RESEARCH PAPER

Limitation of the Cavitron technique by conifer pit aspiration

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Abstract

The Cavitron technique facilitates time and material saving for vulnerability analysis. The use of rotors with small diameters leads to high water pressure gradients (ΔP) across samples, which may cause pit aspiration in conifers. In this study, the effect of pit aspiration on Cavitron measurements was analysed and a modified ‘conifer method’ was tested which avoids critical (i.e. pit aspiration inducing) ΔP. Four conifer species were used (Juniperus communis, Picea abies, Pinus sylvestris, and Larix decidua) for vulnerability analysis based on the standard Cavitron technique and the conifer method. In addition, ΔP thresholds for pit aspiration were determined and water extraction curves were constructed. Vulnerability curves obtained with the standard method showed generally a less negative P for the induction of embolism than curves of the conifer method. Differences were species-specific with the smallest effects in Juniperus. Larix showed the most pronounced shifts in \( P_{50} \) (pressure at 50% loss of conductivity) between the standard (–1.5 MPa) and the conifer (–3.5 MPa) methods. Pit aspiration occurred at the lowest ΔP in Larix and at the highest in Juniperus. Accordingly, at a spinning velocity inducing \( P_{50} \), ΔP caused only a 4% loss of conductivity induced by pit aspiration in Juniperus, but about 60% in Larix. Water extraction curves were similar to vulnerability curves indicating that spinning itself did not affect pits. Conifer pit aspiration can have major influences on Cavitron measurements and lead to an overestimation of vulnerability thresholds when a small rotor is used. Thus, the conifer method presented here enables correct vulnerability analysis by avoiding artificial conductivity losses.

Key words: Embolism, hydraulic conductance, pressure difference, vulnerability, water extraction, xylem pressure.

Introduction

Vulnerability analyses substantially contribute to our knowledge of plant water relations, and, according to Cochard et al. (2005), they may be increasingly applied in ecological studies as well as in breeding programmes (Mayr et al., 2006; Cochard et al., 2008; McDowell et al., 2008; West et al., 2008; Beikircher and Mayr, 2009; Beikircher et al., 2010). A vulnerability curve shows a species’ resistance to drought-induced embolism. It is a plot of per cent loss of hydraulic conductivity (PLC) versus the xylem pressure (\( P \); Tyree and Sperry, 1988, 1989; Tyree and Zimmermann, 2002). Until recently, they have mainly been obtained via hydraulic measurements after inducing a defined \( P \) by dehydration (Sperry et al., 1988; Mayr et al., 2002; Tyree and Zimmermann, 2002), air-injection (Cochard et al., 1992; Sperry and Saliendra, 1994), or centrifugation (Holbrook et al., 1995; Pockman et al., 1995; Alder et al., 1997). By contrast with these methods, the Cavitron technique allows the induction and measurement of PLC in rapid succession while samples are enclosed in the centrifuge, and thus enables the construction of an entire vulnerability curve within a short space of time (Cochard, 2002; Cochard et al., 2005, 2007).

Cavitron technique (Cochard, 2002) is based on the use of the centrifugal force to generate (i) negative xylem pressures (\( P \); MPa) in the sample to induce embolism and (ii) a positive water pressure difference (ΔP; MPa) across...
the case of the conifer method. Therefore, samples are fixed in a custom-built rotor, and sample ends are placed in water reservoirs with outlets at different distances from the rotation axis. The minimum (i.e. most negative) \( P \) in the sample is computed as

\[
P = -0.25 \rho \omega^2 (R^2 + (R-r)^2)
\]

where \( \rho \) is the density of water (kg m\(^{-3}\)), \( \omega \) the angular velocity (rad s\(^{-1}\)), \( R \) the distance (m) from the rotational axis to the downstream reservoir, and \( r \) the difference in water levels (m) between the downstream and the upstream reservoir. From this formula it is evident that the smaller the rotor diameter the higher the \( \omega \) required to obtain a defined \( P \). The high \( \omega \), in turn, strongly influences \( \Delta P \). As the water pressure in the upstream reservoir changes during measurements the time-averaged water pressure difference across the sample (\( \Delta P^* \)) is calculated as

\[
\Delta P^* = 1/6 \rho \omega^2 \left( 3R^2(r_1-r_2) + (R-r_1)^3 - (R-r_2)^3 \right) / (r_1 - r_2)
\]

whereby \( r_1 \) and \( r_2 \) are the differences in water levels between the reservoirs at the times \( t_1 \) and \( t_2 \), respectively. Thus, the smaller a rotor, the higher are the required \( \omega \) (see above) and, in consequence, the higher is \( \Delta P \) (Fig. 1) — unless the water levels in the reservoirs are balanced. In that case \( \Delta P \) is 0 and does not depend on \( \omega \) or \( P \).

To understand the background of the Cavitron technique as well as the interrelationship with pit aspiration presented in this study, it is essential to distinguish carefully between the parameters \( P \) and \( \Delta P \) and their impact on xylem function. The former is a measure of the availability of water at a certain point in a plant or a sample and is physically equivalent to a tension (Tyree and Ewers, 1991). When it exceeds species-specific thresholds, embolism occurs due to air-seeding (Sperry and Tyree, 1990; Tyree and Ewers, 1991; Tyree and Zimmermann, 2002; Cochard et al., 2009). The \( \Delta P \) is the difference in water pressure between two points in a plant or sample, respectively. High \( \Delta P \) is known to cause pit aspiration in conifers, whereby the pressure drop across each pit membrane is relevant (Sperry and Tyree, 1990). The inter-tracheid pits of conifers typically have a torus-margo pit membrane. This architecture allows water to flow between two water-filled tracheids as water passes through the thin and porous margo driven by moderate \( \Delta P \). When a tracheid is filled with air (embolism), the pressure difference between the water-filled (negative pressure) and the air-filled conduit (ambient pressure) increases considerably, the thickened torus is aspirated against the pit opening and seals the air-filled tracheid as long as the pressure difference is lower than the species-specific vulnerability threshold for air-seeding (Chapman et al., 1977; Tyree and Zimmermann, 2002; Hacke et al., 2004; Domec et al., 2006, 2008; Hölttä et al., 2007). Theoretically, this sealing of the tracheid can also happen between two water-filled tracheids when \( \Delta P \) is high enough. In nature this may rarely be the case as the pressure difference over the pit membrane due to normal transpiration is far too small to cause pit aspiration (Gregory and Petty, 1973; Chapman et al., 1977; Hölttä et al., 2007). Although Hammel (1967) and Robson et al. (1988) found that, at the boundary between frozen and unfrozen tracheids, the pressure difference can be sufficient to cause pit aspiration.

So far, the rotor diameter has only been known to influence cavitron measurements of angiosperms negatively (Choat et al., 2010; Cochard et al., 2010a): When mean conduit length exceeds the sample length, water in cut conduits may be thrown out by the rotational motion (Alder et al., 1997) or, due to the lack of filtration of microscopic particles or air bubbles at the pit membranes, embolism may be induced, causing a strongly altered vulnerability curve (Cochard et al., 2005; but also see Li et al., 2008). This should not concern measurements on conifers, as tracheids are mostly only few millimetres long (Tyree and Zimmermann, 2002). Accordingly, Cochard et al. (2005) and Li et al. (2008) demonstrated that vulnerability curves of conifers obtained with conventional methods and the Cavitron technique corresponded well, which would also indicate that pit aspiration does not play any role in Cavitron measurements. However, they used a rotor of about 280 mm in diameter which might be sufficient to avoid critical \( \Delta P \). It is hypothesized that the use of smaller rotors can be problematic when high \( \Delta P \) causes pit closure and, consequently, conductivity losses independent of embolism formation. Species-specific differences in the sensitivity to \( \Delta P \) effects were also expected.

In this study, a small rotor was used with a diameter of 150 mm to measure vulnerability to drought-induced
embolism. The influence of $\Delta P$ on vulnerability curves was analysed, based on two different measurement approaches. First, measurements were carried out at stepwise increasing $\sigma$ and, consequently, increasing $\Delta P$ according to measurement protocols in Cochard (2002), Cochard et al. (2005, 2008), and Li et al. (2008). This method is referred to as the ‘standard method’. Second, $\Delta P$ was kept low by a reduction of $\sigma$ during conductivity measurements and balancing the water levels in the reservoirs before increasing $\sigma$ to attain the desired $P$ (see Materials and methods section). This measuring method is referred to as the ‘conifer method’.

Vulnerability curves obtained with both methods were compared on four conifer species, and $\Delta P$ thresholds determined by measurement of hydraulic conductivity at increasing $\Delta P$. Furthermore, water extraction curves were made with the centrifuge to test if spinning itself affect pits by induced gravity forces.

Materials and methods

Plant material

All measurements were made on the following four conifers: Juniperus communis L., Picea abies (L.) Karsten, Pinus sylvestris L., and Larix decidua Mill. The latter three species were harvested in a natural forest in Natters (47°14’ N, 11°22’ E; 783 m a.s.l., Tyrol, Austria, Central European Alps), while J. communis was growing in the Botanical Garden of Innsbruck (47°16’ N, 11 24’ E; 754 m a.s.l., Tyrol, Austria, Central European Alps). Branches were cut from plants, wrapped in dark plastic bags and transported to the laboratory. There, they were re-cut twice under water and saturated for 24 h.

Vulnerability analyses

Vulnerability analyses were done by plotting the fractional (%) loss of conductivity versus the xylem pressure ($P$). Curves were fitted using an exponential sigmoidal equation (equation 3) given in Pammenter and Vander Willigen (1998):

$$PLC = 100(1+\exp(a(P-P_{06})))$$

where $PLC$ is the per cent loss of conductivity, $P$ is the corresponding xylem pressure (Pa), and $a$ is related to the slope of the curve. $P_{06}$ is the $P$ value corresponding to a 50% loss of conductivity. In addition, $P$ at 10% PLC ($P_{10}$) and 90% PLC ($P_{90}$) were determined. PLC was calculated from the ratio of actual (after inducing a given $P$) to maximum (i.e. first measurement at $-0.25$ MPa) hydraulic conductance ($k$).

Setting of $P$ and measurement of $k$ were done with the Cavitron technique (Cochard, 2002; Cochard et al., 2005; Li et al., 2008) using two different measuring methods (see below). In either case, stem segments were fixed in a custom-built rotor designed by JS Sperry for a Sorvall RC-5 centrifuge (Thermo Fisher Scientific, Waltham, MA, USA; see Li et al., 2008, for details). The rotor was 150 mm in diameter (lumen), and outlets in the upstream and downstream reservoirs were at 1.6 and 0.8 cm, respectively, from the inner rotor wall. The desired $P$ was obtained by setting the rotational speed ($n_{rot}$, rpm) according to equation 4

$$n_{rot} = \frac{P/(0.5p0.1047198*10^{-6})^2}{0.5}$$

where $p$ is the density of water (kg m$^{-3}$), 0.1047198 is the conversion factor of angular velocity ($\sigma$; rad s$^{-1}$) in $n_{rot}$ (1 s$^{-1}$) and $r$ (m) is the radius of the rotor less the distance to the outlet of the downstream reservoir. In the case of the conifer method, where the water reservoirs were emptied, $r$ was the radius of the rotor less the wall thickness of the reservoirs (3 mm).

For conductivity measurements, reservoirs were filled with distilled, filtered (0.22 $\mu$m), and degassed water containing CaCl$_2$ (1 mmol) and KCl (10 mmol). The moving water meniscus was observed using a high resolution camera (Motic MC 2000, Motic China group Co., Ltd.) fixed on a centring telescope (U-CT30, Olympus; Olympus Austria Corporation, Vienna) with scale reticle. This system was calibrated to allow measurement of the actual distance travelled in metres. Whenever the meniscus passed a scale unit, time was registered and the flow rate ($F^*$) calculated as

$$F^* = \frac{\Delta r}{\Delta t}$$

where $\Delta r$ is the actual distance travelled (m), $\Delta t$ the time therefore required, and $s$ is a conversion factor to calculate the shifted volume per distance travelled by the meniscus. Hydraulic conductance ($k$: m$^2$ s$^{-1}$ MPa$^{-1}$) was then calculated using $F^*$ and mean water pressure difference across the sample ($\Delta P^*$; see introduction, equation 2) as

$$k = \frac{F^*}{\Delta P^*}$$

Two different approaches were chosen for Cavitron measurements:

Standard method

For measurements with the standard method the rotational speed ($n_{rot}$) and thus the angular velocity $\sigma$ was set to the target pressure (according to equation 4) and maintained constant for 1 min. This time was assumed to be sufficient to generate the maximum level of cavitation as it has been reported that cavitation is forming within a few seconds after exposure to the respective $P$ (Cochard et al., 2005), and (ii) during the water extraction analyses (see below) it has been observed that water was released only within the first seconds after increasing $P$ and that the meniscus was stable afterwards. After setting $P$, the flow rate ($F^*$) was measured and hydraulic conductance ($k$) was calculated as described above (see equations 5 and 6). The first measurement was done at $-0.25$ MPa and the following at successively lower pressures. In the case that the meniscus was moving too rapidly at high speed, after setting $P$ $n_{rot}$ was lowered for conductivity measurements. However, in contrast to the conifer method (see below), water levels in the reservoirs were never balanced before increasing $n_{rot}$ and thus high $\Delta P$ values were achieved at low pressures. For example, at a $P$ of $-6$ MPa, $\Delta P$ was 1.25 MPa (Fig. 1).

Conifer method

In the conifer method, $\Delta P$ was always kept low by (i) doing all $k$ measurements at $-0.25$ MPa and (ii) balancing the water levels before increasing $P$ as described in the following. First, $P$ was set to $-0.25$ MPa (equals $\Delta P$ of 0.05 MPa for our rotor diameter) and after 1 min, $k$ was measured in a similar way to the standard method. In contrast to the standard method, water levels in the reservoirs were balanced before $P$ was decreased to the next desired value. This was done either by waiting until the water levels were balanced due to the water flow itself, or (as conductivity was rather low in most samples and thus a lot of time, often more than 1 h, was required to balance the water levels) by stopping the centrifuge briefly. By stopping the centrifuge for a few seconds, reservoirs were emptied except for a small amount of water. This remaining water caused a 1 mm thick water film at the reservoir bottom during spinning so that sample ends were still submerged. Then, $n_{rot}$ was set to the next desired $P$ for 1 min. Due to the balanced water levels, $\Delta P$ was thus nearly zero regardless of $n_{rot}$ (also see Introduction). After increasing $P$, $n_{rot}$ was decreased again to $-0.25$ MPa before the reservoirs were refilled for the next conductivity measurement.
Thus, in the conifer method $\Delta P$ never exceeded 0.05 MPa. A problem with this measuring method could have arisen if conduits refilled during spinning at low $n_{rot}$. At least for our study this can be excluded, because $k$ was always constant during the measurement phase. Also Cochard (2002) did not observe a conductance recovery in *Laurus nobilis* when, after spinning at $-3.1$ MPa, samples were spun at $-0.8$ MPa for 4 h.

**Pressure–conductivity analyses**

To analyse at which $\Delta P$ pit aspiration occurs, about 15 cm long, debarked twigs were fixed in a pressure chamber (Model 1000 Pressure Chamber, PMS Instrument Company, Corvallis, OR, USA). The sample end in the chamber was positioned in a plastic bottle filled with CaCl$_2$/KC1 solution (see vulnerability analysis). The outer sample end was connected to a Pasteur pipette via a silicone tube. Flow rates were determined with a FC-connected balance (Sartorius BP61S, 0.0001 g precision, Sartorius AG, Göttingen, Germany) by weight registration every 10 s. Measurement pressure was stepwise increased, starting from 0.1 MPa until flow ceased. The applied pressure thereby corresponds to $\Delta P$ in Cavitron measurements. Specific hydraulic conductivity ($k_s$) was calculated as

\[ k_s = \frac{Q}{A \Delta P} \]  

where $Q$ is the volume flow rate (m$^3$ s$^{-1}$), $A$ is the xylem cross-sectional area (sapwood less heartwood; m$^2$), and $\Delta P$ in this case corresponds to the water pressure (Pa) applied with the pressure bomb.

Curves were fitted similar to vulnerability curves (equation 3) whereby $PLC$ was substituted by percentage loss of $k_s$. $\Delta P$ at 10, 50, and 90% loss of $k_s$ corresponded to 10, 50, and 90% pit aspiration ($\Delta P_{asp10}$, $\Delta P_{asp50}$, $\Delta P_{asp90}$).

This method may cause problems when the solution in the chamber is saturated with gas and, subsequently, gas comes out of the solution in the stem section protruding from the bomb. However, this was only observed at pressures where PLC was already up to 100%. Furthermore, this effect would be similar on all measured species and thus could not explain the variation shown in Fig. 3. Also, Cochard et al. (2010b, who did a similar experiment on *Fagus sylvatica*, observed no decrease in conductance.

**Critical sample length**

Critical sample length was defined as the minimum length at which less than 10% pit aspiration occurred at $P_{50}$. It was calculated by plotting $\Delta P$ versus $P$ for various diameters (as in Fig. 1) and subsequently inserting of the respective $\Delta P_{asp10}$ ($\delta$-value) in the linear equations obtained. The diameter at which the thereby calculated $\delta$-value corresponded to $P_{50}$ was taken as the critical sample length.

**Water extraction curves**

Water extraction curves (also see Cochard et al., 2010b) were made with the Cavitron. Therefore, two reservoirs with outlets at 1.6 cm from the bottom were used. Samples were first spun at $-0.25$ MPa until water levels were balanced and menisci overlapped. Then, $n_{rot}$ was increased successively to different $P$ and the respective distances travelled by the meniscus noted. For the calculation of $P$ (equation 1), $R$ was taken as the distance from the rotational axis to the actual water level. The percentage of extracted water was calculated as the ratio of actual (at a given $P$) to maximum distance travelled. Curves were fitted similar to vulnerability curves (equation 3) whereby $PLC$ was substituted by percentage extracted water. $\Delta P_{ex50}$ is the $\Delta P$ at 50% of total water extraction.

**Number of samples and statistics**

Vulnerability analyses, water extraction curves, and pressure–conductivity analyses were made on 4–10 samples, originating from at least three individual plants per species, and method. For conductivity measurements with the Cavitron at least five $k$ values were taken per measurement and means calculated. Differences between methods were tested with the Welch-test at a probability level of 5%.

**Results**

In all four conifers, vulnerability curves analysed with the standard and conifer methods, respectively, differed. Vulnerability thresholds, i.e. xylem pressure at 10, 50, and 90% loss of conductivity ($P_{10}$, $P_{50}$, $P_{90}$) were higher (i.e. less negative) when measured with the standard method (Table 1; Fig. 2). The greatest differences were found in *L. decidua*, where $P_{50}$ of the standard method was 2.08 MPa higher than that measured with the conifer method. Differences were lower in *P. sylvestris* and *P. abies* and lowest in *J. communis* (0.66 MPa; Table 1). In general, $P_{50}$ measured with the conifer method corresponded well to $P_{90}$ values obtained with conventional hydraulic methods (Table 1).

According to the findings of the vulnerability analysis, *L. decidua* was also most susceptible in the pressure–conductivity analysis. In this species, 10% pit aspiration occurred already at an applied water pressure of 0.32 MPa ($\Delta P_{asp10}$). Table 2 lists $\Delta P$ thresholds for pit aspiration and $\Delta P$ induced with the standard method at a $P$ causing 50% of embolism. Thereby $P_{50}$ was taken from curves obtained with the conifer method as these $P_{50}$ values were similar to the results of conventional methods and thus suggested to be close to the actual $P_{50}$. *J. communis* was the only species which showed the onset of pit aspiration ($\Delta P_{asp10}$) at a $\Delta P$ higher than the $\Delta P$ corresponding to 50% of embolism. In all other species, pit aspiration had already occurred at the $\Delta P$ where embolization took place ($\Delta P_{asp10}$ was lower than $\Delta P$ at $P_{50}$; see Table 2), and thus both pit aspiration and embolism formation contributed to conductivity losses. Accordingly, $\Delta P_{asp50}$ of *L. decidua* is less negative than its $\Delta P$ at $P_{50}$. In *P. sylvestris* both values are identical and in *P. abies*, $\Delta P_{asp50}$ is slightly above $\Delta P$ at $P_{50}$. In *J. communis*, $\Delta P_{asp50}$ was 1.25 MPa higher than $\Delta P$ at $P_{50}$. Critical sample length, i.e. minimum length at which less than 10% pit aspiration occurs at $P_{50}$ ranged from 116 mm for *L. decidua* to 324 mm for *L. nobilis* (Table 2).

Water extraction curves in all four species showed a course very similar to vulnerability curves (Figs 2, 4) and $50%$ water extraction ($P_{ex50}$) in all species was reached close to $P_{50}$ (Table 1). No indications for any direct influence of spinning, like pit aspiration at higher $P$, was substituted by percentage loss of $P_{10}$, $P_{50}$, $P_{90}$, $P_{50}$ was taken as the critical sample length, i.e. minimum length at which less than 10% pit aspiration occurred at $P_{50}$; see Table 2), and thus both pit aspiration and embolism formation contributed to conductivity losses. Accordingly, $\Delta P_{asp50}$ of *L. decidua* is less negative than its $\Delta P$ at $P_{50}$. In *P. sylvestris* both values are identical and in *P. abies*, $\Delta P_{asp50}$ is slightly above $\Delta P$ at $P_{50}$. In *J. communis*, $\Delta P_{asp50}$ was 1.25 MPa higher than $\Delta P$ at $P_{50}$. Critical sample length, i.e. minimum length at which less than 10% pit aspiration occurs at $P_{50}$ ranged from 116 mm for *J. communis* to 324 mm for *L. decidua* (Table 2).

Water extraction curves in all four species showed a course very similar to vulnerability curves (Figs 2, 4) and 50% water extraction ($P_{ex50}$) in all species was reached close to $P_{50}$ (Table 1). No indications for any direct influence of spinning, like pit aspiration at higher $P$, which would alter position and shape of the curve by slowing down the release of water, were observed.

**Discussion**

The main advantage of the Cavitron technique over other methods of vulnerability analysis lies in its time-efficiency and low material requirement: Embolism induction and conductivity measurements are achieved in parallel, and an
Table 1. Vulnerability curve parameters (parameter a, xylem pressure at 10, 50, and 90% loss of conductivity: P_{10}, P_{50}, P_{90}) and xylem pressure at 50% water extraction (P_{ex50}) of Juniperus communis, Picea abies, Pinus sylvestris, and Larix decidua.

Data (mean ± SE) show parameters of vulnerability curves measured with the standard and the conifer method and P_{SO} values measured with a rotor with 280 mm in diameter (published in Cochard, 2006) as well as conventional hydraulic methods (conventional; (1) Beikircher et al., 2010; (2) Mayr et al., 2006; (3) Cochard, 1992; (4) Poyatos et al., 2008). Asterisks indicate significant intraspecific differences between standard and conifer method (P<0.05, Welch test).

<table>
<thead>
<tr>
<th>Parameter a</th>
<th>Juniperus communis</th>
<th>Picea abies</th>
<th>Pinus sylvestris</th>
<th>Larix decidua</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard method</td>
<td>0.77±0.04</td>
<td>1.25±0.09</td>
<td>1.65±0.14</td>
<td>2.00±0.18</td>
</tr>
<tr>
<td>Conifer method</td>
<td>0.77±0.04</td>
<td>1.01±0.06*</td>
<td>2.64±0.32*</td>
<td>1.61±0.09</td>
</tr>
<tr>
<td>P_{10}</td>
<td>–2.10±0.22</td>
<td>–1.18±0.18</td>
<td>–0.83±0.17</td>
<td>–0.36±0.15</td>
</tr>
<tr>
<td>Conifer method</td>
<td>–2.78±0.20</td>
<td>–1.80±0.18*</td>
<td>–2.54±0.15*</td>
<td>–2.18±0.43*</td>
</tr>
<tr>
<td>P_{50}</td>
<td>–4.96±0.07</td>
<td>–2.93±0.06</td>
<td>–2.17±0.05</td>
<td>–1.46±0.05</td>
</tr>
<tr>
<td>Conifer method</td>
<td>–5.62±0.06*</td>
<td>–3.98±0.05*</td>
<td>–3.37±0.05*</td>
<td>–3.54±0.04*</td>
</tr>
<tr>
<td>280 mm rotor diameter</td>
<td>–6.37±0.18</td>
<td>–3.97±0.04</td>
<td>–3.20±0.02</td>
<td>–3.87±0.10</td>
</tr>
<tr>
<td>Conventional</td>
<td>–6.60±0.18</td>
<td>–3.39±0.03</td>
<td>–2.53</td>
<td>–2.78±0.11</td>
</tr>
<tr>
<td>P_{90}</td>
<td>–7.82±0.09</td>
<td>–4.68±0.07</td>
<td>–3.50±0.06</td>
<td>–2.56±0.05</td>
</tr>
<tr>
<td>Conifer method</td>
<td>–8.47±0.09*</td>
<td>–6.16±0.07*</td>
<td>–4.21±0.06*</td>
<td>–4.90±0.27*</td>
</tr>
<tr>
<td>P_{ex50}</td>
<td>–5.61±0.06</td>
<td>–4.08±0.02</td>
<td>–3.24±0.05</td>
<td>–3.93±0.06</td>
</tr>
</tbody>
</table>

The entire curve can be measured on a single sample independent of season and foliage (Cochard et al., 2005, 2007; Li et al., 2008). However, our study clearly demonstrates that pit aspiration can limit measurements on conifers and that rotor diameter may thus be critical.

According to the hypothesis formulated in the Introduction, vulnerability curves made with the standard method, where high ΔP can occur (see Materials and methods), were significantly shifted to less negative P compared to curves made with the conifer method (Fig. 2; Table 1). Effects of ΔP were found to be highly species-specific, with the most pronounced shift in vulnerability thresholds in L. decidua, and slightest differences in J. communis (Table 1; Fig. 2).

With the conifer method, it was possible to avoid high ΔP by equilibration of water contents in the water reservoirs. In all four species, xylem pressure at 50% loss of conductivity (P_{50}) measured with the conifer method was similar to the P_{50} measured using conventional hydraulic methods (Table 1).

The reason for shifts in vulnerability curves was pit aspiration. As demonstrated in our pressure–conductivity analysis (Fig. 3), pit aspiration in three out of the four species studied induced considerable conductivity losses at ΔP corresponding to P inducing embolism in the standard method (Table 2). At P_{50} (actual P_{50}, see Results), the ΔP induced by the 150 mm rotor caused pit aspiration resulting in conductivity losses of c. 60% in L. decidua and c. 40% in P. abies and P. sylvestris. Only in J. communis, pit aspiration contributed hardly to overall PLC (4% at P_{50}) and thus showed the smallest shift in vulnerability curves obtained with the standard and the conifer methods (Fig. 2; Table 1). Critical sample length (i.e., minimum sample length at which less than 10% pit aspiration occurs in the range of P_{50}) showed that a considerably bigger rotor would be required to achieve similarly good conformities between the vulnerability curves for the other measured species (Table 2). Accordingly, in the study of Cochard (2006) P_{50} values achieved with a bigger rotor (280 mm in diameter) correspond well to those achieved with conventional methods (see Table 2).

The differences are very obvious when differences between ΔP_{asp50} and ΔP at P_{50} are compared. Table 2 clearly indicates that the overlapping of the two processes, embolism formation and pit aspiration, was most relevant in L. decidua, followed by P. sylvestris and P. abies and hardly relevant in J. communis. Pit aspiration upon experimentally increased pressure gradients was already demonstrated by Sperry and Tyree (1990). The authors found a decrease in hydraulic conductivity (at a ΔP less than –1 MPa) only in conifers and the effect was reversed when the pressure was decreased again. They hypothesized that observed differences between species may be due to the flexibility of the inter-tracheid pit membrane. The lower the flexibility the higher is the pressure required to get the torus in a sealing position. On the other hand, a high rigidity may also hold the torus in a sealing position even at large pressure differences and thus account for a high resistance to air-seeding (but also see Cochard et al., 2009). Accordingly, species of the genus Juniperus, which has an extraordinarily high resistance to drought-induced embolism (Pockman and Sperry, 2000; Martinez-Vilalta et al., 2004; Mayr et al., 2006; Willson and Jackson, 2006; West et al., 2007; Beikircher and Mayr, 2008; Willson et al., 2008) showed pit aspiration at lowest P in the study of Sperry and Tyree (1990) and this study, respectively. Besides interspecific differences in susceptibility to pit aspiration, variation...
between early- and late-wood pits may also play a role (Gregory and Petty, 1973; Bolton and Petty, 1977; Cochard et al., 2009). Furthermore, pit aspiration is not only of interest in experimental approaches, but also in applied sciences and to the timber industry as it influences the permeability of sapwood (Comstock and Côté, 1968).

An attempt was made to compute the combined effect of $P$-induced embolism formation and $\Delta P$-induced pit aspiration from vulnerability curves (conifer method) and pit aspiration analysis. Interestingly, the calculated effects (data not shown) were much lower than the measured differences (i.e. the standard vulnerability curves deviated much more from the curves obtained with the conifer method as indicated by pit aspiration thresholds (Figs 2, 3), which indicates that both processes were interrelated and, in consequence, effects on conductivity amplified. It is suggested that cavitation of a tracheid might lead to an increase of $\Delta P$ at surrounding pits and induce their aspiration.

Table 2. Parameters of the pressure–conductivity analysis (parameter a, water pressure difference causing 10, 50, and 90% pit aspiration: $\Delta P_{\text{asp}10}$, $\Delta P_{\text{asp}50}$, $\Delta P_{\text{asp}90}$), water pressure difference ($\Delta P$) obtained with the standard method at $P_{50}$ of the conifer method and critical sample length of *Juniperus communis*, *Picea abies*, *Pinus sylvestris*, and *Larix decidua*; mean ± SE.

<table>
<thead>
<tr>
<th></th>
<th><em>Juniperus communis</em></th>
<th><em>Picea abies</em></th>
<th><em>Pinus sylvestris</em></th>
<th><em>Larix decidua</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter a</td>
<td>-2.49 ± 0.20</td>
<td>-6.12 ± 0.84</td>
<td>-11.36 ± 1.20</td>
<td>-6.64 ± 0.58</td>
</tr>
<tr>
<td>$\Delta P_{\text{asp}10}$</td>
<td>1.52 ± 0.11</td>
<td>0.59 ± 0.10</td>
<td>0.50 ± 0.03</td>
<td>0.32 ± 0.04</td>
</tr>
<tr>
<td>$\Delta P_{\text{asp}50}$</td>
<td>2.40 ± 0.04</td>
<td>0.95 ± 0.05</td>
<td>0.69 ± 0.01</td>
<td>0.65 ± 0.01</td>
</tr>
<tr>
<td>$\Delta P_{\text{asp}90}$</td>
<td>3.29 ± 0.04</td>
<td>1.31 ± 0.00</td>
<td>0.88 ± 0.01</td>
<td>0.98 ± 0.02</td>
</tr>
<tr>
<td>$\Delta P$ at $P_{50}$</td>
<td>1.15 0.81</td>
<td>0.69 0.73</td>
<td>204 206</td>
<td>324</td>
</tr>
<tr>
<td>Critical sample length (mm)</td>
<td>116 204 206 324</td>
<td>3390</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Furthermore, pit margo elasticity is known to be strongly increased after pressure treatments (Cochard et al., 2010b), which might also influence xylem vulnerability to air seeding. A rapid increase in $\Delta P$ may cause high local pressure differences and pit aspiration before pressures are equilibrated within the sample. It is also not known if a pit, when already aspirated to one pit opening by $\Delta P$, can prevent air seeding when sealing to the other pit opening would be required.

In our centrifuge experiments, pits might also have been influenced by the spinning procedure itself. It was unclear if the centrifugal forces can cause displacement or even a sealing of the torus. In Cavitron vulnerability measurements (Fig. 2), embolism and pit aspiration influence the measured variable (hydraulic conductivity) and can thus not be distinguished. We therefore made water extraction experiments, where embolism was estimated by the release of water from broken water columns (Fig. 4). As these measurements were made at zero $\Delta P$, $\Delta P$ related pit aspiration played no role. Any other change at the pits would be obvious in differences of shape or position in the water extraction curve compared with the vulnerability curve (with the conifer method or obtained by conventional methods). Figure 4 clearly indicates that the direct effects of spinning forces on pit structures did not play a role. Also Cochard et al. (2010b) reported a close correlation between $P_{50}$ and $P$ at 50% water extraction. Thus, water extraction curves may even be a simplified alternative to vulnerability measurements as long as the amount of extracted water caused by embolism linearly correlates to conductivity losses.

In summary, use of a small rotor, although advantageous for small samples, may be unsuitable for vulnerability analysis of conifers. The high rotational speeds required to reach low $P$ can cause high $\Delta P$ and, in consequence, pit aspiration, when water levels in the reservoirs are unbalanced. In large rotors, spinning velocities and $\Delta P$ are lower. Thus, the effect of pit aspiration overall negatively correlates with rotor size but the remarkably species-specific differences in pit aspiration thresholds have to be considered. We recommend use of the conifer method for Cavitron measurements on conifers unless test experiments prove that rotor diameter and species-specific thresholds for pit aspiration allow the use of the faster standard Cavitron method.

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References


![Fig. 4. Per cent extracted water at different xylem pressure ($P$; MPa) in the Cavitron. Vertical lines show $P$ at which 50% water was extracted ($P_{\text{ex}50}$).](image-url)


