What do we know about growth of vessel elements of secondary xylem in woody plants?

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ABSTRACT

Despite extensive knowledge about vessel element growth and the determination of the axial course of vessels, these processes are still not fully understood. They are usually explained as resulting primarily from hormonal regulation in stems. This review focuses on an increasingly discussed aspect – mechanical conditions in the vascular cambium. Mechanical conditions in cambial tissue are important for the growth of vessel elements, as well as other cambial derivatives. In relation to the type of stress acting on cambial cells (compressive versus tensile stress) we: (i) discuss the shape of the enlarging vessel elements observed in anatomical sections; (ii) present hypotheses regarding the location of intrusive growth of vessel elements and cambial initials; (iii) explain the relationship between the growth of vessel elements and fibres; and (iv) consider the effect of mechanical stress in determining the course of a vessel. We also highlight the relationship between mechanical stress and transport of the most extensively studied plant hormone – auxin. We conclude that the integration of a biomechanical factor with the commonly acknowledged hormonal regulation could significantly enhance the analysis of the formation of vessel elements as well as entire vessels, which transport water and minerals in numerous plant species.

Key words: vessel element, mechanical stress, diurnal strain, vascular cambium, ring-porous wood, diffuse-porous wood, intrusive growth

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I. INTRODUCTION

Processes of growth and differentiation of cambial derivatives, deposited by the vascular cambium – the lateral plant meristem responsible for production of secondary xylem and secondary phloem cells (Crag, Lyons-Sobaski & Wise, 2018) (Fig. 1) – are extremely complex and insufficiently explained. Even such basic issues as functioning of the vascular cambium and the development of derivative cells, the mechanism of increase in circumference of the cambial cylinder (Wloch et al., 2009, 2013; Wilczek et al., 2014;
Fig 1. Scheme showing position of vascular cambium and cambial derivatives deposited on the xylem and phloem side within a tree stem. Cambial initials of the vascular cambium are shown in dark violet. Other undifferentiated cambial cells are shown in light violet. Cells of the vascular cambium are purposely spaced apart in order to indicate the periclinal and anticlinal walls of the cambial initials. Pith and periderm are also shown. Periderm (phellem, phellogen and phellogen) is marked in different hues of brown. Phellogen is responsible for the deposition of phellem and phelloderm. The proportional thickness (number of cells) of different tissues is not taken into account. CI, cambial initials; Pe, periderm; Ph, phloem; Pi, pith; VC, vascular cambium; Xy, xylem.

Miodek et al., 2021), and the course of intrusive growth of cambial initials, vessel elements, and wood fibres (Jura et al., 2006; Wilczek et al., 2018; Gizińska, Miodek & Kojs, 2021) still remain debated. One of the key processes related to cambial activity is xylogenesis, i.e. wood formation, with its underlying mechanisms of growth of vessel elements and vessel pattern formation. Mature vessel elements are dead cells, devoid of protoplasts, connected to each other by perforation plates, and forming long strands – vessels (Evert, 2006; Fig. 2). These are specialised for conduction of water and minerals from the roots to the leaves. Vessels can take various sizes, both in length and diameter (Zimmermann, 1982). Moreover, dicotyledonous trees show distinct distribution patterns of vessels with different tangential and radial dimensions within their annual growth rings (Beck, 2010) which are classified into different wood porosity types. In ring-porous wood earlywood vessels are characterized by much larger transverse dimensions than latewood vessels; diffuse-porous wood vessels have similar diameters throughout the annual growth ring; and semi-ring porosity represents an intermediate condition (IAWA Committee, 1989). Woody plants with different porosity types adapt differently to various environmental conditions, with an example of such adaptation being how trees respond to the onset of a new growth season (e.g. after winter frosts) (discussed in Gizińska et al., 2015). Thus, understanding the processes of growth of vessel elements and pattern formation of water-conducting system is of great interest (Zasada & Zahner, 1969; Zimmermann, 1983; Zakrzewski, 1991; Kutin, Sano & Funada, 2003; Wilczek et al., 2011b; Gizińska et al., 2021).

There is much information available concerning the hormonal regulation of cambial activity and the formation of vessel elements (e.g. Zakrzewski, 1983; Uggl et al., 1996; Uggl, Mellerowicz & Sundberg, 1998; Kramer, 2002, 2006; Schrader et al., 2003; Nilsson et al., 2008; Pettersson et al., 2009; Vanneste & Friml, 2009; Grunewald & Friml, 2010; Huang et al., 2010; Santos et al., 2010; Zhang et al., 2010; Aloni, 2013, 2015, 2021; Rutschow, Baskin & Kramer, 2014; Bhalerao & Fischer, 2017). However, in order to understand how vessel elements increase their transverse dimensions (radial and tangential), it is necessary to pinpoint the factors responsible for determining the course of growth of a vessel element, in addition to the commonly accepted contribution of auxin (Aloni, 2007, 2013, 2015; Vanneste & Friml, 2009; Grunewald & Friml, 2010). One such factor is mechanical stress, which seems to play an important role in most, if not all, plant tissues (Hamant et al., 2008; Hamant & Traas, 2010; Nakayama et al., 2012; Landrein & Hamant, 2013; Nick, 2013; Hamant & Haswell, 2017; Ackermann & Stanislas, 2020; Du & Jiao, 2020; Sampathkumar, 2020; Ramos, Maizel & Alim, 2021; Trinh et al., 2021) including cambium and developing xylem (Hejnowicz, 1980, 1997, 2012; Kwiatkowska & Nakielksi, 2011; Kojs, 2013). The role of mechanical pressure in maintaining the functionality of the cambial tissue and progression of xylogenesis was first shown by Brown & Sax (1962), who found that longitudinal incisions of trees stems, reaching into the xylem led to callus formation due to the removal of mechanical stress. Callus is composed of large cells with thin walls, and is formed not only after injury but also in tissue culture (Beck, 2010). Initial cells of the intact vascular cambium have specific shapes depending on their type. Fusiform initials are axially elongated with small radial dimensions and give rise to the axial system of cells (tracheids, vessel elements, axial parenchyma, and wood fibres). Ray initials have an isodiametric shape and give rise to the radial system of cells (i.e. they are responsible for the formation of rays within the xylem and phloem) (Hejnowicz, 2012; Crang et al., 2018). Brown & Sax (1962) showed that the normal pattern of tissue differentiation was restored at incision sites after the formation of new phellogen and cambium. The important role of mechanical stress in the differentiation of xylem cells was confirmed by the lack of radial growth in in vitro cultures of cambial cells; cells of the cambium deprived of adjoining layers of xylem and phloem, and thus of their specific mechanical environment, take a spherical shape (discussed in Srivastava, 1973). Lintilhac & Vesecký (1984) found that applying external compressive forces induces spatially ordered divisions in in vitro culture of pith tissue.

The influence of mechanical conditions on the growth of vessel elements is extremely interesting, which is reflected in the most recent trends in plant sciences dealing with influence of mechanics on morphogenesis (Robinson et al., 2013; Ackermann & Stanislas, 2020; Du & Jiao, 2020; Sampathkumar, 2020; Ramos et al., 2021) and ongoing
discussion regarding the types of stress occurring in the vascular cambium. There are partially opposing views regarding the type of mechanical stress (compressive versus tensile) that acts upon cambial cells in a radial direction (e.g., Hejnowicz, 1997, 2012; Kojs, 2012; Kojs, Malik & Wistuba, 2012). Cambial cells and their derivatives show two basic types of growth—symplastic or intrusive (Wilczek et al., 2018; Gizińska et al., 2021; Miodek et al., 2021). The main type of growth in radial enlargement of a stem is symplastic (Miodek et al., 2021). During symplastic growth, the surfaces of the walls of adjacent cells do not move relative to each other, so contacts between cells do not change and cells grow in a coordinated manner (Tulik, 2007). When individual regions of cell walls grow at different rates, this can lead to changes in contacts between cells, and such growth is defined as intrusive growth (Sinnott & Bloch, 1939; Gizińska et al., 2021). Intrusive growth plays an important role in the vascular cambium and developing xylem (Jura et al., 2006; Wilczek et al., 2011a, b, 2014a; Hejnowicz, 2012), such as participating in cambial cell rearrangement which translates into the specific structure and properties of the wood. During development of secondary xylem of angiosperm trees two types of cells show intrusive growth—fibres (Wilczek et al., 2018) and vessel elements (Gizińska et al., 2021). Intensive intrusive growth of vessel elements can be observed near the border of the cambium and the zone of xylem development. It is therefore possible that the mechanical conditions in which vessel elements and cambial initials grow intrusively are similar. Furthermore, can intrusive growth in both the vascular cambium and the zone of xylem cell development be explained similarly by specific mechanical conditions? To date, anatomical studies have considered the locations and directions of intrusive growth for specific types of cells separately (i.e., cambial initials, wood fibres, and vessel elements). A consequence of this approach may be a failure to understand the higher levels of organization of the complex adaptive system represented by a tree. It may be speculated that in the control of developmental processes, both the biophysical/biomechanical and biochemical/molecular aspects operate in a complementary manner. Below we attempt to elucidate the relationship between the location of intrusive growth of vessel elements and cambial initials, while taking into account the biomechanical background of these processes. Most importantly, this review aims to highlight the potential role of biomechanical forces in vessel element formation. A full explanation of the functioning of the vascular cambium will be possible only by considering all physiological, anatomical, and biomechanical factors involved.

II. DEFINITIONS

Vascular cambium can be defined in various ways (Hejnowicz, 2012). In the narrow definition the cambium is considered to be a single layer of initials, i.e. meristematic cells that are constantly dividing, periclinal as well as anticlinal (Fig. 3). Their derivatives, which can still divide periclinally (mother cells), are included together with the initials in the ‘cambial zone’ (Wilson, Wodzicki & Zahner, 1966). Application of the narrow definition is rather problematic, as in many cases we are unable to precisely identify cambial initials among periclinally dividing cells. Therefore, many authors use a broader definition, in which cambium comprises all dividing cells, i.e. both initials and mother cells (Wilson et al., 1966; Hejnowicz, 2012). However, application of this broad definition may also be problematic. For instance, as cells divide less frequently on the phloem side,
one can sometimes observe undifferentiated cells resembling cambial cells in this region that do not fit into the broad definition of cambium as they do not undergo periclinal divisions. Moreover, adjoining cells belonging to different radial files may exhibit different stages of development, i.e. cells in one radial file may still divide, while cells in another file may have already started to differentiate. Figure 3 provides an example where the majority of dividing cells are located between the red dotted lines, which approximately encompass the vascular cambium according to the broad definition. Nonetheless, some cells which still undergo divisions reach far into the secondary xylem (cells marked with asterisks in Fig. 3). In this review, we therefore define cambium simply as a zone of undifferentiated cells located between the developing secondary xylem and secondary phloem; these cells do not exhibit strong growth in a radial direction (Fig. 3; zone approximated by black dotted lines). Outside the vascular cambium we can recognize developing xylem and phloem (growing and differentiating cells). Beyond this, after completion of cell development, mature secondary xylem and mature secondary phloem can be observed.

III. MECHANICAL STRESS AND AUXIN

Auxin is a plant hormone present in the vascular cambium (Sundberg, Little & Cui, 1990) and in growing and differentiating derivative cells, where it is distributed according to a radial concentration gradient (Uggl et al., 1996, 1998). Auxin is key to the canalization hypothesis, which explains the basis of vascular strand formation (Sachs, 1991) and the acid growth hypothesis regarding cell wall loosening and cell elongation (Rayle & Cleland, 1992). The concentration of auxin affects the size and number of vessel elements (Aloni & Zimmermann, 1983; Zakrzewski, 1991; Aloni, 2021). It is likely that hormonal regulation and mechanical stress are linked, especially during processes involved in the growth of vessel elements and determination of the course of vessels. Nakayama et al. (2012) confirmed that PIN1 (PIN-FORMED1) proteins, involved in polar auxin transport, show sensitivity to a mechanical stimulus. An increase in mechanical strain led to: (i) an overall increase in the amount of PIN1 transporters in a cell, and (ii) an increase in the fraction of transporters localized to the cell membrane (Nakayama et al., 2012). PIN proteins are cellular efflux transporters of auxin (Rubery & Sheldrake, 1974; Raven, 1975; Friml, 2003; Vanneste & Friml, 2009; Heisler et al., 2010; Nakayama et al., 2012). A possible mechanism by which auxin transport is linked to cell wall strain involves: (i) transfer of strain from the wall to the cell membrane (with extracellular receptors likely participating in this process); and (ii) in the areas of higher strain, auxin efflux transporters accumulate on the cell membrane, as a result of the interaction of auxin with auxin binding protein 1 (ABP1), which interferes with constitutive cycling of PINs between the plasma membrane and endosomes leading to the inhibition of clathrin-mediated endocytosis, and possibly also as a result of increased exocytosis (Robert et al., 2010; Li, Friml & Grunewald, 2012; Nakayama et al., 2012). A recent study (Li et al., 2019) showed that PIN1 relocalization is initiated by mechanical stimulation which causes transient changes in cytoplasmic Ca^{2+} concentration. Therefore, it is likely that a...
combination of biomechanical and hormonal regulation plays a key role during the formation of vessel elements as well as whole vessels.

IV. MECHANICAL CONDITIONS IN THE VASCULAR CAMBium AND DEVELOPING XYLEM

Mechanical stress is known to be extremely important for cell functioning, growth, and differentiation. For example, application of a pressure girdle to *Fagus sylvatica* stems inhibits vessel formation (Bauer & Eschrich, 1997). Zhou et al. (2006, 2007) observed that protoplasts placed under compressive stress elongate in the direction perpendicular to the stress, and Lintilhac & Vesecky (1984) and Lynch & Lintilhac (1997) showed that mechanical stress determines the orientation of the division plane in cell protoplasts embedded in agarose blocks. However, the patterns of mechanical stress present in the vascular cambium are still unresolved. Some authors assume that tensile stress acts in a circumferential (tangential) direction (see Fig. 1) and compressive stress acts in a radial direction (Hejnowicz, 1980, 1997, 2012; Kwiatkowska & Nakielski, 2011). According to this view radial compressive stress would result from the radial growth of xylem (growth of derivative cells following periclinal divisions in the cambium) in the presence of constraints imposed by external tissues, i.e. bark (Hejnowicz, 1980; Kwiatkowska & Nakielski, 2011). Observations of intrusively growing cells have shown that their tips are rounded (Kojs, 2013), as intrusive growth of a cell is enabled when space is created in its vicinity (Romberger, Hejnowicz & Hill, 1993; Hejnowicz, 2012), i.e. cell growth occurs into a previously created space. Vessel elements show strong intrusive growth at the border of the cambium and developing xylem (Figs 4 and 5), with this growth mainly involving their radial (anticlinal) walls (Barnett, 1992). Intrusive growth of vessel elements occurs between periclinal (tangential) walls of cells of adjacent radial files of the axial system, which can be easily deduced from the lack of continuity of these files (Wilczek et al., 2011b; Hejnowicz, 2012; Gizińska et al., 2021).

So how is a space into which a vessel element can grow created between the periclinal walls of cells, assuming that cambium is compressed in a radial direction (Kwiatkowska & Nakielski, 2011)? By using incision experiments, Hejnowicz (1980) identified radial tensile stress in the vascular cambium of two ring-porous species in spring. Under such mechanical conditions, separation of the periclinal walls of cells in radial files adjacent to a growing vessel element seems possible, resulting in the formation of spaces available for intrusive growth (Hejnowicz, 2012; Kojs, 2012). Hejnowicz (1980, 1997) speculated that radial tensile stress in the cambium may result from synchronous collapse of sieve tubes, as well as companion cells, in phloem functioning in the previous growing season. The value of tensile stress acting on the middle lamellae between periclinal walls of cells would depend on the degree of synchronization of such phloem cell collapse (Hejnowicz, 1997). According to this concept radial tensile stress in the cambium would change to compressive stress later in the growing season (Hejnowicz, 1997; Kwiatkowska & Nakielski, 2011). However, the study of *Robinia pseudoacacia* xylogenesis (Miodek et al., 2020) revealed that the phloem within the stem seems to collapse much too early to explain the growth of first broad vessel elements of earlywood (Fig. 6) – phloem showed a reduced radial dimension in March, compared to the radial dimension observed in

![Fig 4. Developing broad vessel elements in ring-porous species. (A, B) *Quercus robur* L. branch (seen under polarized light and bright-field illumination, respectively). (C) *Robinia pseudoacacia* L. stem. Note in B that during intensive growth phases vessel elements with thin primary cell walls may exhibit a radially flattened shape. The rounded shape of broad earlywood vessel elements becomes permanent later in development. In A, the polarized light image allows visualization of the progress of maturation, i.e. deposition of secondary cell walls. White and black arrowheads in A and B indicate a fragment of primary cell wall of the developing vessel element. Green arrows in C indicate regions of strong indentations of the neighbouring rays caused by the growing vessel element. These indentations are associated with the tangential growth of vessel elements. Tangential intrusive growth of a vessel element ceases as soon as it contacts ray cells. R, ray; VC, vascular cambium; VE, vessel element; Xy, xylem.](image-url)
August of the previous year. Moreover, assuming that radial stretching of vascular cambium in ring-porous trees is of a transient nature (occurring only in the spring) (Hejnowicz, 1997; Kwiatkowska & Nakielski, 2011): (i) how can one explain growth of vessel elements in latewood of ring-porous species if the cambium is radially compressed in the later part of the growing season; and (ii) how can growth of vessel elements in diffuse-porous species be explained, especially in the later part of the growing season? Hejnowicz (2012) proposed that the rate of radial growth in tissue containing a developing vessel element may differ from that of nearby vesselless regions. Temporally vesselless cambial sectors grow faster, with such growth meaning that they take over the pressure exerted by outer tissues, allowing simultaneous stretching of the sector containing the growing vessel element.

An alternative viewpoint is a hypothesis based on changes in hydration of tree tissues related to transpiration occurring during the day and its cessation at night (Kojs & Rusin, 2011; Kojs, 2012, 2013; Kojs et al., 2012). Numerous reports have identified diurnal cycles of plant tissue deformation – a phenomenon known as diurnal strain (e.g. Kozlowski & Winget, 1964; Simonneau et al., 1993; Ueda & Shibata, 2001; Yoshida et al., 2003) – that have been used to explain the intrusive growth of cambial cells (Wloch, Mazur & Kojs, 2001; Wloch, Mazur & Beltowski, 2002; Kojs et al., 2004a; Kojs, Wloch & Rusin, 2004b; Jura et al., 2006). The key difference between this hypothesis and that outlined above regarding the predominance of radial compression of vascular cambium in the growing season lies in the timing and frequency of the presence of radial tensile stress within the vascular cambium. In this latter hypothesis it is assumed that there is a regular temporal pattern of radial tensile stress in the vascular cambium. In trees with developed foliage it is hypothesized that during the day radial compression of the vascular cambium would result from water loss due to ongoing transpiration. At night, or during the day in conditions that limit transpiration (e.g. cold, rain) tissues would be rehydrated and strong radial tensile stress would be...
created in the cambium (Kojs & Rusin, 2011; Kojs, 2012, 2013; Kojs et al., 2012) – xylem strain is considerably lower than phloem strain due to the higher content of cells with lignified and thick secondary walls. According to Kojs (2012), this strong radial tensile stress would allow the radial growth of cells. Therefore, radial growth might be viewed as an adaptation to alternating mechanical stress cycles (Kojs et al., 2012; Kojs, 2013). Furthermore, it may be speculated that prior to leaf development the cycle is reversed, i.e. tensile stress in the radial direction required for the growth of vessel elements is created during the day, and compressive stress at night, due to increased tissue hydration (i.e. changes in the osmotic potential of cells) during the day resulting from increased temperature and water availability (P. Kojs, in preparation). The experiments discussed above involving incisions of F. excelsior and R. pseudoacacia cambia (Hejnowicz, 1980) may be consistent with this hypothesis, where they were carried out during the day and before leaf development in ring-porous species: tangential incisions of cambia were made in early May 1977 in F. excelsior, and in 1978 (when earlier phases of cambial reactivation were studied) in F. excelsior and R. pseudoacacia (Hejnowicz, 1980). Bud break and foliage development in ring-porous species is often delayed in comparison to diffuse-porous ones (Friesner, 1942; Aloni & Peterson, 1997; Panchen et al., 2014). Radial tensile stress in the vascular cambium associated with the diurnal cycle of tissue swelling may provide an explanation for the growth of vessel elements in both ring- and diffuse-porous species during the entire growing season. Radial tension, together with shearing stress, would control the formation of microspaces between periclinal walls of cells in the cambium and at the border of the cambium and developing xylem (Kojs, 2012, 2013; see microspaces in Fig. 5). Pectins are the main constituent of the middle lamellae. Pectins show thixotropy, meaning that they can lower their viscosity in response to mechanical force (Woodcock, 1985; Barnes, 1997) in a reversible way (Barnes, 1997). Under shearing mechanical stress pectin gels undergo elastic deformation up to a threshold value, beyond which liquid-crystal domains are created that can flow in relation to other parts of the pectin gel (Kerst et al., 1990; Mujumdar, Beris & Metzner, 2002). Thus mechanical stress operating in/near the cambial tissue may lead to a reduction in viscosity of the middle lamellae, separation of parts of primary walls and thus to the formation of spaces. Mother cells of vessel elements may grow intrusively into these spaces. Kojs (2013) observed the formation of large microspaces at the border of the cambium and xylem between 3 a.m. and 5 a.m. Since sampling and anatomical investigations are not usually carried out at night, it may be that this phenomenon has been overlooked. As mentioned above, anatomical observations indicate that intrusive growth of vessel elements occurs near the border of these two tissues (Figs 4 and 5).

There is currently no consensus regarding the mechanical conditions under which cambial cells and their derivatives function and grow. Consideration of physiological aspects involving tissue swelling and shrinkage (Ueda & Shibata, 2001; Yoshida et al., 2003), as well as anatomical investigations concerning the precise course of intrusive growth of cambial initial cells (Wloch & Polak, 1994; Wloch et al., 2001, 2002, 2009, 2013; Kojs et al., 2004a,b; Jura et al., 2006; Karcewskas et al., 2009; Wilczek et al., 2018; Miodek et al., 2021), have raised serious objections against earlier assumptions concerning mechanical stress in the vascular cambium.

V. SHAPE OF ENLARGING VESSEL ELEMENTS

Before attaining their final transverse dimensions, broad vessel elements observed in cross section are often flattened in a radial direction (Figs 4 and 5). Such vessel elements, as revealed by polarization microscopy, have a primary cell wall (or at least parts of this wall) not thickened by secondary cell wall layers (Figs 4 and 6). Initially, this flattened shape was explained as resulting from differential growth rates of vessel elements in different directions. From their examination of Quercus rubra tissues Zasada & Zahner (1969) stated that vessel elements first rapidly increase their tangential dimension, followed by a gradual increase in radial dimension. Hejnowicz (2012) presented a scheme showing successive stages of vessel element growth in which tangential and radial diameters become equivalent only in the final stage of vessel element development. According to Hejnowicz (2012), the occurrence of tensile stress in the radial direction would allow separation of periclinal walls of cells in radial files adjacent to a vessel element.

We suggest that in the presence of such strong radial tensile stress in the vascular cambium, the frequently observed initially flattened shape of vessel elements may not necessarily be a result of differential growth rates in the radial and tangential directions. Assuming that growth of a vessel element starts into a microspace created after radial tensile stress-induced separation of neighbouring periclinal cell walls, and that cells grow when tissue hydration is high during the diurnal cycle (Kojs & Rusin, 2011; Kojs, 2012, 2013), it may be the case that sampling procedures affect the mechanical conditions within the tissue leading to artefacts in vessel shape. The collected tissue will no longer be subjected to strong tensile stress, some cells may lose their turgidity, and the thin primary walls of developing vessel elements could easily collapse. In Figs 4 and 8 it can be observed that broad earlywood vessel elements assume a permanent rounded, but sometimes radially elongated, shape during their later development. Figure 8 shows a radial gradient of growth and differentiation, inwardly from the vascular cambium. It is clear that the developing vessel element has a semi-circular shape on the xylem side associated with asymmetrical secondary cell wall deposition, but an irregular shape on the cambial side (closer to the cambial initial surface) which is associated with the occurrence of a thin primary cell wall that may be prone to deformation. Additionally, in cases where the secondary walls of developing vessels are thin due to gene
modification (Turner & Somerville, 1997; Hao & Mohnen, 2014), or to metabolic (Damari-Weissler et al., 2009), or environmental (Aloni, 2021) stresses, collapsed and deformed vessels are formed. Note that diurnal cycles of changes in vivo tissue hydration will also include periods of tissue shrinkage due to loss of water (Simonneau et al., 1993; Kojs & Rusin, 2011). Thus, it can be speculated that there will be periods in a diurnal cycle when mechanical conditions are not suitable for growth of vessel elements. In these periods, vessel elements with fragments of non-lignified primary cell wall could temporarily assume a slightly flattened shape. If this flattening persists over time it may result in the formation of a peculiarly shaped (deformed) mature vessel (A. Miodek, unpublished data). Such tissue shrinkage due to loss of water (Simonneau et al., 1993; Kojs & Rusin, 2011) may also explain the formation of deformed/collapsed vessel elements with thin secondary cell walls. Burggraaf (1973) reported that the sampling method is critical for the observed shape of vessel elements, noting that while in earlier analyses (Burggraaf, 1972) the observed shape of vessel elements was as described by Zasada & Zahner (1969), a different sampling technique produced only circular-shaped developing vessel elements (Burggraaf, 1973). The sampling technique proposed by Burggraaf (1973) included careful tissue collection and preparation of thick sections of developing earlywood. Thus, the techniques used for sample collection and preparation may be of great importance in the context of observations of vessel elements at early stages of their growth.

An alternative model of vessel element development was presented by Wilczek et al. (2011b), who suggested that: (i) vessel elements grow intrusively between periclinal cell walls; (ii) the whole tissue undergoes symplastic growth in a radial direction; and (iii) at each stage of growth, tangential and radial diameters of the growing vessel element are similar. In their model, growth of a vessel element takes place in periods when the whole cambial tissue is under radial tensile stress. If a relatively large space is created between periclinal walls as a result of strong radial tensile stress, a vessel element mother cell may start to grow to fill this space.

VI. INTRUSIVE GROWTH OF VESSEL ELEMENTS AND CAMBIAL INITIALS

Enlarging vessel elements grow intrusively in close vicinity to cambial initials. Therefore, it is of interest to consider whether cambial initials and enlarging vessel elements are subject to similar mechanical conditions and have a common mechanism of intrusive growth. There is inconsistent information available regarding the location of intrusive growth of cambial initials. Some sources state that intrusive
growth of cambial initials is localized between the radial cell walls (e.g. Wloch, 1976; Evert, 2006), or imply this location by arguing that an increase in circumference of non-storied cambium occurs through the intrusive growth of initials after oblique (pseudotransverse) anticlinal divisions (Butterfield, 1973; Zagórska-Marek, 1981; Romberger et al., 1993; Evert, 2006; Hejnowicz, 2012). In order to increase the circumference of the cambial cylinder, microspaces into which cambial initial cells can grow intrusively must be created between the radial walls of cells. Thus, the cells between which an initial is growing become separated and move apart in a tangential direction (Fig. 9A). Following this concept, it was stated that cambial cells are stretched in the tangential direction (Hejnowicz, 1980, 1997, 2012; Romberger et al., 1993; Kwiatkowska & Nakielni, 2011). Contrary to this pattern, intrusive growth of vessel elements is thought to proceed in a different way, and under different mechanical conditions. Intrusive growth of vessel elements occurs between periclinal walls of cells as a result of radial tensile stress (Hejnowicz, 2012), while intrusive growth of initials occurs between radial walls of cells under conditions of tensile stress in the tangential direction and compressive stress in the radial direction (Hejnowicz, 1997). This disparity is confusing, since there seems to be no clear reason for these two types of cells to grow by different mechanisms.

By contrast, recent reports show that intrusive growth of cambial initials occurs between the periclinal walls of initial cells and their closest derivatives (Kojs et al., 2004a,b; Karcewska et al., 2009; Wloch et al., 2013; see Fig. 9B), thus in the same way as intrusive growth of enlarging vessel elements (Wilczek et al., 2011b; Gizińska et al., 2021). The location of intrusive growth of cambial initials between periclinal cell walls is consistent with evidence that intrusive growth does not affect the circumference of the cambial cylinder (Jura et al., 2006; Wloch et al., 2009, 2013; Wilczek et al., 2011a; Miolek et al., 2021; see also Fig. 5). It should be noted that growth of vessel elements is often associated with substantial changes in the arrangement of adjacent cells (Evert, 2006). As for cambial initials (Jura et al., 2006), the location of intrusive growth of vessel elements between periclinal walls of adjacent cells of the axial system should not increase the tangential dimension of surrounding tissue. This is logical as intrusive growth of a vessel element in the tangential direction is related to disruption of continuity of adjacent radial files (Wilczek et al., 2011b; Hejnowicz, 2012; Gizińska et al., 2021), or shortening of the periclinal walls of cells of the axial system (Gizińska et al., 2021). Interestingly, a tangential increase in diameter of a vessel element can take place not only intrusively, but also symplastically (Gizińska et al., 2021). This refers to specific situations, when a growing vessel element is adjacent to a ray/rays. Partial/complete separation of periclinal walls can be observed in radial files of axial system cells that are separated from a vessel element by a ray/rays (Gizińska et al., 2021). It can be deduced that growing vessel elements do not contribute to an increase in circumference of the developing secondary xylem.

While considering this recent evidence for the location of intrusive growth of initials between periclinal cell walls, the issue of cell elimination is pertinent. Cell elimination in the vascular cambium is now thought to occur as a consequence of intrusive growth of an initial cell between an initial depositing a radial file which will be eliminated, and its closest derivative (Kojs et al., 2004a; Jura et al., 2006; Wloch

![Fig 9. Schemes of transverse sections showing the types of stress occurring in the vascular cambium, according to two hypotheses of intrusive growth of cambial initials. These mechanical stresses may either allow the initial cell to grow intrusively between anticlinal cell walls of neighbouring initials (A) or between periclinal walls of neighbouring initials and their closest derivatives (B). (A) An initial cell located above/below the visible transverse plane would grow intrusively in an axial direction into a microspace created by separation of the anticlinal walls (marked as a thick black line). (B) An initial cell located above/below the visible transverse plane would grow intrusively in an axial direction into a microspace created by separation of the periclinal walls (marked as a thick black line). The direction in which separation occurs is indicated by red arrows. The type of stress (compressive versus tensile) is indicated by large, grey arrows. Cambial initials are shaded green; mother cells are blue.](image-url)
et al., 2009; Hejnowicz, 2012); elimination occurs as a result of intrusive cell growth between periclinal walls. During elimination, a given initial cell loses its initial status, is removed from the initial surface and differentiates on the xylem/phloem side (Wloch & Polap, 1994; Jura et al., 2006; Wloch et al., 2013; Wilczek et al., 2014a). It is speculated that cambial initials do not actively intrude between walls of neighbouring cells (Hejnowicz, 1980, 1997), but instead grow into microspaces created between cells (Kojs, 2013; Wloch et al., 2013). Therefore, elimination of cells due to intrusive growth of other cells would require the occurrence of radial tensile stress, as is the case for growing vessel elements. Thus, according to recent reports, growth of both vessel elements and cambial initials as well as cell elimination in the vascular cambium are all associated with intrusive growth between periclinal walls (Jura et al., 2006; Karczewska et al., 2009; Wilczek et al., 2014a,b; Wloch et al., 2013; Gizińska et al., 2021; Miodek et al., 2021). Therefore, it seems that intrusive growth along periclinal cell walls is key to our understanding of events occurring in the cambium and its surroundings. Moreover, the mechanical conditions in which these processes take place are also similar—radial tensile stress.

The location of intrusive growth between periclinal cell walls has fundamental consequences in the context of cambial cell rearrangement in proximity to growing vessel elements. For example, intrusive growth/elimination within vesselless radial files of cells, occurring in the neighbourhood of a growing latewood vessel element may be observed in ring-porous species (Fig. 5). It would be difficult to explain such phenomena if the vascular cambium is under constant radial compression (except for during early spring) (Hejnowicz, 1997; Kwiatkowska & Nakielski, 2011). As mentioned earlier, under the assumption that the cambium is under radial compression, it was speculated that a vessel element can grow under local radial tensile stress generated by adjacent, currently vesselless radial files (Hejnowicz, 2012). Such radial files would grow most rapidly, and thus take over the compression force exerted by the bark (Hejnowicz, 2012). However, intrusive growth/cell elimination from the initial surface can be observed in such radial rows—observations that can be easily explained if we assume that cambium is radially stretched (Kojs, 2012; Kojs et al., 2012), which seems to be true for ring-porous trees in the spring (Hejnowicz, 1980). Radial stretching of the cambium may lead to the formation of microspaces (Kojs, 2012, 2013; Wloch et al., 2013) into which initials can subsequently grow intrusively. This would result in simultaneous elimination of initials from the initial surface (Jura et al., 2006), since intrusive growth of one cell and elimination of the neighbouring cell are inherently coupled and should be considered as simultaneous outcomes of cambial cell rearrangement. The cell events shown in Fig. 5 (growth of a vessel element, rearrangement of cambial cells) thus seem to be related to tensional rather than compressive radial stress.

In summary, there remains a need for a unified theoretical framework, providing a comprehensive explanation of the various processes occurring in the vascular cambium and zone of xylem/phloem cell development. There are currently different concepts on how a given cell type (e.g. wood fibre, initial cell, vessel element) grows, with some conflicting explanations partly related to the different assumptions concerning the mechanical conditions in which these cells grow.

VII. INTRUSIVE GROWTH OF VESSEL ELEMENTS AND WOOD FIBRES

Fibres, especially those with thick secondary cell walls, are responsible for the mechanical strengthening of xylem tissue. As a result of axial intrusive growth at their tips, fibres are much longer than the fusiform initials from which they originate (Evett, 2006; Jura-Morawiec et al., 2008; Hejnowicz, 2012). A recently published study concerning intrusive growth of the tips of wood fibres indicates that this takes place between the periclinal walls of adjacent cells within one or two radial rows of cells (Wilczek et al., 2018), in agreement with previous observations (Wenham & Cusick, 1975). Thus intrusive growth of fibres takes place between the periclinal walls as is the case for vessel elements (Wilczek et al., 2011b; Hejnowicz, 2012; Gizińska et al., 2021) and most likely for cambial initials (Jura et al., 2006; Karczewska et al., 2009; Wloch et al., 2013; Miodek et al., 2021).

Wilczek et al. (2018) also noted that wood fibres occurring near vessel elements showed limited intrusive growth, thereby confirming previous observations on the ring-porous species Robinia pseudoacacia L. (Fujita, Tohyama & Harada, 1984, 1986) and the diffuse-porous species Aesculus hippocastanum Willd. (Honjo, Ogata & Fujita, 2006; Yahya et al., 2015). Kojs (2013) suggested that the growth of wood fibres requires both radial tensile stress and shearing stresses. Radial tensile stress within the vascular cambium seems to be a necessary pre-condition for the growth of vessel elements to occur (Hejnowicz, 1980, 2012). Kojs (2013) assumed that the formation of large spaces by separation of the periclinal cell walls, which are immediately filled by growing vessel elements, causes local relaxation of the radial tensile stress accumulated in the surrounding tissues. This implies that the degree of relaxation of radial tensile stress might translate into the range of intrusive growth of fibres, and therefore to decreased fibre length. Wilczek et al. (2018) thus explained their observations by acknowledging the role of mechanical stress. In the present context it is noteworthy that in some species wood fibres do not occur, or occur infrequently, in the direct vicinity of vessel elements, since the latter are surrounded by parenchyma cells (including rays) or groups of tracheids and parenchyma cells (Metalfe & Chalk, 1965). It may be the case that a specific type of cell into which a cambial derivative differentiates as well as its final shape and size are dependent upon the specific mechanical conditions occurring at the moment of its formation.
VIII. FORMATION OF VESSEL ELEMENTS FROM OVERWINTERING CELLS

The existence of overwintering mother cells of vessel elements has been suggested by several studies (e.g. Barnett, 1992; Kitin et al., 1999; Frankenstein, Eckstein & Schmitt, 2005; Fonti, Solomonoff & Garcia-González, 2007). Kitin et al. (2003) stated that it is not possible to predict from which cambial cell a vessel element will differentiate. They also highlighted that it is not known whether positions of vessel cell elements are random, or whether their location is determined by a specific mechanism. However, when radial tensile stress is present, as for example in ring-porous trees in the spring (Hejnowicz, 1980), it can be proposed that mechanical factors may play an important role in the formation of vessel elements. In this scenario radial tensile stress, together with shearing stresses, could lead to the formation of axial/tangential micro-cracks at the border of the cambium and zone of xylem cell development (Kojs, 2013). Shearing stresses lead to increased fluidity of the middle lamellae (Leroux, 2012) between periclinal walls of cambial cells, facilitating their mechanical separation by radial tensile stress, and thereby enabling intrusive growth of vessel elements between the periclinal walls of adjacent cells (Wilczek et al., 2011b; Hejnowicz, 2012; Gizińska et al., 2021). Nonetheless, even if mechanical stress does cause separation of cell walls (allowing growth of selected cells), the issue of whether the future vessel element to be formed is randomly determined is still under debate.

IX. COURSE OF VESSELS

It is widely accepted that the arrangement of the majority of xylem cells reflects the arrangement of initials in the vascular cambium at the moment of their deposition. This assumption is often made in indirect studies of cambial tissue involving analyses of xylem sections, especially for conifers which are characterized by a relatively simple wood structure with smaller variation in cell types compared to angiosperm trees (Hejnowicz & Zagórska-Marek, 1974; Wloch, 1976; Krawczyszyn, 1977; Hejnowicz & Romberger, 1979; Wilczek, Miodek & Gizińska, 2014b). The relationship between the arrangement of cambial initials and vessel elements seems to be more complex, since vessel elements show intensive growth, both symplastic and intrusive, during their development (Gizińska et al., 2021). Moreover, vessels are interconnected to form a network, in which deflections of vessels are often observed in both a tangential and, to some extent, in a radial direction (Burggraaf, 1972; Kitin et al., 2004). Among factors that may affect the course of vessels, the relationship between growing vessel elements and rays is likely to be important. Vessel elements do not grow intrusively between periclinal walls of ray cells (Gizińska et al., 2021), and nearby rays may locally modify expansion of vessel elements (Wilczek et al., 2011b). Adjacent rays may contribute to the anisotropic growth of vessel elements and thus to local deflections of vessels in a tangential direction (Wilczek et al., 2011b; see also Fig. 10A). However, the arrangement of rays on its own is insufficient to explain all tangential deflections of a vessel. As shown in Fig. 10B, C, slight deflections in the vessel course can be observed in the

Fig 10. Examples of local deflections of vessels seen in tangential sections. The course of vessels is indicated by dashed lines. (A) Juvenile wood of Acer pseudoplatanus L. White arrowheads indicate areas where the vessel deviates slightly in the direct vicinity of a ray. (B, C) Juvenile wood of Robinia pseudoacacia L. Black arrowheads indicate areas where vessels change their course slightly in the absence of rays in their direct neighbourhood. In C, note the presence of a second vessel (indicated by grey arrowhead) close to the vessel identified by the dashed line.
absence of rays. In Fig. 10C a change in course of the vessel occurred in the vicinity of a second vessel (visible in the lower part of the micrograph). Interestingly, Kitin et al. (2004) pointed out that radial deflections indicate that both initiation and completion of differentiation of vessel element mother cells might not occur simultaneously. It is plausible that mechanical stress which controls the localization of PIN proteins (Heisler et al., 2010; Li et al., 2012; Nakayama et al., 2012; Li et al., 2019) may be a key factor determining the course of forming vessels. We suggest that future studies of plants should consider the mechanical stresses operating in their developing tissues.

X. CONCLUSIONS

(1) From an anatomical perspective, vessel elements grow intrusively and symplastically. Intrusive growth occurs between the periclinal walls of axial system cells, in a similar way for both wood fibres and cambial initials. It is postulated that intrusive growth of a developing vessel element between the periclinal walls of axial system cells requires the presence of radial tensile stress enabling the separation of the periclinal walls of neighbouring cells.

(2) The mechanical conditions in which vessel elements grow and differentiate have not been fully elucidated. Incision experiments indicated the presence of radial tensile stress in the vascular cambium of ring-porous species in the spring, supporting the hypothesis that radial tensile stress is present in the vascular cambium of angiosperm trees.

(3) Mechanical factors seem to be vital for developmental processes occurring in the vascular cambium.

(4) Clarifying the relationship between mechanical stress and auxin in the vascular cambium is particularly important for a better understanding of developmental processes in plants. Mechanical stress is a strong candidate for a key player involved not only in vessel element growth, but also in determining the course of vessels.

(5) Future research should make use of methods allowing the visualization and quantification of time course of mechanical forces in three dimensions and most importantly associate them with developmental processes occurring within plants.

XI. REFERENCES

Ackermann, F. & Stanislas, T. (2020). The plasma membrane—an integrating compartment for mechanism-signalizing. Plant 9(4), 905.

Aloni, R. (2007). Phytohormonal mechanisms that control wood quality formation in young and mature trees. In The Compendium Wood Workhop 2007 (eds K. Enniste, P. Hanner & J. Walker), pp. 1–22. The Wood Technology Research Centre, University of Canterbury, Christchurch.

Aloni, R. (2013). The role of hormones in controlling vascular differentiation. In Cellulare Aspects of Wood Formation. Plant Cell Monographs 20 (ed. J. Fromm), pp. 99–139. Springer, Berlin, Heidelberg.

Aloni, R. (2015). Ecophysiological implications of vascular differentiation and plant evolution. Trees 29, 1–16.

Aloni, R. (2021). Vascular Differentiation and Plant Hormones. Springer Nature, Cham.

Aloni, R. & Peterson, C. A. (1997). Auxin promotes dormancy callose removal from the phloem of Magnolia kobus and callose accumulation and earlywood vessel differentiation in Quercus robur. Journal of Plant Research 110, 37–44.

Aloni, R. & Zimmerman, M. H. (1983). The control of vessel size and density along the plant axis. A new hypothesis. Differentiation 24, 203–208.

Barnes, H. A. (1997). Thixotropy—a review. Journal of Non-Newtonian Fluid Mechanics 70, 1–33.

Beert, J. R. (1992). Reactivation of the cambium in Aesculus hippocastanum L.: a transmission electron microscope study. Annals of Botany 70, 169–177.

Bauer, T. & Escherich, W. (1997). Mechanical pressure inhibits vessel development of xylegic cambial derivatives of beech (Fagus sylvatica L.). Trees 11, 349–355.

Beek, C. B. (2010). An Introduction to Plant Structure and Development. Cambridge University Press, Cambridge.

Bhalerao, R. P. & Fischer, U. (2017). Environmental and hormonal control of cambial stem cell dynamics. Journal of Experimental Botany 68, 79–87.

Brown, C. L. & Sax, K. (1962). The influence of pressure on the differentiation of secondary tissues. American Journal of Botany 49, 683–691.

Burggraaf, P. D. (1972). Some observations on the course of the vessels in the wood of Fagus excelsior L. Acta Botanica Neerlandica 21, 32–47.

Burggraaf, P. D. (1973). On the shape of developing vessel elements in Fagus excelsior L. Acta Botanica Neerlandica 22, 271–278.

Butterfield, B. G. (1973). Variation in the size of xylem cambial initials and vessel members in Heleola angulifolia Roaw. New Zealand Journal of Botany 11, 391–410.

Crang, R., Lyons-Sobrski, S. & Wise, R. (2018). Plant Anatomy: A Concept-Based Approach to the Structure of Seed Plants. Springer Nature, Cham.

Damari-Weissler, H., Rachamilewitch, S., Aloni, R., German, M. A., Cohen, S., Zwieniecki, M. A., Holbrook, N. M. & Granov, D. (2000). LeFK2 is required for phloem and xylem differentiation and the transport of both sugar and water. Planta 230, 795–805.

Deng, F. & Jiao, Y. (2020). Mechanical control of plant morphogenesis: concepts and progress. Current Opinion in Plant Biology 57, 16–23.

Evert, R. F. (2006). Evan’s Plant Anatomy: Mesophyll, Cells, and Tissues of the Plant Body: Their Structure, Function, and Development. John Wiley and Sons, Hoboken.

Fonti, P., Solomonoff, N. & García-González, I. (2007). Earlywood vessels of Castanea sativa record temperature before their formation. New Phytologist 173, 562–570.

Frankenstein, C., Eckstein, D. & Schmitz, U. (2005). The onset of cambium activity—a matter of agreement? Dendrochronologia 23, 57–62.

Frederik, R. C. (1942). Dendrometer studies on five species of broadleaf trees in Indiana. Butler University Botanical Studies 5, 160–172.

Freml, J. (2008). Auxin transport—shaping the plant. Current Opinion in Plant Biology 11, 74–79.

Fujita, M., Tohyma, M. & Harada, H. (1984). Three-dimensional analysis of the cambium and the derivatives in Robinia pseudoacacia L. In Proceeding of Pacific Regional Wood Anatomy Conference (ed. S. Scio), pp. 91–93. Conference Secretariat, Tsukuba.

Fujita, M., Tohyma, M. & Harada, H. (1986). An approach to three-dimensional analysis of cambial cells and their derivatives in Robinia pseudoacacia L. Bulletin of the Kyoto University Forests 57, 283–289.

Gizynska, A. Mioder, A. & Koen, P. (2021). Rays hamper intrusive growth of vessel elements. Trees 35, 749–760.

Gizynska, A. Mioder, A., Wilezczek, A., Wloch, W. & Iqbal, M. (2015). Wood porosity as an adaptation to environmental conditions. Nature Journal (Fragmenta Naturalis) 48, 54–52.

Grunewald, P. D., Burggraaf, P. D. & Henningsen, K. (2010). The march of the PINs: developmental plasticity by dynamic polar targeting in plant cells. The EMBO Journal 29, 2700–2714.

Hamant, O., Heisler, M. G., Jonsson, H., Krupinski, P., Uttewaal, M., Borov, P., Corson, F., Sahlin, P., Boumdouaou, A., Meyerowitz, E. M., Coulter, Y. & Traas, J. (2008). Developmental patterning by mechanical signals in Arabidopsis. Science 322, 1650–1655.

Hamant, O. & Hassell, E. S. (2017). Life behind the wall: sensing mechanical cues in plants. BMC Biology 15, 59.

Hamant, O. & Traas, J. (2010). The mechanics behind plant development. New Phytologist 185, 369–385.

Hao, Z. & Mohnen, D. (2014). A review of xylan and lignin biosynthesis: foundation for studying Arabidopsis irregular xylan mutants with pleiotropic phenotypes. Critical Reviews in Biochemistry and Molecular Biology 49, 212–241.

Heisler, M. G., Hamant, O., Krupinski, P., Uttewaal, M., Ohno, C., Jonsson, H., Traas, J. & Meyerowitz, E. M. (2010). Alignment between PIN1 polarity and microtube orientation in the shoot apical meristem reveals a tight coupling between morphogenesis and auxin transport. PLoS Biology 8(10), e1000516.
How do vessel elements grow?

Hejnowicz, Z. (1980). Tensional stress in the cambium and its developmental significance. *American Journal of Botany* 67, 1–5.

Hejnowicz, Z. (1997). Graviresponses in herbs and trees: a major role for the redistribution of tissue and growth stresses. *Planta* 203, S136–S146.

Hejnowicz, Z. (2012). *Anatomia i histologia roślin zasypiskowych*. Oxylogy, Poznań, Warszawa.

Hejnowicz, Z. & Romberger, J. A. (1979). The common basis of wood grain figures is the systematically changing orientation of cambial fusiform cells. *Wood Science and Technology* 13, 304–332.

Hejnowicz, Z. & Zagórska-Marer, B. (1974). Mechanism of changes in grain inclination in wood produced by storied cambium. *Acta Societatis Botanicorum Poloniae* 43, 381–398.

Honjo, K., Ogata, Y. & Fujita, M. (2006). Introduction and verification of a novel method for measuring wood fiber length using a single cross section in *Artemisia mongolica*. *Tuu* 20, 356–362.

Huang, F., Zago, M. K., Iqbal, M., Van der Putten, H. W., Chalk, L. & Meyerowitz, E. M. (2019). Calcium signals are necessary to establish auxin transporter polarity in a plant stem cell niche. *Nature Communications* 10, 726.

Lintulnic, P. M. & Vesecky, T. B. (1984). Stress-induced alignment of division plane in plant tissues grown in vitro. *Nature* 307, 363–364.

Ling, T. M. & Lintulnic, P. M. (1997). Mechanical signals in plant development: a new method for single-cell studies. *Developmental Biology* 181, 246–256.

Metcalf, C. R. & Chalk, L. (1965). Anatomy of the Deciduous. Leaves, Stem, and Wood in Relation to Taxonomy with Notes on Economic Uses. Claradon Press, Oxford.

Miodek, A., Gzińska, A., Klisz, M., Wojda, T., Ukalski, A. & Kojs, P. (2020). Direct exposure to solar radiation causes radial growth eccentricity at the beginning of the growing season in *Robinia pseudoacacia*. *IAWA Journal* 41, 61–84.

Miodek, A., Wóch, I., Iqbal, M., Gzińska, A. & Kojs, P. (2021). Controversy over the mode of growth of cambial cylinder. *The Botanical Review* 87, 243–257.

Mojumdar, A., Berin, A. N. & Metzner, A. (2002). Transient phenomena in plant cells. *Molecular Microbiology* 45, 157–178.

Nick, P. (2013). Mectotubules, signalling and abiotic stress. *The Plant Journal* 75, 309–323.

Nakayama, N., Smith, R. S., Mandel, T., Robinson, S., Kimura, S., Boudaoud, A. & Kühlemeier, C. (2012). Mechanical regulation of auxin-mediated cell elongation of tracheary elements. *Current Biology* 22, 1464–1478.

Nilsen, J., Karlberg, A., Antti, H., Lopez-Vernaza, M., Mellerowicz, E., Perkov-Reichenmann, C., Sandberg, G. & Bhalerao, R. P. (2008). Dissecting the molecular basis of the regulation of wood formation by auxin in hybrid aspen. *The Plant Cell* 20, 843–853.

Panacha, Z., A., Prems, R. B., Nordin, B., Ellwood, E. R., Steven, A.-D., Renner, S. S., Willis, C. G., Fahey, R., Whittemore, A., Du, Y. & Davis, C. C. (2014). Leaf out times of temperate woody plants are related to phylogeny, deciduousness, growth habit and wood anatomy. *New Phytologist* 203, 1209–1219.

Petersson, S. V., Johansson, A. I., Kowalczyk, M., Makovecuk, A., Wang, Y. J., Moritz, T., Grebe, M., Benfey, P. N., Sandberg, G. & Ljung, K. (2009). An auxin gradient and maximum in the *Arabidopsis* root apex shown by high-resolution cell-specific analysis of IAA distribution and synthesis. *Current Biology* 19, 230–235.

Ramos, J. R. D., Maizel, A. & Alim, K. (2021). Tissue-wide integration of mechanical cues promotes effective auxin patterning. *The European Physical Journal Plus* 136, 250.

Raven, J. A. (1975). Transport of indoleacetic acid in plant cells in relation to pH and electrical potential gradients, and its significance for polar IAA transport. *New Phytologist* 74, 163–172.

Rayle, D. L. & Cleland, R. E. (1992). The acid growth theory of auxin-induced cell elongation is alive and well. *Plant Physiology* 99, 1271–1274.

Robert, S., Kleine-Vehn, J., Barbez, E., Sauer, M., Pacheco, T., Baster, P., Vanneyte, S., Zhang, J., Simon, S., Čovanová, M., Haváky, K., Dujardin, H., Pacy, Z., Brand, A. Y., Jones, A. M. A., et al. (2010). ABP1 mediates auxin inhibition of clathrin-dependent endocytosis in a plant stem cell niche. *Current Biology* 20, 121–127.

Robert, P. H. & Schiefelbein, J. A. (1994). Carrier-mediated auxin transport. *Planta* 191, 189–201.

Rutschoff, H. L., Baskin, T. I. & Kramer, E. M. (2014). The carrier AUXIN RESISTANT (AUX1) dominates auxin flux into *Arabidopsis* protoplasts. *New Phytologist* 204, 356–354.

Sachs, T. (1991). Cell polarity and tissue patterning in plants. *Development Supplement* 1, 83–93.

Sampaikumar, A. (2020). Mechanical feedback-loop regulation of morphogenesis in plants. *Development* 147, dev177964.

Santos, F., Teale, W., Fleck, C., Volpers, M., Bertoft, B. & Palme, K. (2010). Modelling polar auxin transport in developmental patterning. *Plant Biology* 12, 1–14.

Schrader, J., Baba, K., May S. T., Palme, K., Bennett, M., Bhalerao, R. P. & Sandberg, G. (2003). Polar auxin transport in the wood-forming tissues of hybrid aspen is under simultaneous control of developmental and environmental signals. *Proceedings of the National Academy of Sciences of the United States of America* 100, 401–406.

Simonneau, T., Harib, R., Goutouly, J.-P. & Huguett, J.-G. (1993). Diurnal changes in stem diameter depend upon variations in water content: direct evidence in peach trees. *Journal of Experimental Botany* 44, 615–621.

Li, H., Frimi, J. & Grunwald, W. (2012). Cell polarity: restricting prevents developmental cramps. *Current Biology* 22, R635–R637.

Li, T., Yan, A., Bhatia, N., Atein, A., Artik, E., Durand-Senet, P., Tare, P. T., Schroeder, J. I., Heidler, M. G. & Meyerowitz, E. M. (2019). Calcium signals are necessary to establish auxin transporter polarity in a plant stem cell niche. *Nature Communications* 10, 726.
Sinnott, E. W. & Bloch, R. (1939). Changes in intercellular relationships during the growth and differentiation of living plant tissues. *American Journal of Botany* **26**, 625–634.

Srivastava, L. M. (1973). Cambial activity in trees. *Arborea* **33**, 46–66.

Sundberg, B., Little, C. H. A. & Cel, K. (1990). Distribution of indole-3-acetic acid and the occurrence of its alkali-labile conjugates in the extraxylary region of *Pinus sylvestris* stems. *Plant Physiology* **93**, 1295–1302.

Trehin, D.-C., Alonso-Serret, J., Asaoka, M., Colin, L., Cortés, M., Malwé, A., Takatsui, S., Zhao, F., Traas, J., Trehin, G. & Hamant, O. (2021). How mechanical forces shape plant organs. *Current Biology* **31**, R143–R159.

Tulik, M. (2007). Wzrost intruzyny komórek u roślin drzewiastych. *Syflos* **151**, 3–13.

Turner, S. R. & Somerville, C. R. (1997). Collapsed xylem phenotype of Arabidopsis identity mutants deficient in cellulose deposition in the secondary cell wall. *The Plant Cell* **9**, 689–701.

Ueda, M. & Shibata, E. (2001). Diurnal changes in branch diameter as indicator of water status of Hinoki cypress Chamaecyparis obtusa. *Trees* **15**, 315–318.

Ugglä, C., Millerowicz, E. J. & Sundberg, B. (1998). Indole-3-acetic acid controls cambial growth in Scots pine by positional signaling. *Plant Physiology* **117**, 113–121.

Ugglä, C., Moritz, T., Sandberg, G. & Sundberg, B. (1996). Auxin as a positional signal in pattern formation in plants. *Proceedings of the National Academy of Sciences of the United States of America* **93**, 9282–9286.

Vanneste, S. & Friml, J. (2009). Auxin: a trigger for change in plant development. *Cell* **136**, 1005–1016.

Wenham, M. W. & Cusick, F. (1975). The growth of secondary wood fibres. *New Phytologist* **74**, 247–261.

Wilczek, A., Giżńska, A., Miodek, A. & Włoch, W. (2014a). Nowa hipoteza wzrostu promieniowego i przebudowy kambialnego woskowego rejonu drzewiastych. *Kosmos* **63**, 591–601.

Wilczek, A. B., Iqbal, M., Włoch, W. & Klisz, M. (2018). Geometric analysis of intrusive growth of wood fibres in *Robinia pseudoacacia*. *IWA Journal* **39**, 194–208.

Wilczek, A., Jura-Morawiec, J., Iqbal, M. & Włoch, W. (2011b). Correlation of intrusive growth of cambial initials to rearrangement of rays in the vascular cambium. *IWA Journal* **32**, 313–331.

Wilczek, A., Miodek, A. & Giżńska, A. (2014b). Terminal xylem and initial parenchyma in anatomical investigations – a review of definitions. *Nature Journal (Fragmenta Naturae)* **47**, 31–44.

Wilczek, A., Włoch, W., Iqbal, M. & Koś, P. (2011b). Position of rays and lateral deviation of vessel elements in the stem wood of some dicotyledonous species with storeyed, double-storeyed, and nonstoreyed cambium. *Botany* **89**, 849–860.

Wilson, B. F., Wodzicki, T. J. & Zahniser, R. (1966). Differentiation of cambial derivatives: proposed terminology. *Forest Science* **12**, 438–440.

Włoch, W. (1976). Cell events in cambium, connected with the formation and existence of a whirled cell arrangement. *Acta Societatis Botanicae Poloniae* **45**, 313–326.

Włoch, W., Jura-Morawiec, J., Koś, P., Iqbal, M. & Kruczkowski, J. (2009). Does intrusive growth of fusiform initials really contribute to circumferential growth of vascular cambium? *Botany* **87**, 154–163.

Włoch, W., Mazur, E. & Bieńkowska, M. (2002). Formation of spiral grain in the wood of *Pinus sylvestris* L. *Trees* **16**, 306–312.

Włoch, W., Mazur, E. & Koś, P. (2001). Intensive change of inclination of cambial initials in *Picea abies* (L.) Karst. tumours. *Trees* **15**, 498–502.

Włoch, W. & Polaś, E. (1994). The intrusive growth of initial cells in re-arrangement of cells in cambium of *Tilia cordata* MILL. *Acta Societatis Botanicae Poloniae* **63**, 109–116.

Włoch, W., Wilczek, A., Jura-Morawiec, J., Koś, P. & Iqbal, M. (2013). Modelling for rearrangement of fusiform initials during radial growth of the vascular cambium in *Pinus sylvestris* L. *Trees* **27**, 879–893.

Woodcock, L. V. (1985). Origins of thixotropy. *Physical Review Letters* **54**, 1513–1516.

Yahiya, R., Sundararoyo, A., Inai, T. & Sugiyama, J. (2013). Distance from vessels changes fiber morphology in *Araucaria angustifolia*. *IWA Journal* **36**, 35–43.

Yoshida, M., Ikawa, M., Kaneda, K. & Okiyama, T. (2003). Stem tangential strain on the tension wood side of *Fagus crenata* saplings. *Journal of Wood Science* **49**, 475–483.

Zagórska-Marék, B. (1981). Ontogeneza kambium. *Wisdomoci Botaniczne* **25**, 89–110.

Zarzecki, J. (1983). Hormonal control of cambial activity and vessel differentiation in *Quercus robur*. *Physiologia Plantarum* **57**, 537–542.

Zarzecki, J. (1991). Effect of indole-3-acetic acid (IAA) and sucrose on vessel size and density in isolated stem segments of oak (*Quercus robur*). *Physiologia Plantarum* **81**, 234–238.

Zasada, J. C. & Zahniser, R. (1969). Vessel element development in the earlywood of red oak (*Quercus rubra*). *Canadian Journal of Botany* **47**, 1965–1971.

Zhang, J., Nodzyńska, T., Pęcik, A., Rolčík, J. & Friml, J. (2010). PIN phosphorylation is sufficient to mediate PIN polarity and direct auxin transport. *Proceedings of the National Academy of Sciences of the United States of America* **107**, 918–922.

Zhou, J., Wang, B., Li, Y., Wang, Y. & Zhu, L. (2007). Responses of chrysanthemum cells to mechanical stimulation require intact microtubules and plasma membrane-cell wall adhesion. *Journal of Plant Growth Regulation* **26**, 55–68.

Zhou, J., Wang, B., Zhu, L., Li, Y. & Wang, Y. (2006). A system for studying the effect of mechanical stress on the elongation behavior of immobilized plant cells. *Colloids and Surfaces B: Biointerfaces* **49**, 163–174.

Zimmermann, M. H. (1982). Functional xylem anatomy of angiosperm trees. In *New Perspectives in Wood Anatomy* (ed. P. Baas), pp. 59–70. Martinus Nijhoff/Dr W. Junk Publishers, The Hague.

Zimmermann, M. H. (1983). *Xylem Structure and the Avent of Sap*. Springer-Verlag, Berlin.

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