Do Growth-Limiting Temperatures at the High-Elevation Treeline Require an Adaptation of Phloem Formation and Anatomy?

Dennis Marko Schröter and Walter Oberhuber*

Department of Botany, Leopold-Franzens-University of Innsbruck, Innsbruck, Austria

Low temperatures during the growing season restrict the growth of trees at high elevations and lead to the formation of the high-elevation treeline. To ensure the survival and growth of trees in such extreme locations, sufficient vascular transport capacity – enabled by vascular anatomical characteristics – is required. However, in contrast to the xylem, only little is known about the effects of low temperatures on the anatomy and formation of the phloem as important nutrient- and signal-conducting tissue. In this review, known findings of cold-induced changes in the anatomical and phenological properties of vascular tissues are used as starting points to discuss how low temperatures might affect phloem formation at the treeline and how this conductive tissue might adaptively respond to this growth-limiting environmental variable. Data currently available suggest that low temperatures lead to changes in the anatomy and phenological development of the phloem. In order to ensure the functionality of the phloem and thus the survival of trees at the high-elevation treeline, appropriate adaptations to the prevailing low temperatures are therefore to be expected and are discussed in this review.

Keywords: adaptation, intra-annual phloem formation, low temperature, phloem anatomy, phloem transport, phenology, tree growth, treeline

INTRODUCTION

Phloem is a highly complex vascular tissue that performs numerous functions for the growth and survival of trees. The long-distance transport of both phytohormones and RNAs as information signals between above- and belowground tissues (Lacombe and Achard, 2016; Kondhare et al., 2021) and carbohydrates as nutrients from source to sink (Savage et al., 2016) are considered to be important key functions, enabled by the anatomical properties of phloem (Schulz and Thompson, 2009; Knoblauch and Oparka, 2012; Liesche et al., 2015). The characteristics of the phloem in trees differ from those in herbaceous plants (for more details see De Schepper et al., 2013 and the literature therein). Phloem transport occurs along three functional units (i.e., collection, transport, and release phloem) via sieve cells and sieve tube elements in gymnosperms and angiosperms, respectively (van Bel, 2003; De Schepper et al., 2013). The water required for phloem transport is provided by the radial connection to the xylem via ray parenchyma (Box 1; Hölttä et al., 2009; Pfautsch et al., 2015; Sevanto, 2018). In this process, ray parenchyma cells provide not only the radial transport of water and carbohydrates between xylem and phloem (van Bel, 1990) but also
in and out of the axial parenchyma tissue (Pfautsch et al., 2015). Sieve elements and companion cells are laterally connected via sieve areas consisting of several sieve pores. Axially, sieve elements are connected by sieve areas in gymnosperms, which form a sieve plate with large sieve pores in angiosperms (Liesche et al., 2017). During the growing season, there is a transition from early phloem cells with wide lumens that have a conducting function to late phloem cells with narrow lumens that primarily have a storage function (Gričar et al., 2021). These phloem cells are separated by a band of axial parenchyma cells (Figure 1; Gričar and Ćufar, 2008).

Although both the xylem and phloem arise from radial cell division of the cambium (Larson, 1994), the formation of both tissues is not synchronous and is driven by different factors (Prislan et al., 2013; Gričar et al., 2015, 2021). In contrast to the xylem, the phloem was found to be less susceptible to environmental influences and mainly endogenously controlled (Alfieri and Evert, 1968; Rosner et al., 2001; Gričar and Ćufar, 2008; Prislan et al., 2013; Swidrak et al., 2014; Miller et al., 2020; Gričar et al., 2021). Even before the reactivation of cambial activity after dormancy, phloem formation starts with expansion and differentiation of overwintered phloem cells (Alfieri and Evert, 1968, 1973; Antonova and Stasova, 2006; Gričar and Ćufar, 2008). After the reactivation of the cambium, the onset of phloem cell production precedes xylem production by approximately 2–3 weeks (Alfieri and Evert, 1968; Antonova and Stasova, 2006, 2008; Swidrak et al., 2014; Jyske et al., 2015), due to the short functional life span of sieve cells of 1–2 years before they collapse (Evert, 2006).

Because climate change strongly affects tree growth and the phloem fulfills important functions in carbon allocation, phloem anatomy and phenology gained increasing importance in the last decade (e.g., Liesche et al., 2015, 2017; Miller et al., 2020; Gričar et al., 2021). Surprisingly, no studies were published yet dealing with intra-annual development and anatomy of phloem cells at high-elevation treeline sites, where trees reach their uppermost distributional limit due to temperature constraints on tree growth. Globally, trees at high-elevation treelines are exposed to seasonal mean air temperature between 5.5 and 7.5°C (Körner, 1998). In this brief review, we will evaluate possible adaptations in phloem phenology and anatomy under low temperatures.

## TEMPERATURE LIMITATION OF TREE GROWTH AT HIGH ELEVATIONS

Low temperatures are considered to be the main reason for the upper distributional boundary of trees (e.g., Körner and Paulsen, 2004; Rossi et al., 2007; Caccianiga et al., 2008; Körner, 2015). Only a few taxa are able to reach the uppermost high-elevation limit for trees – the treeline – which is a conspicuous climate-driven ecological boundary and occurs worldwide at high elevations (Körner, 2012). In the temperate climate zone, conifers of the genera Pinus, Larix, and Picea, as well as tree species belonging to Betulaceae and Fagaceae, are among the most successful tree taxa dominating at the high-elevation treeline (e.g., Körner and Paulsen, 2004; Caccianiga et al., 2008; Körner, 2012; Leuschner and Ellenberg, 2017).

At high elevations, the meristem activity, i.e., cell division, enlargement, and differentiation, is more sensitive to low-temperature conditions prevailing during the growing season than photosynthesis. The “carbon-sink-limitation hypothesis” is currently the most widely accepted hypothesis for explaining the formation of treelines (e.g., Körner and Paulsen, 2004; Rossi et al., 2007; Körner, 2015). While photosynthesis is still active at temperatures near the freezing point (e.g., Häsler, 1982; Weng et al., 2005), the exponential increase in cell doubling time below c. 5°C causes the upper elevational limit for the occurrence of trees (Körner, 2003, 2015). Numerous studies revealed that both aboveground and belowground growth is strongly reduced at temperatures below 5°C (Häsler et al., 1999; Alvarez-Uria and Körner, 2007, 2011; Rossi et al., 2007, 2008; Schenker et al., 2014). All growth processes, i.e., root, radial stem, and shoot growth as well as budburst, and subsequent leaf and flower development require water, carbon compounds, nutrients, and hormones via the vascular system, i.e., phloem and xylem (Savage and Chuine, 2021). Several findings have already demonstrated the effects of low temperatures on the xylem, i.e., reductions in wood formation, cell diameter, and cell wall thickness (e.g., Rossi et al., 2007, 2008; Castagneri et al., 2017; Babushkina et al., 2019). Therefore, it is necessary to consider low-temperature influences at high elevations on the anatomical characteristics and phenology of the phloem to comprehensively understand temperature-induced growth limitation.

### ARE CARBON AVAILABILITY AND PHLOEM TRANSPORT REDUCED AT THE HIGH-ELEVATION TREELINE?

A continuous supply of carbohydrates is essential for tree growth (Cuny et al., 2015; Deslauriers et al., 2016), especially under extreme climatic conditions prevailing at the treeline. Several authors have shown that photosynthesis is less limited by low temperatures than growth (Grace et al., 2002, for a review, see Körner, 2015). Larcher (2003) reported that photosynthetic production in evergreen conifers commences at temperatures between −5 and −3°C. An excess of these assimilates are stored as non-structural carbohydrates (Box 1; NSC) like starch and other carbon compounds (sugar, alcohol, lipids, proteins) which can be later mobilized for growth, development, and regeneration (Palacio et al., 2014). Several studies have shown an increasing trend in the content of NSC in leaves, stem, and root along an elevational transect, i.e., decrease in air temperature, up to the treeline (e.g., Hoch et al., 2002; Hoch and Körner, 2003, 2009; Gruber et al., 2011). An increase in NSC storage is considered as an immediate physiological response to low temperatures and to compensate for the imbalance between carbon production and demand (Hoch and Körner, 2012).

As mentioned above, root growth decreases at temperatures below 5°C (Alvarez-Uria and Körner, 2007, 2011; Schenker et al., 2014). However, at the treeline, reduced nutrient availability due to low nutrient mineralization rates (e.g., Meentemeyer, 1977)
functional even at the world’s highest treeline formed by nutrient uptake at the treeline. Hence, phloem is able to remain phloem transport in favor of root growth to ensure an adequate This implies both a sufficient carbon availability and an efficient lowered by self-shading, which underpin the previous results. Pinus cembra et al. (2017) demonstrated an accelerated fine root turnover and resulting in a higher root to shoot ratio. Additionally, Kubisch Picea abies fine root biomass with elevation in Norway spruce (Hoch and Körner, 2005). In this regard Hertel and Schöling (2011) found an increase of appropriate adaptive responses for an efficient nutrient uptake. In this regard Heid and Schöling (2011) found an increase of fine root biomass with elevation in Norway spruce (Picea abies) resulting in a higher root to shoot ratio. Additionally, Kubisch et al. (2017) demonstrated an accelerated fine root turnover and higher production rate in Pinus cembra under soil temperatures lowered by self-shading, which underpin the previous results. This implies both a sufficient carbon availability and an efficient phloem transport in favor of root growth to ensure an adequate nutrient uptake at the treeline. Hence, phloem is able to remain functional even at the world’s highest treeline formed by Polylepis sp. at c. 5,000 m a.s.l. (Hoch and Körner, 2005).

**EXPECTED ADAPTATIONS OF PHLOEM ANATOMY AND PHENOLOGY AS A RESPONSE TO EXTREME CLIMATE CONDITIONS AT THE HIGH-ELEVATION TREELINE**

In field experiments, several authors found that phloem formation and anatomy are less responsive to variability in environmental conditions compared to the xylem (Alfieri and Evert, 1968; Rosner et al., 2001; Gričar et al., 2014; Swidrak et al., 2014; Miller et al., 2020). However, extreme climate conditions might impair intra-annual phloem formation and/or phloem anatomy which would adversely affect carbon transport and

---

**BOX 1 | Glossary of terms.**

- **Axial parenchyma**: Longitudinally arranged parenchyma cells for vertical transport and storage of carbohydrates.
- **Callose**: Polysaccharide (β-1,3-glucan) found in cell walls of higher plants; involved in phloem transport and deposited as response to injury.
- **Companion cells**: Specialized nucleated parenchyma cells regulating transport of substances in and out of sieve elements.
- **Early phloem**: First formed phloem cells with large diameter primarily for transport of substances.
- **Initial early phloem**: Undifferentiated overwintering cells formed at the end of the previous growing season, which differentiate to early phloem in spring.
- **Late phloem**: Formed after early phloem with narrow diameter primarily for storage.
- **Non-structural carbohydrates**: Comprise primarily soluble sugars and starch; provide substrates for growth, metabolism and osmoregulation.
- **Sieve area**: Are developed in angiosperms in transverse cell walls between axial connected sieve elements (cf., Figure 1).
- **Sieve plate**: Are developed in angiosperms in transverse cell walls between axial connected sieve elements (cf., Figure 1).
- **Sieve pores**: Pores lined with callose within sieve areas and sieve plates (cf., Figure 1).
- **Ray parenchyma**: Radially arranged parenchyma cells for storage and lateral transport of water and carbohydrates between xylem and phloem.

---

as well as cold-related physical constraints in water and nutrient uptake from soil to root (Weih and Karlsson, 2001, 2002) require appropriate adaptive responses for an efficient nutrient uptake. In this regard Hertel and Schöling (2011) found an increase of fine root biomass with elevation in Norway spruce (Picea abies) resulting in a higher root to shoot ratio. Additionally, Kubisch et al. (2017) demonstrated an accelerated fine root turnover and higher production rate in Pinus cembra under soil temperatures lowered by self-shading, which underpin the previous results. This implies both a sufficient carbon availability and an efficient phloem transport in favor of root growth to ensure an adequate nutrient uptake at the treeline. Hence, phloem is able to remain functional even at the world’s highest treeline formed by Polylepis sp. at c. 5,000 m a.s.l. (Hoch and Körner, 2005).

Does the Temperature-Induced Reduction in Cambial Activity Also Affect Phloem Anatomical Features?

Based on several studies on coniferous species, temperature-induced changes in xylem anatomy affect cell size, as well as cell wall thickness, the latter being related to carbon mobilization and deposition rates (e.g., Rossi et al., 2008; Lupi et al., 2010; Körner, 2012). Castagneri et al. (2017) and Babushkina et al. (2019) demonstrated along an elevational gradient, that low temperatures especially at high elevations are one of the key climatic factors affecting xylem cell enlargement and cell wall deposition, which subsequently are reflected in a reduction of cell diameter and cell wall thickness. Similar effects on phloem anatomy are to be expected at the treeline, where low temperatures prevail during the growing season (Figure 1). Moreover, it should be noted that possible low-temperature effects on phloem formation and anatomy may impair the carbon transport in some tree species. Liesche et al. (2015) demonstrated in their study a lower phloem transport rate in narrow sieve cells of gymnosperms compared to angiosperms, which supports this assumption. Additionally, Pfautsch et al. (2015) reported that phloem sap becomes increasingly viscous with decreasing temperature, which additionally increases resistance in phloem transport. Because at high elevation xylem cells were found to be highly vulnerable to cavitation (Mayr et al., 2002, 2003), impaired phloem transport could also have an impact on the repair of embolized xylem cells (cf., Pfautsch et al., 2015).

How Can Phloem Transport Efficiency Be Ensured Under Low Temperatures Prevailing at the High-Elevation Treeline?

Several findings conducted at mid-elevation sites (1,200 m a.s.l., Gričar et al., 2014, 2015, 2021; Miller et al., 2020)

---

| Sieve pores: Pores lined with callose within sieve areas and sieve plates (cf., Figure 1). | Ray parenchyma: Radially arranged parenchyma cells for storage and lateral transport of water and carbohydrates between xylem and phloem. |
FIGURE 1 | Schematic illustration of known and assumed low-temperature effects on phloem anatomy and formation along an elevational transect. Upper graphic: Current studies dealing with environmental influence on the phloem conducted at different elevations. Below: General phloem structure at low-elevation sites, alterations caused by decrease in temperature at mid-elevation sites, and effects and possible phloem adaptations at the high-elevation treeline. Low elevation, top: Annual phloem ring exemplified by Picea abies after Gričar and Čufar (2008) consist of early phloem (EP) with initial early phloem (IEP), the continuous tangential band (BAP) of axial parenchyma cells (AP), late phloem (LP) with adjacent cambial zone (CZ) and latewood tracheids (LW); bottom: phloem cells (axial view) with lateral sieve areas (LSA) and sieve pores (SPo), as well as axial sieve areas (SA) in conifers (right) and sieve plates (SP) with larger sieve pores in angiosperms (left), respectively.

indicate plasticity in phloem formation, i.e., the ability to adapt structurally to ensure optimal function under local environmental conditions. Phloem cells show a tip-to-base widening (Jyske and Höltä, 2015; Ryan and Robert, 2017), and to compensate for the smaller conducting area the number of conduits per unit phloem area increases (Jyske and Höltä, 2015). Due to aerodynamic coupling of the shoot to the macroclimate, height growth at the treeline is
more strongly restricted than radial stem growth (Hoch and Körner, 2005; Körner, 2012; Wang et al., 2012). Therefore, it can be assumed that phloem responds with an increase in cell density resulting in an increased ratio of early to late phloem cells for better conductivity. The priority of the early phloem as the conductive part over the late phloem (storage) is shown by its robustness to environmental influences and underpins its greater importance for the survival of trees (Larson, 1994; Gričar et al., 2015, 2021). Temperature-related changes in radial dimension (lumen) of initial early phloem (Box 1; i.e., first tangential rows of cells at the phloem growth ring boundary formed in the previous year) and late phloem cells (Gričar et al., 2015), as well as a seasonal variability in length of sieve elements and axial parenchyma (Jyske et al., 2015), suggest an adaptive change in cell size for an efficient conductivity under low temperatures. Furthermore, several authors reported a decrease in cell wall thickness in earlywood and latewood tracheids at mid elevations (1,260 m a.s.l., Gindl et al., 2001; 1,200 m a.s.l., Gričar et al., 2015), and a similar effect in sieve cell wall thickness could be conceivable as another way to increase the sieve cell lumen diameter at high elevations. However, the conductive capacity of the phloem at the treeline may be favored not only by adaptations in the aforementioned anatomical traits but also by an increase in sieve pore size and frequency along the pathway (Mullendore et al., 2010). Sieve pores can be lined with callose or nearly occluded with it, which may play a significant role in transport regulation through the symplast (Neuberger and Evert, 1975; Knoblauch and van Bel, 1998). Callose is deposited as a response to cell injury, senescence, or dormancy (Evert, 1990) whereby not all pores are sealed with it (Montwe et al., 2019; Prislan et al., 2019). Due to the prevailing extreme climate conditions at the high-elevation treeline, less callose might accumulate as a temperature-dependent response to increase sieve pore size and create less resistance to ensure phloem transport capacity (cf., McNairn, 1972). Because of the severely time-limited lifespan of sieve elements, the priority for tree survival is in phloem production while xylem thickness can be reduced or become discontinuous (Evert, 2006). Especially under extreme environmental conditions, this is reflected in a lower xylem to phloem ratio (Larson, 1994; Plomion et al., 2001; Gričar et al., 2009).

Phenological adaptations can also influence anatomical properties of vascular tissues in favor of functional capacity, as demonstrated by Cuny et al. (2019). It became apparent, that in cold environments xylem differentiation rate (i.e., cell elongation and cell wall thickening) decreased but to mitigate the influence of low temperatures the duration of these processes was extended. Miller et al. (2020) reported a delay in the transition from early to late phloem in *Picea abies* along an elevational gradient extending from 450 m to 1,250 m a.s.l., which was related to the onset of cambial activity, cell production rate, and the number of cells in the early phloem. Findings in this study also provide the first evidence that in contrast to xylem formation (e.g., Rossi et al., 2006) maximum phloem cell production rate does not coincide with the summer solstice. These results suggest that at the high-elevation treeline the maximum phloem growth rate may shift to the warmest period instead of being controlled by photoperiod or endogenous factors. Furthermore, it is conceivable that more overwintering phloem cells are formed or that phloem cells remain functional for more than one to two growing seasons, which would correspond to the observations of Holdheide (1951). Hence, adaptations in intra-annual phloem formation and anatomy in response to low temperatures and climate extremes (e.g., the occurrence of frost drought) prevailing at the high-elevation treeline are to be expected (Figure 1; cf., Agusti and Greb, 2013; Gričar et al., 2015, 2021; Miller et al., 2020). This would also entail a shift from endogenous to exogenous control of phloem formation.

**AREAS OF FUTURE RESEARCH**

Expected adaptations of phloem phenology and anatomy induced by extreme climatic conditions prevailing at the high-elevation treeline are shown in Figure 1. In order to understand the effects of low temperatures on the phloem at high elevations, several questions need to be clarified, e.g., (i) How does phloem phenology and anatomy adapt to climate extremes (e.g., cold summer with shortened growing period, early or late frost events) at the treeline? (ii) To what extent is phloem transport capacity and efficiency affected by low temperatures? (iii) Is tip-to-base widening occurring at treeline (cf., Jyske and Hölttä, 2015; Ryan and Robert, 2017)? (iv) Are there organ specific differences in phloem formation and anatomy, e.g., in branches (twigs) vs. roots due to exposure to different temperature regimes, i.e., high vs. low daily temperature variability? (v) How sensitive is phloem formation to environmental stress (e.g., frost drought) within the treeline ecotone? (vi) Is there a year-to-year variability in phloem formation depending on weather conditions or does endogenous control of phloem formation also predominate at the high-elevation treeline? A combination of high measurement frequency and long-term observations in the field together with field chilling experiments, laboratory manipulations, and histo-anatomical analyses will be necessary to answer these questions. As there are differences in phloem anatomy between gymnosperms and angiosperms (Jensen et al., 2012; Liesche et al., 2015, 2017) the question arises if species-specific adaptations regarding phloem anatomy and/or phenology at the high-elevation treeline occur.

**CONCLUSION**

Phloem fulfills important functions for tree growth and development (Hölttä et al., 2009; Pfautsch et al., 2015;
However, compared to the xylem much less is known about the influence of growth-limiting temperatures at high elevations on the anatomy and intra-annual formation of the phloem. Therefore, to advance our knowledge about the characteristics of phloem tissue at the high-elevation treeline, field and experimental studies are urgently needed. This review is intended to highlight the existing gaps in knowledge regarding phloem adaptation to extreme climate conditions prevailing at the high-elevation treeline and to serve as a stimulus to open up a new area of research.

REFERENCES

Agusti, J., and Greb, T. (2013). Going with the wind–Adaptive dynamics of plant secondary meristems. Mech. Dev. 130, 34–44. doi: 10.1016/j.mod.2012.05.011
Alfieri, F. J., and Evert, R. F. (1968). Seasonal development of the secondary phloem in Pinus. Am. J. Bot. 55, 518–528. doi: 10.2307/2440583
Alfieri, F. J., and Evert, R. F. (1973). Structure and Seasonal development of the secondary phloem in the Pinaceae. Bot. Gaz. 134, 17–25. doi:10.1086/336674
Alvarez-Uria, P., and Körner, C. (2007). Low temperature limits of root growth in deciduous and evergreen temperate tree species. Funct. Ecol. 21, 211–218. doi: 10.1111/j.1365-2427.2007.01231.x
Alvarez-Uria, P., and Körner, C. (2011). Fine root traits in adult trees of evergreen Scots pine stems. Russ. J. Dev. Biol. 37, 306–320. doi: 10.1134/S1062360409050043
Alvarez-Uria, P. and Sta. S. V. (2008). Seasonal development of phloem in Siberian larch stems. Russ. J. Dev. Biol. 39, 207–218. doi:10.1134/S1062360408040024
Babushkina, E. A., Belokopytova, L. V., Zhimova, D. F., and Vaganov, E. A. (2019). Plasticity in variation of xylem and phloem cell characteristics of Norway spruce under different local conditions. Front. Plant Sci. 6:730. doi: 10.3389/fpls.2015.00730

AUTHOR CONTRIBUTIONS

Both authors listed have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

FUNDING

The publication of this manuscript was financially supported by the University of Innsbruck.
Hoch, G., and Körner, C. (2012). Global patterns of mobile carbon stores in trees at the high-elevation tree line. Glob. Ecol. Biogeogr. 21, 861–871. doi: 10.1111/j.1466-8238.2011.00731.x

Hoch, G., Popp, M., and Körner, C. (2002). Altitudinal increase of mobile carbon pools in Pinus cembra suggests sink limitation of growth at the Swiss treeline. Ökos 98, 361–374. doi: 10.1034/j.1600-0706.2002.980301.x

Holdheide, W. (1951). “Anatomie mitteleuropäischer Gehölzrinden (mit mikrophotographischem Atlas),” in Handbuch der Mikroskopie in der Technik, 5th Edn, ed. H. Freund (Frankfurt-am-Main: Umschau Verlag), 193–367.

Hölzt, M., T., Mencuccini, M., and Nikinmäki, E. (2009). Linking phloem function to structure: analysis with a coupled xylem-phloem transport model. J. Theor. Biol. 259, 325–337. doi: 10.1016/j.jtbi.2009.03.039

Jensen, K. H., Liesche, J., Bohr, T., and Schulz, A. (2012). Universality of phloem transport in seed plants. Plant Cell. Environ. 35, 1065–1076. doi: 10.1111/j.1365-3040.2011.02472.x

Jyske, T., and Hölttä, T. (2015). Comparison of phloem and xylem hydraulic architecture in Picea abies stems. New Phytol. 205, 102–115. doi: 10.1111/nph.12973

Jyske, T. M., Suuronen, J.-P., Pranovich, A. V., Laakso, T., Watanabe, U., Kuroda, M., and Hölttä, T. (2015). Comparison of long-distance signalling in plants. J. Exp. Bot. 72, 4218–4236. doi: 10.1093/jxb/erab048

Körner, C. (1998). A re-assessment of high elevation treeline positions and their explanation. Oecologia 115, 445–459. doi: 10.1007/s004420050540

Körner, C. (2002). Altitudinal increase of mobile carbon stores in trees at the high-elevation tree line. Glob. Ecol. Biogeogr. 11, 74–80. doi: 10.1034/j.1399-3054.2002.1150108.x

Meinmeyer, V. (1977). “Climatic regulation of decomposition rates of organic matter in terrestrial ecosystems,” in Proceedings of the Environmental Chemistry and Cycling Processes. Department of Energy Symposium Series CONF-760429, eds D. C. Adams and I. L. Brubin (Washington, DC: U.S. Department of Energy), 779–789.

Miller, T. W., Stangler, D. F., Larysch, E., Seifert, T., Spiecker, H., and Kahle, H. P. (2020). Plasticity of seasonal xylem and phloem production of Norway spruce along an elevational gradient. Trees 34, 1281–1297. doi: 10.1007/s00468-020-01997-6

Monte, D., Hacke, U., Schreiber, S. G., and Stanfield, R. C. (2019). Seasonal vascular tissue formation in four boreal tree species with a focus on callose deposition in the phloem. Front. For. Glob. Change 2:8. doi: 10.3389/ffgc.2019.00058

Mullendore, D. L., Windt, C. W., Van As, H., and Knoblauch, M. (2010). Sieve tube geometry in relation to phloem flow. Plant Cell 22, 579–593. doi: 10.1105/tpc.109.097094

Neuner, S. D., and Evert, R. F. (1975). Structure and development of sieve areas in the hypocotyl of Pinus resinosa. Protoplasma 84, 109–125. doi: 10.1007/BF02075947

Palacio, S., Hoch, G., Sala, A., Körner, C., and Millard, P. (2014). Does carbon storage limit tree growth? New Phytol. 201, 1096–1100. doi: 10.1111/nph.12602

Pflaust, S., Hölttä, T., and Mencuccini, M. (2015). Hydraulic functioning of tree stems—fusing ray anatomical, radial transfer and capacitance. Tree Physiol. 35, 706–722. doi: 10.1093/treephys/tpv058

Plomion, C., Leprovost, G., and Stokes, A. (2001). Wood formation in trees. Plant Physiol. 127, 1513–1523. doi: 10.1104/pp.1010816

Piršlan, P., Gríčar, I., de Luis, M., Smith, K. T., and Čufar, K. (2013). Phenological variation in xylem and phloem formation in Fagus sylvatica from two contrasting sites. Agric. For. Meteorol. 180, 142–151. doi: 10.1016/j.agrformet.2013.06.001

Piršlan, P., Mrak, P., Žnidarič, N., Strus, J., Humar, M., Thaler, N., et al. (2019). Intra-annual dynamics of phloem formation and ultrastructural changes in sieve tubes in Fagus sylvatica. Tree Physiol. 39, 262–274. doi: 10.1093/treephys/tpy102

Rosner, S., Baier, P., and Kikut, S. (2001). Osmotic potential of Norway Spruce (Picea abies (L.) Karst.) secondary phloem in relation to anatomy. Trees 15, 472–582. doi: 10.1007/s00468-001-0131-9

Rossi, S., Deslauriers, A., Anfodillo, T., and Carraro, V. (2007). Evidence of threshold temperatures for xylegenesis in conifers at high altitudes. Oecologia 152, 1–12.

Rossi, S., Deslauriers, A., Anfodillo, T., Morin, H., Saracino, A., Motta, R., et al. (2006). Conifers in cold environments synchronize maximum growth rate of tree-ring formation with day length. New Phytol. 170, 301–310. doi: 10.1111/j.1469-8137.2006.01660.x

Savio, S., Deslauriers, A., Gríčar, J., Seo, J. W., Rathgeber, C. B. K., Anfodillo, T., et al. (2008). Critical temperatures for xylegenesis in conifers of cold climates. Glob. Ecol. Biogeogr. 17, 696–707. doi: 10.1111/j.1466-8238.2008.00417.x

Ryan, M. G., and Robert, E. M. R. (2017). Zero-calorie sugar delivery to roots. Nat. Plants 3, 922–923. doi: 10.1038/s41477-017-0070-0

Salmon, Y., Dietrich, L., Sevanto, S., Hölttä, T., Dannoura, M., and Epron, D. (2019). Drought impacts on tree phloem: from cell-level responses to ecological significance. Tree Physiol. 39, 173–191. doi: 10.1093/treephys/tpy153

Savage, J. A., and Chuihe, I. (2021). Coordination of spring vascular and organ phenology in deciduous angiosperms growing in seasonally cold climates. New Phytol. 230, 1700–1715. doi: 10.1111/nph.17289
Savage, J. A., Clearwater, M. J., Haines, D. F., Klein, T., Mencuccini, M., Sevanto, S., et al. (2016). Allocation, stress tolerance and carbon transport in plants: how does phloem physiology affect plant ecology? *Plant Cell. Environ.* 39, 709–725. doi: 10.1111/pce.12602

Schenker, G., Lenz, A., Körner, C., and Hoch, G. (2014). Physiological minimum temperatures for root growth in seven common European broad-leaved tree species. *Tree Physiol.* 34, 302–313. doi: 10.1093/treephys/tpu003

Schulz, A., and Thompson, G. A. (2009). *Phloem Structure and Function: Encyclopedia of Life Sciences (eLS)*. 2nd Edn, Wiley. doi: 10.1002/9780470015902.a0001290.pub2

Sevanto, S. (2018). Drought impacts on phloem transport. *Curr. Opin. Plant Biol.* 43, 76–81. doi: 10.1016/j.pbi.2018.01.002

Swidrak, I., Gruber, A., and Oberhuber, W. (2014). Xylem and phloem phenology in co-occurring conifers exposed to drought. *Trees* 28, 1161–1171. doi: 10.1007/s00468-014-1026-x

van Bel, A. J. E. (1990). Xylem-phloem exchange via the rays: the undervalued route of transport. *J. Exp. Bot.* 41, 631–644. doi: 10.1093/jxb/41.6.631

van Bel, A. J. E. (2003). The phloem, a miracle of ingenuity. *Plant Cell. Environ.* 26, 125–149. doi: 10.1046/j.1365-3040.2003.00963.x

Wang, Y., Ćufar, K., Eckstein, D., and Liang, E. (2012). Variation of maximum tree height and annual shoot growth of Smith fir at various elevations in the Sygera Mountains, southeastern Tibetan Plateau. *PLoS One* 7:e31725. doi: 10.1371/journal.pone.0031725

Weih, M., and Karlsson, P. S. (2001). Growth response of mountain birch to air and soil temperature: is increasing leaf-nitrogen content an acclimation to lower air temperature? *New Phytol.* 150, 147–155. doi: 10.1046/j.1469-8137.2001.00708.x

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher’s Note: All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Copyright © 2021 Schröter and Oberhuber. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.