The SHORT-ROOT-like gene *PtSHR2B* is involved in *Populus* phellogen activity

Andreia Miguel\(^1,2\), Ana Milhinhos\(^1,2,\ast\), Ondřej Novák\(^3\), Brian Jones\(^4,5\) and Célia M. Miguel\(^1,2,\dagger\)

\(^1\) Instituto de Biologia Experimental e Tecnológica (IBET), Apartado 12, 2781-901 Oeiras, Portugal
\(^2\) Instituto de Tecnologia Química e Biológica António Xavier, Universidade Nova de Lisboa (ITQB-UNL), Avenida da República, 2780-157 Oeiras, Portugal
\(^3\) Laboratory of Growth Regulators, Centre of the Region Haná for Biotechnological and Agricultural Research, Faculty of Science, Palacký University & Institute of Experimental Botany AS CR, Šlechtitelů 27, 783 71, Olomouc, Czech Republic.
\(^4\) Umeå Plant Science Centre, Department of Plant Physiology, Umeå University, Umeå, Sweden
\(^5\) Faculty of Agriculture, Food and Natural Resources, University of Sydney, Sydney, Australia

\(\ast\) Present address: Department of Plant Sciences, University of Oxford, Oxford OX1 3RB, UK.
\(\dagger\) Correspondence: cmiguel@itqb.unl.pt

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Abstract

SHORT-ROOT (SHR) is a GRAS transcription factor first characterized for its role in the specification of the stem cell niche and radial patterning in *Arabidopsis thaliana* (*At*) roots. Three *SHR*-like genes have been identified in *Populus trichocarpa* (*Pt*). *PtSHR1* shares high similarity with *AtSHR* over the entire length of the coding sequence. The two other *Populus SHR*-like genes, *PtSHR2A* and *PtSHR2B*, are shorter in their 5' ends when compared with *AtSHR*. Unlike *PtSHR1*, that is expressed throughout the cambial zone of greenhouse-grown *Populus* trees, *PtSHR2Bprom:uidA* expression was detected in the phellogen. Additionally, *PtSHR1* and *PtSHR2B* expression patterns markedly differ in the shoot apex and roots of *in vitro* plants. Transgenic hybrid aspen expressing *PtSHR2B* under the 35S constitutive promoter showed overall reduced tree growth while the proportion of bark increased relative to the wood. Reverse transcription–quantitative PCR (RT–qPCR) revealed increased transcript levels of cytokinin metabolism and response-related genes in the transgenic plants consistent with an increase of total cytokinin levels. This was confirmed by cytokinin quantification by LC-MS/MS. Our results indicate that *PtSHR2B* appears to function in the phellogen and therefore in the regulation of phellem and periderm formation, possibly acting through modulation of cytokinin homeostasis. Furthermore, this work points to a functional diversification of *SHR* after the divergence of the *Populus* and *Arabidopsis* lineages. This finding may contribute to selection and breeding strategies of cork oak in which, unlike *Populus*, the phellogen is active throughout the entire tree lifespan, being at the basis of a highly profitable cork industry.

Key words: Cytokinin, lateral meristem, phellogen, *Populus tremula* × *Populus tremuloides*, secondary growth, SHORT-ROOT.

Introduction

Plant growth occurs from specialized regions called meristems. Mitotic divisions in the meristems produce the cells that eventually differentiate into the organs and tissues that comprise the body of the plant. Four main meristems exist in woody perennials, the root and shoot apical meristems, that provide cells for shoot and root growth, respectively, and the
vascular cambium and cork cambium (phellogen) that generate cells for the secondary, or radial, growth of the stem, branches, and roots.

During secondary growth, the epidermis is replaced in the stems and roots by the periderm. This protective tissue is derived from the activity of the phellogen or cork cambium that forms a continuous ring of meristematic cells around the stem and roots. Periclinal divisions of the phellogen initials give rise to cells that differentiate into phelloderm towards the inside of the stem and into phellem towards the outside (Evert, 2006), in a way similar to the differentiation of vascular tissues from the vascular cambium. In a few species, of which the cork oak is an extreme example, the activity of the phellogen contributes to a significant enlargement of the trunk, producing a thick layer of phellem or cork. In adult cork oak trees, cork is regularly stripped from the tree and used for several industrial applications due to its exceptional impermeability, insulation, density, high energy absorption, resilience, and elasticity properties (Silva et al., 2005).

Several lines of evidence have indicated that although the structure of the meristems differs, there are commonalities in the molecular mechanisms underlying their function (Schrader et al., 2004; Baucher et al., 2007; Du and Groover, 2010). Examples of genes known to be involved in the regulation of different meristems include the Class III HD-ZIP (Emery et al., 2003; Schrader et al., 2004; Du et al., 2011; Robischon et al., 2011), the Populus Class I KNOX homeobox genes ARBORKNOX1 and ARBORKNOX2 (ARK1 and ARK2) which are orthologues of Arabidopsis SHOOTMERISTEMLESS (STM) and BREVIPEDICELLUS (BP), respectively (Long et al., 1996; Mele et al., 2003; Groover et al., 2006; Du et al., 2009; Liebsch et al., 2014), SHORT-ROOT (SHR) (Benfey et al., 1993; Fukaki et al., 1998; Helariutta et al., 2000; Schrader et al., 2004; Dhondt et al., 2010; Wang et al., 2011), and Arabidopsis WUSCHEL (WUS) (Mayer et al., 1998; Brand et al., 2000; Schoof et al., 2000) and WUSCHEL-RELATED HOMEOBOX 5 (WOX5) (Haecker et al., 2004; Schrader et al., 2004; Baucher et al., 2007; Sarkar et al., 2007; Tucker and Laux, 2007; Stahl and Simon, 2009; Stahl et al., 2009).

Although considerable attention has been paid to the functioning of the apical meristems and the vascular cambium, little is known about the functioning of the phellogen, despite its critical importance in establishing a protective layer on stems, branches, and roots. Given the existing evidence, it can be hypothesized that the mechanisms underlying phellogen function will overlap those of the other meristems (Soler, 2008).

The SHR transcription factor belongs to the GRAS family of plant-specific proteins that are characterized by a variable N-terminal domain but a highly conserved C-terminal domain (Helariutta et al., 2000; Bolle, 2004). The Arabidopsis SHR (ArSHR) has been well characterized, being a key regulator, along with the related GRAS protein SCARECROW (SCR), of radial patterning and stem cell niche specification in the roots (Benfey et al., 1993; Di Laurentzio et al., 1996; Helariutta et al., 2000; Nakajima et al., 2001). It is essential for the asymmetric cell divisions of the cortex/endodermal initial (CEI) (Benfey et al., 1993; Helariutta et al., 2000; Nakajima et al., 2001), and for the periclinal divisions of cortex cells in a maturing root (Paquette and Benfey, 2005). Fukaki et al. (1998) demonstrated that AtSHR is also involved in radial patterning in the shoot. More recently, Dhondt et al. (2010) demonstrated that, similarly to the Arabidopsis root, SHR functions in association with its downstream target, SCR, in the regulation of cell proliferation and vascular differentiation in leaves. The SHR/SCR mechanism therefore appears to have been co-opted to regulate cell proliferation and differentiation in multiple organs.

SHR modulates the expression of genes involved in a wide range of processes during Arabidopsis root development, including transcriptional regulation, signalling, and response to hormones, and in the regulation of cell cycle genes (Levesque et al., 2006; Sozzani et al., 2010). In Arabidopsis, correct patterning of the central vascular cylinder is mediated through movement of the SHR protein from the stele into the endodermis (Nakajima et al., 2001), where it activates its target, SCR, that together activate miR165a and miR166b (Carlsbecker et al., 2010). The regulation of vascular patterning by SHR in the Arabidopsis root involves the modulation of cytokinin (CK) homeostasis through the direct regulation of the cytokinin-degrading enzyme, CYTOKININ OXIDASE 3 (CKX3) (Cui et al., 2011; Hao and Cui, 2012).

SHR has also been studied in tree species such as Pinus radiata, where it was suggested to have roles in root meristem formation and maintenance, and in the cambial region of hypocotyls (Solé et al., 2008). The putative Populus orthologue of AtSHR, PtSHR1, is expressed in the cambial zone (Schrader et al., 2004; Wang et al., 2011) and functions as a regulator of cell division and meristem activity in the shoots (Wang et al., 2011). Partial suppression of the PtSHR1 transcript in transgenic lines leads to taller trees with a larger vascular cambium due to an increase in cell proliferation in the cambial zone (Wang et al., 2011). In both Arabidopsis and Populus, it has been suggested that SHR regulates growth through the control of cell divisions, in a concentration-dependent manner (Paquette and Benfey, 2005; Wang et al., 2011; Koizumi et al., 2012).

Whereas there is only one SHR gene in Arabidopsis, three SHR-like genes have been identified in the Populus genome, PtSHR1, PtSHR2A, and PtSHR2B (Wang et al., 2011). Based on sequence similarity and on functional studies with the PtSHR1 coding sequence, driven by the AtSHR promoter, PtSHR1 is considered to be the putative ortholog of the Arabidopsis SHR (Wang et al., 2011). In this study, PtSHR2B was investigated in order to better understand its role in meristem function in hybrid aspen. We show that PtSHR2B is expressed in the phellogen, pointing to a regulatory role in this meristem during secondary growth. Overexpression of PtSHR2B in hybrid aspen not only affected overall plant growth, but altered the ratio between the amount of wood and bark tissues in the stem. We further present experiments that indicate that PtSHR2B may act, at least partially, through the regulation of CK homeostasis.
Materials and methods

Plant material

Hybrid aspen (Populus tremula L.×Populus tremuloides Michx.; Clone T89) was propagated in vitro on half-strength basal MS medium (Murashige and Skoog, 1962) and maintained in a growth chamber at 21 °C in a 16 h light/8 h dark photoperiod. For greenhouse experiments, in vitro established wild-type and transgenic plants were placed in a soil:peat:perlite (1:3:1) potting mix and acclimatized in a growth chamber, gradually decreasing the humidity from 95% to 70% over 5 weeks before transfer to the greenhouse, where plants were grown for a minimum of 10 weeks prior to analysis. The position of all pots within the greenhouse was changed weekly to minimize positional bias experimental error.

After 10 weeks in the greenhouse, the five youngest fully expanded leaves were collected (Supplementary Fig. S1A at JXB online). Bark was isolated by peeling off the stem across the cambial zone. The remaining stem tissues (xylem and pith), hereafter termed ‘wood’, were collected together. All samples were immediately frozen in liquid nitrogen and stored at –80 °C until further processing. Intact stem sections were also collected and fixed in FAA (5% formaldehyde, 5% acetic acid, 50% ethanol) for anatomical analysis.

Gene constructs and genetic transformation

To detect promoter activity of PtSHR1, PtSHR2A, and PtSHR2B, transgenic lines were generated using constructs incorporating ~2.5 kbp sequence upstream of the start codon of each of the genes, fused to the coding sequence of uidA encoding β-glucuronidase (GUS) (PtSHRprom:uidA), using the pKGWFS7.0 binary vector (Karimi et al., 2002). The sequences (eugene3.01860017, eugene3.00070144, and eugene3.00640143, for PtSHR1, PtSHR2A, and PtSHR2B, respectively) were retrieved from the Populus trichocarpa genome version 3.1 (http://genome.jgi-psf.org/Poptri_1/Poptri_1.home.html) and were previously described by Wang et al. (2011). In Phytovaze v9.1 (http://www.phytovaze.net/), the corresponding loci are Potri.007G063300, Potri.007G132000, and Potri.017G019900 for PtSHR1, PtSHR2A, and PtSHR2B, respectively. To express PtSHR2B ectopically, its coding sequence was cloned downstream of the 35S Cauliflower mosaic virus (CaMV) promoter in the pK7WG2.0 vector (35S:PtSHR2B) (Karimi et al., 2002). Hybrid aspen stem sections were transformed according to Nilsson et al. (1992). Ten independent transgenic lines were generated for each construct. Representative PtSHR1prom:uidA and PtSHR2Bprom:uidA lines and three independent overexpression lines, 2B_7, 2B_8, and 2B_12, were chosen for further analysis, after checking for the presence of single insertion events by Southern blot analysis. In this study, the nomenclature and the gene sequence information followed Wang et al. (2011).

Histochemical GUS assay

GUS assays were performed in transformed hybrid aspen leaves, roots, and shoot apex collected from 6-week-old in vitro grown shoots, and 6-month-old and 1-year-old stems of greenhouse-grown trees. Tissues were placed in ice-cold 90% acetone for 30 min and then washed in water prior to immersion in the GUS staining solution [10 mM sodium phosphate buffer pH 7.0, 0.5% Triton X-100, 2 mM potassium ferricyanide, and 2 mM X-Gluc (5-bromo-4-chloro-3-indolyl β-D-glucuronide), vacuum infiltrated, and incubated overnight in the dark at 37 °C]. After washing in water, tested leaves, roots, and shoot apex of in vitro plants were gradually dehydrated to 70% ethanol and stem sections of the greenhouse-grown plants were fixed in FAA and then included in Technovit 7100 resin (Heraeus Kulzer), according to the manufacturer’s instructions, with minor modifications: after vacuum infiltration, samples were left for 2 d at 4 °C in the pre-infiltration solution. The solution was then replaced and samples left for another 7 d at 4 °C. The material was subsequently placed in the infiltration solution and left for 1–3 weeks at 4 °C, followed by polymerization at room temperature.

Steremicroscope observations were performed with a Nikon SMZ800, and images were captured using an Olympus SC30 camera and software. Microscope observations were made with a Nikon Inverted Microscope Eclipse TE300, and images taken with a Nikon DS-F1i camera using the NIS-Elements F3.0 software.

Anatomical analysis and growth measurements

Stem pieces of 1- and 2-year-old shoots of wild-type hybrid aspen were collected and fixed in ice-cold FAA, as previously described, vacuum infiltrated, and left overnight in a desiccator at 4 °C. After gradual dehydration to 100% ethanol, tissues were embedded in resin as described above, and 6-8 µm thick sections were stained with Toluidine Blue O. Several growth parameters were analysed in 10-week-old greenhouse-grown 35S:PtSHR2B trees (Supplementary Fig. S1A). Tree height and total length between the 10th (EN10) and the 17th (EN17) internodes from the shoot tip were recorded. Stem diameters were measured at the reference internode (EN14), showing fully differentiated secondary vascular tissues and chosen for comparison between transgenic and wild-type plants, and at the stem base (ENbase), corresponding to an internode between the 20th and 25th internodes depending on the tree. Measurements of distances between different stem tissues were taken at a minimum of four positions around the circumference of the stem sections using ImageJ software (Abràmoff et al., 2004; Schneider et al., 2012) (Supplementary Fig. S1B). The total lamina area of the five leaves surrounding EN14 was determined using a leaf area meter (LI-3000A, LI-COR Inc.).

Reverse transcription–quantitative PCR (RT-qPCR)

The first five fully expanded leaves down from the shoot tip, and bark and wood from the ENbase were collected and ground to a fine powder (Supplementary Fig. S1A) using either a mortar and pestle (leaves and bark) or a grinder mill (M 20 Universal mill, Ika). Isolation of bark and wood tissues was done as described in the ‘Plant material’ section. Total RNA from leaves and bark was isolated as described by Reid et al. (2006) with minor modifications (Marum et al., 2012). Total RNA from wood samples was extracted using the protocol as described in Chang et al. (1993), and all RNA samples were treated with TURBO DNase (Ambion) according to the manufacturer’s instructions. cDNA synthesis was performed from 1.5 µg of DNase-treated RNA, using a Transcriptor High Fidelity cDNA Synthesis Kit (Roche) with anchored oligo(dT)18 primers. Quantitative real-time PCR (qPCR) was carried out in 96-well plates in a LightCycler 480 (Roche) using SYBR Green I Master Mix (Roche). Primers for amplifying a transcript fragment of PtSHR2B, 5′-CAGCAATACCCCTTGGACACAG-3′ and 5′-ACCAGCTTCCCTTGGTG-3′, were designed using the P. trichocarpa genome version 3.1 (http://genome.jgi-psf.org/Poptr1_1/Poptr1_1.home.html) and the gene sequence (eugene3.00640143) described by Wang et al. (2011). For amplification of PtCKX3 (Potri.004G152500.1) transcripts, specific primers 5′-TCAGATCCAAACCCTTGATTTC-3′ were designed. Quantitative real-time PCR (qPCR) was carried out in 96-well plates in a LightCycler 480 (Roche) using SYBR Green I Master Mix (Roche). Primers for amplifying a transcript fragment of PtSHR2B, 5′-CAGCAATACCCCTTGGACACAG-3′ and 5′-ACCAGCTTCCCTTGGTG-3′, were designed using the P. trichocarpa genome version 3.1 (http://www.phytowome.net/). For amplification of PtRR7 transcripts (Potri.016G038000.1), primers were as previously described (Niinemnen et al., 2008). PtCYP2 (Potri.004G168800.1) was used as a reference gene (Brummer et al., 2004; Milhinhos et al., 2013). The PCR program used was 95 °C for 10 min, 45 cycles of 10 s at 95 °C, 20 s at 60 °C for PtRR7 and PtCKX3 or 20 s at 63 °C for PtSHR2B, and 10 s at 72 °C. The annealing temperature for the reference gene primers was 60 °C or 63 °C, depending on the experiment. Three technical replicates were used for each of the three biological samples in each experiment. To normalize values obtained from different plates, a calibrator sample consisting of cDNA synthesized
from RNA from leaves of a transgenic line was used in each plate. Normalized relative quantities were obtained through the ΔΔC_T method (Livak and Schmittgen, 2001; Pfaffl, 2001; Hellemans et al., 2007) and the amplification efficiency determined using Real-Time PCR Miner (Zhao and Fernald, 2005).

Quantification of cytokinins
To quantify the CK levels in bark and wood tissues, samples were ground into a fine powder. CKs were extracted and isolated from ~10 mg of frozen tissues from bark and wood as previously described (Svačinová et al., 2012), including modifications described by Antoniadi et al. (2015). To each extract, the stable isotope-labelled CK internal standards (0.1 pmol of CK bases, ribosides, N-glucosides, 0.25 pmol of O-glucosides, and 0.5 pmol of nucleotides) were further added as a reference. Purified samples were analysed by an LC-MS/MS system consisting of an ACQUITY UPLC® System (Waters) and a Xevo™ TQ-S (Waters) triple quadrupole mass spectrometer. Quantification was obtained using a multiple reaction monitoring (MRM) mode of selected precursor ions and the appropriate product ion. Five independent biological replicates were analysed for each sample.

Statistical analysis
The assessment of statistical significance in transcript profiles and phenotypic parameters was carried out using non-parametric analysis, Mann–Whitney U-test. A significance level of _P_ = 0.05 was used. An ANOVA was performed to assess the statistical significance in the quantification of CKs. Statistics were performed using the Statistica (StatSoft Inc., http://www.statsoft.com) software package.

Results
In this work, the _Populus SHR_-like gene PtSHR2B was characterized and compared with the putative _Populus_ orthologue of the _Arabidopsis SHR_ gene, PtSHR1 (Wang et al., 2011). _AtSHR_ and _PtSHR1_ genes have been previously implicated in the regulation of primary apical meristems and vascular cambium activity. Transverse sections of hybrid aspen stems at different developmental stages showed a distinct phellogen meristem layer, already present at the end of the first year of growth and characterized by rectangular cells that are flattened radially and divide mostly by periclinal division (Supplementary Fig. S1C). The characteristic layer of suberized phellem cells in the periderm could be observed. The periderm in 2-year-old stems had only a slight increase in phellem layer thickness compared with 1-year-old stems (Supplementary Fig. S1C, D).

PtSHR1 and PtSHR2B show different expression patterns
While tissues from wild-type controls were always negative to GUS histochemical assay (Fig. 1A–D), analysis of hybrid aspen plants carrying either the _PtSHR1_ or _PtSHR2B_ promoter driving _uidA_ expression (_PtSHR2B_prom:_uidA_) indicated different patterns of promoter activity. Analysis of _PtSHR1_ promoter activity in greenhouse-grown trees that had undergone substantial secondary growth showed GUS throughout the cambial zone and in xylem rays (Fig. 1E, F). Additionally, _GUS_ expression was found in the leaf vasculature and in the root stele of _in vitro_ plants (Fig. 1G, H), similarly to its homologue in _Arabidopsis (AtSHR)_ (Helariutta et al., 2000; Dhondt et al., 2010; Wang et al., 2011). In the shoot apex, _GUS_ staining was observed in the apical meristem and vasculature (Fig. 1I). In the case of the _PtSHR2B_ promoter, the greenhouse-grown trees showed _GUS_ staining strongly localized in the phellogen cell layer (Fig. 1J, K), suggesting a specific function for the modified version of _SHR_ in this meristem. In leaves from _in vitro_ plants, _GUS_ expression was not detected (Fig. 1L), but in the roots _GUS_ staining showed a stark contrast to that observed with the _PtSHR1_ promoter, with _GUS_ expression being observed at the root tip (Fig. 1M). Differences in the expression driven by each of the two promoters were also found in _in vitro_ developing shoot tip where _PtSHR2B_prom:_uidA_ staining was restricted to apical and axillary meristems (Fig. 1N). Since no _GUS_ signal was ever detected in any of the analysed tissues from plants carrying the _PtSHR2B_prom:_uidA_ construct, the study only proceeded with the analysis of _PtSHR2B_.

Profiling of _PtSHR2B_ transcript levels by RT–qPCR in the tissues of wild-type greenhouse-grown hybrid aspen revealed the highest levels in the bark, with significantly lower levels in wood and leaf tissues (Fig. 2). These results corroborate the _GUS_ staining observations, confirming that expression is predominantly in the phellogen, although not restricted to it (Figs 1J, K, 2).

Ectopic expression of _PtSHR2B_ reduces overall tree growth
To explore the role of _PtSHR2B_ in hybrid aspen, the _PtSHR2B_ coding region was isolated and transformed into hybrid aspen under the control of the constitutive CaMV 35S promoter. RT–qPCR confirmed the ectopic expression of _PtSHR2B_ in leaves and stem tissues of _in vitro_ 35S::_PtSHR2B_ plants (Fig. 3A). However, in the greenhouse-grown trees, the increased accumulation of _PtSHR2B_ transcript was observed in leaves and bark, but not in wood tissues (Fig. 3B). No obvious phenotype could be observed in _in vitro_ grown plants. However, 10-week old greenhouse-grown transgenic trees showed a reduced growth compared with the wild type (Fig. 4; Supplementary Fig. S2). Tree height in all of the transgenic lines was significantly reduced when compared with the wild-type plants (Fig. 4A, B). Both control and transgenic trees experienced growth deceleration between weeks 7 and 10, possibly due to a greenhouse temperature increase during this period. Stem diameter was significantly reduced compared with the wild type at the base of the transgenic trees (Fig. 4C) although no change could be observed in the reference internode (EN14). The reduction in height in the transgenic trees was primarily the result of a reduced internode length, as the mean internode length between EN10 and EN17 was significantly shorter in the transgenic trees compared with the wild type (Fig. 4D). The total number of internodes was slightly reduced, but this was only significant in one of the transgenic lines (Fig. 4E). The
total average fresh weight and lamina area of the five leaves surrounding the reference internode was also significantly reduced in the transgenic trees (Fig. 4F, G).

Altered stem anatomy in 35S:PtSHR2B ectopic expression plants

Transverse sections taken from stems of the 35S:PtSHR2B transgenic trees were analysed by light microscopy. The secondary xylem [radial distance from the pith side of the lignified xylem to the cambial zone (Supplementary Fig. S1B)] was significantly reduced in the transgenic trees compared with the wild type in two of the three transgenic lines (Fig. 5A), whereas the bark layer was wider than in the wild type (Fig. 5A). The difference in the proportion of wood and bark tissues was more evident at the base of the stem, where secondary growth is far more extensive (Fig. 5B). Additionally, the ratio between the phellem and the stem radius was reduced in two out of the three transgenic lines (Fig. 5C, D). Some variation was seen between the independent transformant lines, which is commonly observed and consistent with previous work using Populus (Robischon et al., 2011; Milhinhos et al., 2013).

Fig. 1. Localization of PtSHR1 and PtSHR2B promoter-driven uidA expression. (A, C–E, F, I–K, L–N, Q–S) GUS histochemical assay in 6-week-old hybrid aspen in vitro grown plants. (B, G, H, O, P) GUS histochemical assay in stems from 6-month-old greenhouse-grown hybrid aspen. Images A, F, G, L–N and O were taken under the microscope (scale bars=100 µm). Images B–E, H–K, and P–S were taken under the stereomicroscope (scale bars=475 µm). AX, axillary meristems; Pg, phellogen; SAM, shoot apical meristem; VC, vascular cambium.

Fig. 2. Relative expression levels of PtSHR2B in tissues of wild-type hybrid aspen grown in the greenhouse for 10 weeks. Results are expressed relative to the expression in the bark, and values are the means ±SE of at least four biological replicates, except for wood where a mix of different wild-type plants was used, and three technical replicates. Asterisks indicate the significance of the difference between each tissue and the bark. (This figure is available in colour at JXB online.)
Quantification of cytokinins in stem tissues

To clarify the link between PtSHR2B and CK homeostasis, we quantified the levels of CKs present in wood and bark tissues of the transgenic trees. Naturally occurring CKs are adenine derivatives with either isoprenoid or aromatic side chains. Isoprenoid CKs can be distinguished as isopentenyladenine (iP)-, trans-zeatin (tZ)-, cis-zeatin (cZ),- or dihydrozeatin (DHZ)-type derivatives, depending on the hydroxylation and reduction of the side chain (Ha et al., 2012). Total CK levels were altered in both tissues, as was the distribution of its derivatives (Fig. 7; Table 1). All 35S:PtSHR2B transgenic trees presented high levels of total CKs in bark but lower levels in wood tissues (Fig. 7).

Discussion

The vascular cambium functioning and the development of vascular tissues have been the subject of numerous reports (Baucher et al., 2007; Du and Groover, 2010; Schuetz et al., 2012; Sanchez et al., 2012). In contrast, despite the important role that the phellogen plays in providing cells for the development of the protective layer on stems, branches, and roots, it has received minimal attention. Under our growth conditions, a surrounding meristematic phellogen and a phellem (cork) layer had formed in 1-year-old hybrid aspen stems. The thickness of this layer was almost unchanged between 1- and 2-year-old hybrid aspen stems. Consistency in the phellem layer thickness during the lifespan of the close relative Populus tremuloides Michx. has long been known (Kaufert, 1937), indicating that in Populus species phellogen cell divisions are matched by a shedding of cork layer cells.

Populus is recognized as a model species in the study of angiosperm tree function. It has a high annual rate of secondary growth, and like other model species has the advantages of being relatively easily genetically modified and cultured. There are also ample molecular, genomic, and bioinformatics resources available for various Populus species. Importantly, the P. trichocarpa genome has been fully sequenced (Tuskan et al., 2006). The sequence data indicate that two whole-genome duplication events occurred in ancestors of the species (Tuskan et al., 2006). Although duplicated genes are often lost over evolutionary time, higher gene retention is often found, particularly for specific classes of genes, such as: (i) genes with regulatory functions, namely transcription factors and developmental regulators (Blanc and Wolfe, 2004; Seoighe and Gehring, 2004; Carretero-Paulet and Fares, 2012); and (ii) for genes derived from a previous round of genome duplication (Seoighe and Gehring, 2004). The biased retention is most probably because multiple copies of the retained genes impart specific beneficial effects for the organism (Seoighe and Gehring, 2004; Carretero-Paulet and Fares, 2012). One of the ways in which multiple genes can be beneficial is through speciation, leading to divergent function and expression patterns (Blanc and Wolfe, 2004; Tuskan et al., 2006; Rodgers-Melnick et al., 2012). In Populus, it has been hypothesized that after duplication, gene preservation is influenced by a combination of subfunctionalization and
selection favouring retention of genes that encode proteins with a large number of interactions (Rodgers-Melnick et al., 2012). The presence of three SHR-like genes in the Populus genome indicates that SHR fits these criteria. In this work we show that the patterns of GUS expression driven by Populus PtSHR1 and PtSHR2B promoters differ markedly both in the shoot apex and in the roots. Commensurate with AtSHR expression in Arabidopsis, GUS staining in the roots of the PtSHR1prom:uidA plants was confined to the stele. In contrast, the PtSHR2B promoter activity was detected in the root tip, suggesting that the two genes have different functions in the root. In greenhouse-grown plants with extensive secondary growth, GUS expression driven by both PtSHR1 and PtSHR2B promoters was strongly associated with the lateral meristems. However, while GUS staining was detected in the vascular cambium of the PtSHR1prom:uidA plants, the

Fig. 4. Phenotypic characterization of transgenic 35S:PtSHR2B hybrid aspen lines grown in the greenhouse for 10 weeks. (A) The wild type (WT) and plants from independent transgenic lines (2B_7; 2B_8 and 2B_12). (B) Tree height. Values are the means ±SE of at least eight biological replicates. (C) Stem diameter at the reference internode (EN14) and at the stem base (ENbase) corresponding to the 20th–25th internode. Values are the means ±SE of at least six biological replicates. (D) Mean internode length between the 10th and the 17th internodes (EN10 and EN17). Values are the means ±SE of at least eight biological replicates. (E) Total number of internodes. Values are the means ±SE of at least eight biological replicates. (F) Total weight of the five leaves surrounding EN14. Values are the means ±SE of at least five biological replicates. (G) Total area of the leaves surrounding EN14. Values are the means ±SE of at least three biological replicates. Asterisks indicate the significance level between each individual line and the WT (*P<0.05, **P<0.01, and ***P<0.001, Mann–Whitney U-test). (This figure is available in colour at JXB online.)
PtSHR2B promoter drove GUS expression in the phellogen. PtSHR1 has previously been reported to be expressed in the vascular cambium, and to regulate its activity (Schrader et al., 2004; Wang et al., 2011). Furthermore, AtSHR and PtSHR1 have been shown to have broad activity in meristems in the roots and shoots (Schrader et al., 2004; Wang et al., 2011). However, this is the first work reporting an SHR-like gene promoter activity in the phellogen, suggesting that PtSHR2B fulfills an important function in this lateral meristem.

This finding led us to explore further the function of PtSHR2B by ectopically expressing it in hybrid aspen. Compared with the wild type, overall growth was reduced in transgenic trees as evaluated by several parameters to assess

![Fig. 5. Anatomical characterization of the 35S:PtSHR2B hybrid aspen stems at the 14th internode (EN14) and at the stem base (ENbase) corresponding to an internode between the 20th and 25th internodes. (A and B) Ratio between the width of the wood or the bark layer and the stem radius at EN14 (A) and ENbase (B). (C and D) Ratio between the width of the phellem layer and the stem radius at EN14 (C) and ENbase (D). Values are the means ±SE of at least three biological replicates, and asterisks indicate the significance of the difference between each individual line and the wild type (*P<0.05, **P<0.01, and ***P<0.001, Mann–Whitney U-test). For each tree cross-section, the measurements were taken at a minimum of four equidistant positions around the circumference of the stem. (This figure is available in colour at JXB online.)](https://academic.oup.com/jxb/article-abstract/67/5/1545/2885132/)

![Fig. 6. Expression of genes involved in cytokinin function, determined at the stem base (ENbase) corresponding to an internode between the 20th and 25th internodes, in the 35S:PtSHR2B hybrid aspen plants grown in a greenhouse for 10 weeks. (A, B) Relative transcript levels of (A) the cytokinin primary response gene, PtRR7, and (B) the cytokinin oxidase gene, PtCKX3, in bark and wood tissues of 35S:PtSHR2B trees. Values are the means ±SE of at least three biological replicates, and asterisks indicate the significance of the difference between each individual line and the wild type (*P<0.05, **P<0.01, and ***P<0.001, Mann–Whitney U-test). (This figure is available in colour at JXB online.)](https://academic.oup.com/jxb/article-abstract/67/5/1545/2885132/)

![Fig. 7. Cytokinin (CK) content in the bark and wood tissues of 35S:PtSHR2B hybrid aspen plants grown in a greenhouse for 10 weeks. Values are the means ±SE of at least three biological replicates, and asterisks indicate the significance between each individual line and the wild type (*P<0.05, **P<0.01, and ***P<0.001, one-way ANOVA). (This figure is available in colour at JXB online.)](https://academic.oup.com/jxb/article-abstract/67/5/1545/2885132/)
primary and secondary growth. Total tree height and mean internode length between EN10 and EN17 were amongst the most significantly reduced growth parameters in all the transformed lines. The total number of internodes was only slightly reduced, further indicating that stem elongation was inhibited in the transgenic lines. The stems of the transgenic trees were less tapered than those of the wild-type plants since both groups had similar stem diameters in the upper region of the stem (at EN14), but the stems of the transgenics were appreciably narrower at the base (EN20–EN25). When the stem tissues were analysed in more detail, we verified a relative reduction in the wood layer, while an increase in the proportion of bark, composed of phloem, phelloderm, and phellem, was observed. We could not correlate the increase in the bark width with an increase of the phellem given that the shedding of cork layer cells in *Populus* occurs simultaneously with cell division of phellogen initials (Kaufert, 1937).

Altogether, the phenotypic analysis of these plants suggests that, in addition to the effect on secondary growth patterns, *PtSHR2B* ectopic expression affects overall plant growth. When studying the function of *PtSHR1* in hybrid aspen, Wang et al. (2011) have shown that its partial suppression enhanced overall plant growth, and suggested that the protein acts as a dose-dependent negative regulator of meristem activity. Our results do not allow a conclusion to be drawn on the dose-dependent action of *PtSHR2B* but support that it might interfere in plant growth through similar mechanisms to those of *PtSHR1*.

In order to investigate the possible mechanisms by which *PtSHR2B* may exert its function, we have analysed the expression of the CK-related genes *PtRR7* and *PtCKX3*, which encode a CK primary response and a CK degradation protein, respectively. As inducers of cell division and differentiation, CKs are central regulators of plant development (Miyawaki et al., 2006; Sakakibara, 2006). In fact, the importance of CK signalling in the regulation of cambial activity in *Populus* stems has been previously demonstrated (Nieminen et al., 2008). In our work, we found that the expression of both genes was significantly up-regulated in the bark tissues of all the transgenic lines, but their expression levels were either similar or only slightly increased in the wood tissues when compared with the wild type. *PtRR7* transcript levels were reported to be positively correlated with the amount of CK present across the stems of hybrid aspen trees (Nieminen et al., 2008). Cytokinin oxidase gene expression has also been shown to be up-regulated in response to increased CK levels (Motyka et al., 1996; Jiao et al., 2003). Additionally, in *Arabidopsis*, the *CKX3* transcript levels were significantly reduced in the *shr* roots (Cui et al., 2011). Our transcript quantification results would seem consistent with increased CK endogenous levels in the tissues where expression is higher. In fact, through quantification of CKs we were able to confirm that endogenous total CK levels were significantly higher in the bark tissues, in two out of the three transgenic lines. This trend was not verified in wood tissues that, despite the lower CK levels in two out of three lines, showed a slight up-regulation of *PtRR7*. Differences in CK biosynthesis and homeostasis regulation, or even their crosstalk with auxin in the vascular tissues may contribute to explain this observation (Kieber and Schaller, 2014), although additional studies would be required to clarify this issue.

The mechanisms by which the overexpression of *PtSHR2B* might lead to increased CK levels in the bark tissues were not addressed in our study. However, it is tempting to speculate that similarly to *AtSHR* (Sebastian et al., 2015), *PtSHR2B* may interfere with CK homeostasis. It is possible that the increased CK levels in the bark (containing the phellogen layer) result from an increased amount of *PtSHR2B* transcripts which may regulate target genes with specific domains of expression in these tissues. The decrease in the wood layer, indicative of modified cambium activity, might also be explained by the slightly reduced levels of total CKs in the wood tissues. This will need to be tested in future studies with tissue-specific promoters, since secondary effects of ectopic expression of *PtSHR2B* cannot be ruled out.

The data presented here suggest that *PtSHR2B* is involved in lateral meristem functioning. Although it cannot be excluded that *PtSHR2B* plays a role alongside *PtSHR1* in regulating vascular cambium activity, our data suggest that speciation and functional diversification have led to these two *AtSHR* homologues playing different roles in *Populus* stems. *PtSHR2B* appears to function principally in the phelogen

### Table 1. Distribution of the cytokinin derivatives between bark and wood tissues, in 10mg of tissue, of 3SS:*PtSHR2B* hybrid aspen plants grown in the greenhouse for 10 weeks

|        | Cks  | Total | iP types | tZ types | DHZ types | cZ types |
|--------|------|-------|----------|----------|-----------|----------|
| Bark   |       |       |          |          |           |          |
| WT     | 61.2±5.8 | 13.8±1.3 | 7.8±1.4 | 2.7±0.4 | 36.9±3.9 |
| 2B_7   | 75.9±18.8 | 6.9±0.1* | 10.1±1.8 | 5.9±0.4* | 52.9±16.5* |
| 2B_8   | 79.3±5.6* | 9.7±0.3 | 11.0±0.4* | 6.7±0.7*** | 51.9±5.1* |
| 2B_12  | 132±22.8** | 5.9±1.4** | 14.2±6.6* | 6.7±2.9* | 86.3±14.6*** |
| Wood   |       |       |          |          |           |          |
| WT     | 29.2±2.0 | 12.5±0.7 | 2.7±0.6 | 0.5±0.1 | 13.5±1.0 |
| 2B_7   | 17.5±6.5 | 4.4±1.9** | 2.3±1.0 | 0.8±0.3 | 9.9±3.8 |
| 2B_8   | 20.8±1.8* | 5.2±0.8** | 2.6±0.3 | 0.7±0.008** | 12.4±0.7 |
| 2B_12  | 12.2±3.7*** | 4.3±1.8*** | 1.8±0.6 | 0.7±0.3 | 5.7±1.2*** |

Values are means ±SE of at least three biological replicates, and asterisks indicate the significance between each individual line and the wild-type (*P<0.05, **P<0.01, and ***P<0.001, one-way ANOVA). CK, cytokinin; iP, isopentenyladenine; tZ, trans-zeatin; DHZ, dihydrozeatin; cZ, cis-zeatin.
and therefore in the regulation of phellem and periderm formation. This finding may contribute to strategies aimed at the selection and improvement of species such as cork oak in which, unlike *Populus*, the phellogen is active throughout the entire tree lifespan, being at the basis of a highly profitable cork industry.

**Supplementary data**

Supplementary data are available at *JXB* online.

**Figure S1.** Schematic representation of tissue sampling, stem section measurements, and anatomical aspects of hybrid aspen stem.

**Figure S2.** Growth curves of the wild-type and 35S::PtSHR2B hybrid aspen plants.

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