Axillary buds are dwarfed shoots that tightly regulate GA pathway and GA-inducible 1,3-β-glucanase genes during branching in hybrid aspen

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Abstract

Axillary buds (AXBs) of hybrid aspen (Populus tremula×P. tremuloides) contain a developing dwarfed shoot that becomes para-dormant at the bud maturation point. Para-dormant AXBs can grow out after stem decapitation, while dormant AXBs pre-require long-term chilling to release them from dormancy. The latter is mediated by gibberellin (GA)-regulated 1,3-β-glucanases, but it is unknown if GA is also important in the development, activation, and outgrowth of para-dormant AXBs. The present data show that para-dormant AXBs up-regulate GA receptor genes during their maturation, but curtail GA biosynthesis by down-regulating the rate-limiting GIBBERELLIN 3-OXIDASE2 (GA3ox2), which is characteristically expressed in the growing apex. However, decapitation significantly up-regulated GA3ox2 and GA4-responsive 1,3-β-glucanases (GH17-family; α-clade). In contrast, decapitation down-regulated γ-clade 1,3-β-glucanases, which were strongly up-regulated in maturing AXBs concomitant with lipid body accumulation. Overexpression of selected GH17 members in hybrid aspen resulted in characteristic branching patterns. The α-clade member induced an acropetal branching pattern, whereas the γ-clade member activated AXBs in recurrent flushes during transient cessation of apex proliferation. The results support a model in which curtailing the final step in GA biosynthesis dwarfs the embryonic shoot, while high levels of GA precursors and GA receptors keep AXBs poised for growth. GA signaling, induced by decapitation, reinvigorates symplasmic supply routes through GA-inducible 1,3-β-glucanases that hydrolyze callose at sieve plates and plasmodesmata.

Key words: Apical dominance, axillary branching, bud dormancy, callose, gibberellin, 1,3-β-glucanase, para-dormancy, plasmodesmata, Populus, strigolactone.

Introduction

The architecture and three-dimensional shape of a tree arise gradually during multiple seasons. Branches compete for resources, whereas the development of secondary, tertiary, and higher order branches exponentially enhances the complexity of a tree (Turnbull, 2005; Costes et al., 2014; Costes and Gion, 2015). Branching patterns are constrained by...
phyllotaxis as branches grow out of axillary meristems (AXMs) that arise in the axes of leaves. Although the molecular mechanisms that regulate branching are thought to be conserved between annuals and woody perennials, there are some distinct differences in the initiation of AXMs, axillary bud (AXB) formation, and branching. For example, in the annual Arabidopsis, AXM initiation is delayed, and the AXBs they produce are simple and lack scales (Grabic and Bleecker, 2000; Long and Barton, 2000; Greb et al., 2003). In contrast, deciduous woody perennials produce AXMs by default, in conjunction with nascent leaves, and in continuity with the shoot apical meristem (SAM) (Garrison, 1955). The first products of these AXMs are bud scales which confine an emerging dwarfed shoot (Romberger, 1963; Brunner et al., 2014; Rinne et al., 2015). Differences in the timing of AXB outgrowth give rise to distinct branching styles. In the proleptic branching style, AXBs must pass through a dormancy phase before they can grow out (Halle et al., 1978; Barthélémy and Caraglio, 2007). In contrast, in sylleptic branching, incomplete AXBs give rise to branches (Wu and Hinckley, 2001). Whereas syllepsis is highly variable due to its environmental responsiveness, the more robust phenomenon of prolepsis reflects strong apical dominance (Ceulemans et al., 1990; Cline, 1997; Wu and Stettler, 1998).

Apical dominance is the phenomenon whereby AXBs are held captive in a state of para-dormancy by a proliferating apex (Phillips, 1975; Cline, 1991, 1997). AXBs can be released from this repressed state by decapitation, a procedure that has been widely used to study branching. Early studies attributed the dominance of the apex (i.e. its developmental supremacy over the AXBs) to the local production of auxin and its basipetal transport (Thimann and Skoog, 1934; Phillips, 1975; Cline, 1991). In a recent model for Arabidopsis, the apex monopolizes the polar auxin transport stream (PATS), saturating its transport capacity and hampering access of AXBs to the PATS (Domagalska and Leyser, 2011). The concept of auxin-based apical dominance is relative in the sense that a proliferating apex does not always prevent branching (Dun et al., 2006). Moreover, in caulescent plants, AXBs may be activated ahead of the decapitation-induced depletion of the PATS (Morris et al., 2005), a phenomenon that was attributed to the sudden availability of sucrose (Mason et al., 2014). This conclusion is reminiscent of the early hypothesis that the growing apex deprives the AXBs of nutrients (Cline, 1991), and suggests that resource allocation is a factor in apical dominance. The emerging picture is that the role of auxin in branching is not straightforward as branching is sensitive to the overall balance of systemic and local processes. In brief, the systemic networks involve auxin, the branch inhibitor strigolactone, the branch facilitator cytokinin, as well as competition-driven shifts in sink–source relationships (Ferguson and Beveridge, 2009; Domagalska and Leyser, 2011; Mason et al., 2014). In addition to the systemic networks, AXBs may self-regulate through internally produced agents. In Arabidopsis, such a local agent is BRANCED1 (BRC1), a homolog of the maize transcription factor teosinte branched1 (Doebely et al., 1997). BRC1 acts downstream of strigolactone to suppress AXB outgrowth (Aguilar-Martinez et al., 2007; Niwa et al., 2013). Activation of an AXB involves initiation of auxin biosynthesis, production and polarization of PINFORMED1 (PIN1) auxin efflux carriers in the bud-to-stem path, and differentiation of functional vascular connections to the main stem (Li and Bangerter, 1999; Balla et al., 2011; Domagalska and Leyser, 2011).

Deciduous woody perennials may recruit similar mechanisms as their response to decapitation is essentially the same (Cline, 1991, 1997; Rinne et al., 1993), and their genomes contain genes homologous to those involved in herbaceous branching (Czarnecki et al., 2014; Waldie et al., 2014). For example, in hybrid aspen (Populus tremula×P. tremuloides), BRC1 and MAX1 were identified (Rinne et al., 2015). Both genes appeared to be highly expressed in AXBs and down-regulated upon decapitation. Moreover, xylem feeding of the synthetic strigolactone analog GR24 inhibited AXB activation in internode cuttings (Rinne et al., 2015). Together this indicates that they can function locally in AXBs. In trees, such local agents might be particularly important considering the extended transport paths.

In the hybrid aspen clone T89, apical dominance prevents branching in current year AXBs, but it does not prevent the development of an embryonic shoot, which is essentially a dwarfed side shoot. The newly formed AXM follows a developmental program, in which it first produces five primordia that develop into ‘perfect’ scales to protect the subsequent 10 primordia that develop into embryonic leaves. AXB development is completed at the so-called bud maturation point (BMP) (Rinne et al., 2015), and results in a para-dormant ‘embryonic’ or ‘pre-formed’ shoot (Romberger, 1963; Brunner et al., 2014; van der Schoot et al., 2014; Rinne et al., 2015). During branching, the tightly packed embryonic leaves expand, the compressed internodes elongate, and subsequently new leaves are initiated. Thus, in proleptic branching, cell division and morphogenesis of the branch are temporarily separated from expansion by a waiting period of minimally one season.

The dwarfed stature of the embryonic shoot suggests that AXBs are gibberellic acid (GA) deficient. Nonetheless, the role of GA in apical dominance and para-dormancy has not received much attention. One reason might be that the phenotypes of GA biosynthesis mutants (Murfet and Reid, 1993; Silverstone et al., 1997) and overexpressors of GA catabolism genes often show enhanced branching, in both herbaceous plants and trees (Agharkar et al., 2007; Lo et al., 2008; Mauriat et al., 2011; Jaworski and Busov, 2014). This seems to suggest that GA actually inhibits branching (Scott et al., 1967; Rameau et al., 2015). On the other hand, overexpression of GA catabolism genes has variable effects on branching (Busov et al., 2003; Mauriat et al., 2011). This might reflect the complex feed-back structure of the GA pathway (Hedden and Thomas, 2012), with branching depending on the balance of GA biosynthesis, catabolism, and receptor abundance (Ueguchi-Tanaaka et al., 2005).

Most overexpression studies focus on GA levels in the stem, while the stem and AXBs might respond independently. The importance of GA levels in AXBs themselves is demonstrated by the up-regulation of GA biosynthesis genes in AXBs of hybrid aspen that are released from dormancy by...
chilling (Rinne et al., 2011). In support of this, GA supply also induces branching in a number of woody species (Saure, 1985; Rinne et al., 2011; Ni et al., 2015). AXBs of hybrid aspen resemble short day (SD)-induced terminal buds (TBs), both morphologically and molecularly (Rinne et al., 2015), although their vascularization might initially differ (Pizzolato and Larson, 1977). In TBs, dwarving and eventual cessation of development involve the narrowing of plasmodesmata (PD) and their subsequent closing during dormancy establishment (Rinne and van der Schoot, 1998; Ruonala et al., 2008). This is achieved by local activation of 1,3-β-glucon synthase (glycosyl transferase GT48-family), an enzyme that deposits callose in dormancy sphincter complexes at the PD of the SAM (Ruonala et al., 2008). Such precisely regulated disruption of symplasmic circuitry by ‘circuit breakers’ (Paul et al., 2014a) effectively prevents metabolic and electric coupling, and the movement of transcription factors and morphogens that sustain SAM function (Rinne and van der Schoot, 1998; Rinne et al., 2001; Kim et al., 2003; Urbanus et al., 2010). The central role of PD is also evident from experiments in Arabidopsis, in which artificially induced callose at the PD of the SAM compromised or terminated SAM function (Daum et al., 2014). It is likely, therefore, that symplasmic alterations are also instrumental in regulating AXB development and branching.

Symplasmic permeability is under homeostatic regulation by 1,3-β-glucon synthases and callose-hydrolyzing 1,3-β-gluconases (glycosyl hydrolase GH17) at PD and sieve plate pores (Rinne and van der Schoot, 2003; Levy et al., 2007; Levy and Epel, 2009). During dormancy establishment, the balance shifts toward net callose deposition, whereas chilling-induced release reverses it. In hybrid aspen, the central 1,3-β-gluconases involved are GA responsive (Rinne et al., 2011). Xylem feeding of GA and chilling of dormant AXBs affect the local expression of these callose-degrading enzymes, indicating that GA and 1,3-β-gluconases are players in dormancy release and branching.

GH17-families are relatively large in both Arabidopsis (~50 members) and Populus trichocarpa (~100). Family members are grouped into three clades (Doxey et al., 2007; Rinne et al., 2011), 10% of which have a cell wall-related function (Geisler-Lee et al., 2006; Doxey et al., 2007). Shoot elongation mostly involves α-clade members, whereas γ-clade members function prominently in defense and stress responses (Maule et al., 2011; Rinne et al., 2011; Benitez-Alfonso et al., 2013; Sager and Lee, 2014). Many α-clade members possess a glycosylphosphatidylinositol (GPI) anchor for attachment to the exoleaflet of the plasma membrane, and/or a carbohydrate-binding module 43 (CBM43) that binds cell wall callose (Doxey et al., 2007; Levy and Epel, 2009; Simpson et al., 2009). The more distant γ-clade members lack known signals and may associate with lipid bodies (Rinne et al., 2011). In Arabidopsis, members of the α-clade localize to PD (Levy et al., 2007; de Storme and Geelen, 2014; Gaudioso-Pedraza and Benitez-Alfonso, 2014; Knox and Benitez-Alfonso, 2014). In hybrid aspen, both α- and γ-clade members can localize to PD, potentially targeting callose deposits at distinct parts of the PD (Paul et al., 2014b).

Here, we investigate the putative roles of GA pathway and GH17-family genes in embryonic shoot dwarfing, para-dormancy, dormancy, and branching. The results suggest that AXBs are GA deficient but highly sensitive to GA owing to low expression of the rate-limiting GA biosynthesis gene GIBBERELLIN 3-OXIDASE2 (GA3ox2) and high expression of two GIBBERELLIN INSENSITIVE DWARF1-like (GID1-like) GA receptor genes. The rate-limiting GA3ox2 is significantly up-regulated in AXBs after decapitation, but not in decapitation-insensitive dormant AXBs, showing that SDs block its expression whereas chilling de-represses it. Expression analyses of GA-responsive α- and γ-clade members of the GH17-family indicate that these enzymes modulate symplasmic permeability during AXB transitions. Functional studies in which representative members of the α- and γ-clade were overexpressed in hybrid aspen support the conclusion that GA biosynthesis, and its downstream effects on GH17-family members, are crucial in AXB formation and activation.

Materials and methods

Plant material and designs for experiments

Hybrid aspen (Populus tremula × P. tremuloides) clone T89 was micropropagated in vitro, planted in soil as described previously (Ruonala et al., 2008), grown in a greenhouse under long days (LDs) (18 h light) at ~18 °C and 75–80% relative humidity, and watered twice a day. Natural light was supplemented to a level of 200 μmol m−2 s−1 at 400–750 nm (Osram). After 6 weeks, when the plants had reached a height of 70–80 cm, and elongation and leaf production rates were constant, the plants were divided into three groups. Group one was kept in LDs as a control group. Group two was moved to an SD regime with a 10 h photoperiod for 5 weeks to induce dormancy. Group three was kept in LDs and decapitated just above the BMP to remove apical dominance. Kinetics of AXB outgrowth and dormancy were described previously (Rinne et al., 2015).

Initiation of SD-induced TBs is strictly controlled by photoperiod length and requires repeated photoperiodic cycles. To compare AXBs and TBs of the same developmental age, it was necessary to determine how many SD cycles were required to initiate TBs. As no molecular markers of TB initiation are known, and the early morphological changes are hidden from sight by a cluster of newly initiated leaves, we used two indirect growth analyses methods. First, we counted the number of nascent leaves and primordia around the SAM of an LD plant, and assessed how many of these leaves unfolded under SDs before the TB became visible, which took 21–28 d. With the second growth analysis, we established where in this trajectory the first signs of irreversible TB formation could be detected in plants that were returned to LDs.

AXB anatomy and staining of lipid and callose

AXBs were fixed overnight at 4 °C in 2% (v/v) glutaraldehyde and 3% (v/v) paraformaldehyde in 100 mM phosphate citrate buffer, as described earlier (Rinne et al., 2001). Briefly, samples were infiltrated gradually with LR White Resin (LRW) of increasing concentration (3% (v/v) paraformaldehyde in 100 mM phosphate citrate buffer, as described earlier (Rinne et al., 2001). Briefly, samples were infiltrated gradually with LR White Resin (LRW) of increasing concentration (30–70%), and kept for 4 d in 100% LRW. Polymerization was conducted at 55 °C for 24h. Lipid body accumulation was studied by staining longitudinally cut 1–3 μm thick sections with Sudan II (black) (1% w/v in 70% ethanol), filtered prior to use. The sections were stained in a continuously stirred solution at 50 °C, subsequently cleared with 70% ethanol for 2 min, and mounted in water for light microscopic
observation. For callose staining, longitudinal fresh hand sections were made under a dissection microscope through AXB–node units that were submersed in 10 mM 2-deoxy-d-glucose (2-DDG; Sigma-Aldrich) to inhibit formation of cutting-induced callose (Rinne et al., 2005). Subsequently, the sections were incubated in the dark for ~1 h in 0.1× K$_2$HPO$_4$ buffer (pH 9.5) containing aniline blue (0.01%) and 2-DDG (100× stock). Callused border epidermis cells were observed under a fluorescence microscope as described previously (Rinne et al., 2005), and photographed with a digital camera (Nikon Coolpix 995).

RNA extraction and quantitative RT–PCR analysis

The apex and every second one of the subtended AXBs up to node 30 (counted from the apex) were collected from LD plants. In parallel, apices and maturing AXBs at the even nodal positions 2–14 were collected from SD plants at SD weeks 2, 3, and 5. Two types of decapitation experiments were carried out. In the first experiment, the five AXBs immediately under the cut were collected at 8 d post-decapitation. In the second experiment, the first AXB under the cut was collected after 1, 2, 3, 5, and 7 d. For each data point, RNA was extracted from six plants, and divided into two biological replicates, each containing material from three individual plants. To exclude possible diurnal variations in gene expression, sampling was carried out at exactly the same time of day for all analyses.

RNA was extracted from 0.2 g of frozen tissue and ground in a mortar with 750 μl of extraction buffer (Qiagen RTL buffer, containing 1% PVP-40). After addition of a 0.4 vol. of K$_2$CO$_3$ at pH 6.5 and further grinding, the solution was transferred to a 2 ml tube, incubated on ice for 15 min, and centrifuged at 12 000 rpm at 4 °C for 15 min. The supernatant was transferred to a 1.5 ml tube, and a 0.5 vol. of 100% EtOH was added. The mix was transferred to two RNeasy-spin columns and further processed in accordance with instructions of the Qiagen Plant RNA isolation kit. RNA was DNase (Ambion) treated, cleaned using the total RNA purification system ‘Purelink RNA mini kit’ (Invitrogen), and reverse transcribed with SuperScriptIII reverse transcriptase (Invitrogen). Quantitative reverse transcription–PCR (qRT–PCR) analyses were performed with the ABI Prism 7500 Fast sequence detection system using SYBR Green PCR master mix (Applied Biosystems). Transcript levels were normalized using an actin gene. Gene-specific primer sequences for the analyses were designed using Primer3 (http://frodo.wi.mit.edu/ primer3) (Supplementary Table S1 at JXB online).

Transformation and in vitro culture of hybrid aspen

For vector construction and Agrobacterium-mediated transformation, genomic clones of T89 GH17 44 and GH17 102 were amplified and subsequently cloned into the pMDC32 destination vector (Curtis and Grossniklaus, 2003) using the Gateway system (Invitrogen), replacing the ccdB gene downstream of the dual Cauliflower mosaic virus (CaMV) 35S promoter. The overexpression vectors were transformed into the Agrobacterium tumefaciens strain GV3101 (pMP90). Hybrid aspen (clone T89) was first grown in vitro under sterile conditions for 4–5 weeks (photoperiod 18 h, light intensity 28 μmol m$^{-2}$ s$^{-1}$, temperature 20 °C). Explants of these plants were used for Agrobacterium-mediated transformation (Häggman et al., 2003). Briefly, 3–5 mm long internodal stem segments were cut and placed on solid callus production medium, hereafter referred to as MS [half-strength Murashige and Skoog (1/2× MS) medium; Duchefa, M0225], which contained 2% sucrose, 0.5 μM 6-benzylaminopurine (BAP; Sigma, B3408), 4 μM 2,4-dichlorophenoxyacetic acid (2,4-D; Fluka, 31518), and 0.7% agar (pH 5.6). Explants were incubated under light for 3 d prior to co-cultivation with Agrobacterium. Fresh cultures of A. tumefaciens strain GV3101, containing the binary plasmids Pro35S::GH17 44 or Pro35S::GH17 102, were grown in Luria broth (LB) medium (1.0% tryptone, 0.5% yeast extract, and 1.0% NaCl) containing antibiotics (20 μg ml$^{-1}$ rifampicin, Sigma, R3501), 30 μg ml$^{-1}$ gentamicin (Sigma, G6896), and 100 μg ml$^{-1}$ kanamycin (Sigma, K4378). The cultures were grown until OD$_{600}$ (optical density) of ~0.5. Subsequently, the cultures were centrifuged for 10 min. at 3000 rpm, washed once in distilled water, and re-suspended in a MS1 solution which was supplemented with 2% sucrose to an OD$_{600}$ of ~0.5. Acetylsyringone (Sigma, D134406) was added to the culture in a final concentration of 20 μM, and cultures were further grown at room temperature for 1 h with shaking (60 rpm). The explants were co-cultured with pre-inoculated A. tumefaciens cells for 4 h (room temperature, 60 rpm), and then incubated on MS1 plates for 48 h in the dark. Thereafter Agrobacterium cells were removed by rinsing the explants three times in 1/2× MS liquid medium containing 2% sucrose, and twice in 1/2× MS liquid medium containing 2% sucrose, 300 mg 1$^{-1}$ vancomycin (Duchefa, V0155), and 500 mg 1$^{-1}$ clorofuran (cefotaxime sodium, Duchefa, C0111), for 15 min per wash (room temperature, 60 rpm). The explants were blotted on a sterile filter paper and transferred to MS1 plates with antibiotic selections (15 μg ml$^{-1}$ hygromycin (Sigma, H9773) and 250 μg ml$^{-1}$ clorofuran) to initiate callus growth. At a size of ~5 mm, the calluses were transferred to the shoot regeneration medium MS2 [1/2× MS medium containing 2% sucrose, 0.1 μM thidiazuron (TDZ; Duchefa, T9016), and 0.7% agar at pH 5.6], with antibiotic selections (15 μg ml$^{-1}$ hygromycin and 250 μg ml$^{-1}$ clorofuran). Approximately 5 cm tall plantlets were transferred to the rooting medium MS3 [1/2× MS medium supplemented with 100 mg 1$^{-1}$ myo-inositol, 2.85 μM indole acetic acid (IAA; Sigma, I2886), and 0.8% agar at pH 5.6] without any antibiotic selection. Rooted cuttings were transferred to soil for greenhouse growing. Expression of overexpressed genes in different lines was analyzed by qPCR in leaves, stems, and AXBs. The number, position, and length of sylleptic branches was monitored after the plants had grown in soil in the greenhouse for 2 months.

Bioinformatics

BLAST searches in GenBank and the P. trichocarpa genome v2.0 (Tuskan et al., 2006) databases (http://www.ncbi.nlm.nih.gov/BLAST; http://www.phytozome.net) was used to identification of GH17 and GA biosynthesis and signaling genes. ClustalW (http://www.ebi.ac.uk/Tools/msa/clustalw2) was used to perform multiple sequence alignments. The PLACE database was used to compare plant cis-acting regulatory DNA elements in the putative promoter region (1000 bp upstream) of GH17 genes of P. trichocarpa (http://togodb.biociscencedbc.jp/togodb/view/place_main) (Higo et al., 1999). The expression map of Arabidopsis genes, e-FP browser, was used to search the expression patterns of orthologous genes in the GH17-family (http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi). Gene-specific primer sequences for the qPCR analysis were designed using Primer3 (http://frodo.wi.mit.edu/primer3).

Results

Time frame of axillary and terminal bud formation

AXB formation in hybrid aspen is a default process (Fig. 1). Branching of the T89 clone is proleptic, meaning that under normal growth conditions AXBs only give rise to branches after a period of (winter) dormancy. This makes it an excellent system to investigate AXB development, para-dormancy, and decapitation-induced branching, as spontaneous bud burst is absent. AXBs contain a dwarfed shoot system, the development of which is completed at the BMP (Fig. 1). In our experimental conditions, reaching this point takes ~4 weeks (Rinne et al., 2015). By then, the AXB contains ~10–12 embryonic leaves. In comparison, TB formation is a non-default process that is under strict control by SDs, and results in dormancy establishment in ~5 weeks. To enable comparison of AXBs and TBs at a similar developmental stage,
we first aimed to establish the point in time at which the SAM transitions to a TB. Because molecular markers for scale initiation, an early sign of the transition, have not been identified, an indirect growth analysis method was used (Materials and methods). The analysis showed that under SDs two leaves (±0.9) were produced before leaf production ceased and TB formation started. Under the assumption that the plastochron did not change, the reprogramming of leaf primordia to scale primordia took 3 d. In an alternative procedure, plants were exposed to SDs for a restricted number of days before returning them to LDs, to assess the earliest time point at which the apex would become morphologically affected. A very short exposure to SDs, for 2–4 d, did not produce visible signs of TB formation, but newly formed internodes around the BMP (not shown), but weakening of apical dominance directly preceded TB initiation (i.e. changes in the SAM and scale initiation), the results indicate that the SD response at the SAM was as early as 3 d after the start of SD exposure. The time frame to form a dormant TB at week 5 is thus roughly similar to the 4 weeks needed to form a completed AXB.

**Axillary buds amass lipid bodies in long and short days**

Structural analyses previously showed that the developmental trajectories of TBs and AXBs converge on a shared morphogenetic program (Rinne et al., 2015). We here investigate if para-dormant and dormant AXBs also share the unique cellular features that characterize dormant TBs. Cytohistological studies, using the lipid stain Sudan II (black), showed that the lipid bodies that amass in the SAM and rib meristem of SD-induced dormant TBs (van der Schoot et al., 2011) are also prominent in para-dormant and dormant AXBs. A direct comparison showed that the amount of lipid bodies was very similar (Fig. 2). In all cases, intensely black lipid bodies crowded the cytoplasm of the SAM and the rib meristem cells. In contrast, the apex of growing LD plants contained very few lipid bodies that stained light-blueish (Fig. 2A). In para-dormant AXBs, lipid bodies were particularly prominent in the upper cell layer of the central zone (L1), and in the rib meristem (Fig. 2B). In conclusion, lipid body accumulation appears to be an integral part of bud development as such, preceding para-dormancy as well as dormancy. Another significant feature that is shared by all buds is the loss of cellular water. Under LDs, AXBs desiccated considerably, lowering their water content in developing AXBs from 80% to ~60% around the BMP, and then further during para-dormancy to 50% (Supplementary Fig. S2). Taken together, lipid body accumulation and desiccation accompany the formation of TBs as well as AXBs.

**Expression profiles of GA pathway genes in developing buds**

Xylem feeding of GA\(_4\) activates dormant AXBs and induces them to grow out (Rinne et al., 2011), suggesting that the dwarfing of the embryonic shoot is caused by a deficiency in GA signaling. We investigated this possibility by analyzing the expression of genes that are central to GA catabolism, biosynthesis, and signaling in AXBs of different developmental stages and activity states. The results show that the GA pathway genes that function in dormancy cycling at the shoot apex have expression patterns characteristic of developing, para-dormant, and dormant AXBs (Fig. 3).

**GA catabolism**

The \(GA2-OXIDASE\)-family (\(GA2ox\)) of GA-catabolizing genes (Gou et al., 2011) functions in reducing the levels of bioactive GA. Several \(GA2ox\) members were expressed in AXBs (Fig. 3A). The transcript levels typically increased in developing AXBs (i.e. until they reached the mature stage around the BMP). \(GA2ox2\) expression was below the detection limit (not shown), but \(GA2ox1\), \(GA2ox3\), and \(GA2ox4\)
were significantly up-regulated during AXB development, from 10- to 400-fold. \textit{GA2ox6} was very modestly up-regulated (not shown). \textit{GA2ox1} and \textit{GA2ox3} expression was very low in proliferating apices compared with developing AXBs, suggesting a comparatively diminished catabolism of GA in the apex (Fig. 3A). This is in agreement with the consen-
sus view that GA is necessary to facilitate shoot elongation growth. Consistent with this, the SD-induced cessation of apical growth and TB formation resulted in the up-regulation of three of the four GA catabolism genes (Fig. 3A). \textit{GA2ox4} stood out because it was strongly up-regulated not only in dormant TBs but also in the AXBs that were still developing when SD exposure started. Because these developing AXBs also become dormant (Rinne et al., 2015), the GA catabolism gene \textit{GA2ox4} might be central in dormancy establishment.

\textbf{GA biosynthesis}

The two selected members of the \textit{GA3-OXIDASE}-family (\textit{GA3ox}) that function in the last biosynthesis step of biologically active GA showed opposite expression patterns (Fig. 3B). \textit{GA3ox1} positively reflected AXB development. It was hardly expressed in growing apices, gradually up-regulated in developing AXBs, and maintained at a relatively steady level below the BMP. In contrast, \textit{GA3ox2} was characteristic of apical growth and elongation. It was highly expressed in the apex, but considerably down-regulated during AXB development (Fig. 3B). Consistent with these distinct patterns, SDs induced up-regulation of \textit{GA3ox1} and down-regulation of \textit{GA3ox2} in TBs and developing AXBs. The members of the \textit{GA20-OXIDASE}-family (\textit{GA20ox}), which produce precursors for the \textit{GA3ox}-family, were also differentially regulated. \textit{GA20ox8} showed very little change during AXB development and under SDs. As this gene is chilling regulated (Rinne et al., 2011), it might function predominantly in dormancy release. In contrast, \textit{GA20ox4} (Fig. 3B) and \textit{GA20ox3} (not shown) were up-regulated during AXB development under LDs, reaching a steady expression level around the BMP. \textit{GA20ox4} was further up-regulated to a much higher level in developing TBs and AXBs that established dormancy under SDs (Fig. 3B), suggesting that during para-dormancy and dormancy high levels of biologically inactive precursors are produced. This would allow a rapid production of biologically active GA if needed, because only the final GA3ox enzyme has to be produced.

\textbf{GA signaling}

Expression of \textit{DE1}, a \textit{DELLA}-like gene, was low and stable in AXBs up to the BMP, and slightly down-regulated in older AXBs. Under SDs, it was exclusively up-regulated in TBs (Fig. 3C). Because DELLA proteins interact with a GA receptor, thereby affecting tissue sensitivity to GA, we also analyzed the expression of two putative homologs of the rice gene \textit{GID1}, encoding the GID1 receptor (Ueguchi-Tanaka et al., 2005). These genes, \textit{GID1A} and \textit{GID1B}, align with Arabidopsis \textit{GID1A} and \textit{GID1B}, respectively. Both genes were substantially up-regulated during AXB development, up to 10- and 100-fold, respectively, and further up-regulated during dormancy. This is congruent with the hypothesis that para-dormant and dormant buds are GA deficient as well as GA sensitized.

\textbf{Unique expression patterns of PD callose-related GH17 genes}

The involvement of GA-regulated, PD-targeting, and callose-
hydrolyzing members of the 1,3-β-glucanase-family (GH17)
was previously investigated in relation to TB dormancy cycling (Rinne et al., 2011). We here analyzed their comparative expression during AXB development, para-dormancy, and dormancy. The α-clade members (Fig. 4A) are post-transcriptionally modified for excretion to the cell wall. They possess a GPI anchor (GH17_102, GH17_65, and GH17_33) and/or a CBM43 (GH17_102, GH17_79, and GH17_98). The CBM43 module facilitates their targeting to cell wall callose that is present around PD orifices (Doxey et al., 2007; Levy and Epel, 2009; Simpson et al., 2009). Under LDs,
expression of the α-clade genes *GH17_102* and *GH17_79* gradually declined from apices toward the AXB at the BMP, with apices having a twice as high expression level (detailed in Supplementary Fig. S3). In contrast, expression of *GH17_65*, *GH17_33*, and *GH17_98* was more equal in the apex and all subtending AXBs (Fig. 4A). Under SDs, *GH17_102* and
**GH17_79** were also down-regulated in the dormant TB. In contrast, **GH17_65** and **GH17_33** were up-regulated in dormant AXBs and TBs, while **GH17_98** was only up-regulated in dormant AXBs (Fig. 4A). Regulation of the γ-clade genes (**GH17_44**, **GH17_101**, **GH17_39**, and **GH17_37**) was distinctly different from that of the α-clade (Fig. 4B). Their expression levels increased strongly but transiently during AXB development up to the BMP, closely correlating with the accumulation of lipid bodies, where they might be stored for later recruitment (Paul et al., 2014b). In the dormant TBs and AXBs, all four γ-clade genes were substantially up-regulated, although less so in the case of **GH17_37** (Fig. 4B).

**Decapitation induces GA biosynthesis and GH17 genes, and activates AXBs**

Although in the T89 clone AXBs do not branch out in the year in which they are formed, they can be induced to do so by decapitation. Branching depends on functional symplasmic connections between the stem and the AXB, which is expected to require both GA and GA-regulated GH17 members that diminish callose at PD and sieve plate pores (Levy and Epel, 2009; Rinne et al., 2011). We therefore analyzed the expression of GA pathway genes in conjunction with the GA-induced GH17 members. Para-dormant AXBs possessed considerable amounts of callose at PD, especially in the SAM, provascular tissue, and sieve tubes plus surrounding ground tissue of the embryonic stem, a wedge-like extension that connects the SAM to the main stem (Supplementary Fig. S4A–C). Activated AXBs of single-node cuttings showed that in 3 d callose deposits were strongly diminished (Supplementary Fig. S4D), before signs of bud burst occurred (Rinne et al., 2015). For gene expression analyses, plants were decapitated directly above the BMP, at nodal position 12, and gene expression was analysed after 8 d in five successive AXBs proximal to the cut (insets of Figs 3 and 4). Although all five AXBs might become activated, those closest to the cut will usually win the competition and become the new leading shoot (Rinne et al., 2015). Therefore, the most proximal AXB was chosen to analyze the expression of a number of selected genes in more detail. The data show that significant changes in gene expression took place within 24 h. In all cases, decapitation substantially affected the expression of GA catabolism, biosynthesis, and signalling genes (Fig. 3).

The GA-catabolizing **GA2ox** genes (**GA2ox1** and **GA2ox3**), which were significantly up-regulated during AXB development, were down-regulated 1 week after decapitation in all five proximal AXBs, but most significantly in the most proximal one (Fig. 3A, insets). The GA-catabolizing enzyme, **GA2ox4**, which was somewhat up-regulated during AXB development and more substantially during dormancy establishment, was reduced only in the AXB closest to the cut (Fig. 3, inset). The GA-catabolizing gene **GA2ox5** did not show a decapitation-induced down-regulation in the proximal AXB, but was up-regulated instead (Fig. 3A, insets). A time-resolved analysis of one member in this family, **GA2ox3**, showed that in the uppermost AXB, the gene was down-regulated within a single day to the low levels characteristic of a proliferating apex (Fig. 3D).

The GA biosynthesis gene families **GA3ox** and **GA20ox** were both affected by decapitation. The genes **GA3ox1** and **GA20ox4**, which were up-regulated during AXB development, were down-regulated in the proximal AXB. In contrast, the four lower AXBs, which commonly did not produce branches, up-regulated the expression (Fig. 3B, insets). In sharp contrast to **GA3ox1** and **GA20ox4**, the gene **GA3ox2** was up-regulated in the uppermost AXB to ~10- and 70-fold during the first and second day post-decapitation, respectively (Fig. 3D). By day 5, it reached a 100-fold expression level, which subsequently declined to the level typical of growing apices. In the remaining four AXBs, the expression of **GA3ox2** was lower at 8 d post-decapitation (Fig. 3B, inset). Notably, this gene is characteristically expressed in the growing apex of non-decapitated plants, while it is repressed in dormancy (Fig. 3B). Thus, **GA3ox2** appears to be crucial in activation of the proximal AXB and may enable it to become the new leading shoot.

GA signaling was also severely affected by decapitation. At 8 d post-decapitation, expression of the **DELLA1**-like gene **DEI** was strongly reduced in the three uppermost AXBs (Fig. 3C, inset). The GA receptor genes **GID1A** and **GID1B**, which were highly expressed at the BMP in intact plants, were considerably down-regulated in the AXB proximal to the cut 8 d post-decapitation (Fig. 3C, inset). However, a day-by-day analysis of **GID1B** showed that in this AXB the expression shortly peaked at the second day after decapitation (Fig. 3D). The **GID1A** and **GID1B** repression occurred when the AXB had become more elongate and the GA biosynthesis had increased. In contrast, in the lower AXBs, **GID1A** and **GID1B** were up-regulated 2- to 3-fold, in agreement with the low expression of the GA biosynthesis gene **GA3ox2** (Fig. 3A, B, insets). The expression ratio of **GID1A** and **GID1B** to **GA3ox2** was also high during AXB and TB development, reflecting their inhibited state.

As anticipated, decapitation also affected the GA-regulated genes that encode the PD-related GH17-family proteins. The expression of growth- and GA-γ-regulated α-clade GH17 genes (**GH17_102**, **GH17_65**, **GH17_33**, **GH17_79**, and **GH17_98**) was up-regulated during the first day post-decapitation, up to 10-fold in the case of **GH17_102**, and remained high during AXB activation (Fig. 4A). The γ-clade GH17 genes **GH17_39**, **GH17_101**, **GH17_37**, and **GH17_44** that were strongly up-regulated during AXB development were down-regulated by decapitation, especially in the proximal AXB (Fig. 4B, insets).

**Overexpression of GH17 members in hybrid aspen induces distinct branching phenotypes**

The distinct expression patterns of the α-clade and γ-clade GH17 members during apical growth, AXB development, AXB para-dormancy, and decapitation-induced AXB activation warrant the hypothesis that these enzymes have unique roles in developmental processes. To probe their putative function, we constitutively overexpressed a representative of each clade in the T89 clone of hybrid aspen (Fig. 5A, B). **GH17_102** was selected from the α-clade because its
expression was highest in the apex (Fig. 4A) and it was closely co-expressed with PIN1 and the meristem-related gene WUS (Rinne et al., 2015). From the γ-clade we selected GH17_44, encoding a lipid body-related protein, because it was virtually absent from the proliferating apex, but significantly up-regulated in AXB (Fig. 4B). Other reasons for selecting these genes relate to the differences in their promoter regions.

Promoter analysis, using a publicly available database of plant cis-acting regulatory DNA elements http://togodb.biosciencedbc.jp/togodb/view/place_main, showed that the promoter region (1000bp upstream) of GH17_102 was enriched with multiple central elements that imply auxin and GA regulation (Supplementary Table S3). GH17_102 was the only one among the studied GH17-family genes which has the UP2 motif in its promoter region. In Arabidopsis, this motif relates to decapitation-induced AXB activation (Tatematsu et al., 2005). In addition, GH17_102 possesses the same target sequences for the meristem-specific transcription factors LEAFY and WUS as are found in the intron of the AGAMOUS gene (Lohmann et al., 2001). In contrast, lipid body-related GH17 genes possess the sugar-responsive cis-element SRE in their promoter region (Supplementary Table S3), which in Arabidopsis is also present in a number of genes that are down-regulated by decapitation (Tatematsu et al., 2005).

Since GH17_102 was predominantly expressed in proliferating meristems, we anticipated that its overexpression would release the repressed para-dormant state of AXBs, thereby changing the proleptic branching habit of the T89 clone to a sylleptic one (Figs 5G, 6A; Supplementary Table S2).
Consistent with this, GH17_102 overexpression lines (Fig. 5B) showed AXB burst and formation of sylleptic branches (Fig. 5F, H). Notably, AXB branching was not sporadic but patterned. Branches gradually emerged in an acropetal pattern, suggesting that once the AXBs had reached a certain level of development and maturation they were able to burst and produce a branch. We observed this in tissue culture plants and in younger soil-grown plants (Fig. 5F, H). When plants grew taller than 0.5 m this acropetal branching pattern diminished. All lines were SD responsive and made TBs in the same time frame as the wild type.

Because GH17_44 transcripts were virtually absent from the proliferating apex, we expected that its overexpression would interfere with apex function. Indeed, all overexpression lines had apical deviations. Two of the GH17_44 overexpression lines (lines 1 and 2) produced large terminal buds and pronounced AXBs under LDs, and as early as the tissue culture stage (Fig. 5C). Although the large AXBs could burst after decapitation, the apical meristems of the emerging branches would again become arrested. Other, more severely affected GH17_44 overexpression lines showed fasciation and SAM termination, whereas some lines continued growth repetitively from the uppermost AXB in a sympodial manner (Fig. 5D). At least eight of the GH17_44 overexpression lines (lines 8–12 and 16–18), those with less enhanced expression levels, were able to grow in soil. Contrary to expectation, these lines also produced branches (Figs 5J, 6), not from mature AXBs like the GH17_102 overexpression lines but from numerically younger AXBs. Remarkably, branching occurred in distinct recurrent flushes, even when plants became older (Supplementary Table S2). Line 9 displayed the most severe branching phenotype (Fig. 6B), reminiscent of the behavior of wild-type plants that were exposed for 6–12 d to SD conditions, and often branched from the uppermost AXB (Supplementary Fig. S1). As overexpression of GH17_44 could induce spontaneous bud formation in some lines, it seems likely that recurrent branching in less severe GH17_44 lines was due to the recurrent formation of a TB-like structure, which then causes syllepsy indirectly.

Discussion

The ‘architectural model’ of a tree species describes its branch geometry and patterns of reiterated development (Hallé et al., 1978). However, the shape of a crown is developmentally plastic, and influenced by internal competition between branches as well as interactions with the environment (Tomlinson, 1983; Turnbull, 2005; Barthélémy and Caraglio, 2007; Costes et al., 2014; Costes and Gion, 2015). How the crown maintains its operational hierarchy over multiple seasons is virtually unknown. In a recent study, we compared AXB and TB development as well as the expression of meristem-specific and branching-related genes, including BRC1 and MAX1 (Rinne et al., 2015). This comparison showed that in terms of structure, development, and gene expression patterns, AXBs and TBs are very much alike, despite distinct differences in how and where they are formed. Furthermore, AXBs appeared morphogenetically active up to the BMP, where they become para-dormant, a phase where morphogenesis has simply ceased (Rinne et al., 2015). The present results support a model in which GA pathway genes, and genes encoding GA-inducible 1,3-glucanases that degrade PD callose during dormancy release, are subject to a comparable regulation in development and activation of para-dormant AXBs (Fig. 7).

Developing AXBs are sinks that accumulate lipid bodies

Dormant TBs of deciduous perennials can contain substantial amounts of lipid bodies. Known examples include birch (Rinne et al., 2001) and hybrid aspen (Rinne et al., 2011; van der Schoot et al., 2011), and the evergreen Rhododendron (Lynch and Rivera, 1981). That dormant AXBs also possess
lipid bodies is therefore not surprising (Fig. 2D). However, the finding that AXBs that develop under LDs also contain them (Fig. 2B) shows that lipid bodies are not exclusive indicators of dormancy, but rather an integral part of bud development. That such carbon reserves accumulate in young developing AXBs reflects that they are sinks that import from the phloem. Commonly, AXBs are not regarded as sinks; however, in hybrid aspen developing AXBs are not very different from the proliferating apex, except for the fact that cell enlargement and shoot elongation are postponed to the next season. That only AXBs above the BMP become dormant might relate to the fact that developing hybrid aspen buds are sinks, which are presumably accessible to photoperiodically regulated signals (Fig. 1), such as FLOWERING LOCUS T (FT) (Böhlenius et al., 2006).

Lipid bodies are known to be abundant in seeds, but they are constituents of most plant cells, although in vegetative cells they are small and few in number (van der Schoot et al., 2014). This changes during bud development when lipid bodies amass in quantities resembling those in seeds (Fig. 2; Rinne et al., 2011; van der Schoot et al., 2011). Like in seeds, which can also overwinter, lipid bodies in AXBs and TBs may serve as energy stores and confer cryo-protection during dehydration (Shimada et al., 2008). Although less severe than in seeds, developing TBs and AXBs also dehydrate during development, irrespective of whether or not they establish dormancy (Faust et al., 1991; Welling et al., 1997; Supplementary Fig. S2). 1,3-β-Glucanases of the γ-clade can localize at lipid bodies, which target the plasma membrane and PD, as shown with immunolabeling and by eGFP tagging of 1,3-β-glucanases and the lipid body marker protein oleosin (Rinne et al., 2001, 2011; Paul et al., 2014b). The accumulated lipid bodies may function during chilling-induced release from dormancy as vehicles that shuttle 1,3-β-glucanases to PD to remove callose and restore the symplasmic organization of the SAM (Rinne et al., 2011; Paul et al., 2014b). The fact that lipid bodies and 1,3-β-glucanases of the γ-clade GH17-family also accumulate in developing AXBs suggests that they have a similar role as in TBs, which makes sense because AXBs as a rule overwinter in proleptic hybrid aspen before they can branch. Considering that lipid bodies are rapidly mobilized by xylem feeding of GA (Rinne et al., 2011), it seems possible that the γ-clade GH17 members stored at lipid bodies may also have a role in activation of para-dormant AXBs, while subsequent outgrowth relies on GA₄-induced 1,3-β-glucanases of the GH17 α-clade.

GA pathway genes in AXB inhibition and activation

The present findings that AXBs do not express the growth-related GA biosynthesis gene GA₃ox2, unless decapitated (Fig. 3B, D), while they do express the GA precursor genes (Fig. 3B), suggests that AXBs can rapidly produce biologically active GA by simply activating the last biosynthesis step. The implication is that para-dormant AXBs are poised for activation and outgrowth (Fig. 7). It is unclear how in para-dormant AXBs GA₃ox2 expression is maintained at a low level, but it could relate to a lack of auxin. Various GA biosynthesis genes are known to be induced by auxin (Figerio et al., 2006), including a putative Pisum ortholog of the poplar gene GA₃ox2 (Ross et al., 2000). While decapitation of hybrid aspen up-regulated the proximal AXB GA₃ox2 (Fig. 3D) and PIN1-like genes (Rinne et al., 2015), the catabolism gene, GA₂ox₃, was down-regulated (Fig. 3A). Because GA catabolism appears to be restricted to a narrow zone under the SAM, protecting the SAM from the disturbing influence of GA (Sakamoto et al., 2001; Jasinski et al., 2005), this could direct GA-induced cell elongation to the rib zone. Considering that GA increases and stabilizes PIN₁ abundance (Willige et al., 2011), GA might stimulate auxin transport from the AXM to the rib zone, where it could promote its own auxin-mediated biosynthesis.

An additional crucial component in GA signaling is the GA receptor GID1 (Ueguchi-Tanaka et al., 2005), the levels of which reflect GA sensitivity, as demonstrated in rice through overexpression of GIDI1 (Ueguchi-Tanaka et al., 2005). The substantial up-regulation of two Populus homologs of GIDI during AXB development (Fig. 3C) may thus enhance the sensitivity of the system to GA, while it is already poised for activation by high expression of GA precursor genes. The data indicate that AXBs in general are considerably more sensitive to GA than the growing apex, as GIDI expression levels were ~10- (GIDI1) and...
70- (GID1B) fold higher in LDs, and >200-fold higher in SDs (GID1B) (Fig. 3C, D). Thus, para-dormant AXBs might be poised for activation by tight regulation of GA pathway genes (Fig. 3B, C), as well as by the GA-responsive and rib meristem-resident gene CENTRORADIALISLIKE1 (CENL1) (Rinne et al., 2015). This poised state is balanced by high expression of GA-deactivating genes (Fig. 3A) and the branch inhibitor genes BRC1 and MAX1 (Rinne et al., 2015).

In theory, in a GA-deficient but highly GA-sensitized system, a slight increase in GA biosynthesis could bring about AXB activation and outgrowth. The assumption that the AXBs represent such a system is supported by the finding that xylem feeding of the growth-related GA4 induces canonical bud burst, even when the AXBs are dormant (Rinne et al., 2011). On the other hand, GA3, which is synthesized via a parallel pathway from the branch point GA12 precursor (Hedden and Thomas, 2012) and which is involved in chilling-induced release from dormancy, did not promote burst of dormant buds (Rinne et al., 2011). The reason for this difference remains unclear, but could relate to different binding affinities of GA3 and GA4 for the GID1 receptor (Ueguchi-Tanaka et al., 2005), or to the fact that they regulate different genes/paralogs. For example, in hybrid aspen, GA3 up-regulates genes encoding the bud development-related γ-clade members, while GA4 up-regulates growth-related α-clade members of the GH17-family (Rinne et al., 2001, 2011). Together, this might explain why the capacity of applied GA to replace vernalization or chilling in different species has been inconsistent (Mutasa-Göttgens and Hedden, 2009).

The present conclusion that GA signaling has a pivotal role in regulating AXB activation is seemingly contradicted by the observation that GA biosynthesis mutants and plants that overexpress GA metabolism genes are often highly branched. Such observations, for example in Pisum, Arabidopsis, and turfgrass, seem to suggest that GA is not required for AXB outgrowth, and that GA may actually inhibit branching (Murfet and Reid, 1993; Silverstone et al., 1997; Agharkar et al., 2007; Lo et al., 2008; Rameau et al., 2015). Silencing of the GA catabolism gene GA2ox in tomato increased levels of GA4 in AXBs, but reduced branching (Martinez-Bello et al., 2015). Considering the locally restricted expression of some GA2ox genes as discussed above, silencing may result in GA spilling over into the SAM. Silencing of GA2ox genes might then inhibit branching through experimentally induced mislocalization of GA, compromising SAM integrity. In Populus, the reduction of GA levels by activation tagging of GA2ox, or the use of different promoters in overexpression studies, reduced or accelerated branching (Busov et al., 2003; Mauriat et al., 2011). The present suggestion that AXBs are GA deficient but GA sensitized could throw some light on these phenomena. In GA-deficient AXBs, overexpression of GA catabolism genes is unlikely to reduce GA levels much further, while it will affect the main shoot (see above; Willige et al., 2011), decreasing the PATS and increasing branching due to reduced apical dominance.

**GA pathway genes in dormant AXBs**

Judged from gene expression profiles, dormant AXBs may accumulate even more precursors of GA than para-dormant AXBs (Fig. 3B). Nonetheless, previous experiments with internode cuttings showed that this would be to no avail. As long as they do not receive sufficient chilling prior to decapitation, dormant AXBs fail to up-regulate GA3ox2, the gene that catalyzes the final GA biosynthesis step (Rinne et al., 2011). Only after sufficient chilling at 5 °C and subsequent return to 18 °C is the gene up-regulated prior to bud burst (Rinne et al., 2011). Together with the current data, this suggests that SDs repress GA3ox2, and that de-repression requires chilling followed by growth-promoting temperatures. This would be a mechanism that safeguards the AXBs from freezing damage, and ensures that growth is initiated at the proper time in spring (Rinne et al., 2011). Coupling of photo- and thermo-periods in scheduling developmental events is also characteristic of other dormancy related genes, including FT1 and FT2 (Hsu et al., 2006, 2011; Ruonala et al., 2008; Rinne et al., 2011; Brunner et al., 2014). In all cases, these alterations might be expected to involve shifts in methylation and chromatin remodeling (Paul et al., 2014a; Considine and Considine, 2016). Interestingly, in seeds of various species, light- and cold-inducible GA3ox genes are central to GA-regulated developmental events (Yamauchi et al., 2004).

**The rib meristem as a target of branch regulators**

A major function of GAs is to stimulate growth through cell elongation (Hedden and Thomas, 2012). This involves activation of the rib meristem and elongation of the descendent cells in the rib zone (Ruonala et al., 2008). A role for GA3ox2 in cell elongation is supported by the strong decapitation-induced up-regulation in the proximal AXB before it starts to telescope out (Fig. 3B). Interestingly, although reduced expression of the growth-related gene GA3ox2 might be a major cause of dwarfing and hindrance to AXB outgrowth, expression of the non-growth-related paralog GA3ox1 increased during AXB development (Fig. 3B). This suggests that in AXBs of hybrid aspen, GA3ox1 and GA3ox2 might have distinct expression domains as they function during bud formation and branching, respectively.

A hormone that opposes GA function as a branch regulator is strigolactone (Shinohara et al., 2013). Although the roles of strigolactone in trees are as yet uncharted, a genuine strigolactone pathway has been identified in poplar (Czarnecki et al., 2014). In hybrid aspen, MAX1 genes are highly expressed in AXBs themselves, and down-regulated during decapitation, indicating that strigolactone could counteract GA locally in AXBs (Rinne et al., 2015). For example, auxin-recruited GA might stimulate cell division and enlargement in the rib meristem and rib zone, as well as the establishment of an auxin export path to the main stem, while strigolactone opposes it by destabilizing PIN1 proteins (Willege et al., 2013). As in hybrid aspen AXBs develop while MAX1 genes are up-regulated, it can be concluded that strigolactone does not impede cell division and morphogenesis, but it could contribute to
inhibition of cell elongation. Such a role for strigolactone has been proposed for the dicots *Pisum* and Arabidopsis (Agusti et al., 2011; de Saint-German et al., 2013). On the other hand, in the monocot rice, strigolactone does not affect cell elongation but it negatively regulates cell division (Hu et al., 2010). Based on the present data, a plausible scenario is that during AXB formation, GA₃ can promote cell division because the simultaneous production of strigolactone and other inhibitors prevents elongation.

**AXB outgrowth requires high capacity delivery conduits**

AXB activation and outgrowth can be thought of as conceptually distinct, but both probably depend on a functional stem–AXB connection. Activation of a para-dormant AXB triggers the re-initiation of morphogenesis, involving patterning processes at the AXM (SAM) and cell division activity in the rib meristem and rib zone. In the light of current understanding, this is regulated by cytokinin, auxin production in leaf primordia, and export to the stem (Müller and Leyser, 2011). Sugar is proposed to be the activation signal in caulescent *Pisum* (Mason et al., 2014). In hybrid aspen, sugar import via the phloem is unlikely due to the cessation of sink activity in para-dormant AXBs, and the accumulation of callose at PD and sieve plates (Supplementary Fig. S4C). Similarly, xylem import of root-produced cytokinin may also not be the initial event given the desiccated state of the AXB and the hard bud scales that prevent evaporation.

Thus, the persistent problem in understanding AXB activation is that the AXM (SAM) of the embryonic shoot must sense a change in the shoot system, as a putative signal must be relayed to it from the stem. We hypothesize that this signal travels via reinvigorated symplasmic connections. This would require first the function of 1,3-β-glucanases that reduce callose at PD and sieve plates (Levy et al., 2007, 2009), opening up symplasmic connections between the stem and AXB. Auxin in the stem could promote removal of dormancy callose in the phloem, as shown for some woody perennials (Aloni et al., 1991; Aloni and Peterson, 1997). In contrast, in the hypocotyl of the herbaceous annual Arabidopsis, auxin can stimulate callose deposition, enabling polar auxin transport during phototropic bending (Han et al., 2014). A role for bud-produced auxin as well as GA-induced 1,3-β-glucanases in AXB activation is supported by the elevated expression of several callose-hydroryzing α-clade GH17 members in decapitation-activated AXBs (Fig. 4A), the reduced callose deposits in activated AXBs on internode cuttings (Supplementary Fig. S4D), and the spontaneous branching of transgenic plants overexpressing the α-clade 1,3-β-glucanases (Fig. 5H).

Analysis of cis-elements in the promoter region of the decapitation-inducible α-clade GH17_102 further supports the conjecture that this gene may be under regulation of GA and auxin (Supplementary Table S3). Its promoter has target sequences that are similar to those present in the intron of the AGAMOUS gene for LEAFY and WUS binding (Lohmann et al., 2001). The WUS protein, which moves through PD from the WUS domain to the overlying CLV3 domain (Daum et al., 2014), might facilitate its own movement by recruiting GH17_102 or another α-clade GH17 enzyme to dilate the PD transport channel. This seems feasible as in Arabidopsis the WUS expression domain (Yadav et al., 2009) overlaps with that of a gene homologous to GH17_102 (Supplementary Fig. S5). Similarly, α-clade genes may play a direct role in AXB activation as the phenotype of GH17_102 overexpressors display sylleptic-like branches at lower nodes (Fig. 5H). Although the branches in the GH17_102 lines were short, the phenotype demonstrates that simply increasing PD permeability can overcome an arsenal of inhibitory factors that are present in mature AXBs. The present data suggest that re-functionalization of symplasmic conduits between stem and dwarfed shoot is a determinant of AXB activation and outgrowth (Fig. 7). How this is co-ordinated at the molecular level remains to be established.

**AXB development is accompanied by γ-clade GH17 gene expression**

The enhanced expression of γ-clade GH17 genes during AXB development reflects the concurrent accumulation of lipid bodies (Figs 2, 4B). The encoded γ-clade members typically lack known membrane localization signals, and instead they may be stored at lipid bodies, attached to the half-membrane via electrostatic interactions, hydrophobic binding, or accessory molecules (Paul et al., 2014; van der Schoot et al., 2014). They do not play a major role in growth, as decapitation decreases the expression of γ-clade GH17 genes in the proximal AXBs (Fig. 4B). However, as they are accumulating and precociously stored in all AXBs, it seems possible that they function in the initial re-invigoration of symplasmic connections between AXB and stem, prior to outgrowth. The promoter region of the γ-clade member GH17_44 contains the sugar-responsive cis-element SRE that in Arabidopsis is negatively regulated by sugar (Tatematsu et al., 2005). Notably, GH17_44, is up-regulated in maturing AXBs that cease sugar import, and down-regulated during AXB activation when sugar import is restarting (Fig. 4B).

The relationship between γ-clade GH17 gene expression and AXB development is not just correlative, as evidenced by overexpression of GH17_44, which could induce bud development. Strong overexpressors developed pronounced TBs and AXBs already at the tissue culture stage under LDs (Fig. 5C). The reason is unclear, but possibly overproduction of the enzymes depletes the membrane lipid pool by directing lipid synthesis towards lipid body formation thereby inhibiting elongation and promoting bud formation. Moderate transgenic GH17_44 lines showed an acrotropic sylleptic branching pattern, with sporadic flushes from the uppermost AXBs (Fig. 6B). These flushes could be due to the tendency to start a TB that temporarily weakens apical dominance, as observed after a brief SD exposure in wild-type plants (Supplementary Fig. S1). The branching of GH17_44 lines resembles that of tropical trees in which rhythmic meristem activity is accompanied by branching at the end of each flush (Halle et al., 1978). Our data suggest that transient reduction in symplasmic connectivity and sink activity at the apex of the main stem may contribute to
sylleptic branching from young AXBs that are still sinks. What the possible connection is between \textit{GH17.44} and the cytokinin-induced gene \textit{EARLY-BUD BREAK1} (EBB1), overexpression of which induces somewhat similar sylleptic branches in \textit{P. deltoides} (Yordanov et al., 2014), remains to be established. It is tempting to speculate that EBB1-like proteins act in the wake of GA-inducible 1,3-\beta-glucanases that re-functionalize symplasmic input conduits.

Conclusions

The present findings add to previous work (Rinne et al., 2015) showing that AXB-intrinsic controls are crucial in branching of trees. Novel intrinsic controls include GA pathway genes and GA-regulated GH17-family members that reduce callose depositions at PD and sieve plate pores. Collectively, they enable and facilitate transport and communication between stem and bud.

Accession numbers

The \textit{P. trichocarpa} gene model identifiers (Tuskan et al., 2006) and/or sequence accessions used for real-time qPCR analysis are listed in Supplementary Table S1.

Supplementary data

Supplementary data are available at \textit{JXB} online.

Figure S1. Reversal of dormancy development and subsequent branching.

Figure S2. Developmental desiccation of AXBs.

Figure S3. Expression analyses of \textit{GH17.102} and \textit{GH17.79}.

Figure S4. Callose in para-dormant and activated AXBs.

Figure S5. Arabidopsis eFP browser.

Table S1. \textit{Populus trichocarpa} genes, identifiers, and primer pairs.

Table S2. Syllepsis-like branching of \textit{GH17} overexpressors.

Table S3. Analysis of putative promoter regions of \textit{GH17} genes.

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References

Aloni R, Peterson CA. 1997. Auxin promotes dormancy callose removal from the phloem of \textit{Magnolia kobus} and callose accumulation and early wood vessel differentiation in \textit{Quercus robur}. Journal of Plant Research 110, 37–44.

Aloni R, Raviv A, Peterson CA. 1991. The role of auxin in the removal of dormancy callose and resumption of phloem activity in \textit{Vitis vinifera}. Canadian Journal of Botany 69, 1825–1832.

Agharkar M, Lomba P, Altpeter F, Zhang H, Kenworthy K, Lange T. 2007. Stable expression of \textit{AtGA2ox1} in a low-input turfgrass (\textit{Paspalum notatum} Flugge) reduces bioactive gibberellin levels and improves turf quality under field conditions. Plant Biotechnology Journal 5, 791–801.

Aguilar-Martínez JA, Poza-Carrón C, Cubas P. 2007. Arabidopsis \textit{BRANCH1} acts as an integrator of branching signals within auxillary buds. The Plant Cell 19, 459–472.

Agusti J, Herold S, Schwarz M, et al. 2011. Strigolactone signaling is required for auxin-dependent stimulation of secondary growth in plants. Proceedings of the National Academy of Sciences, USA 108, 20242–20247.

Balla J, Kalousek P, Reinöhl V, Friml J, Procházka S. 2011. Competitive canalization of PIN-dependent auxin flow from auxillary buds controls pea bud outgrowth. The Plant Journal 65, 571–577.

Barthélémy D, Caraglio Y. 2007. Plant architecture: a dynamic, multilevel and comprehensive approach to plant form, structure and ontogeny. Annals of Botany 99, 375–407.

Benítez-Alfonso Y, Faulkner C, Pendle A, Miyashima S, Helaïruttia Y, Maule A. 2013. Symplastic intercellular connectivity regulates lateral root patterning. Developmental Cell 26, 136–147.

Böhlenius H, Huang T, Charbonnel-Campaa L, Brunner AM, Jansson S, Strauss SH, Nilsson O. 2006. CO/FT regulatory module controls timing of flowering and seasonal growth cessation in trees. Science 312, 1040–1043.

Brewer PB, Dun EA, Gui R, Mason MG, Beveridge CA. 2015. Strigolactone inhibition of branching independent of polar auxin transport. Plant Physiology 68, 1820–1829.

Brunner AM, Evans LM, Hsu CY, Sheng X. 2014. Vernalization and the chilling requirement to exit bud dormancy: shared or separate regulation? Frontiers in Plant Science 5, 732.

Busov VB, Meilan R, Pearce DW, Ma C, Rood SB, Strauss SH. 2003. Activation tagging of a dominant gibberellin catabolism gene (GA 2-oxidase) from poplar that regulates tree stature. Plant Physiology 132, 1283–1291.

Ceulemans R, Steller RF, Hinckly TM, Isebrands JG, Heilman PE. 1990. Crown architecture of \textit{Populus} clones as determined by branch orientation and branch characteristics. Tree Physiology 7, 157–167.

Cline MG. 1991. Apical dominance. Botanical Review 57, 318–358.

Cline MG. 1997. Concepts and terminology of apical dominance. American Journal of Botany 84, 1064–1069.

Considine MJ, Considine JA. 2016. On the language and physiology of dormancy and quiescence in plants. Journal of Experimental Botany 67, 3189–3203.

Costes E, Crespel L, Denoyes B, Morel P, Demene M-N, Lauri E-P, Wenden B. 2014. Bud structure, position and fate generate various branching patterns along shoots of closely related Rosaceae species: a review. Frontiers in Plant Science 5, 666.

Costes E, Gion J-M. 2015. Genetics and genomics of tree architecture. In: Adam-Blondon A-F, Plomion C, eds. Land plants—trees. Amsterdam: Elsevier Ltd Academic Press, 157–200.

Curtis MD, Grossniklaus U. 2003. A gateway cloning vector set for high-throughput functional analysis of genes in plants. Plant Physiology 133, 462–469.

Czarnecki O, Yang J, Wang X, Wang S, Muchero W, Tuskan GA, Chen JG. 2014. Characterization of \textit{MORE AXILLARY GROWTH} genes in \textit{Populus}. PLoS One 9, e102757.

Daum G, Medzihradzky A, Suzuki T, Lohmann JU. 2014. A mechanistic framework for non-cell autonomous stem cell induction in \textit{Arabidopsis}. Proceedings of the National Academy of Sciences, USA 111, 14619–14624.

de Saint-Germain A, Ligerot Y, Charbonnel-Campaa L, Brunner AM, Isebrands JG, Heilman PE. 2011. Strigolactone signaling at plasmodesmata: molecular regulations and developmental relevance. Frontiers in Plant Science 5, 138.

Doebely J, Stec A, Hubbard L. 1997. The evolution of apical dominance in maize. Nature 386, 485–488.

Domagalska MA, Leyser O. 2011. Signal integration in the control of shoot branching. Nature Reviews Molecular Cell Biology 12, 211–221.
Doxey AC, Yaish MWF, Moffatt BA, Griffith M, McConkey BJ. 2007. Functional divergence in the Arabidopsis β-1,3-glucoamylase gene family inferred by phylogenetic reconstruction of expression states. Molecular Biology and Evolution 24, 1045–1055.

Dun EA, Ferguson BJ, Beveridge CA. 2006. Apical dominance and shoot branching. Divergent opinions or divergent mechanisms? Plant Physiology 142, 812–819.

Faust M, Liu D, Millard MM, Stutte GW. 1991. Bound versus free water in dormant apple buds—a theory for endodormancy. HortScience 26, 887–890.

Ferguson BJ, Beveridge CA. 2009. Roles for auxin, cytokinin, and strigolactone in regulating shoot branching. Plant Physiology 149, 1929–1944.

Frigerio M, Alabadi D, Pérez-Gómez J, García-Cárcel L, Phillips AL, Hedden P, Blázquez MA. 2006. Transcriptional regulation of gibberellin metabolism genes by auxin signaling in Arabidopsis. Plant Physiology 142, 553–563.

Garrison R. 1955. Studies in the development of axillary buds. American Journal of Botany 42, 257–266.

Gaudioso-Pedraz R, Benitez-Alfonso Y. 2014. A phylogenetic approach to study the origin and evolution of plasmodesmata-localized glycosyl hydrolases family 17. Frontiers in Plant Science 5, 212.

Geisler-Lee J, Geisler M, Coutinho PM, et al. 2006. Poplar carbohydrate-active enzymes. Plant Physiology 140, 946–962.

Gou J, Ma C, Kadmiel M, Gai Y, Strauss S, Jiang X, Busov V. 2011. Tissue-specific expression of Populus C19 GA-2-oxidases differentially regulate above- and below-ground biomass growth through control of bioactive GA concentrations. New Phytologist 192, 626–639.

Grbic V, Bleecker AB. 2000. Axillary meristem development in Arabidopsis thaliana. The Plant Journal 21, 215–223.

Greif T, Clarens O, Schäfer E, Müller D, Herrero R, Schmitz G, Han X, Hyun TK, Zhang M, Kumar R, Koh EJ, Kang BH, Lucas WJ, Häggman H, Frey AD, Aronen T, Ryynänen L, Julkunen-Tiitto R, Tiimonen H, Pihakaski-Maunsbach K, Jokipii S, Chen X, Tiimonen J, Kallio PT. 2007. Expression of Vitreoscilla hemoglobin in hybrid aspen (Populus tremula×tremuloides). Plant Biotechnology Journal 1, 287–300.

Hallé AF, Oldeman RAA, Tomlinson PB. 1978. Tropical trees and forests: an architectural analysis. Berlin: Springer.

Han X, Hyun TK, Zhang M, Kumar R, Koh EJ, Kang BH, Lucas WJ, Kim JY. 2014 Auxin–callose-mediated plasmodesmal gating is essential for tropic auxin gradient formation and signaling. Developmental Cell 28, 132–146.

Hedden P, Thomas SG. 1993. Developmental mutants: In: Casey R, Davies DP, eds. Peas, genetics, molecular biology and biotechnology. Wallingford, UK: CAB International, 186–216.

Higuchi M, Jang Y, Inouye K, Tsurumaki H, Yamamoto N, Nishiyama Y, Fujita Y, Ohashi Y, Shinoda K, Takano SA, et al. 2005. KNOX action in Arabidopsis is mediated by coordinate regulation of cytokinin and gibberellin activities. Current Biology 15, 1560–1565.

Kim JY, Yua Z, Jackson D. 2003. Developmental regulation and significance of KNOX protein trafficking in Arabidopsis. Development 130, 4351–4362.

Knox JP, Benitez-Alfonso Y. 2014. Roles and regulation of plant cell walls surrounding plasmodesmata. Current Opinion in Plant Biology 22, 93–103.

Kohlen W, Charnikova T, Liu Q, et al. 2011. Strigolactones are transported through the xylem and play a key role in shoot architectural responses to phosphate deficiency in nonarbuscular mycorrhizal host Arabidopsis. Plant Physiology 155, 974–987.

Levy A, Epel BL. 2009. Cytology of the (1,3)-β-glucan (callose) in plasmodesmata and sieve plate pores. In: Basic A, Fincher GB, Stone BA, eds. Chemistry, biochemistry and biology of (1,3)-β-glucans and related polysaccharides. Burlington, MA: Academic Press, Elsevier, 439–463.

Levy A, Erlanger M, Rosenthal M, Epel BL. 2007. A plasmodesmata-associated β-1,3-glucanase in Arabidopsis. The Plant Journal 49, 689–692.

Li CJ, Bangerth F. 1999. Autoinhibition of indoleacetic acid transport in the shoots of two-branched pea (Pisum sativum) plants and its relationship to correlative dominance. Physiologia Plantarum 106, 415–420.

Lo SF, Yang SY, Chen KT, et al. 2008. A novel class of gibberellin2-oxidases control semidwarfism, tillering, and root development in rice. The Plant Cell 20, 2603–2618.

Lohmann JU, Hong RL, Hobe M, Busch MA, Parcy F, Simon R, Weigel D. 2001. A molecular link between stem cell regulation and floral patterning in Arabidopsis. Cell 105, 793–803.

Long J, Barton MK. 2000. Initiation of axillary and floral meristems in Arabidopsis. Developmental Biology 218, 341–353.

Martínez-Bello L, Moritz T, López-Díaz I. 2015. Silencing C19-GA 2-oxidases induces parthenocarpic development and inhibits lateral branching in tomato plants. Journal of Experimental Botany 66, 6879–6910.

Masclaux-Daubresse C, Russel JJ, Babs B, Woenckhaus BN, Beveridge CA. 2014. Sugar demand, not auxin, is the initial regulator of axillary dominance. Proceedings of the National Academy of Sciences, USA 111, 6092–6097.

Maule AJ, Benitez-Alfonso Y, Faulkner C. 2011. Plasmodesmata—membrane tunnels with attitude. Current Opinion in Plant Biology 14, 683–690.

Mauriat M, Sandberg LG, Moritz T. 2011. Proper gibberellin localization in vascular tissue is required to control auxin-dependent leaf development and bud outgrowth in hybrid aspen. The Plant Journal 67, 805–816.

Morris SE, Cox MCH, Ross JJ, Kristiansen S, Beveridge CA. 2005. Auxin dynamics after decapitation are not correlated with the initial growth of axillary buds. Plant Physiology 138, 1665–1672.

Muratet C, Reid JB. 1993. Developmental mutants: In: Casey R, Davies DP, eds. Peas, genetics, molecular biology and biotechnology. Wallingford, UK: CAB International, 186–216.

Mutasa-Göttgens E, Hedden P. 2009. Gibberellin as a factor in floral regulatory networks. Journal of Experimental Botany 60, 1979–1989.

Ni J, Gao C, Chen M-S, Pan B-Z, Ye K, Xu X-F. 2015. Gibberellin promotes shoot branching in the perennial woody plant Jatropha curcas. Plant and Cell Physiology 56, 1655–1666.

Niwa M, Daimon Y, Kuratani K, et al. 2013. BRANCH1 interacts with FLOWERING LOCUS T to repress the floral transition of the axillary meristems in Arabidopsis. The Plant Cell 25, 1228–1242.

Paul NK, Rinne PLH, van der Schoot C. 2014a. Shoot meristems of deciduous woody perennials: self-organization and morphogenetic transitions. Current Opinion in Plant Biology 17, 86–95.

Paul NK, Rinne PLH, van der Schoot C. 2014b. Refurbishing the plasmodesmal chamber: a role for lipid bodies? Frontiers in Plant Science 5, 40.

Phillips IDJ. 1975. Apical dominance. Annual Review of Plant Physiology 26, 341–367.

Pizzolato T, Larson PR. 1977. Auxin flow development in Populus deltoides. II Late ontogeny and vascularization. American Journal of Botany 64, 849–860.

Rameau C, Bertheloot J, Leduc N, Andrieu B,oucher F, Sakr S. 2015. Multiple pathways regulate shoot branching. Frontiers in Plant Science 5, 741.

Rinne P, Tuominen H, Sundberg B. 1993. Growth patterns and endogenous indole-3-acetic-acid concentrations in current-year coppice shoots and seedlings of two Betula species. Physiologia Plantarum 88, 403–412.
Rinne PLH, Kaikuranta PM, van der Schoot C. 2001. The shoot apical meristem restores its symplasmic organization during chilling-induced release from dormancy. The Plant Journal 26, 249–284.

Rinne PLH, Paul JK, Vahala J, Ruonala R, Kangasjärvi J, van der Schoot C. 2015. Long and short photoperiod buds in hybrid aspen share structural development and expression patterns of marker genes. Journal of Experimental Botany 66, 6745–6760.

Rinne PLH, van den Boogaard R, Mensink MGJ, Kopperud C, Kormelink R, Goldbach R, van der Schoot C. 2005. Tobacco plants respond to the constitutive expression of the toposovirus movement protein M5I(M) with a heat-reversing sealing of plasmodesmata that impairs development. The Plant Journal 43, 689–707.

Rinne PLH, van der Schoot C. 1998. Symplasmic fields in the tunica of the shoot apical meristem coordinate morphogenetic events. Development 125, 1477–1485.

Rinne PLH, van der Schoot C. 2003. Plasmodesmata at the crossroads between dormancy, development and defense. Canadian Journal of Botany 81, 1182–1197.

Rinne PLH, Wellin A, Vahala J, Ripel L, Ruonala R, Kangasjärvi J, van der Schoot C. 2011. Chilling of dormant buds hyperinduces FLOWERING LOCUS T and recruits GA-inducible 1,3-β-glucanases to reopen signal conduits and release dormancy in Populus. The Plant Cell 23, 130–146.

Romberger JA. 1963. Meristems, growth and development in woody plants. USDA Technical Bulletin no. 1293. Washington, DC: US Department of Agriculture.

Ross JJ, O’Neill DP, Smith JJ, Kerckhoffs LHH, Elliott RC. 2000. Evidence that auxin promotes gibberellin A(1) biosynthesis in pea. The Plant Journal 21, 547–552.

Ruonala R, Rinne PLH, Kangasjärvi J, van der Schoot C. 2008. CENL1 expression in the rib meristem affects stem elongation and the transition to dormancy in Populus. The Plant Cell 20, 59–74.

Sager R, Lee J-Y. 2014. Plasmodesmata in integrated cell signalling: insights from development and environmental signals and stresses. Journal of Experimental Botany 65, 6337–6358.

Sakamoto T, Kobayashi M, Itoh H, Tagiri A, Kayano T, Tanaka H, Iwahori S, Matsuoka M. 2001. Expression of a gibberellin 2-oxidase gene around the shoot apex is related to phase transition in rice. Plant Physiology 125, 1508–1516.

Saure MC. 1985. Dormancy release in deciduous fruit trees. American Horticultural Society of Science 7, 239–300.

Scott TK, Case DB, Jacobs WP. 1987. Auxin–gibberellin interaction in apical dominance. Plant Physiology 82, 1329–1333.

Shimada TL, Shimada H, Takahashi H, Fukao Y, Haranishimura L. 2008. A novel role of oleosins in freezing tolerance of oilseeds in Arabidopsis thaliana. The Plant Journal 55, 798–809.

Shinohara N, Taylor C, Leyser O. 2013. Strigolactone can promote or inhibit shoot branching by triggering rapid depletion of the auxin efflux protein PIN1 from the plasma membrane. PLoS Biology 11, e1001474.

Silverstone AL, Mak PY, Martinez EC, Sun TP. 1997. The new RGA locus encodes a negative regulator of gibberellin response in Arabidopsis thaliana. Genetics 146, 1087–1099.

Simpson C, Thomas C, Findlay K, Bayer E, Maule AJ. 2009. An Arabidopsis GPI-anchor plasmodesmal neck protein with callose binding activity and potential to regulate cell-to-cell trafficking. The Plant Cell 21, 581–594.

Tatematsu K, Ward S, Leyser O, Kamiya Y, Nambara E. 2005. Identification of cis-elements that regulate gene expression during initiation of axillary bud outgrowth in Arabidopsis. Plant Physiology 138, 757–766.

Thimm KV, Skoog F. 1934. On the inhibition of bud development and other functions of growth substances in Vicia faba. Proceedings of the Royal Society of London B: Biological Sciences 114, 317–339.

Tomlinson PB. 1983. Tree architecture. American Scientist 71, 141–149.

Turnbull CGN. ed. 2005. Plant architecture and its manipulation. Oxford: Blackwell Publishers Ltd.

Tuskan GA, Difazio S, Hansson S, et al. 2006. The genome of black cottonwood, Populus trichocarpa (Torr. & Gray). Science 313, 1596–1604.

Ueguchi-Tanaka M, Ashikari M, Nakajima M, et al. 2005. GIBBERELLIN INSENSITIVE DWARF1 encodes a soluble receptor for gibberelin. Nature 437, 693–698.

Urbanus SL, Martenelli AP, Dinh QD, Aizza LCB, Dornelas MC, Angenent GC, Immink RGH. 2010. Intercellular transport of epidermis-expressed MAD5 domain transcription factors and their effect on plant morphology and floral transition. The Plant Journal 6, 60–72.

van der Schoot C, Paul LK, Paul SB, Rinne PLH. 2011. Plant lipid bodies and cell–cell signaling. A new role for an old organelle? Plant Signaling and Behavior 6, 1732–1738.

van der Schoot C, Paul LK, Rinne PLH. 2014. The embryonic shoot a lifeline through winter. Journal of Experimental Botany 65, 1699–1712.

van der Schoot C, Rinne PLH. 2011. Dormancy cycling at the shoot apical meristem: transitioning between self-organization and self-arrest. Plant Science 180, 120–131.

Waldie T, McCulloch H, Leyser O. 2014. Strigolactones and the control of plant development. The Plant Journal 79, 607–622.

Welling A, Kaikuranta P, Rinne P. 1997. Photoperiodic induction of dormancy and freezing tolerance in Betula pubescens. Involvement of ABA and dehydrins. Physiologia Plantarum 100, 119–125.

Willige BC, Isono E, Richter R, Zourelioud M, Schwechheimer C. 2011. Gibberelin regulates PIN-FORMED abundance and is required for auxin transport-dependent growth and development in Arabidopsis thaliana. The Plant Cell 23, 2184–2195.

Wu R, Hinckley TM. 2001. Phenotypic plasticity of sylleptic branching: genetic design of tree architecture. Critical Reviews in Plant Science 20, 467–485.

Wu R, Stettler RF. 1998. Quantitative genetics of growth and development in Populus, III. Phenotypic plasticity of crown structure and function. Heredity 81, 299–310.

Yadav RK, Girke T, Pasala S, Xie M, Reddy GV. 2009. Gene expression map of the Arabidopsis shoot apical meristem stem cell niche. Proceedings of the National Academy of Sciences, USA 106, 4941–4946.

Yamauchi Y, Ogawa M, Kuwahara A, Hanada A, Kamiya Y, Yamaguchi S. 2004. Activation of gibberellin biosynthesis and response pathways by low temperature during imbibition of Arabidopsis thaliana seeds. The Plant Cell 16, 367–378.

Yordanov YS, Ma C, Strauss SH, Busov VB. 2014. EARLY BUD-BREAK 1 (EBB1) is a regulator of release from seasonal dormancy in poplar trees. Proceedings of the National Academy of Sciences, USA 111, 10001–10006.

Zawaski C, Busov VB. 2014. Roles of gibberellin catabolism and signaling in growth and physiological response to drought and short-day photoperiods in Populus trees. PLoS One 9, e86217.