Heterogeneous distribution of pectin epitopes and calcium in different pit types of four angiosperm species

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Summary

- Intervessel pits act as safety valves that prevent the spread of xylem embolism. Pectin–calcium crosslinks within the pit membrane have been proposed to affect xylem vulnerability to cavitation. However, as the chemical composition of pit membranes is poorly understood, this hypothesis has not been verified.
- Using electron microscopy, immunolabeling, an antimonate precipitation technique, and ruthenium red staining, we studied the distribution of selected polysaccharides and calcium in the pit membranes of four angiosperm tree species. We tested whether shifts in xylem vulnerability resulting from perfusion of stems with a calcium chelating agent corresponded with the distribution of pectic homogalacturonans (HG) and/or calcium within interconduit pit membranes.
- No HG were detected in the main part of intervessel pit membranes, but were consistently found in the marginal membrane region known as the annulus. Calcium colocalized with HG in the annulus. In contrast to intervessel pits, the membrane of vessel-ray pits showed a high pectin content.
- The presence of two distinct chemical domains, the annulus and the actual pit membrane, can have substantial implications for pit membrane functioning. We propose that the annulus could affect the observed shift in xylem vulnerability after calcium removal by allowing increased pit membrane deflection.

Introduction

Water and nutrients move through a complex network of xylem conduits that, when mature, are dead and void of cellular content. Water movement is driven by a gradient in negative pressure that is a result of evaporation from the tiny menisci localized in the cell walls of leaf parenchyma. Consequently, xylem sap is in a metastable state and hydraulic failure can readily occur (Tyree & Zimmermann, 2002). As the xylem pressure decreases owing to increased evaporative demand or lack of soil moisture, there is a higher risk of water columns essentially ‘breaking’ and xylem conduits becoming air-filled (i.e. embolized). Embolized conduits are temporarily or permanently disconnected from the transpiration stream and hydraulic conductivity ($k_h$) decreases.

Interconduit pits connect adjacent conduits to permit water flow while preventing the spread of air from embolized conduits to adjacent functional ones (Sperry & Tyree, 1988; Sperry & Hacke, 2004; Choat et al., 2008). According to the air-seeding hypothesis, microporous pit membranes will support an air—water interface until the pressure difference across the membrane overcomes the capillary forces (Sperry et al., 1996). The air-seeding threshold is inversely related to the pore diameter, such that higher pit membrane porosity will result in more vulnerable xylem (Jarbeau et al., 1995; Choat et al., 2008). A consequence of the role of interconduit pits as safety valves is that they provide a significant constraint to water flow. In both conifer tracheids and angiosperm vessels, interconduit pits contributed, on average, more than half of the total conduit resistivity (Pittermann et al., 2005; Hacke et al., 2006).

Pits are also found between (axial and ray) parenchyma cells and xylem conduits, thus providing an interface between the living and dead components of the xylem. Controlled fluxes of various solutes including ions (De Boer & Volkov, 2003; Nardini et al., 2010), sugars (Sauter et al., 1973; Salles et al., 2009) and amino acids (Sauter & Van Cleve, 1992) occur between ray cells and vessels. These exchange processes are most likely influenced by the properties of vessel-ray pit membranes. In addition, vessel-ray pits may play a role in embolism refilling as water appears to
enter refilling vessels through this interface (Braun, 1970; Brodersen et al., 2010).

The permeability of pit membranes in both intervessel and vessel-ray pits is critical for their function. The porosity and permeability of pit membranes are likely to be affected by the chemical composition of the membrane. However, our knowledge of pit membrane chemistry is limited. Pit membranes develop from the compound middle lamella (cml) (Evert, 2006), and it is usually assumed that their chemical composition resembles that of a typical primary cell wall. However, this assumption awaits further testing because of the extensive remodeling and hydrolysis of the original primary wall that takes place during pit membrane differentiation (Schmid & Machado, 1968; O’Brien, 1970; Morrow & Dute, 1998). According to the current paradigm, pit membranes are composed of multiple layers of cellulose microfibrils embedded in a matrix of hemicellulose and pectins (Choat et al., 2008). Structural proteins that are common in the primary cell wall (Cassab, 1998; Valentin et al., 2010) may also occur in pit membranes (Harrak et al., 1999).

Experimental evidence suggests that pectins play important roles in pit membrane functioning (Wisniewski et al., 1991; Zwieniecki et al., 2001; Boyce et al., 2004; van Ieperen, 2007; Cochard et al., 2010; Nardini et al., 2010). Pectins are a highly complex and heterogeneous group of polysaccharides rich in galacturonic acid (GalA). Different domains of pectins can be distinguished based on their biochemical properties; namely homogalacturonan (HG), rhamnogalacturonan-I (RG-I) and rhamnogalacturonan-II (Willats et al., 2001). Homogalacturonans are linear polymers of 1,4-linked α-D-GalA and are the most abundant pectic domain in the primary cell wall comprising up to 60% of total pectin (O’Neill & William, 2003). The pectic polymers extensively interact with each other as well as with the other cell wall components (Ryden et al., 2000; MacDougall et al., 2001a; Valentin et al., 2010), but the complexity of these connections is not fully understood.

Because of the presence of numerous carboxyl groups, pectins exhibit properties of polyelectrolytes and have an overall negative charge (Valentin et al., 2010). Calcium pectin gels tend to swell when there is an imbalance in the distribution of mobile counterions between the gel and surrounding solution, which results from the Donnan effect (MacDougall et al., 2001a). Crosslinking of the polymer network constrains the swelling tendencies (MacDougall et al., 2001a). The hydrogel behavior of pectins has been used to explain why the $k_b$ of stem segments is higher when measured with salt solution (typically 10–100 mM KCl solution) compared with perfusion with distilled water (Zwieniecki et al., 2001; Lopez-Portillo et al., 2005; Gasco et al., 2006; van Ieperen, 2007).

In addition to the ionic effect, calcium-mediated crosslinks between pectins have been proposed to influence the vulnerability of xylem to embolism (Sperry & Tyree, 1988). Perfusion of stems with oxalic acid and calcium solution induced greater vulnerability in sugar maple, perhaps by disrupting calcium-mediated crosslinks in the pectins of the pit membrane, which could make the membrane more flexible and allow for transient pore widening (Sperry & Tyree, 1988). Herbet & Cochard (2010) also found that embolism vulnerability increased after perfusing stems with calcium chelating solution. In the presence of $>10$ consecutive unmethyl-esterified GalA residues, calcium may interact with the negative charges of the GalA residues to form stable gels based on the ‘egg-box’ model (Cöffal & Mohnen, 2009). While other linkages exist in the pectin network, calcium crosslinking of HG is known to contribute to wall strength (MacDougall et al., 2001a; Micheli, 2001; Cöffal & Mohnen, 2009). The presence of low methyl-esterified HG in intervessel pit membranes may therefore be associated with shifts in vulnerability to embolism and may even explain differences in vulnerability between various species (Herbet & Cochard, 2010).

Both the ionic effect and the magnitude of the shift in vulnerability vary greatly among species (Cochard et al., 2010; Herbet & Cochard, 2010; Jansen et al., 2011). While the factors influencing the ionic effect have been previously investigated (Gortan et al., 2011; Jansen et al., 2011), the factors affecting the magnitude of the vulnerability shift after calcium removal are poorly understood. Herbet & Cochard (2010) noted that the shift was greater in more embolism-resistant species. Of the 13 species tested in their study, two (Salix alba and Betula pendula) remained unaffected by the treatments. It seems possible that pectins were not present in the intervessel pit membranes of these species. Alternatively, if pectins were present, they may not have been capable of forming calcium mediated crosslinks, possibly because of a high degree of esterification. The fact that monoclonal antibodies did not recognize HG in intervessel pit membranes of hybrid poplar saplings (Plavcová et al., 2011) supports the hypothesis that some highly vulnerable species do not possess pectinaceous intervessel pit membranes. The absence of HG would also prevent an increase in vulnerability after calcium removal. To test this hypothesis, we measured the shift in vulnerability in two relatively vulnerable species, Betula papyrifera and Populus balsamifera, and two more resistant species, Prunus virginiana and Amelanchier alnifolia. We hypothesized that HG will be absent in the intervessel pit membranes of the two vulnerable species, and that the vulnerability of these species will therefore remain unaffected by calcium removal. We expected the opposite pattern in the two resistant species.

One of the most powerful ways to study pectin in its physiological context is by using anti-pectin antibody probes (Wisniewski & Davis, 1995; Willats et al., 2001). However, this tool has rarely been used on intervessel pit...
membranes of woody angiosperms. In this present study, we used immunogold labeling with monoclonal antibodies raised against primary cell wall polysaccharides (HG, RG-I and xylolglucan) to probe the chemical composition of pit membranes. We wished to determine whether the pattern of HG labeling corresponded with the presence of calcium in pit membranes. To do this, we used an antimonate precipitation technique and compared the distribution of HG and calcium at the transmission electron microscopy level. Our final objective was to compare the structure and chemistry of different pit types. Although intervessel and vessel-ray pits may occur in the same vessel element, they have very different functions and may therefore differ in their chemical composition.

**Materials and Methods**

**Plant material**

Branches (approx. 1 cm in diameter) of *R. papyrifera* Marsh., *P. balsamifera* L., *P. virginiana* L., and *A. alnifolia* Nutt. were collected in the vicinity of the University of Alberta campus in Edmonton (53°50’ N 113°52’ W). These species were selected to cover a wide range of xylem vulnerability. The trees used for sampling were mature individuals growing in a river valley. For the measurements of vulnerability to embolism, branches were sampled from different randomly selected individuals. The plant material was used immediately or wrapped in plastic bags with wet paper towels and stored at 4°C for no longer than 3 d before the measurements were carried out. For electron microscopy work, xylem samples were collected from the same trees used for the hydraulic measurements. In this case, samples were processed immediately. Samples for hydraulic measurements and transmission electron microscopy (TEM) immunolabeling were collected in October and November 2010, samples for scanning electron microscopy (SEM) observations and TEM calcium localization were collected in February 2011 and samples for ruthenium red staining were taken in May 2011. In addition, a limited set of hydraulic measurements was carried out in February to verify that vulnerability to embolism did not undergo seasonal shifts.

**Shifts in vulnerability to embolism**

To evaluate the effect of calcium removal on vulnerability to embolism, vulnerability curves were obtained for control stems and stems perfused with a calcium chelating sodium phosphate solution (Herbette & Cochard, 2010). Stem segments were trimmed under water to a length of 14.2 cm. Segments were flushed with the treatment (pH 10) or control (pH 4) 10 mM NaPO₄ solution for 30–40 min at 20 kPa. At least 5 ml of solution were perfused through each stem. At pH 10, the phosphate occurs mainly in the form of HPO₄²⁻ anions that readily precipitate Ca²⁺ cations. By contrast, at pH 4 the predominant ionic form of phosphate is H₂PO₄⁻, which does not bind calcium. The same solutions used for flushing were used during the kₜ measurements. After flushing, stem segments were fitted to a tubing apparatus and the maximal hydraulic conductivity (kₜmax) was measured as the flow rate per pressure gradient (Plavcová et al., 2011). The standard centrifuging method (Alder et al., 1997; Li et al., 2008) was used to generate vulnerability curves. Curves were constructed by plotting the negative xylem pressure against the per cent loss of conductivity as described previously (Plavcová et al., 2011). Curves were fitted with a Weibull function and the xylem pressure corresponding to 50% loss of conductivity (P₅₀) was calculated for each stem segment. Six stems per species were measured for each NaPO₄ solution.

**Scanning electron microscopy**

Scanning electron microscopy was used to evaluate whether perfusion with the calcium chelating solution caused changes in pit membrane structure. To prepare the samples for SEM observation, stem segments 5 cm long were perfused with the treatment (pH 10) or control (pH 4) NaPO₄ solution at a pressure of 20 kPa for 1 h. Samples were then soaked in the same solution for 24 h at room temperature. Subsequently, segments were rinsed with distilled water and dehydrated through a gradual ethanol series 30–50–70–90% (30 min each) and placed in 100% ethanol overnight. Finally, segments were air-dried for 24 h, split with a razor blade and mounted on aluminum stubs using conductive silver paste. Samples were sputter-coated with chromium and carbon and observed with a field-emission scanning electron microscope (6301F, JOEL, Tokyo, Japan) using 2 kV acceleration voltage. Four different stems of *P. balsamifera* were observed for each NaPO₄ solution (pH 4 and 10, respectively). *P. balsamifera* was selected for these SEM observations, because both intervessel and vessel-ray pits were relatively easy to find on a tangential surface plane in comparison with the other three species studied. A semiquantitative elemental analysis via SEM coupled X-ray spectrometry (Princeton Gamma-Tech Inc., Princeton, NJ, USA) was used to assess the chemical composition of conspicuous particles that were abundant in the specimens treated with NaPO₄ at pH 10. An acceleration voltage of 50 kV was used for this analysis.

**Polysaccharide localization with transmission electron microscopy**

For immunolabeling, 1 mm³ blocks of xylem tissue sampled from the outer part of the stems were fixed in a mixture of 0.2% glutaraldehyde and 3.7% paraformaldehyde in 25 mM piperazine-N,N’-bis(2-ethanesulfonic acid) (PIPES) for 4.5 h at room temperature. Specimens were
then buffer washed, dehydrated and embedded in LR White resin (London Resin Co., London, UK). Our embedding procedure closely followed the steps previously described by Chaffey (2002), with the modification that we used heat polymerization at 60°C for 24 h. Ultrathin sections (70–90 nm) were cut using an ultramicrotome (Ultracut E, Reichert-Jung, Vienna, Austria) equipped with a diamond knife. Sections were collected on pioioform-coated nickel grids and immunolabeled. Four monoclonal antibodies (JIM5, JIM7, LM6 and LM15; PlantProbes, Leeds, UK) raised against different cell wall polysaccharide epitopes were used in this study. JIM5 and JIM7 bind to HG with a low and a high degree of methyl-esterification, respectively (Knox et al., 1990; Willats et al., 2001; Guillemin et al., 2005). LM6 recognizes arabian side-chains of RG-I (Ermel et al., 2000; Willats et al., 2001; Guillemin et al., 2005). LM15 is targeted to the XXXG motif of xyloglucan (Marcus et al., 2008). Immunolabeling was performed by floating the grids on drops of successively changing solutions. Sections were preincubated for 10 min on a drop of 0.05 M Tris-buffered saline (pH 7.6) with 0.1% Tween 20 and 0.1% BSA, blocked for 20 min with goat serum (Sigma-Aldrich) diluted 1 : 30 (v : v) in the same buffer, treated with the primary antibody (JIM5, JIM7 and LM5 diluted 1 : 4, LM15 diluted 1 : 3) overnight at 4°C, buffer-washed four times and stained with a secondary antibody, goat-anti rat IgG conjugated with 10 nm gold particles (Sigma-Aldrich) for 1 h. The grids were then extensively washed with buffer and filtered water and finally contrasted with 4% uranyl acetate for 25 min and with Reynolds’ lead citrate for 2 min (Reynolds, 1963). Sections were examined under a transmission electron microscope (Morgagni 268; Fei Company, Hillsboro, OR, USA).

The immunolabeling experiment was conceived as a qualitative study aiming to describe the pit membrane chemical composition as well as differences between species. Given our interest in the occurrence of HG in pit membranes, we assessed the labeling density of JIM5 and JIM7 antibodies in the pit membrane annulus. We counted the gold particles in this region. The labeling density in pit membrane annuli was estimated based on 6–15 pits per species and antibody. In addition, the length of the annulus was measured as the distance from the edge of the pit (following the plane of the cml) to the area where the annulus transitioned into the main portion of the pit membrane. This transition zone could be clearly identified based on the difference in the electron density of the membrane. In addition to these measurements, pit membrane diameter was calculated from the pit area assessed in tangential sections with a light microscope at 1000× magnification assuming a circular shape of pit membranes.

To verify the immunolocalization pattern, a histological detection of pectins was performed via ruthenium red staining based on methods described previously (Micheli et al., 2002; Gorton et al., 2011). Xylem tissue was fixed in Karnovsky’s fixative containing 0.1% (w : v) ruthenium red for 1.5 h and post-fixed in 1% buffered osmium tetroxide with 0.1% ruthenium red for 1.5 h at room temperature. Subsequently, samples were dehydrated and embedded in Spurr’s resin (EMS, Hatfield, PA, USA). As osmium tetroxide post-fixation provides some contrast to the tissue, control samples were prepared using the same solutions without ruthenium red. Sections were mounted on coated copper grids and examined without further staining.

### Calcium localization

Calcium was localized within the wood tissue using an antimoniate precipitation technique (Wick & Hepler, 1980; Slocum & Roux, 1982). This method has been successfully used to localize cell wall-bound calcium in the cambial zone of poplar (Baier et al., 1994; Guglielmino et al., 1997) and European ash (Funada & Catesson, 1991). In this study, small blocks (1 mm3) of mature xylem were fixed in a mixture of 2% glutaraldehyde, 2.5% formaldehyde, 0.1% tannic acid and 2% potassium antimoniate in 0.1 M potassium phosphate buffer at pH 7.6 for 6 h at room temperature in the dark. Specimens were then washed twice for 15 min in antimoniate buffer (2% KSB(OH)4 in 0.1 M potassium phosphate at pH 8), postfixed in 1% OsO4 in antimonate buffer for 2 h, washed three times in antimonate buffer for 10 min, washed for 30 min in 0.01 M potassium phosphate buffer without antimonate at pH 7.6, and finally gradually dehydrated and embedded in Spurr’s resin. Ultrathin sections were collected on coated copper grids and observed without further staining.

### Statistical analysis

The statistical environment R (R Development Core Team, 2009) was used to perform the statistical analysis. Analysis of variance followed by Tukey’s HSD post hoc comparison test was conducted to evaluate differences among the species and perfusion solutions. For all tests, differences were considered statistically significant at $P \leq 0.05$.

### Results

#### Shifts in xylem vulnerability to embolism

Vulnerability to embolism was significantly higher in stems perfused with NaPO4 solution at pH 10 compared with the same solution at pH 4 in three out of four species studied ($P < 0.001$) (Fig. 1). The $P_{50}$ values at pH 4 did not differ from values previously measured in our laboratory for these species using a regular measuring solution (20 mM KCl and 1 mM CaCl2) instead of NaPO4 (data not shown). Therefore, perfusion with NaPO4 at pH 4 does not affect xylem vulnerability and the $P_{50}$ at pH 4 reflects the ‘native’
Xylem vulnerability. The difference in P50 (ΔP50) in stems infiltrated with solutions of different pH was largest in the most resistant species, *A. alnifolia* (ΔP50 = 2.12 ± 0.16 MPa). The magnitude of the shift in P50 was proportional to the native vulnerability of species, with more resistant species shifting more (see the Supporting Information, Fig. S1). Although their P50 values at pH 4 did not differ (P = 0.945), *P. balsamifera* exhibited a significant shift in vulnerability (P = 0.020) while *B. papyrifera* did not (P = 0.998). The correlation between ΔP50 and the P50 at pH 4 closely followed the relationship observed by Herbette & Cochard (2010) although we used different species and a different centrifuge method to measure the vulnerability curves.

**Scanning electron microscopy**

Scanning electron microscopy was used to test if perfusion with calcium chelating solution caused visible changes in pit membrane structure. Spherical particles were often found in the vessel lumens of samples treated with NaPO4 at pH 10, but were rare in samples treated with NaPO4 solution at pH 4 (Fig. 2). Semiquantiitative elemental analysis with an X-ray analysis system coupled with the SEM microscope revealed that the particles were rich in phosphorus, calcium, sodium and magnesium. At pH 10, but not at pH 4, these particles were consistently found in high concentrations on the surface of vessel-ray pit membranes (Fig. 2b). At pH 10, precipitate was occasionally found on the surface of intervessel pit membranes (images not shown). However, compared with vessel-ray pit membranes, intervessel pit membranes had far fewer particles, and precipitate was only sporadically observed in different samples. The distribution of these particles within the intervessel pit membrane seemed fairly random. However, in several instances more particles were found around the periphery of the pit membrane, that is, the annulus region.

**Distribution of high and low methyl-esterified homogalacturonan**

None of the antibodies used in this study showed a strong signal in the entire pit membrane of interconduit pits.
Nevertheless, several labeling patterns could be distinguished (Table 1). JIM7 labeling was more or less evenly distributed in the cml (Fig. 3a). By contrast, the JIM5 epitope was only rarely found in the cml between two cells, but was more frequently present in the cml of cell corners. This pattern indicates that the HG in the cml was largely esterified. Pit membranes of all species showed a distinct annulus, that is, an electron-dense area near the periphery of the pit membrane. The annulus was substantially shorter in B. papyrifera in comparison with the other species (Table 2). Importantly, this region was strongly labeled with both JIM5 and JIM7 (Fig. 3a,b,c). Whereas JIM7 labeling was usually evident throughout the entire annulus region (Fig. 3a), the JIM5 epitope was often confined to the tip of the annulus (Fig. 3b,c). The number of gold particles per annulus for JIM5 and JIM7 varied, on average, from 5 to 15 between the species (Table 2). These counts provide an estimate of the ratio between low and high methyl-esterified HG in the annulus. This ratio was highest in B. papyrifera.

The pit annuli of imperfect tracheary elements in P. virginiana, A. alnifolia and B. papyrifera showed the same HG labeling pattern that was observed in intervessel pits (Fig. 4a). We use the term ‘imperfect tracheary element’ in the all-inclusive sense outlined by Carlquist (1986). Pits of imperfect tracheary elements in P. virginiana and A. alnifolia were distinct in that they often had pseudotori thickenings. The attachment of the cap-like thickenings to the membrane varied. Sometimes

Table 1 The intensity and localization of immunogold labeling for polysaccharide-specific antibodies JIM5, JIM7, LM6 and LM15

<table>
<thead>
<tr>
<th>Epitope</th>
<th>JIM5</th>
<th>JIM7</th>
<th>LM6</th>
<th>LM15</th>
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<tbody>
<tr>
<td>Intervessel pit membrane</td>
<td>Low methyl-esterified/nonesterified HG</td>
<td>Methyl-esterified HG</td>
<td>(1-5)-α-L-arabinan of RG-I</td>
<td>XXXG motif of XG</td>
</tr>
<tr>
<td>Annullus</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Amorphous layer of ray cells</td>
<td>++</td>
<td>++</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>cml-between two cells</td>
<td>−/−</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>cml-cell corners</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Pseudotori</td>
<td>−/−</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Membrane under pseudorus</td>
<td>++</td>
<td>++</td>
<td>−/−</td>
<td>−</td>
</tr>
</tbody>
</table>

++, strong signal/enrichment in labeling; +, weak signal; −, no signal; cml, compound middle lamella; HG, homogalacturonan; RG-I, rhamnogalacturonan I; XG, xyloglucan.

Fig. 3 Transmission electron micrographs of the pit membrane annulus region showing the distribution of homogalacturonan (HG) (a,b,c) and calcium precipitate (d) as revealed by immunogold labeling (a,b,c) and an antimonate precipitation technique (d). Labeling with JIM7 antibody is indicative of high methyl-esterified HG and was evident throughout the entire annulus region (a). Labeling with JIM5 antibody shows the distribution of low methyl-esterified HG and was often restricted to the tip of the annulus (b,c). Electron-dense calcium precipitates in the annulus (d) closely resembled the JIM5 labeling pattern (compare b and d). Micrographs show pits of Populus balsamifera labeled with JIM7 (a), Prunus virginiana labeled with JIM5 (b), Betula papyrifera labeled with JIM5 (c) and Amelanchier alnifolia after antimonate precipitation (d). Arrow heads point to annuli; cml, compound middle lamella; pm, pit membrane; scw, secondary cell wall. Bars, (a–c) 0.2 μm, (d) 0.5 μm.
the pads were closely attached to the membrane. In those cases, strong labeling with JIM7 was found throughout the inner membrane layer, that is the part of the membrane located between the pads (images not shown). In other cases, the thickenings were somewhat disconnected and formed hollow horseshoe- or cap-like structures that overarched the membrane (Fig. 4a). Strong labeling with both JIM5 and JIM7 was seen on the inner surface of the cap-like thickenings (Fig. 4a, arrows).

In contrast to intervessel pit membranes, the membranes of vessel-ray pits consistently exhibited strong labeling for both JIM5 and JIM7 (Fig. 4c). JIM5 was localized closer to the surface of the membrane facing the vessel lumen (Fig. 4c), whereas the density of JIM7 labeling was more homogeneous across the pit membrane. JIM5 and JIM7 labeling was also detected in the amorphous cell wall layer of ray parenchyma cells. In *A. alnifolia*, JIM5 labeling was enriched in a darker band that traversed the amorphous layer (Fig. 4c).

The HG distribution patterns, as detected with antibodies, were verified using ruthenium red staining. Electron-dense regions indicated a positive staining reaction with ruthenium red. In agreement with the immunolabeling results, staining was consistently observed in the annulus of intervessel pits (Fig. 5a,b), vessel-ray pit membranes, the amorphous layer (Fig. 5c) and in pseudotori (Fig. 5d). The pattern described above was not found in control samples prepared without ruthenium red, indicating that osmium tetroxide alone was not responsible for the differential contrast. The main part of intervessel pit membranes could be distinguished in samples of all species as a very faint, almost electron-transparent granular layer. No increase in electron density of the main part of intervessel pit membranes was apparent in samples stained in ruthenium red.

### Calcium localization with TEM

The antimonate technique was used to localize electron dense calcium precipitate in TEM-samples. In agreement with the calcium chelating experiments described above (Fig. 2b), the precipitates often formed a thin layer of clumps along the inner vessel walls. By contrast, precipitates were less common in fiber lumens. A distinct layer of precipitate was consistently observed in the annulus region of interconduit pits (Figs 3d, 4b). This pattern matched the labeling with JIM5. The distribution of calcium precipitate also matched the JIM5 labeling patterns found in pseudotori (compare Fig. 4a,b). Precipitate was consistently found on the surface of vessel-ray pit membranes (Fig. 4d, arrows). While the distribution of calcium precipitate closely matched the distinct JIM5 labeling patterns described above, there was less agreement with JIM7 labeling. Calcium precipitate was not found throughout the entire annulus or the entire vessel-ray pit membrane. Instead, the precipitate often formed a lining around the annulus (Fig. 3d) and was restricted to the surface of vessel-ray pit membranes (Fig. 4d).

### Distribution of rhamnogalacturonan I and xyloglucan

The signal for the other antibodies, LM6 and LM15, tended to be weaker and more variable than the patterns seen for JIM5 and JIM7. Nevertheless, both antibodies were localized in the cml. Weak labeling of anti-RG-I LM6 occurred in vessel-ray pit membranes. The LM6 epitope was most abundant in the amorphous layer of ray cells (Fig. S2a), where it provided a strong signal in all species. While LM6 labeled pseudotori (Fig. S2c), the LM6 signal was not enriched in annuli. Intervessel pit membranes of *P. virginiana* showed weak labeling with LM6, but this pattern could not be confirmed in the other species.

The LM15 xyloglucan epitope was consistently found in the cml (Fig. S2b) and in the outer layer of cell corners (Fig. S2d). Weak labeling was also found in intervessel pit membranes of *P. virginiana* and *A. alnifolia* (Fig. S2b, dark grey arrow). LM15 labeling was not observed in pseudotori, vessel-ray pit membranes, and in the amorphous layer (Table 1).

### Table 2

<table>
<thead>
<tr>
<th>Species</th>
<th>Pit membrane diameter ($D_m$) (µm)</th>
<th>Annulus length (nm)</th>
<th>JIM5</th>
<th>JIM7</th>
<th>JIM5 : JIM7</th>
</tr>
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<tbody>
<tr>
<td><em>Betula papyrifera</em></td>
<td>2.3 ± 0.03 (86)a</td>
<td>141 ± 8 (24)a</td>
<td>12 ± 1 (13)a</td>
<td>5 ± 1 (13)a</td>
<td>2.49</td>
</tr>
<tr>
<td><em>Populus balsamifera</em></td>
<td>6.8 ± 0.06 (85)b</td>
<td>275 ± 24 (20)b</td>
<td>9 ± 2 (16)ab</td>
<td>10 ± 2 (6)ab</td>
<td>0.82</td>
</tr>
<tr>
<td><em>Prunus virginiana</em></td>
<td>4.6 ± 0.05 (87)c</td>
<td>331 ± 27 (20)b</td>
<td>5 ± 0 (6)b</td>
<td>10 ± 2 (15)b</td>
<td>0.48</td>
</tr>
<tr>
<td><em>Amelanchier alnifolia</em></td>
<td>4.8 ± 0.05 (76)c</td>
<td>351 ± 23 (20)b</td>
<td>6 ± 1 (15)b</td>
<td>15 ± 1 (9)b</td>
<td>0.44</td>
</tr>
</tbody>
</table>

The intensity of the labeling was assessed by counting the number of gold particles localized in the annulus. The JIM5 : JIM7 antibody ratio is indicative of the ratio of low methyl-esterified to high methyl-esterified homogalacturonan. Values represent mean ± SE, the number of individual pits and annuli used for these measurements is indicated in brackets. Different letters indicate that the means were significantly different between the species, one-way ANOVA followed by Tukey HSD test ($P < 0.05$).
Discussion

A significant increase in xylem vulnerability after perfusion with calcium chelating solution was found in three out of four species studied (Fig. 1). The magnitude of the shift was proportional to the native vulnerability of the species, with the more resistant species showing larger shifts (Fig. S1). These results are in agreement with previous findings of Herbette & Cochard (2010). In addition, our data indicate that at least some highly vulnerable species can become even more vulnerable after calcium removal, as shown by the 0.6 MPa shift seen in *P. balsamifera*. The shift in vulnerability after calcium removal was highly reproducible, suggesting that it was caused by a 'controlled' change in pit membrane properties rather than a complete loss of integrity of the pit membranes. Sperry & Tyree (1988) and Herbette & Cochard (2010) suggested that the shift in vulnerability is caused by disruptions of calcium–HG crosslinks in intervessel pit membranes that would change the rigidity and stretching properties of the membranes.

The pectin localization experiments conducted in this study showed that the main part of intervessel pit membranes in all four species contained very little or no HG. Thus, our original hypothesis that there is a link between the presence or absence of HG in the pit membrane and the magnitude of the vulnerability shift was not supported. The only portion of the membrane that was rich in HG was the marginal annulus region. The pit membrane annulus is a conspicuous feature of the pit membrane and has often been noted by wood anatomists. A distinct annulus occurs both in interconduit pits of angiosperms (Schmid, 1965; Schmid & Machado, 1968; Jansen *et al.*, 2009; Gortan *et al.*, 2011) as well as in torus–margo pits of gymnosperms.

Fig. 4 Corresponding patterns of immunolabeling with JIM5 antibody (a,c) and calcium localization (b,d) in pseudotori (a, b) and vessel-ray pits (c,d) of *Amelanchier alnifolia* as observed with transmission electron microscopy. Gold particles and calcium precipitate were localized on the inner surface of pseudotori (arrows in a and b) of imperforate tracheary elements. Precipitates were also evident in the annulus (arrowheads in b). JIM5 epitopes were abundant in the vessel-ray pit membrane (pm in c) and in the amorphous layer (al) of ray cells (c). The greatest labeling density within the vessel-ray pit membrane was found close to the surface of the membrane, near the vessel lumen (v, arrows; panel c only). Correspondingly, a distinct layer of precipitates was observed lining the outer vessel-ray pit membrane surface in samples treated with antimonate (d). Precipitates were also found around the periphery of the vessel lumen. al, amorphous layer; cml, compound middle lamella; pm, pit membrane; pt, pseudotorus; r, ray cell; scw, secondary cell wall; v, vessel lumen. Bars, 0.5 μm.
(Liese, 1965; Dute et al., 2008; Pittermann et al., 2010). Under TEM, the annulus typically appears more electron-dense than the rest of the membrane. However, the opposite pattern has also been found in a few species (Schmitz et al., 2007; Jansen et al., 2009; Gortan et al., 2011). To the best of our knowledge, it is not known whether the annulus plays a physiological role or if it simply relates to pit development (e.g. the enzymes that remodel the pit membrane may have restricted access to the marginal membrane region).

A high HG content in the annulus was consistently observed in all four species studied and agrees with our previous observations on hybrid poplar saplings (Plavcová et al., 2011) as well as earlier reports (O’Brien, 1970). Furthermore, the anti-RG-I antibody did not display an increased labeling density in the annulus, indicating that it is specifically the HG domain of pectins that is enriched in the annulus. Homogalacturonans are known for their calcium-binding capacity (MacDougall et al., 2001b; Proseus & Boyer, 2008) and this premise was confirmed by our calcium localization experiment. Hence, it is possible that the effect of calcium removal on xylem vulnerability is realized through the disruption of the HG-calcium superstructure within the annulus. The fact that the magnitude of the shift in the four species studied was proportional to the length of the annulus further supports this hypothesis.

Based on our findings, we propose that the pit membrane should not be viewed as an isotropic material. Instead, there are two chemically and structurally distinct domains (the pit membrane annulus and the main part of the membrane) that likely exhibit different mechanical properties. A growing body of evidence suggests that the large pressure difference that is exerted on the pit membrane before air seeding results in pit membrane stretching, and that the extent of membrane deflection influences the cavitation threshold. Features that minimize pit membrane deflection such as vestures (Choat et al., 2004) or shallow pit chambers (Haque & Jansen, 2009; Lens et al., 2011) were found in embolism-resistant xylem. Although calcium crosslinks can substantially enhance cell wall strength (Parre & Geitmann, 2005; Cybulská et al., 2011), cellulose microfibrils most likely represent the main load-bearing component of pit membranes and limit the degree of membrane deformation (Pett, 1972; Sperry & Hacke, 2004). The orientation of microfibrils is critical for mechanical properties. The wall is more pliable in the direction perpendicular to the prevailing orientation of the microfibrils. In the pit membrane, the microfibrils seem to be oriented randomly. However, it is not clear whether the same pattern of microfibril orientation is maintained in the annulus. Observations of developing pits suggest that microfibrils are deposited in a circular fashion near the pit border (Wardrop, 1954; Imamura & Harada, 1973; Chaffey et al., 1997). This deposition pattern is usually interpreted as an initial step in the formation of the pit border. However, Imamura & Harada (1973) state: ‘...the circularly oriented microfibrils have been laid down making the periphery of the pit area, called the pit annulus’.

If microfibrils in the annulus are oriented in a circular fashion, the annulus would extend more than the rest of the membrane during air seeding as the cellulose microfibrils would move apart. When stems were perfused with the...
calcium chelating agent, the annulus region may have become looser and even more extensible. Thus, it is possible that calcium removal led to the formation of small pores or micro-cracks within the annulus or the annulus-membrane interface through which air could penetrate and cause embolism. Alternatively, calcium removal from the annulus may allow for increased pit membrane deflection, which in turn would lead to a widening of membrane pores. A more extensible annulus would allow the pit membrane to deflect further and aspirate sooner against the pit border. As the membrane continues to deflect through the pit aperture, the pores will start to enlarge even more and air seeding will occur at less negative xylem pressure (Sperry & Hacke, 2004).

*Betula papyrifera* was the only species in this study that did not show a shift in vulnerability after calcium removal. Although HG and calcium were present in the pit membrane annulus, annuli of *B. papyrifera* were significantly shorter than those of the other three species. This may explain why the effect of calcium removal was less pronounced in *B. papyrifera*. In addition, intervessel pits of *B. papyrifera* were much smaller than those of the other species. With all other parameters being equal, smaller pits should show less membrane deflection than larger ones (see Eqn 8 in Sperry & Hacke, 2004). Membrane deflection in *B. papyrifera* may therefore be minimal. However, pit membrane diameter alone does not have a strong effect on xylem vulnerability (Jansen et al., 2009; Lens et al., 2011). Pit membrane thickness can also play a role. More detailed knowledge of the overall pit geometry and mechanical properties is necessary to decipher if and how pit membrane deflection influences the air seeding threshold. Our results also indicate that HG in annuli of *B. papyrifera* exhibited a relatively low degree of methyl-esterification (Table 2). Low-esterified HG binds calcium more tightly than does high methoxyl pectin (Tibbits et al., 1998). It is possible that calcium was less susceptible to sequestration in *B. papyrifera* and that the annulus consequently retained its original strength. However, the data on labeling density shown in Table 2 should be interpreted with caution as they originate from only one or two independent immunolabeling experiments per species.

The proposed hypothesis that changes in the extensibility of the annulus can induce increased vulnerability is speculative. An alternative explanation would be that a low amount of HG and calcium is present in interconduit pit membranes, and that this small amount was not detected by the methods employed in this study. Following this argument, one would expect that HG and calcium were completely removed during pit development in *B. papyrifera* (which did not shift), but not in the other species, which did shift. This seems unlikely and none of our data supported this view. The different methods employed in this study all indicated that the annulus is the region of the pit membrane with the highest HG and calcium content. It seems improbable that the treatment with calcium chelating agents would not have any effect on it.

Our results highlight the differences in interconduit and vessel-ray pit membrane chemistry. In contrast to intervessel pits, the entire surface of vessel-ray pit membranes was rich in pectins (Figs 4c, 5c) and calcium (Figs 2, 4d). Vessel-ray pit membranes lacked an annulus. The transitional region between cm1 and pit membrane did not show enhanced HG-labeling, further supporting that the annulus might have a specific role in interconduit pit functioning. Pectins were previously found in vessel-ray pit membranes of *Prunus persica* (Wisniewski & Davis, 1995). It has been suggested that pectinaceous vessel-ray pit membranes effectively isolate intracellular water from extracellular ice, thereby conferring the ability of the tissue to undergo supercooling.

This study also provides information on pseudotori. The function, if any, of these peculiar structures is unknown. Pseudotori occur in tracheary elements of several woody taxa, including Rosaceae, Ericaceae, and Oleaceae (Jansen et al., 2007; Rabaey et al., 2008). Pseudotori develop as secondary thickenings following the formation of plasmodesmata-associated primary thickenings. Later in development, autolytic enzymes remove the primary thickenings leaving the pseudotori intact, and the tracheary cell undergoes programmed cell death (Rabaey et al., 2008). Our immunolabeling experiments complement this description with some chemistry data. We often observed a nonhydrolysed portion of the pit membrane that was protected by pseudotorus caps. This finding further highlights the substantial remodeling and hydrolysis that affects most of the noncellulosic components during the development of interconduit pit membranes (O’Brien, 1970; Czaninski, 1972). In addition, the colocalization of HG and calcium (Fig. 4a,b) demonstrates that calcium abundantly occurs in association with HG in the xylem.

A different pattern in pectin distribution was recently found in pit membranes of four Lauraceae species (Gortan et al., 2011). In these species, intervessel pit membranes reacted strongly with ruthenium red, while annuli and vessel-ray pit membranes showed a negative reaction. Pectins were also present in the pit membranes of grapevine (Sun et al., 2011) and conifers (Hafren et al., 2000). Thus, it is obvious that substantial variability in pit membrane chemistry exists across different plant taxa. Future research will likely highlight the functional and ecological significance of these differences in pit chemistry. A comparative study using angiosperm species with high and low pectin content in their pit membranes would provide further insights into the role of pectins in pit membrane functioning. For example, it would be helpful to see whether a similar shift in vulnerability after calcium removal occurs in Lauraceae species and whether the high pectin content in pit membranes of Lauraceae is associated with high pit resistance and with...
the ability to trap solutes within refilling vessels (Hacke & Sperry, 2003; Nardini et al., 2011).

In addition, the relationship between shifts in vulnerability and the ion-mediated increase in hydraulic conductivity (Zwieniecki et al., 2001; Gasco et al., 2006) needs to be evaluated as both phenomena are attributed to the presence of HG in the pit membranes. However, there are also distinct differences that may partly uncouple these two phenomena. For example, a broader spectrum of pectin types can be responsible for the ionic effect while only the long linear stretches of GalA that are typical for the HG domain are capable of substantial calcium crosslinking. All four species in this study showed a relatively weak ionic effect between 5% and 16% (data not shown). The magnitude of the ionic effect did not correspond with ΔPs. It is not clear whether the hydrogel behavior of pectins in the annulus could influence ΔPs. However, it is possible that the tension of the annulus, as determined by the ion interactions, transmits to the cellulosic matrix of the pit membrane, thereby stretching and relaxing the membrane pores.

In conclusion, this study enhances our knowledge of the chemical composition of pit membranes, which is a prerequisite for a better understanding of the role of these microvalves in xylem functioning. Our results suggest that there is no link between the general presence or absence of HG in intervessel pit membranes and the shift in vulnerability after calcium removal. We provide evidence that pectic HG are not homogeneously distributed in intervessel pit membranes. In the four species studied, the main part of the membrane contained very little or no pectins whereas the annulus showed substantial enrichment in HG-labeling. Calcium precipitation experiments confirmed that calcium colocalizes with HG in the annulus. We therefore hypothesize that the disruption of HG-calcium crosslinks within the annulus can lead to increased vulnerability to cavitation. Our results also highlight differences in chemical composition between interconduit and vessel-ray pit membranes. Observed differences in chemistry likely reflect the different biological functions of these pit types. While intervessel pits facilitate water flow, vessel-ray pits may be designed to provide a selective barrier between living ray cells and dead vessel elements.

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References


Supporting Information

Additional Supporting Information may be found in the online version of this article:

**Fig. S1** Relationship between vulnerability to cavitation and the magnitude of vulnerability shift in four angiosperm species.

**Fig. S2** Immunogold labeling of cell walls with anti-RG-I antibody LM6 and anti-xylanlucan antibody LM15.

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