Specific and dynamic lignification at the cell-type level controls plant physiology and adaptability

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Abstract

Lignins, abundant phenolic cell wall polymers that accumulate in vascular tissue, were essential for plant terrestrialization as they enable sap conduction and mechanical support. Although lignification is currently understood as a random process, different cell types accumulate lignins with different compositions. The biological significance of these cellular differences is however still unknown. We performed single cell analyses to decipher the specific roles of different lignins and their residues on sap conduction and mechanical strengthening in plant xylem, using inducible pluripotent cell cultures and genetically modified whole plants. We show that specific lignins dynamically accumulate in each cell type and their morphotypes using distinct genetic programs, and that different lignin residues have non-redundant roles on plant biomechanical and hydraulic properties. Lignin is therefore a dynamic polymer changing composition to tailor the load bearing and sap conduction properties of each cells, in order for plants to adapt to developmental and environmental constraints.

Keywords: Arabidopsis, poplar, inducible pluripotent stem cell cultures, cell wall, lignin, controlled lignification, biomechanics, vessel collapse, cell morphotypes, drought resistance, single-cell quantitative image analyses, irregular xylem, structural equation modelling

Introduction

Complex multicellular organisms require adjustable vascular systems to maintain tissue hydration and prevent embolism independently and irrespective of developmental and environmental constraints. Vertebrate animals have a closed vascular system that functions under a self-adjustable positive pressure caused by the heart muscles and modulation of blood conduit diameters. In contrast, plants have an open system under environmentally-controlled negative pressure with non-adjustable conduits of different diameters. These conduit cells, the tracheary elements (TEs), are part of a tissue called xylem that both provides the load-bearing skeleton and the sap conduction network. The dual role of TEs is enabled by the reinforcement of their primary cell walls (PCWs) with secondary cell walls (SCWs), and the subsequent removal of the intracellular content, except for the cell walls, by programmed cell death (Ménard et al. 2015; Derbyshire et al. 2015). As the plant grows, new TEs form, die and connect to older TEs to form a contiguous hollow conductive network that channels the sap throughout the plant (Ménard and Pesquet 2015). Because TEs have to undergo cell death to become functional, they have been long considered to have a static role in plant sap conduction. Yet, to adapt to developmental and environmental changes, the organisation of the continuously growing xylem dynamically changes. This affects not only the proportion and organisation of the different TE morphotypes – narrow protoxylem (PX) with annular or spiral patterns or wide metaxylem (MX) with reticulate or pitted patterns, but also the cell types surrounding TEs – un lignified xylem parenchyma (XP) and lignified xylary fibres (XF) (Fig. S1; Chaffey et al. 2002; Derbyshire et al. 2015).

In the conductive networks formed by connected hollow TEs, the sap only ascends due to a negative pressure pull caused by the water potential gradients (Ψ) along the soil/plant/atmosphere continuum; the sap water is eventually released in the air as vapour by the evapotranspiration of leaves. Ψ along the continuum depends on differences in capillary, osmotic, mechanical, gravitational and vapour pressures, causing Ψ in soil and air to fluctuate with environmental conditions (Fig. S1). In contrast, plants control their Ψ using both stomatal movements to regulate leaf transpiration rates, and the intracellular osmolarity of XPs and XFs to alter osmotic pressure (Bentrup 2017; Holbrook et al. 1995; Pockman et al. 1995). Al-
though the contribution of the mechanical pressure exerted by TE cell walls (the matric potential) is still unclear (Boyer 1967), we know TEs cannot withstand very large Ψ changes, such as the ones due to extreme drought, and would collapse inwardly, disrupt sap transport resulting in embolism (Fig. S1; Brodribb and Holbrook 2005; Zhang et al. 2016; Coleman et al. 2008; Kitin et al. 2010; Voelker et al. 2011). However, the regulation of plant hydration by modulating structural features of TE morphotypes and xylem organisation remains poorly understood.

The xylem sap conduction properties, mechanical rigidity and toughness are determined by the accumulation in TEs of lignins, phenolic polymers with complex composition and structure (Barros et al. 2015; Pesquet et al. 2019). Unlike other cell wall polymers, lignin formation continues post-mortem in dead TEs by cooperating with neighbouring living XFs, XP and the xylem sap content which supply monomers (Pesquet et al. 2010; Pesquet et al. 2013; Derbyshire et al. 2015; Blaschek, Champagne, et al. 2020). This post-mortem lignification is catalyzed by highly stable and cell wall embedded oxidative enzymes, filling the gaps in between polysaccharidic cell wall polymers with lignins. The predominant lignin monomers are C6,C3 phenylpropanoids varying in their C6 meta groups, such as monomethoxylated guaiacyl (G) and dimethoxylated syringyl (S), and their C3 functions, such as alcohol (XCHOH) or aldehyde (XCHO) (Dixon and Barros 2019). Other lignin subunits with C6 phenyl residue (P) and C6,C1 benzaldehydes can also be incorporated in plant lignins. Due to the low spatial resolution, standard lignin analyses have led to consider lignin polymerisation as a random process only regulated by monomer supply (Barros et al. 2015). However, recent works challenged this model by showing that lignification is a genetically controlled process regulating the amounts, linkages and composition of lignin in each cell type and their different cell wall layers (Blaschek, Nuoendagula, et al. 2020; Blaschek, Champagne, et al. 2020; Yamamoto et al. 2020). Indeed, different TE morphotypes have been shown to contain lignins with different amounts and differently positioned C CHO (Blaschek, Champagne, et al. 2020; Yamamoto et al. 2020). Yet, the biological roles of lignin dynamics and diversity in each TE morphotype are still not understood. Herein, we investigated the biological roles of the xylem organisation, TE morphotypes...
and their specific lignins as well as their *post-mortem* lignification capacity. We used inducible plant pluripotent cell suspension cultures (iPSCs) and genetically engineered herbaceous and woody plants with modified lignins to show that each TE morphotype exhibits specific lignin amounts, composition and structures that change during *post-mortem* lignification. We demonstrated that the different lignin subunits are essential and non-redundant to fine-tune the biomechanical and hydraulic properties in each TE to sustain extreme Ψ changes. Instead of being a “random” polymer, we demonstrated that lignin structure is specifically tailored during the maturation of each TE morphotype to enable and dynamically adapt their conductive function in changing developmental and environmental conditions.

**Results**

*Post-mortem* lignification enables isolated TE morphotypes to resist extreme Ψ differentials. To define the role of lignins in TEs capacity to withstand extreme negative pressure, iPSCs were hormonally induced to produce intact and isolated TEs (Ménard et al. 2017). iPSCs synchronously formed all TE morphotypes, which underwent cell death 5–7 days after induction, followed by *post-mortem* lignification (Pesquet et al. 2010; Pesquet et al. 2013; Derbyshire et al. 2015). We monitored these *post-mortem* changes in TEs at the nanoscale using scanning electron microscopy (SEM) coupled with energy-dispersive X-ray spectroscopy (EDS) by measuring elemental changes in cell wall composition (Fig. 1A, B). Ratios of carbon (C) to coating chromium (Cr) content showed gradual *post-mortem* increases of C-rich compounds only in SCWs, which plateaued by day 50 (Fig. 1C). Ratios of carbon (C) to oxygen (O) content revealed that the compounds accumulating *post-mortem* in TE SCWs were lowly oxygenated as expected from lignin (9 C:3–5 O for lignin-units compared to 6 C:6 O for cellulose-units; Fig. 1C). Altogether we showed that TE SCWs continue lignifying for more than 40 days after cell death.

The impact of *post-mortem* lignification on TEs was then tested using atomic force microscopy (AFM) on 5–20 µm² areas of isolated 10- and 50-day-old PX and MX TEs (Fig. 1D). AFM analysis showed that *post-mortem* lignification had no impact on PCW or on SCW surface adhesion but altered the biomechanical properties of SCW for each TE morphotype by significantly increasing stiffness and decreasing deformability (Fig. 1E). The role of these biomechanical changes in TE SCWs due to *post-mortem* lignification was evaluated in response to extreme Ψ changes by exposing isolated 10- and 50-day-old TEs to two different drying methods and then observing them using SEM. Critical-point drying (CPD), which minimizes Ψ changes during the drying process, was compared to air drying, which involves extreme Ψ changes. Parenchymatic cells showed no collapse after CPD but were completely flattened by air drying (Fig. 2A, B) whereas TEs were similarly unaffected by CPD but withstood air drying (Fig 2A, B). The proportion of TEs fully or partially collapsed after air drying decreased as *post-mortem* lignification progressed, with the majority of 50-day-old TEs remaining completely intact (Fig. 2C, D). Both PX and MX TEs (Fig. 2E, F) withstood collapse in air drying better as *post-mortem* lignification progressed, although PX were consistently more sensitive than MX TEs (Fig. 2C, D). To ensure that lignins were the factor determining the observed increased resistance, lignification of TE SCWs was prevented with piperonylic acid (PA) treatment (Van de Wouwer et al. 2016; Decou et al. 2017); as a result, un lignified TEs completely collapsed with air drying (Fig. 2G). We thus demonstrated for the first time that *post-mortem* lignification of TEs is a unique dynamic maturation process made to biomechanically reinforce TEs against Ψ differentials for optimal sap conduction.

Specific lignin concentration and composition set the resistance of each TE morphotype to negative pressure in herbaceous whole plants. We then investigated TE lignification and resistance to collapse in herbaceous whole plants using *Arabidopsis thaliana*. Three TE morphotypes are present in 8-week old stems of wild-type (WT) plants, identifiable by their distance to the cambium: protoxylem TEs (PX), primary metaxylem TEs (PMX), and secondary metaxylem TEs (SIX) (Fig. 3A). All TEs were surrounded by ~35% of other TEs whereas the remaining neighbouring cell types varied in

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**Figure 2** | Gradual post-mortem lignification enables all TE morphotypes to resist extreme Ψ differentials. **A** Scanning electron micrograph of 30-day-old isolated TEs and parenchyma cells produced from iPSCs and prepared using critical point drying (CPD). Note that both TEs and parenchyma are intact as indicated by blue and white arrows respectively. **B** Scanning electron micrograph of 30-day-old isolated TEs and parenchyma cells produced from iPSCs and prepared using air drying. Note that parenchyma cells (black arrow) are completely flattened whereas TEs were either fully collapsed (red arrow), partially collapsed (yellow arrow) or intact (blue arrow). **C** Relative proportion of 10- to 50-day-old TEs from iPSCs that were fully collapsed, partially collapsed, or intact after CPD; n = 7–50 individual cells per cell type and time-point. **D** Relative proportion of 10- to 50-day-old TEs from iPSCs that were fully collapsed, partially collapsed, or intact after air drying. Error bars represent ± SD of 2 independent experiments; n = 18–50 individual cells per cell type and time-point. **E** Scanning electron micrograph of a 30-day-old PX TE after air drying. **F** Scanning electron micrograph of a 30-day-old MX TE after air drying. **G** Scanning electron micrograph of 30-day-old unlignified TEs after air drying.

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the relative proportion of XP and XF (Fig 3B and S2). PX and SMX had smaller lumen diameters compared to PMX (Fig. 3C and S2). MXs presented higher lignin levels than PX (Fig. 3D). All TEs exhibited similar S/G residue proportions (Fig. 3E) but showed different C6/C5 benzaldehyde proportions, higher in PX than in MXs (Fig. 3F). Relative coniferyl alcohol (G CHO) levels were higher in the SMX than in PX or PMX, whereas relative terminal coniferaldehyde (G CHO) residues were present in similar amounts in all three morphotypes (Fig. 3G, H). Therefore each TE morphotype has not only specific dimensions and neighbouring cells but also specific lignin composition, amounts, structures and organisation.

TE collapse, also called irregular xylem (irx), has been previously observed in response to drought (Brodribb and Holbrook 2005; Zhang et al. 2016) but also when modifying lignification with genetic mutations (Turner and Sommerville 1997; Brown et al. 2005) or drugs (Amrhein et al. 1983; Smart and Amrhein 1985). We thus evaluated, for each different TE morphotype, the importance of lignin for the resistance to collapse using cross-sections of nine loss-of-function mutants altered in lignin concentration and/or composition. We measured area, perimeter, circularity, convexity, as well as lignin structure using in situ quantitative imaging (Blaschek, Nuendagula, et al. 2020; Blaschek, Champagne, et al. 2020; Yamamoto et al. 2020). The perimeter or neighbouring cell types remained similar for all TE morphotypes between mutants (Fig. S2). TE circularity (indicating TE general deformation) and convexity (indicating TE inwards collapse), were however altered for specific TE morphotypes in distinct mutants. Plants mutated in CCR1 exhibited significantly reduced circularity and convexity in all TE morphotypes (Fig. 4A and S2). Mutants mutated in both 4CL1 and 4CL2 showed TE collapse only in PX and PMX, mutants in CCaOMT only affected PMX and mutants in both CAD4 and CAD5 were only compromised in PX (Fig. 4A and S2). Different TEs in each mutant exhibited significant differences in lignin structure (Fig. S3). These results revealed that specific changes in lignin, altering the C6 and C5 parts of lignin residues, differently control the biomechanical properties of each TE morphotype.

We computed structural equation models to link the morphological and biochemical changes for each TE morphotype. PX collapse was prevented by increases in lignin amount, G CHO and terminal G CHO contents and TE perimeter (Fig. 4B). Only S residue content escalated collapse, whereas G CHO position and the proportion of neighbouring TEs had no effect (Fig. 4B). In contrast, PMX collapse was prevented by increased lignin amount, total and terminal G CHO levels but was negatively affected by increases in S and neighbouring TEs (Fig. 4C). This result complemented recent observations that showed collapse of PMX when genetically increasing their S content (Sakamoto et al. 2020). TE perimeter and G CHO content did not affect PMX collapse (Fig. 4C). SMX collapse was prevented by increases in G CHO but enhanced by increases in benzoaldehydes (Fig. 4D). TE perimeter, number of neighbouring TEs, S and G CHO contents did not affect the collapse of SMX (Fig. 4D). We also performed interaction analyses to determine the interdependence between morphological and biochemical features (Fig. S4). In the PX, the synergistic strengthening effects of G CHO and G CHO against TE collapse were dependent on TE perimeter (Fig. S4). In the PMX, the effects of G CHO were similarly synergistic in preventing collapse with total lignin, but depended on vessel adjacency (Fig. S4). Overall, TE susceptibility to collapse was prevented for each TE morphotype by specific changes and interactions between cell/tissue morphology and lignin amount, composition (S/G and G CHO/G CHO compositional ratios) and structure (G CHO terminal to total). Our results demonstrate that the different lignin residues have non-redundant roles and need to be specifically coordinated to determine the biomechanical properties of each TE morphotype.

TE resistance to negative pressure in woody plants depends on tissue organisation and post-mortem incorporation of specific subunits. We similarly evaluated TE lignification and resistance to collapse in woody whole plants of hybrid poplar by focusing in cross-sections of poplar stems on different TE types, primary (PV) and secondary xylem TE/vesels (SV), and different developmental states, young and old SVs defined within or beyond the 50% distance to the cambium (Fig. 5A). All TEs in WT plants had similar surrounding cell types but varied in the lumen area, which was smallest in young SVs (Fig. 5B, C). In situ analysis showed differences in lignin
levels (lowest in PVs) and S/G composition (highest in young SVs and gradually decreasing in old SVs and PVs) (Fig. 5D, E). Benzaldehyde and GCHO levels (higher in PVs than SVs), and terminal GCHO levels (gradually increasing from young SVs to PVs) revealed changes between TE types as well as during their development (Fig. 5F–H). Complete analysis of TE maturation across wood sections showed gradual increases of lignin levels but also changes in S/G and GCHO/GCHO (Fig. S5). These results show that dynamic post-mortem lignification of TEs occurs in whole woody plants, and lead to different lignin structures.

To assess the link between lignin and TE collapse, we used two genotypes with reduced lignin amounts and composition by down-regulating C4H and CCR. Lignin-reduced poplar plants had TEs with both significantly reduced circularity and convexity only in SVs (Fig. S1 and S6) but major changes in lignin composition and amounts were also observed for all TEs (Fig. S6). Structural equation models were computed to define the parameters affecting the collapse of each TE type as well as during SV post-mortem maturation. As PVs did not collapse, none of the measured parameters had any effect, thereby showing unique resilience of PVs (Fig. 5I). In contrast, old SVs collapse was prevented by increases of GCHO and lignin levels but promoted by increases in terminal GCHO and P residues as well as increases in neighbouring TEs (Fig. 5K). The collapse of young SVs was prevented by increases of GCHO but promoted by increases of S, terminal GCHO and P residues, TE perimeter and neighbouring TEs (Fig. 5L). Similar to Arabidopsis, some of these effects were interdependent on each other. The effects of P and GCHO residues in old SVs were modulated by their adjacency to other vessels, while the negative effect of GCHO depended on high levels of P residues (Fig. S5). In young SVs, the effects of S, GCHO and P values were similarly adjusted by each other and vessel perimeter/adjacency (Fig. S5).

Altogether, these results show that the non-redundant combinatorial effects of lignin structure and cell neighbours on the biomechanical properties of specific TE types are dynamically modulated during their post-mortem maturation.

Lignin residues have non-redundant roles in controlling specific mechanical properties. To link the observed changes in lignin to whole plant mechanical properties, we performed flexure measurements using three point bending experiments on Arabidopsis plant stems (Fig. 6A and S7). We first evaluated the influence of turgor pressure and sap content on stem biomechanical properties by comparing flexure measurements of stem segments incubated in air, pure water or 1 M sorbitol for several hours. Reducing water content in stems with sorbitol significantly reduced stem flexural strength and flexibility but not stiffness (Fig. 6B–C and S7). We then used two Arabidopsis mutants to investigate how stem mechanical properties were affected by changes in lignin composition and concentration, specifically in the residues affecting TE collapse: the fah1 mutant, devoid of S and instead accumulating GCHO (Meyer et al. 1998; Blaschek, Nuoendagula, et al. 2020; Yamamoto et al. 2020), and the cad4×cad5 mutant enriched in total and terminal GCHO (Blaschek, Champagne, et al. 2020; Sibout et al. 2005; Yamamoto et al. 2020). Biochemical analyses showed minor changes in lignin amounts, lower in fah1 and cad4×cad5 compared to WT plants, but major changes in S/G and GCHO/GCHO as well as terminal/total GCHO (Fig. 6D and S7). Flexural strength and stiffness were significantly increased ~2-fold in the S devoid mutant, but significantly decreased ~3-fold in the GCHO enriched mutant, compared to the WT (Fig. 6E and S7). In contrast, flexibility was similar between fah1 and WT but significantly increased ~3-fold in cad4×cad5 (Fig. 6F). Altogether, our results showed that S/G controlled stem flex-
Figure 5 | Different TE morphotypes depend on specific post-mortem accumulated lignins for their resistance against collapse. A. Scheme of the three TE types in the xylem of poplar stems, oriented on the pith–cambium axis: primary vessels (PV) in yellow, old secondary vessels (old SV) in purple and young secondary vessels (young SV) in blue. B. Relative proportion of adjacent TEs to each TE types. Note that the proportion of TEs neighboring other TEs is independent of TE type and very similar to the proportions in A. thaliana. C. Area of each TE type determined from cross sections. D–H Relative amounts of cell wall lignin (D), S/G compositional ratio (E), relative benzaldehyde levels (F), GCHO levels (G) and terminal GCHO levels (H) for each TE morphotype determined by Raman microspectroscopy. Letters in panels B–H indicate significant differences according to a Tukey-HSD test (per panel; \( \alpha = 0.05 \); \( n = 56–72 \) individual cells from 5 individual plants per TE type). I. Representation of TE perimeter for each TE type in transverse cross-sections from stems of Populus tremuloides RNAi plants altering lignin biosynthesis. J–L Structural equation models of the factors influencing TE convexity and circularity in the PV (J), old SV (K) and young SV (L). Blue arrows and positive standardized coefficients indicate significant positive effects, red arrows and negative standardized coefficients indicate significant negative effects. Dashed arrows indicated predictors that were included and improved the model, but whose specific effects were not statistically significant. Greyed out variables did not improve the model and were excluded. See also supplementary figures S4–S6 and supplementary tables S1 and S2.

Increased TE flexibility due to lignin composition enables plants to better resist drought. We then evaluated the role of TE flexibility due to \( \text{GCHO} \) residues on plant sap conduction capacity in response to normal watering or extreme \( \Psi \) changes using simulated drought. Under normal condition, 4–5-week-old WT plant rosettes had evapotranspiration rates at 7.5 mg water loss per min, which was significantly reduced by \( \sim 25\% \) under simulated drought with 10% and 20% PEG\textsubscript{6000} solution for 72h (Fig. 7A–C and S7). Leaf wilting and chlorosis were visible under simulated drought and accentuated by PEG\textsubscript{6000} treatment (Fig. 7A, B). Recovery experiments by transferring plants to normal watering for 96 h showed that plants in 20% PEG\textsubscript{6000} did not recover whereas only one out of 8 plants in 10% PEG\textsubscript{6000} fully recovered (Fig. 7D). In the \textit{cad4}×\textit{cad5} mutant, evapotranspiration rates gradually decreased with increasing PEG\textsubscript{6000} levels (Fig. 7A–C and S7). However, the mutant plants showed less wilting and chlorosis than the WT, resembling the untreated plants (Fig. 7A, B). Recovery experiments to normal watering showed that 35–47% of \textit{cad4}×\textit{cad5} plants fully recovered after both 10% and 20% PEG\textsubscript{6000} treatments (Fig. 7D). Overall our results showed that the increased flexibility conferred by \( \text{GCHO} \) residues to TEs enables plants to better resist the extreme \( \Psi \) changes caused by drought. We therefore demonstrated that lignin structure directly controls the hydraulic properties of TEs to cope with environmental changes.

Discussion

Plants are unique in our biosphere in having an open vascular system that is driven by environmentally controlled water potential gradients (\( \Psi \)). Despite this dependence on external conditions, plants possess an impressive capacity to survive and adapt in challenging habitats. Lignin in plant vascular tissues has until now been treated as a randomly assembled polymer with a structural role but no specific cellular or physiological function. Lignin residue diversity was however acquired several times convergently by different vascular plant genera during plant evolution (Weng et al. 2008). We recently questioned this “random” concept by showing that different xylem cell types, and their morphotypes, accumulated lignin \( \text{GCHO} \) residues using cell-specific genetic, cooperative and oxidative processes (Blaschek, Nuoendagula, et al. 2020; Blaschek, Champagne, et al. 2020; Yamamoto et al. 2020). We also showed that \textit{post-mortem} lignification of TEs occurs in multiple plant species (Pesquet et al. 2010; Pesquet et al. 2013). Yet the physiological roles of TE \textit{post-mortem} lignification and of specific lignin in each TE morphotype remained unclear. We herein demonstrated for the first time that each TE morphotype accumulates specific lignins (composition, concentration
and structure) which are dynamically modified during its post-mortem cell wall maturation. We showed that the different lignin residues had non-redundant effects on TE biomechanical properties. Lignin S/G affected the stiffness and flexural strength whereas G CHO/G CHO modulated the flexibility, together with fine-tuning due to G CHO position, benzaldehydes and P residues as well as neighboring TE proportion and TE perimeter (Fig. 2–6 and S7). While increased G CHO had previously been suggested to compromise stem biochemical properties (Özparpucu et al. 2017; Özparpucu et al. 2018), we suggest that lignin composition finely regulates the trade-off between stiffness and flexibility.

Figure 6 | Distinct lignin monomers control non-redundantly specific mechanical properties. A Arabidopsis thaliana stem segment undergoing three-point-bending. Note that flexural behavior is presented in movie S1. B, C The flexural stiffness (B) and sustained elastic deformation before irreversible breaking, i.e. flexibility, (C) of WT stem segments incubated in water, air, or sorbitol determined by three-point-bending; n = 5–8 segments from 3 individual plants per condition. Letters indicate significant differences according to a Tukey-HSD test (per panel; α = 0.05). Note that turgor pressure contributes slightly to the extent of reversible elastic deformation, but not to its stiffness. D Total lignin content measured by thioglycolic acid derivatization, S/G and G CHO/G CHO ratios measured by pyrolysis-GC/MS for the three genotypes. Values are averages ± SD of three independent replicates. E, F The flexural stiffness (E) and sustained elastic deformation before irreversible breaking (F) of stem segments from WT, S depleted fah1 and G CHO over-accumulating cad4×cad5 mutant plants determined by three-point-bending; n = 10–24 individual stems per genotype, cut into multiple segments. Letters indicate significant differences according to a Tukey-HSD test (per panel; α = 0.05). See also supplementary figure S7.

Our work moreover revises the current paradigm of plant xylem sap hydraulics, which considered TEs as static structures with no adaptive influence on plant sap conduction. We herein demonstrate that the mechanical properties of TEs due to specific lignins directly control their hydraulic properties. Altogether we show that post-mortem lignification of TEs is essential for vascular plants to dynamically adjust their vascular system to varying environmental conditions. The exact identity and function of the different cooperating cell types, the distance by which they can cooperate with TEs as well as the metabolites used and regulating molecular actors still need to be identified. Our study thereby completely revises the "random" lignification of plant cell to a genetically controlled process at the level of each cell morphotype to enable plants to thrive and dynamically adapt to changes in climate.

Figure 7 | Coniferaldehyde induced flexibility of TE lignin improves plant resistance to, and recovery from, extreme ψ differentials. A Top view of 4- to 5-week-old A. thaliana WT and G CHO over-accumulating cad4×cad5 mutant plants after being irrigated with water, 10% PEG or 20% PEG for 3 days. B Top view rosette area after being irrigated with water, 10% PEG or 20% PEG for 3 days. Letters indicate significant differences according to a Tukey-HSD test (per panel; α = 0.05). C Evapotranspiration rates of WT and cad4×cad5 plants after being irrigated with water, 10% PEG or 20% PEG for 3 days. Small grey dots represent individual measurements, larger coloured dots represent the average per plant. Letters indicate significant differences according to a Tukey-HSD test (per panel; α = 0.05). D Proportion of WT and cad4×cad5 plants that did not, partly or fully recover after being treated (with water, 10% PEG or 20% PEG for 3 days) by a 4 days recovery period in water-saturated soil. See also supplementary figure S7.

Materials & methods

Inducible pluripotent cell suspension cultures (iPSCs). Arabidopsis thaliana iPSCs were produced and induced to differentiate into isolated vessel elements as described by Pesquet et al. (2010) and Ménard et al. (2017). Cell suspensions were induced by adding hormones to 30 mg ml⁻¹ of 9 days old cells (fresh weight) in 1x Murashige and Skoog (MS) medium (Duchefa, M0222.0025) at pH 6.0 with 10 µM of morpholinio-ethanesulfonate (Sigma-Aldrich, M8250) and 3% sucrose. Xylogenic induction was triggered by adding 3.2 nM α-naphthalenediacetic acid (Sigma Aldrich, N0640), 44.4 nM 6-benzyl-aminopurine (Sigma Aldrich, B3408) and 4 µM 24-epibassinolide (Sigma Aldrich, E1641). The inhibition of lignin monomer biosynthesis was performed by adding 12.5µM of piperonylic acid (Sigma-Aldrich, P49805) at the time of hormonal induction of TE differentiation in suspension cultures as described by Van de Wouwer et al. (2016) and Decou et al. (2017).

Plant material. Arabidopsis thaliana and hybrid poplar plants were grown in climatic growth chambers under a 16/8 h long day light regime with 150 µmol m⁻² s⁻¹ illumination using (Aura T5 Eco Saver Long Life HO light tubes; AuraLight, Seweden) and 22°C/18°C in 60% humidity. Arabidopsis thaliana mutants in the Columbia Col-0 background used included ccoaomt1 (SALK_151507; Kai et al. 2008), fah1 (EMS mutant; Meyer et al. 1998), omt1 (SALK_135290; Tohge et al. 2007), del-

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Atomic Force Microscopy (AFM). AFM imaging was performed on cell samples air dried for less than 1 hour using a Dimension Icon AFM (Bruker, Nanoscope controller, Santa Barbara, CA, USA). The measurement was conducted under Peak-Force QNM mode in air condition by using the probe TESPA-V2 (Bruker). The force set-point was 0.15V. The height, peak-force error, DMT (Derjaguin-Muller-Toporov) modulus, adhesion, and deformation images were recorded after calibrating the probes on Mica. The images were processed by NanoScope Analysis 1.5 software (Bruker) and quantification performed using ImageJ distribution Fiji (Schindelin et al. 2012).

Scanning electron microscopy (SEM) coupled with Energy-dispersive X-ray spectroscopy (EDS). Cell samples for electron imaging and chemicals analysis were dispersed and sedimented on glass coverslips, then either (i) dehydrated in a series of graded ethanol and critical point dried (CPD) using a Leica EM CPD300 critical point dryer or (ii) submitted to air drying, and finally coated with 5 nm chromium using Quorum Technologies Q150T ES metal coater. The samples morphology was analyzed by field-emission scanning electron microscopy (SEM; Carl Zeiss Merlin) using an in-lens secondary electron detector at accelerating voltage of 4 kV and probe current of 100 pA. Elemental composition measurements were performed using an energy-dispersive X-ray spectrometer (EDS; Oxford Instruments X-Max 80 mm2) at accelerating voltage of 10 kV and probe current of 300 pA, where the elemental composition percentage is an average of multiple line and point analyses.

Histological preparation and analyses. Eight week old stem samples were cleared in 70% ethanol, rinsed in water and embedded in 10% agarose prior to sectioning to 50 µm with a VT1000S vibratome (Leica, Sweden). Semi-quantitative Raman microspectroscopy was performed as described by Blaschek, Nuoodagula, et al. (2020) on the different vessel types using a confocal Raman microscope (RAMANplus, Nanophoton, Japan and LabRAM HR 800, Horiba, France) with a 532 nm laser. Averaged spectra were obtained from three to seven cell walls per TE morphotype and plant, in one to three plants per genotype for Arabidopsis and from 17 to 71 cell walls per TE morphotype and plant, in two to six plants for poplar. Quantitative Wiesner test was performed as described by Blaschek, champagne, et al. (2020) using an Olympus BX60 brightfield microscope equipped with an Olympus UPFLN 40X objective (NA 0.75), an Olympus XC30 CCD colour camera. TE morphological features (distance from cambium, lumen area, perimeter, circularity, neighbouring cell types) were measured from microscopy images using the ImageJ distribution Fiji (Schindelin et al. 2012). TE convexity was determined as 4π(area/perimeter3), and TE circularity was determined as 4π(area/perimeter2), and TE convexity as area/area of convex hull. Fiji macros are available at https://github.com/leonardblaschek/fiji.

Three point flexural test. The stiffness and strength of stems were assessed using three-point flexural tests with an Instron 5966 universal testing machine (Instron, USA) equipped with a 100 N load cell in a humidity and temperature controlled room (50% RH and 23°C). 4–5 cm long stem segments from 25–35 cm stems of 6–7 week-old plants were placed on two supporting pins that were separated at an average span-to-diameter ratio of 38–39±4. Treatment to alter stem water content included incubation for several hours prior to bending in pure distilled water or 1M sorbitol (Sigma, S1876) solutions, all other measurements were performed in air. After manually lowering the loading pin until just in contact with the sample, the probe was lowered automatically at a constant displacement rate of 2 mm min-1 until a final displacement of 7 mm. The flexural strength σmax (MPa) was calculated as the maximum flexural stress using Eq. 1.

\[
\sigma_{\text{max}} = \frac{8F_{\text{max}}L}{\pi d^3}
\]  

In this equation, \(F_{\text{max}}\) (N) is the maximum force the specimen can withstand before kinking, \(L\) (mm) is the span length between the supporting pins, and \(d\) (mm) is the diameter of the circular cross-section of the specimen that was determined using optical microscopy imaging. The flexural stiffness \(E\) (MPa) was calculated from the slope of the initial linear part of the flexural stress-strain curve using Eq. 2.

\[
E = \frac{4FL^3}{3\pi Dd^4}
\]

In this equation, \(D\) (mm) is the maximum deflection of the centre of the stem. The flexibility of stems was defined as the strain at maximum stress, i.e. the amount of deformation a stem could endure before irreversibly breaking.

Evapotranspiration and simulated drought. Simulated drought treatments were made by watering plants with 0, 10 or 20% polyethylene glycol (PEG) 6000 (Sigma-Aldrich, 8,07491) in tap water for 72h in growing conditions (150µE light, 25°C, 60% relative humidity). Evapotranspiration was measured using mass difference for 20 minutes on a LA-124i microbalance (VWR) directly connected to a computer and monitored using the i-Weight software (VWR). Plant recovery were made by placing plants directly in water for 72h. Light intensity, temperature and relative humidity were constantly monitored during the course of the measurements (Fig. S7). Images of the rosettes were acquired with a Nikon D750 camera equipped with a 50mm F1.4 DG HSM lens. Image segmentation and rosette area measurements were performed in Fiji (Schindelin et al. 2012).
Lignin biochemical analysis. Lignin concentration in cell wall was determined after cell wall isolation according to Yamamoto et al. (2020) and the thioglycolic acid lignin derivatization as described by Suzuki et al. (2009) on purified cell walls. Absorbance was measured at 280 nm and calibrated using a regression curve obtained using different quantities of alkaline spruce lignin (Sigma Aldrich, 471003). Pyrolysis-GC/MS was used to measure S/G and G CHO/G CHO, according to Gerber et al. (2012) on 60 µg (±10 µg) of 8-week-old stem samples. Thioacidolysis-GC/MS-FID was used to determine the terminal/total positional ratio of β-O-4 linked G CHO residues on 5 mg (±1 mg) of isolated cell wall from 8-week-old stem samples as described by Yamamoto et al. (2020).

Data analyses and structural equation modelling. Data analysis and visualisation was performed in R (v4.0.4), using the tidyverse collection of packages (v1.3.0). The structural equation models were built using the piecewisesem package (v2.1.0; Lefcheck 2016). The included variables were measured for each individual vessel, except for the Wiesner test intensity in A. thaliana, for which the average per individual plant and vessel type was used. The multiple linear regression models comprising the structural equation models were selected using a bidirectional step-wise optimization approach, excluding interaction terms for clarity. Significantly contributing interaction terms were identified separately and visualized using the interactions R package (v1.1.1). Model fits and coefficients are summarized in table S1 and S2. R code used in this study is available at https://github.com/leonardblaschek/Rscripts/blob/master/irx_pub_figs.rmd.

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Author Contributions

EP conceived the study. EP and DM designed the experiments. DM, LBl, KK, CZ, N, CCL, ZB and EP performed the experiments. LBl, KK, CCL, DM and EP analysed the data. LBe, AM, ZB, SK and EP ensured financial support and scientific expertise. EP wrote the article. All co-authors revised the manuscript.

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