceeds in this fashion, then it would be expected that \( P_{50} \)s produced by the two
techniques would agree. The optical technique emphasizes the importance of connections between
collectors more than the size of individual vessels, and due to the nature of cavitation propagation,
this is likely to accurately capture the dynamics of flow restriction. Although the slopes of
vulnerability curves produced by the cavitron tended to be shallower than those using the optical
method, this is may be explained by the smaller diameter branches used on the optical versus
cavitron technique. Small (3-6mm) diameter branches were used for the optical measurements to
ensure cavitations could be visualized from all depths in the stem. Larger diameter stem samples
used in the cavitron measurements are likely to incorporate more than one year of growth in the
sampled branch, particularly considering the slow growth of many of the conifer species used here
for comparison. Thus the cavitron curves reflect the integrated vulnerability of a much larger sample
of tracheids than the < one year old stems measured by the optical method, likely leading to a
shallower slope (Torres Ruiz et al., 2016).

A significant discrepancy between \( P_{50} \) in optical and centrifuge methods was only observed in stems
of the angiosperm Rosmarinus. Although both methods recorded extremely high cavitation
resistance in this species, the cavitron produced a more negative \( P_{50} \). Further examination of this
species and other highly resistant angiosperms will be needed to determine whether this
disagreement is due to artefacts or some systematic bias of one of the two methods. One possible
contributing factor is the long travel time from Hobart to France prior to measurement of this
individual. Samples of the same species measured locally with the cavitron yielded values much
closer to the OV value (Herve Cochard, pers. comm). This very resistant end of the vulnerability
spectrum is of particular interest as it appears as a critical adaptation in both conifer (Larter et al.;
Brodribb et al., 2014) and angiosperm (Blackman et al., 2012) tree species inhabiting semi-arid
woodland.
The success of the optical method in providing a time resolved map of cavitation in water stressed stems, while yielding an accurate measure of vulnerability in terms of $P_{50}$, opens the door to new applications. The simplicity and low cost of the technique makes it highly appealing for ecological and genetic research where large sample sizes are required. In addition the technique provides a means of viewing cavitation in tissues that have been difficult to measure. Flowers have recently been successfully measured using the optical method to show embolism relative to leaves in herbs and woody species (Zhang and Brodribb, 2017), while roots present an obvious future target. The optical method is ideally suited to explore how cavitation moves within and between plant tissues as water stress intensifies, and has the potential to provide an integrated view of cavitation in major plant organs as cavitation propagates within an individual.

MATERIALS AND METHODS

Plant Material

Thirteen species of conifers from four conifer families (Table I) were sampled from a potted conifer collection growing in glasshouses at the University of Tasmania. All plants were >10 years old and were growing in 20L pots under well watered conditions in partially open glasshouses such that light and temperature were close to ambient conditions in Hobart (Australia). Samples for centrifuge analysis were collected and measured in 2012 while samples for optical analysis were made in 2016 on the same individuals or clones. All species were represented by three replicates collected as cuttings from different individuals in the wild, or wild collected seeds. In addition we collected two angiosperms with contrasting water stress tolerance to represent opposite ends of the angiosperm vulnerability spectrum, but which had relatively short vessels such that they could be measured using the centrifuge technique. These two species (Rosmarinus officinalis and Betula pendula) were both collected at the end of a wet spring (2016) from single garden plants in Hobart.

Cavitron stem vulnerability

We carried out measurements on one or two branches from three to 16 trees per species. Transpiration losses were prevented by removing the needles or leaves immediately after sampling and wrapping the branches in moist paper to keep them humid and cool (5°C) until the measurement of embolism resistance (within three weeks of sampling). All samples were sent via an international express shipping company to France within three days. Vulnerability to drought-induced embolism was then determined at the Caviplace (University of Bordeaux, Talence, France; http://sylvain-delzon.com/caviplace) with the Cavitron technique (Cochard, 2002; Cochard et al., 2005). The bark was removed from conifer branches before sampling, to prevent resin contamination, and all branches were recut with a razor blade, under water, to a standard length of 0.27 m. The percentage loss of conductance (PLC) was determined at different speeds (i.e. different xylem pressures) to obtain a vulnerability curve for each sample. These vulnerability curves show the percentage loss of xylem conductance as a function of xylem pressure (see Delzon et al., 2010 for details). For each branch, the relationship between PLC and xylem water pressure was fitted with the following sigmoidal equation (Pammenter and Van der Willigen, 1998):
where $P_{50}$ (MPa) is the xylem pressure inducing a 50% loss of conductivity and $S$ (% MPa$^{-1}$) is the slope of the vulnerability curve at the inflection point. Mean values of embolism vulnerability parameters ($P_{50}$ and $S$) correspond to the average values of three to 16 samples per species.

Additionally, we used our VCs to calculate P12 and P88, which are respectively the 12% and 88% loss of hydraulic conductivity. P12 and P88 are physiologically significant indexes because they are thought to respectively reflect the initial air-entry tension producing embolisms and the irreversible death-inducing xylem tension (Urli et al., 2013).

### Optical stem vulnerability

The same individuals or clones of trees collected in 2012 for cavitron determination of $P_{50}$ were revisited and sampled using the optical vulnerability method. Branches in the order of 1m long were cut from trees early in the morning and transferred in plastic bags to the laboratory about 50m away. Branches were generally allowed to equilibrate in moist plastic bags in the dark for a period of 60 minutes to ensure stomata were closed before preparing the stem for imaging. The optical method cannot quantify existing embolism in the wood and is only able to measure new cavitations. For this reason great care was taken to ensure that samples were not exposed to any form of water stress or freezing in the months before measurement.

A stem psychrometer (ICT Australia) was fitted as close as possible to the region of stem being scanned for embolism formation. In fitting the psychrometer a small square of bark was removed avoiding damage to the wood. The psychrometer was partially insulated with polystyrene and set to log leaf water potential every 10 minutes. The cooling time for the psychrometer was increased from 5s to 30s as stems dried, ensuring a stable reading of the wet-bulb temperature. Reference leaf water potentials were taken during the drying period using a Scholander pressure chamber, to ensure that leaf and stem water potentials were equilibrated, as would be expected due to stomatal closure prior to the commencement of cavitation (Brodribb and Holbrook, 2003). However, after stem cavitation had begun Scholander and psychrometer values often tended to diverge as would be expected due to hydraulic disconnection between leaves and stems.

A stem approximately 3-6mm in diameter and approximately 80-120 cm in length for conifers or 1-2m in length for the angiosperms, was selected for scanning. Branches that were actively elongating or expanding leaves were avoided to be sure that the xylem was mature (non-living). The depth of xylem that could be reliably visualized for cavitation was approximately 1mm, so a selection of stems were sectioned before-hand to determine the approximate branch thickness that would yield 1mm of xylem above the pith. A leafless region of the stem, approximately 15mm in length was prepared so that xylem on one side of the pith could be imaged. A region of bark approximately 15-20mm in length was carefully removed from one side of the stem to expose the underlying wood without causing damage the xylem. The easiest way of doing this was to run two parallel axial cuts along the bark either side of the desired window, avoiding damage to the underlying xylem, and to use a needle or fingernails to peel the bark gently back from the cuts. Once a window was created, it was firmly secured either onto a flatbed scanner (Perfection 800, Epson) or a microscope stage (Leica M205) using padded clamps to ensure no movement of the sample during drying. Once secured, the

$$PLC = \frac{100}{1 + e^{\left(\frac{S}{25(P_{i} - P_{50})}\right)}}$$
scanner or camera was set to capture images at a rate of one per minute, and the sample left to dry slowly until cavitations were no longer recorded (typically in the order of 48-120 hours). During drying, the target region of the stem was mostly darkened except for the light of the microscope (a ring illumination using LED lighting) or scanner. The rest of the stem was exposed to laboratory lighting, and ambient conditions of 22°C and 55% RH. In the case of the scanner, images were collected in normal reflective mode rather than the transmission mode used for leaves. Samples were allowed to dry until no further cavitations could be seen in the xylem for a period of 12 hours. In some samples, a thin layer of hydrogel (Tensive Gel, Parker USA) was applied to the exposed xylem surface to improve light transmission and reduce evaporation from the surface. This had no appreciable effect on the value of $P_{50}$ when compared between samples (unpublished data) but care was necessary to avoid reflections of movements as the gel shrinks during the drying process.

Once completed, image sequences were analysed to identify cavitation, which was easily seen as changes in the reflection of the exposed xylem. Analysis by image difference using ImageJ (NIH), was carried out by are subtracting successive images to reveal fast changes in contrast produced by cavitation. These rapid changes were easily identified in image subtractions, and could be filtered from slow movements caused by drying. Thresholding of image differences allowed automated counting of cavitation events using the “analyze stack” function in ImageJ. Full details including an overview of the technique, image processing as well as scripts to guide image capture and analysis are available at http://www.opensourceov.org.

A time-resolved count of cavitations in each stem, quantified as a number of pixels per event during stem drying was compiled and this was converted to a % of total pixels cavitated. The psychrometer output was then used to determine a fitted function that described the change in stem water potential over time. Typically this was a linear function once stomata were closed, but occasionally polynomial functions were fitted to account for variation in the slope $\frac{d\psi_{stem}}{dt}$. Combining the cavitation count with the function describing $\frac{d\psi_{stem}}{dt}$ allowed the cumulative number of cavitations to be expressed as a function of $\psi_{stem}$. The $P_{50}$ for each sample stem was taken directly from this plot. One value of $P_{50}$ was measured for each of three stems, allowing a mean and SD to be presented for each species.

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Table I- Species list

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<thead>
<tr>
<th>Species</th>
<th>Family</th>
<th>Class</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Agathis robusta</em> (C.Moore ex F.Muell.) F.M.Bailey</td>
<td>Araucariaceae</td>
<td>Conifer</td>
</tr>
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<td><em>Araucaria bidwillii</em> Hook.</td>
<td>Araucariaceae</td>
<td>Conifer</td>
</tr>
<tr>
<td><em>Araucaria cunninghamii</em> Mudie.</td>
<td>Araucariaceae</td>
<td>Conifer</td>
</tr>
<tr>
<td><em>Wollemia nobilis</em> W.G.Jones, K.D.Hill &amp; J.M.Allen</td>
<td>Araucariaceae</td>
<td>Conifer</td>
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<td><em>Diselma archeri</em> Hook.f.</td>
<td>Cupressacae</td>
<td>Conifer</td>
</tr>
<tr>
<td><em>Acmopyle pancheri</em> (Brongn. &amp; Gris) Pilg.</td>
<td>Podocarpaceae</td>
<td>Conifer</td>
</tr>
<tr>
<td><em>Afrocarpus falcatus</em> (Thunb.) C.N.Page</td>
<td>Podocarpaceae</td>
<td>Conifer</td>
</tr>
<tr>
<td><em>Dacrycarpus imbricatus</em> (Blume) de Laub.</td>
<td>Podocarpaceae</td>
<td>Conifer</td>
</tr>
<tr>
<td><em>Lagarostrobos franklinii</em> (Hook.f.) Quinn</td>
<td>Podocarpaceae</td>
<td>Conifer</td>
</tr>
<tr>
<td><em>Phyllocladus aspleniifolius</em> (Labill.) Hook.f.</td>
<td>Podocarpaceae</td>
<td>Conifer</td>
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<tr>
<td><em>Prumnopitys ladei</em> (F.M.Bailey) de Laub.</td>
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<tr>
<td><em>Retrophyllum comptonii</em> (J.Buchholz) C.N.Page</td>
<td>Podocarpaceae</td>
<td>Conifer</td>
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<tr>
<td><em>Retrophyllum rosiglosii</em> (Pilg.) C.N.Page</td>
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<td>Conifer</td>
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<tr>
<td><em>Betula pendula</em> Roth</td>
<td>Betulacae</td>
<td>Angiosperm</td>
</tr>
<tr>
<td><em>Rosmarinus officinalis</em> L.</td>
<td>Lamiaceae</td>
<td>Angiosperm</td>
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</table>
FIGURE CAPTIONS

Figure 1. A- Cumulative area of cavitated xylem in a sample stem of *Callitris rhomboidea* is shown to increase rapidly approximately 1 day after a hydrated branch was excised (time zero) and allowed to dry. After a rapid rise in cavitation the rate of new xylem cavitated (quantified as number of pixels) falls back to zero approximately 3 days after excision. The insert graph shows that the size of newly cavitated regions visualized in the stem reaches a maximum during the steepest part of the curve (insert). During this period, very large blocks of tracheids were cavitating in the 2 minute interval between scans. B- Cumulative area of cavitated xylem expressed as a function of stem water potential showing a classic sigmoidal vulnerability curve. C- A mosaic of colour maps showing the spatial progression of cavitation through time in this 20mm long branched sample, the same stem sample as B and C. Sequential blocks of 280 images have been stacked together (frame numbers shown at the lower portion of each tile), with cavitated pixels coloured according to the water potential at which cavitation occurred. In this sample the smaller branches proved to be more resistant to cavitation than the main branch.

Figure 2. Similar plots as in Fig.1 showing the progression of cavitation in a stem of the angiosperm *Rosmarinus officinalis*. Despite the extreme resistance to cavitation in this stem the vulnerability curve shows a very steep transition from 12 to 88% cavitation. The reason for this steep transition can be clearly seen as due to a number of large and long cavitations between frames 279 and 333.

Figure 3. A comparison of vulnerability curve shape produced by the cavitron (black circles) and the optical method using branches from the same three individuals of the conifer *Lagarostrobus franklinii*. Although the mean $P_{50}$ is very similar in both species, the slope of the curves between 12% and 88% were steeper using the optical method.

Figure 4. Mean $P_{50}$ (±sd) for stems of the same individuals measured with the optical and cavitron methods. Very close agreement was found in the conifer sample between methods (regression slope 0.98; $r^2=0.93$). Among the two angiosperms sampled, good agreement was found in one species, while the cavitron method produced a more negative $P_{50}$ in the second. Slopes produced by the two techniques (insert graph) were correlated ($r^2=0.35$; p<0.05), but the optical technique produced a steeper slope in 14/16 species (1:1 shown as dotted line in each plot).


Brodribb TJ, McAdam SA, Jordan GI, Martins SC (2014) Conifer species adapt to low-rainfall climates by following one of two divergent pathways. Proceedings of the National Academy of Sciences 111: 14489-14493


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Figure 2. Similar plots as in Fig. 1 showing the progression of cavitation in a stem of the angiosperm *Rosmarinus officinalis*. Despite the extreme resistance to cavitation in this stem the vulnerability curve shows a very steep transition from 12 to 88% cavitation. The reason for this steep transition can be clearly seen as due to a number of large and long cavitations between frames 279 and 333.
Figure 3. A comparison of vulnerability curve shape produced by the cavitron (black circles) and the optical method using branches from the same three individuals of the conifer *Lagarostrobus franklinii*. Although the mean P50 is very similar in both species, the slope of the curves between 12% and 88% were steeper using the optical method.
Figure 4. Mean P50 (±sd) for stems of the same individuals measured with the optical and cavitron methods. Very close agreement was found in the conifer sample between methods (regression slope 0.98; r² = 0.93). Among the two angiosperms sampled, good agreement was found in one species, while the cavitron method produced a more negative P50 in the second. Slopes produced by the two techniques (insert graph) were correlated (r²=0.35; p<0.05), but the optical technique produced a steeper slope in 14/16 species (1:1 shown as dotted line in each plot).


Larter M, Pfautsch S, Domec J-C, Trueba S, Nagalingum N, Delzon S Aridity drove the evolution of extreme embolism resistance and the radiation of conifer genus Callitris. New Phytologist: n/a-n/a


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