CELLULOSE SYNTHASE INTERACTING 1 is required for wood mechanics and leaf morphology in aspen

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SUMMARY

Cellulose microfibrils synthesized by CELLULOSE SYNTHASE COMPLEXES (CSCs) are the main load-bearing polymers in wood. CELLULOSE SYNTHASE INTERACTING1 (CSI1) connects CSCs with cortical microtubules, which align with cellulose microfibrils. Mechanical properties of wood are dependent on cellulose microfibril alignment and structure in the cell walls, but the molecular mechanism(s) defining these features is unknown. Herein, we investigated the role of CSI1 in hybrid aspen (Populus tremula × Populus tremuloides) by characterizing transgenic lines with significantly reduced CSI1 transcript abundance. Reduction in leaves (50–80%) caused leaf twisting and misshaped pavement cells, while reduction (70–90%) in developing xylem led to impaired mechanical wood properties evident as a decrease in the elastic modulus and rupture. X-ray diffraction measurements indicate that microfibril angle was not impacted by the altered CSI1 abundance in developing wood fibres. Instead, the augmented wood phenotype of the transgenic trees was associated with a reduced cellulose degree of polymerization. These findings establish a function for CSI1 in wood mechanics and in defining leaf cell shape. Furthermore, the results imply that the microfibril angle in wood is defined by CSI1 independent mechanism(s).

Keywords: aspen, Populus, cell wall, wood mechanics, cellulose, transgenic trees, cellulose interacting 1, CSI1, pavement cell.

INTRODUCTION

Xylem supports the upright growth of trees, providing mechanical resistance against gravity and wind, and facilitates the effective transport of water and nutrients to aerial tissues of plants (Groover et al., 2010). Xylem, or more commonly wood, is also a natural nanocomposite material, which potentiates sustainable applications that may drive a low carbon economy. The material and mechanical properties of wood is a function of the cellular architecture and the molecular interactions in the xylem cell walls. Wood of angiosperm trees consists of three main cell types. The water and nutrient transporting vessels, the ray cells involved in nutrient storage and radial transport, and the load-bearing xylem fibres that provide structural support (Groover et al., 2010). Dimensions of fibres and wood density are important determinants of the mechanical properties of wood (Beery et al., 1993), as is the angle at which cellulose microfibrils (CMFs) are laid down in the fibre walls (Evans and Elic, 2001; Barnett and Bonham, 2004). In wood, the CMFs are surrounded by heteropolysaccharide hemicelluloses and the complex phenolic lignin, which also are important for the mechanical performance of wood (Gibson, 1992). Understanding how wood formation and cell wall biosynthesis define the mechanical performance is a key aim in wood biology. Identification of the genes, the regulatory network and the master regulators of these processes are important and
relevant for tree breeding and biotechnological approaches focusing on improving the properties of timber and other wood-derived products.

Wood formation initiates in the cambial cell division zone. During cell expansion, the xylem fibre cell wall consists of a middle lamella between adjoining cells and a primary cell wall, which collectively combine to form the compound middle lamella. Plant cell expansion and the final cell dimensions are dictated by turgor-driven anisotropic extension of the primary cell wall, which largely depends on CMF orientation and their interactions with other matrix polymers (Cosgrove, 2005). In addition to anisotropic expansion, xylem fibres grow by intrusive tip growth where the tip of the cell grows in between neighbouring cells (Gorshkova et al., 2012). Upon reaching their final size, xylem fibres initiate the synthesis of secondary cell walls, which at maturity typically contain two to three layers (Kerr and Bailey, 1934; Bailey and Vestal, 1937). The layer structure of the secondary fibre walls originates from altering orientation of the CMFs. The layers of the fibre secondary walls are classified as the outermost S1 layer (adjacent to the primary wall), the middle S2 layer representing the bulk of the total wall thickness, and the innermost S3 layer (Harada, 1962; Fengel and Stoll, 1973; Fengel et al., 1988). The mechanical properties of wood along the longitudinal fibre cell axis are particularly affected by the orientation of parallel CMFs in the S2 layer (Preston, 1974).

The orientation of the CMFs in relation to the longitudinal axis of the fibre cell is defined as the cellulose microfibril angle (MFA). Generally, a low average MFA confers stiffness and strength to the wood, while a higher average MFA results in flexibility and toughness (Reiterer et al., 2001). The molecular underpinnings defining the MFA in xylem fibres are not known, but immunofluorescence microscopy observations support a role for cortical microtubules (MTs) during both primary and secondary wall biosynthesis (Funada et al., 1997; Barnett and Bonham, 2004). Immunostaining of developing xylem fibres using a tubulin antibody revealed parallel-arranged MTs, which were associated with secondary wall thickening in hybrid aspen (Populus tremula × tremuloides) and horse chestnut (Aesculus hippocastanum) (Chaffey et al., 2000; Chaffey et al., 2002). Chaffey et al. (2002) observed variation in MT orientation between neighbouring fibres undergoing secondary cell formation and hypothesized a connection to the different CMF alignments in the S2, S1 and S3 layers. Experiments in Arabidopsis using MT-disrupting chemicals and MT defect mutants support a functional link between MTs and CMFs (Roberts et al., 2004; Oda et al., 2005; Wightman and Turner, 2008). Live cell imaging of fluorescent protein-tagged cellulose synthases (CESAs) in Arabidopsis established that cellulose synthase complexes (CSCs) track along the cortical MTs, which align with CMFs (Paredez et al., 2006). This observation was made first during primary cell wall biosynthesis in etiolated hypocotyls and subsequently during secondary cell wall formation in protoxylem vessel-induced Arabidopsis seedlings (Paredez et al., 2006; Wightman and Turner, 2008; Watanabe et al., 2015; Li et al., 2016; Watanabe et al., 2018).

Genetic and protein interaction studies aimed at finding linker proteins between MTs and CSCs identified an armadillo/beta-catenin-like repeat containing protein called CELULOSE SYNTHASE INTERACTING 1 (CSI1)/POM2 (Gu et al., 2010). CSI1 can guide the CSCs along the cortical MTs during primary wall biosynthesis (Li et al., 2012; Bringmann et al., 2012a), and performs this function by interacting with both CSCs and MTs (Li et al., 2012). Arabidopsis csi1 mutants are impaired in cell expansion and display twisting epidermal cell files and leaves (Bringmann et al., 2012b; Landrein et al., 2013). Initial analysis of csi1 mutants did not show defects in secondary cell walls (Gu and Somerville, 2010) but subsequently fluorescent CSC imaging established a role for CSI1/POM2 during the initial phase of secondary cell wall pattern establishment in xylem vessels (Schneider et al., 2017).

CSI1 analyses during secondary cell wall biosynthesis have been limited to Arabidopsis xylem vessels, which lack the lamellar wall structure observed in xylem fibres. Here, we investigated the function of CSI1 in developing wood, and in particular xylem fibre formation in hybrid aspen (Populus tremula × tremuloides) using RNA interference of functional orthologues of the Arabidopsis CSI1.

RESULTS

**Populus CSI1s are functional orthologues of the Arabidopsis CSI1**

The *Populus* genome encodes two putative orthologues of the Arabidopsis CSI1 (Figure 1a). PtcCSI1A and PtcCSI1B are 95% similar to each other at the amino acid sequence level (Figure S1). To first test whether the *Populus* CSI1A can perform the same function as the Arabidopsis CSI1 we expressed the hybrid aspen (Populus tremula × tremuloides) PttCSI1A in the Arabidopsis csi1 null mutant pom2-4. The PttCSI1A construct complemented the pom2-4 silique and inflorescence length phenotypes confirming that PttCSI1A is indeed a functional orthologue of the Arabidopsis CSI1 (Figure S2). Based on publicly available transcriptome data, PtcCSI1A and PtcCSI1B are expressed in both leaves and stem (popgenie.org and http://aspwood.popgenie.org/aspwood-v3.0/). In the stem, the transcript abundance of both genes increase from cambium across the primary wall and cell expansion zone, peak at the onset of the secondary cell wall formation, and then decrease rapidly at the onset of cell death and xylem maturation (Figure 1b). Developing aspen wood also contains transcripts of the Arabidopsis CSI3 homolog, but the transcript
level does not show obvious changes during wood formation (Figure 1b). These developing wood transcript profiles suggested a function for PtCSI1A and PtCSI1B during xylem cell expansion and secondary cell wall formation. To study the role of PttCSI1 in trees, transgenic hybrid aspen containing a 35S promoter driven PttCSI1RNAi construct targeting both CSI1A and CSI1B were generated and grown under greenhouse conditions for 2 months, to a height of approximately 1.5 m. Quantitative polymerase chain reaction (qPCR) analysis of PttCSI1A and PttCSI1B showed significant reduction in the transcript abundance of both genes in leaves and developing wood of three independent transgenic lines (Figure 1c–f).

**CSI1RNAi causes leaf twisting and defects in pavement cell shape**

The greenhouse grown CSI1RNAi lines showed a modest growth phenotype (Figure 2a). The average stem diameter was reduced in lines 1 and 3, and the stem height was slightly reduced in all transgenic lines (Table 1). The most obvious visual phenotype was a reduction in leaf size and the appearance of occasional twisting of the leaves (Figure 2b and Table 1). This was reminiscent of the twisting rosette leaves and other tissues observed in the Arabidopsis csi1/pom2 null mutants (Bringmann et al., 2012a; Landrein et al., 2013). Interestingly, light microscopy inspection of the CSI1RNAi leaf epidermis revealed that the pavement cells lacked the multi-lobed jigsaw puzzle shape observed in the wild-type (WT) cells (Figure 2c). CMFs are known to mediate directional cell growth and the leaf epidermis plays an important role in defining leaf shape and size by bearing the stress caused by leaf growth. Hence, it seems likely that the leaf area reduction and twisting in the CSI1RNAi lines could be due to misaligned CMFs manifesting in defects in pavement cell and leaf shape formation.

**CSI1RNAi wood is mechanically weaker**

The phenotypic changes in the CSI1RNAi lines and the reduced CSI1 transcript levels in the developing wood indicated that the CSI1RNAi lines could be used to investigate CSI1 function during wood formation and secondary growth. To this end, wood from WT, CSI1RNAi lines 1 and 3 was analysed in detail. Light microscopy of wood cross-
sections in the CSI1RNAi lines showed no obvious differences in the overall cellular anatomy among the transgenic lines and WT trees (Figure 3a). However, we observed that the 20-μm thick wood cross-sections derived from CSI1RNAi wood were more prone to cracking during sample preparation. The cracks occurred across the secondary cell walls of both fibres and vessels indicating that the mechanical integrity of the wall was affected in both cell types (Figure 3b). Transmission electron microscopy (TEM) was used to produce images of the secondary cell wall layer structure of xylem fibre walls, but no obvious changes in the thickness of the S1 or S2 layers or the structure of the fibre walls was observed (Figure 3c).

To examine the cell wall structure and mechanical properties further, we compared WT and CSI1RNAi wood in micromechanical tensile tests. Longitudinal-tangential wood strips with dimensions of 30 × 2 × 0.1 mm were strained to failure using a microtensile testing stage. The stiffness (modulus of elasticity, MOE) in longitudinal direction and the maximum force required to break the sample (modulus of rupture, MOR) were determined from the stress-strain curves. Significant differences were observed in both MOE and MOR between PttCSI1RNAi lines and WT trees. The MOE was reduced approximately by one-third, from 2.95 GPa in the WT to 2 GPa in the transgenics (Figure 4a). The MOR was also reduced in both lines although the difference was only significant for line 3 (Figure 4b).

**CSI1RNAi wood density and average MFA are not changed**

To investigate the origin of the observed mechanical weakness in CSI1RNAi wood we compared the cell wall composition in the transgenic lines and WT. No consistent differences were observed in lignin, cellulose or hemicellulosic sugar content (Table S2). These results indicated that the mechanical wood phenotype was not linked to changes in the composition of the cell wall matrix.

Based on the leaf twisting phenotype, the defects in the pavement cell shape and the CSI1 function in connecting CSCs and MTs, we hypothesized that the cellulose MFA could be affected in the CSI1RNAi wood. The average wood MFA of the S2 layer of xylem fibre walls was determined using X-ray diffraction. Surprisingly, no consistent difference between WT and the transgenic lines was observed in normal wood (Table 2). In line 3, there was a tendency towards a slight MFA increase, but this was not statistically significant. The MFA agrees with those reported in the literature for the S2 layer in juvenile wood of *Populus* sp. (Barnett and Bonham, 2004). We also sampled stem areas with apparent tension wood fibres and gelatinous layer (G-layer) formation where a bimodal orientation distribution of CMFs is seen with very small MFAs for the G-layers and larger angles (>30°) for the S2 layer of tension wood (Müller et al., 2006). However, this mechano-gravitropism induced MFA shift was near-identical between the WT and the CSI1RNAi lines (Table 2). These MFA results showed that the mechanical wood phenotype cannot be explained by a change in the orientation of the CMFs in the secondary cell wall layers, and indicated that CSI1 may not be needed for the alignment of CMFs during secondary cell formation in wood fibres in aspen trees.

**CSI1RNAi reduced the xylem fibre area and the degree of cellulose polymerization**

The longitudinal xylem fibre area measured from light microscopy images of macerated wood showed a consistent reduction by approximately 20% (Table 2). These data suggested that CSI1RNAi reduced the extent of fibre cell expansion. This is in line with the leaf data and suggested a defect during primary wall cellulose biosynthesis also in
In addition to guiding the CSCs along the MTs, CSI1 mutations have been shown to reduce the average speed of CSCs in the plasma membrane in Arabidopsis (Gu et al., 2010). A similar slowdown of CSCs during wood formation could lead to shorter cellulose chains if the lifetime of the CSCs at the plasma membrane does not change. To compare the degree of cellulose polymerization between WT and CSI1RNAi lines, cellulose was extracted from stem wood using peracetic acid (PAA) extraction. PAA is effective for lignin removal while minimizing the impact on CMF degradation (Poljak, 1948; Kumar et al., 2013). After the PAA extraction and solvent exchange cellulose was dissolved in a lithium chloride/N,N-dimethylacetamide (LiCl/DMAc) solution, which is a non-degrading solvent for cellulose (Potthast et al., 2015). To measure the absolute molecular weight of the cellulose fraction we used size exclusion chromatography (SEC) coupled to a laser light scattering (LS) detector, which allows the direct measurement of the molecular weight of polymers in solution (Einsteins, 1910; Wyatt, 1993; Podzimek, 1994). The SEC/LS data indicated a slight decrease in the number average molar mass ($M_n$) and particularly in the weight average molar mass ($M_w$) of cellulose in the CSI1RNAi lines (Table 2). The $M_n$ values are the arithmetic average molecular weight of all cellulose molecules, whereas the $M_w$ is the mass averaged molecular weight ($M_w = \sum M_i^2 / \sum M_i$). Thus, $M_w$ is more sensitive than $M_n$ to decreases in the amount of the higher molecular weight cellulose molecules. This shift in the molecular weight distribution of CMFs is best illustrated by

![Figure 4](image-url)

**Figure 4.** Mechanical properties of wild-type (WT) and CSI1RNAi hybrid aspen (Populus tremula × tremuloides) wood. (a) Modulus of elasticity (b) modulus of rupture (c) density. Error bars represent ± SD (n = 5 (WT and CSI1RNAi-1) and n = 6 (CSI1RNAi-3) biological replicates. Means not sharing a common letter are significantly different at P < 0.05, as determined by Tukey’s test after one-way ANOVA.

| Line          | Average cellulose MFA (°) | Fibre area (µm²) |
|---------------|---------------------------|-----------------|
| WT            | 17.9 ± 2.7³                 | 9655 ± 65³      |
| CSI1RNAi-1    | 16.4 ± 1.8³                 | 7810 ± 314⁴     |
| CSI1RNAi-3    | 20.7 ± 4.4³                 | 7409 ± 576⁵     |

X-ray diffraction determined average MFA: Mean ± SD. Normal wood: WT and CSI1RNAi-1 five trees and CSI1RNAi-3 six trees, three to 10 wood sections per tree. Tension wood: WT two trees, CSI1RNAi-1 five trees and CSI1RNAi-3 three trees, one to nine wood sections per tree. Fibre area and fibre-tip length: mean ± SD, n = 5 (WT and CSI1RNAi-1 and CSI1RNAi-3; molecular weight of cellulose: mean ± SD, n = 4–5 (WT and CSI1RNAi-1 and CSI1RNAi-3). Means not sharing a common letter are significantly different at P < 0.05, as determined by Tukey’s test after one-way ANOVA.
comparing the SEC/LS graph of WT and the CSI1RNAi line 3 (Figure 5a). The $M_w$ and $M_n$ values were not significantly different between WT and transgenics (Table S3). However, a multivariate analysis using orthogonal projections to latent structures discriminant analysis (OPLS-DA) showed that the cellulose degree of polymerization (DP) in line 3 is significantly different from WT, while line 1 lies between the two (Figure 5b). OPLS-DA is suitable for determining the difference between groups (Bylesjö et al., 2006). The horizontal component of the OPLS-DA score scatter plot shows the variation between the genotypes while the vertical dimension captures within genotype variation. The reduction in molecular weight distribution correlated with the reduction in ultimate stress in lines 1 and 3. These results support a role for CSI1 in defining the cellulose DP in wood, and represent a structural change associated with the mechanical phenotype.

DISCUSSION

Molecular mechanisms defining the mechanical properties of wood are poorly understood. The mechanical properties derive from cellular and molecular interactions at tissue and cell wall level. CMFs are the main load-bearing and tension resisting components in the cell walls. CSI1 was identified in Arabidopsis as a linker between cortical MTs and CSC with a role in guiding cellulose biosynthesis in the primary cell walls. We identified functional orthologues of CSI1 in aspen and show that they control the CMF length that, in turn, impacts the mechanical properties of the wood. In the developing wood of aspen CSI1A and CSI1B are expressed during primary cell wall biosynthesis and cell expansion as well as secondary cell wall formation (Figure 1). Reduction of CSI1A and CSI1B transcript level in the leaves to 50–20% of WT results in leaves that twisted, likely due to the defects in pavement cell expansion and consequently cell shape (Figure 1). This is fitting with the results from Arabidopsis showing that the leaf pavement cell shape depends on cortical MT-dependent wall reinforcements in the neck regions of the cells (Fu et al., 2005). Live imaging of fluorescently labelled MT has shown that the cortical MTs align along the maximal tensile stress, which then regulates pavement cell shape by controlling cellulose biosynthesis (Sampathkumar et al., 2014). Hence, the leaf results of the CSI1RNAi lines suggest that CSI1 is involved in guiding CSCs in the pavement cells and consequently the directional cell expansion and leaf growth in aspen. Wood growth is mainly radial, while primary growth typically expands in three directions at the same time. The mechanical forces shaping primary growth and epidermal cell walls differ from forces in developing wood, which grows against the pressure established by the phloem tissues (including bark). It is also worth noting that the secondary cell wall layers of xylem fibres and vessels are formed before the stresses they experience after maturation and cell death. Hence, the patterns in xylem secondary cell walls appear genetically hardwired and less plastic than in the primary walls. An example of this developmental cell wall pattern program is seen in suspension-cultured plant cells, which can be induced to form protoxylem vessels in the absence of any tissue-derived mechanical signals (Fukuda, 1996). The differences between primary and secondary cell wall growth suggests that the mechanosensing processes guiding primary cell wall cellulose biosynthesis may not explain CMF patterning in secondary cell walls. The reduction of CSI1A and CSI1B transcript abundance in the developing wood of the CSI1RNAi lines was shown to be 10–30% of WT (Figure 1), and effectively allowed us to test the functional role of CSI1-mediated CSC guiding during wood formation and secondary growth in aspen. In the developing wood CSI1A and CSI1B transcript levels are abundant after cell expansion, suggesting a functional role in secondary cell wall formation (Figure 1). In support of this, the CSI1RNAi stem sections

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Figure 5. Size exclusion chromatogram analysis of wild-type (WT) and CSI1RNAi hybrid aspen (Populus tremula × tremuloides) stem wood cellulose. (a) Molecular weight distribution of cellulose from WT and CSI1RNAi line 3. (b) Molecular weight distribution of cellulose analysed using orthogonal projections to latent structures discriminant analysis (OPLS-DA). Scores plot shows the separation between WT (triangles) and CSI1RNAi line 1 (pentagons) and line 3 (squares). Each symbol represents one replicate tree. OPLS-DA R2X (cum) 0.82. Weight fraction (WF), derivative of logarithmic molecular weight (dLog MW).
were more brittle perpendicular to the direction of the fibre (Figure 3), and mechanically weaker in the longitudinal direction (Figure 4). We observed no significant differences in the chemical composition of wood or wood density that could explain the mechanical weakness (Figure 4 and Table S2). The fibre cells were approximately 20% smaller in the CSI1RNAi lines (Table 2). The reduced fibre area points to a defect in fibre cell expansion in agreement with the pavement cell defects in leaves and published results from Arabidopsis csi1 mutants (Gu et al., 2010). However, a correlation with the CSI1RNAi fibre size and the mechanical phenotype cannot currently be substantiated in the literature, as mechanical changes associated with wood fibre size change typically also coincide with other changes such as density and cell wall chemistry.

CMFs are the main load-bearing elements in the longitudinal direction, and based on the CSI1 function in CMF alignment during primary wall biosynthesis, we hypothesized a CSI1RNAi effect on MFA in wood. Experiments with both wood and isolated fibre samples have shown that the changes in the S2 MFA correlate with changes in the resistance to tensile stress and elastic deformation of the fibre wall or woody tissue (Preston, 1974). We used X-ray diffraction to measure the average MFA in the S2 layer of fibre walls and in areas showing characteristics of a typical tension wood G-layer MFA angle. These measurements did not reveal consistent differences in the average MFA of CSI1RNAi and WT (Table 2). The MFA in the tension wood S2 layer is similar or higher compared with normal wood S2, while in the G-layer the MFA in the same cells changes to near parallel to the fibre axis (Clair et al., 2010). Hence, the lack of a MFA phenotype in CSI1RNAi G-layers suggests that CSI1 is not involved in the dynamic secondary cell wall MFA response to mechanical and gravity cues (Table 2). The lack of MFA phenotype in the CSI1RNAi trees is in line with the Arabidopsis results showing that CSI1 is not needed for the maintenance of the CMF pattern in xylem vessels (Schneider et al., 2017). We cannot exclude the possibility that the CSI1RNAi lines still contained sufficient residual CSI1 activity during secondary cell wall formation. However, the transcript levels in developing wood were reduced more than in the twisting leaves providing some support against remaining CSI1 activity as the reason for no apparent MFA phenotype in wood.

We discovered a shift in the cellulose DP distribution towards shorter CMFs in the transgenics (Figure 5). The reduction in cellulose DP could manifest from one of several possibilities, including but not limited to, CSI1RNAi reduced half-life of CSCCs in the plasma membrane and/or reduced rate of cellulose biosynthesis. The latter possibility is supported by experiments in Arabidopsis showing reduced velocity of YFP-CESA6 particles in the hypocotyls of csi1 mutants (Gu et al., 2010; Lei et al., 2013). In a subsequent study, Lei et al. (2015) showed that CSI1 is also involved in the formation of endocytosis-derived CSC containing vesicles in the vicinity of the plasma membrane. Thus, it is possible that CSC endocytosis defects in the CSI1RNAi could cause shortening of the CMFs. We hypothesize that CMF length contributes to the mechanical strength of wood, and that the reduction in cellulose DP contributed towards the mechanical wood phenotype. Interestingly, the reduction in cellulose DP and MOR in longitudinal fibre direction appeared linked (Figures 4b and 5b) indicating that the length of the celluloseic glucan chains increases the strength of wood. However, it is important to note that any kind of chemical treatment and extraction is likely to decrease the cellulose DP. Hence we cannot exclude the possibility that the CSI1RNAi cellulose DP is equal to WT in situ, but the CMFs have structural defects making them more sensitive than WT microfibrils to the PAA extraction and LiCl/DMAc solubilization. Despite the analytical limitations in the SEC-LS procedure, it can be concluded that CSI1RNAi affects cellulose biosynthesis in wood. We hypothesize that CSI1RNAi caused reduced cellulose DP and/or structural changes in the CMFs, which may affect CMF interaction with the other matrix components, thereby contributing to weakening of the wood. To dissect the aspen CSI1 function further it will be informative to create CSI1 null mutants using the CRISPR genome editing technology.

**EXPERIMENTAL PROCEDURES**

**Plant material and growth conditions**

Transgenic and WT hybrid aspen (Populus tremula × tremuloides) trees were micropropagated in vitro for 4 weeks and then transferred to the greenhouse for further growth in commercial soil and fertilizer mixture (Hasselfors Garden Planteringsjord, https://www.hasselforsgarden.se) under an 18-h light/6-h dark photoperiod at a temperature of 22°C/15°C (light/dark) and 50%–70% humidity. The trees were fertilized using 150 ml 1% Rika-Å (N/P/K, 7:1:5; Weibull Horto, SW Horto AB, Hammenhög, Sweden) once a week for the first 3 weeks of greenhouse growth.

The trees were harvested after 8 weeks of growth in the greenhouse. Stem diameter was measured 10 cm above the soil using digital calipers, while stem height was determined as the distance between the soil surface and the shoot tip. Wood samples were collected for gene expression analysis, wood anatomy, wood chemistry and mechanical tests were collected from a stem section 10–60 cm above the soil. Samples for gene expression analysis were immediately frozen in liquid nitrogen, while the samples for chemistry and mechanical tests were placed on dry ice and stored at −80°C. Stem wood anatomy sections were prepared from material (stored at −80°C until use) cut 10 cm above the soil, while samples used for fibre maceration and cellulose degree of polymerization analysis originated from 40 to 50 cm above soil and were dried and stored at room temperature until use. Fully expanded leaves for transcript abundance analysis were collected, frozen in liquid nitrogen and stored at −80°C until use. The area of fully expanded leaves was determined using a LI-3000C Portable Leaf Area Meter (LI-COR Bioscience,
www.licor.com) on the leaf number 17, counting down from the first fully emerged leaf at the top of the tree.

**CS11RNAi vector construction, hybrid aspen transformation and qPCR**

The RNA interference cassette was created in the pBluescript SK+ vector using a 171-bp fragment targeting PtCSI1A (Potri.007G087200) and PtCSI1B (Potri.005G080100). Hybrid aspen was transformed as described by Nilsson et al. (1992). Quantitative real-time PCR was used to determine PtCSI1A and PtCSI1B transcript abundance. **UBIQUITIN** transcript abundance was used as a reference gene. Primers for PtCSI1A and PtCSI1B are listed in Table S1.

**Complementation of Arabidopsis pom2-4**

Total RNA extracted from *Populus tremula × tremuloides* wood was transcribed into cDNA using SuperScriptIII Reverse Transcriptase Kit. CS1A coding sequence (6456 bp) was PCR amplified from the corresponding cDNA using Phusion High-Fidelity DNA polymerase (Thermo Fisher Scientific, www.thermofisher.com) with primers specifically containing Gibson cloning sequence (Table S1) and cloned into Gateway entry vector pDONR207 using HiFi DNA Assembly cloning Kit (E5520S; New England Biolabs, www.neb.uk.com) following the manufacturer’s instructions. The presence of the CS1A sequence was confirmed by sequencing and then recombinated into the destination vector pH2GW7 using Gateway LR Clonase Enzyme mix (Thermo Fisher Scientific). The CS1A construct, under the control of the 35S promoter, was introduced into the Arabidopsis pom2-4 mutant by Agrobacterium tumefaciens using the floral dipping method (Clough and Bent, 1998). Transgenic lines were selected on agar plates containing *Agrobacterium* using the floral dipping method (Clough and Bent, 1998). Transgenic lines were described by Nilsson et al. (1992). Quantitative real-time PCR was used to determine PtCSI1A and PtCSI1B transcript abundance. **UBIQUITIN** transcript abundance was used as a reference gene. Primers for PtCSI1A and PtCSI1B are listed in Table S1.

**Genome expression analysis**

Leaves and developing wood scrapings were homogenized in liquid nitrogen using a mortar and pestle. Total RNA was isolated using TRIZOL® Reagent. In short, 5 µg of total RNA was transcribed into cDNA and amplified using a Maxima First Strand cDNA Synthesis Kit containing dsDNase (Thermo Scientific). qPCR was performed with video extensometry using a stereomicroscope and a Leica DC300 camera (www.leicamicrosystems.com) on the leaf number 17, counting down from the first fully emerged leaf at the top of the tree.

For light microscopy, 20-μm thick wood cross-sections were prepared using a cryotome (Table S1) and stained with 2% aqueous uranyl acetate for 60 min and then with lead citrate (Reynolds, 1963) for 6 min before examination with TEM (JEM 1230; JEOL, Tokyo, Japan). Pictures were taken using a GAtan MSC 600CW 2k × 2k CCD camera.

**Wet chemical analysis of wood**

Cell wall chemical composition of WT and transgenic trees was measured using a subsample of isolated stem sections. Cell wall carbohydrates and total lignin were measured by first grinding the solid xylem tissue in a Wiley Mill to pass a 40 mesh and then extracted with hot acetone in Soxhlet apparatus for a minimum of 12 h. Cell wall carbohydrates and total lignin (acid-soluble and -insoluble lignin, in combination forming total lignin) were determined as described in Huntley et al. (2003) using a modified Klason method. Cell wall carbohydrates were quantified with a high-performance liquid chromatography system using a DX-600 (Dionex, Sunnyvale, CA, USA) equipped with a PA1 (Dionex) column, detector with a gold electrode and SpectraAS3500 auto injector (Spectra-Physics, Santa Clara, CA, USA). Carbohydrate amounts were quantified relative to monomeric cell wall-associated carbohydrates (glucose, xylose, mannose, galactose, rhamnose and arabinose). The amounts of KIason lignin and cell wall sugars represent percentages, relative to the initial weight of dry wood sample analysed.

**Mechanical tests**

Mechanical properties of the wood specimens were measured using a microtensile testing stage described by Burgess et al. (2003). The samples were prepared from specimens isolated 10–20 cm above soil. Longitudinal sections were prepared using a scalpel to obtain a wood block of approximately 25 × 2 × 0.1 mm (length × width × thickness). Tangential longitudinal sections of 100 μm thickness were then cut using a cryotome at −20°C. The cambium region was first removed, and then 10 consecutive longitudinal tangential sections were cut and considered as technical replicates and stored in double-distilled (dd)H2O until physical testing. The values of the technical replicates were used to calculate the mean of the biological replicates. Strain measurement was performed with video extensometry using a stereomicroscope and a CCD camera. The test length was set to 12 mm and the samples were subjected to a constant strain of 10 μm sec−1. The force was recorded with a 50-N load cell (Sensotec Sensors, Honeywell, http://www.honeywell.com). Wood density was determined on the same wood samples as used for the mechanical tests. Density was measured based on wet volume (length, width and thickness) and air dried (48 h at room temperature) weight, using the formula $P = \frac{m}{V}$. **MFA**

CMF angles were determined by measuring CMF orientation of all mechanically tested specimens by wide-angle X-ray diffraction. A Nanostar (Bruker AXS, Ettingen, Germany) was used equipped with a 2D detector (HySIntron, www.bruker.com) and a CuKα radiation source with a wavelength of 0.154 nm. The X-ray beam diameter was set to about 300 μm and the sample-detector distance was set to 8.5 cm. For each sample, one diffraction image was taken with 800-sec exposure time. From the diffraction images, azimuthal intensity profiles of the (200)-Bragg peak of cellulose were calculated with a step size of 1° by radial integrating the intensity within the q-range of the (200)-Bragg peak. The
The samples were finally diluted to a concentration of 1 mg ml\(^{-1}\). This, the samples were washed with ddH\(_2\)O until a neutral pH was obtained after 2 min at room temperature to dissolve hemicellulose. After H\(_2\)O\(_2\). The maceration solution was removed and samples washed with water at two subsequent times. Fibre images were recorded using a Zeiss Axioplan2 light microscope and a Zeiss AxioCam HRC Camera (www.ZEISS.com). The fibre area was measured from the images using IMAGEJ software (http://rsbweb.nih.gov).

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**AUTHOR CONTRIBUTIONS**

AB, OS, AM and MR planned and performed experiments and analysed data. TN, SM and SP wrote and analysed data. AB, OS, AM and MR planned and performed experiments. TN, SM and SP planned experiments.

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**DATA AVAILABILITY STATEMENT**

All data are contained within the manuscript. All materials used in the study will be available upon request.

**SUPPORTING INFORMATION**

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

**Table S1.** Primer and the RNAi sequences.

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