

Tip-to-base conduit widening remains consistent across cambial age and climates in *Fagus sylvatica* **L.**

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Water transport, mechanical support and storage are the vital functions provided by the xylem. These functions are carried out by different cells, exhibiting significant anatomical variation not only within species but also within individual trees. In this study, we used a comprehensive dataset to investigate the consistency of predicted hydraulic vessel diameter widening values in relation to the distance from the tree apex, represented by the relationship $D_h \propto L^{\beta}$ (where D_h is the hydraulic vessel diameter, L the distance from the stem apex and *β* the scaling exponent). Our analysis involved 10 Fagus sylvatica L. trees sampled at two distinct sites in the Italian Apennines. Our results strongly emphasize that vessel diameter follows a predictable pattern with the distance from the stem apex and *β* ∼ 0.20 remains consistent across cambial age and climates. This finding supports the hypothesis that trees do not alter their axial configuration represented by scaling of vessel diameter to compensate for hydraulic limitations imposed by tree height during growth. The study further indicates that within-tree variability significantly contributes to the overall variance of the vessel diameter-stem length exponent. Understanding the factors that contribute to the intraindividual variability in the widening exponent is essential, particularly in relation to interspecific responses and adaptations to drought stress.

Key words: allometry, architecture, diffuse-porous, hydraulic, quantitative wood anatomy, vessels, xylem traits.

Introduction

Water uptake and transport in vascular plants play a pivotal role in several developmental processes, including transpiration, photosynthesis and nutrient uptake. As long-living organisms, trees undergo structural and functional changes during ontogeny, which require steady adjustments in xylem hydraulic architecture in response to ever-increasing distances over which water must be transported (Meinzer et al. 2011). Regardless of their differences in growth rate and plant habit, trees encounter a common challenge as they grow taller in order to maintain water transport efficiency and conductance per unit leaf area. This challenge arises from flow resistance, proportional to the total path length, as trees increase in height. To maintain a constant conductance per unit leaf area and supply a branching canopy, vessels must widen proportionally from the tip to the base of the stem as individual plants grow taller. However, while a set of large xylem conduits can have lower overall carbon costs than smaller conduits with the same hydraulic conductance, this efficiency comes with the potential cost of increased vulnerability to embolism (Hacke et al. 2006, 2023; Jacobsen and Pratt 2023, but see Hajek et al. 2016). Consequently, to ensure water transport through plants, xylem conductance, embolism vulnerability and the construction costs of the conducting system must be carefully balanced as trees grow taller. Because of the importance of this equilibrium in plant hydraulic evolution and forest responses to climate change, optimality models have been developed over the past few decades to predict how and why plants vary vessels diameters throughout their bodies.

Several conflicting theories have been proposed to describe the coordination between tree size and vascular architecture. Some models implicitly assume constant vessel diameter (e.g. Shinozaki et al. 1964a, b), while others predict that they widen from tip to base (e.g. West et al. 1999; Savage et al. 2010). Among them, the theoretical West, Brown and Enquist (WBE, West et al. 1997, 1999) model posits that the fractal-like geometry found in many biological systems is the fundamental basis for the allometric scaling observed in the vascular plant transport system. Empirical measurements strongly indicate that vessel lumen areas will narrow axially from tree base to stem apex and branches with the power function as the distance from the tree apex increases, as observed in a wide range of species, across individuals of different heights as well as across different taxa (Anfodillo et al. 2006; Prendin et al. 2018; Koçillari et al. 2021; Petit et al. 2023 among the others). This consistent axial scaling is a common trait across a wide range of tree species, suggesting that natural selection primarily shapes the distribution of xylem vessels diameter along the axis through uniform biophysical principles governing plant functional integrity, rather than through niche differentiation and divergent ecological strategies (Anfodillo et al. 2013; Olson et al. 2014; Williams et al. 2019; Zhong et al. 2019). Along the xylem pathway, the vessels' widening pattern is well approximated by the power function $y = \alpha \cdot L^{\beta}$, where *y* represents the vessel hydraulic diameter, *α* is the allometric constant, *L* is the distance from stem apex and β is the tip-to-base vessel widening exponent, which describes the rate of basipetal widening. A zero-value *β* indicates vessels

Received: December 7, 2023. **Revised:** June 17, 2024. **Accepted:** July 1, 2024 © The Author(s) 2024. Published by Oxford University Press. All rights reserved. For permissions, please e-mail: journals.permission@oup.com. of uniform diameter as the pipe model implicitly assumes (Shinozaki et al. 1964a, b), whereas larger *β* values indicate greater rates of axial widening. According to the theoretical WBE model (West et al. 1999), a vascular design that fully compensates for the progressive increase in hydraulic resistance with tree height should have an ideal vessel diameter widening exponent $\beta = 0.25$ (for the idealized plant of the WBE model) and translates into $\beta \sim 0.2$ when the scaling is between vessel diameter and distance from the stem apex (Becker et al. 2000; Anfodillo et al. 2006; Petit and Anfodillo 2009). However, from several studies it has emerged that the rate of vessel widening (*β*) varies across tree species, habit and growth environment, typically ranging from ≈ 0.13 to ≈ 0.34 (Petit et al. 2010, 2011; Pfautsch et al. 2018; Williams et al. 2019), for reasons that are currently not entirely clear. Even well-established models contain assumptions that require further testing and await the incorporation of novel empirical observations. In fact, since the publication of the WBE model, several criticisms have been highlighted that question its basic framework, assumptions and generality and reveal empirical patterns that contradict parts of the WBE model. Savage et al. (2010) tried to improve the theory's representation of vascular architecture, highlighting how hydraulic safety and efficiency may have shaped the evolution of vascular networks (but see Sopp and Valbuena 2023). That is, the packing rule contradicts the premise of the WBE model that vessel frequency remains constant as vessel radii taper. Other studies over the past few years have reported new empirical observations that are waiting to be incorporated into hydraulic models. For instance, taller plants have broader vessels at the base because of tip-to-base vessel widening, as well as wider terminal vessels (e.g. Olson et al. 2018) thought to be the result of selection maintaining a constant leaf-specific sapwood conductance with height increase. Lechthaler et al. (2019a, 2020) found for *Acer pseudoplatanus* L. that vessels in leaf veins enlarge from their termination toward the petiole base, twice to how they behave in stems. Further, in profiles of diameter from tip to base, vessels close to the stem base are frequently found to be narrower than expected (Petit and Anfodillo 2009; Olson et al. 2018; Pfautsch et al. 2018, but see Koçillari et al. 2021).

From the above it has emerged that there are still many outstanding issues regarding the variability of xylem anatomical traits as an individual tree grows taller, which currently limits a full understanding of vessel scaling patterns along the stem flow path (see Rosell et al. 2017 and Olson et al. 2021 for an overview). Among them, the vessel diameter is expected to widen with distance from the stem tip according to a power law $(D_h \propto L^{\beta})$; however, not all observations confirmed a precise value or a strict power-like pattern of *β*, the vessel widening exponent. Furthermore, although tipto-base vessel widening has been demonstrated in certain species, there is limited empirical evidence to support the notion that vessels should expand uniformly throughout the stem with height growth (Weitz et al. 2006; Prendin et al. 2018; Petit et al. 2023). Studies that commonly included the analyses of vertical patterns of wood anatomical traits were restricted to only a few outermost tree-rings at different points along the main stem (Anfodillo et al. 2006; Petit et al. 2010; Pfautsch et al. 2018; Williams et al. 2019) or branches and petioles (Chen et al. 2012; Ray and Jones 2018). The presence of tip-to-base widening intraspecific variability is an essential prerequisite for empirically testing the adaptive versus non-adaptive hypothesis where natural selection narrows a wider field of evolutionary possibilities (see Figure 2 in Olson et al. 2021).

In this work, we evaluated the consistency of theoretical predictions of vessel widening function in *Fagus sylvatica* L., a diffuse-porous tree species with wide niche across Palearctic temperate forests. Our primary objectives were to investigate: (i) whether the widening exponents of this tree species are consistent with the theoretical predictions of the WBE model and (ii) whether the scaling exponent remains invariant with age, size and growth environment. Various hypotheses can be conceptualized regarding the anatomical changes a tree undergoes during its growth to counteract the effects of height increase: (i) uniform tip-to-base vessel widening (*β*) with height growth (Prendin et al. 2018), but with the allometric constant (α) increasing with tree height (Olson et al. 2014); (ii) tip-to-base vessel widening not being ontogenetically stable, but with the scaling exponent (β) increasing with tree height, resulting in a slight increase in vessel diameter for a given tree height (Rosell et al. 2017); and (iii) both tip-to-base vessel widening (*β*) and the allometric constant (α) remain relatively constant with height growth (Prendin et al. 2018; Petit et al. 2023) (see Fig. 1). To test these hypotheses, we have generated a comprehensive dataset encompassing the within-tree variation in vessel diameter. This extensive dataset was constructed through a retrospective reconstruction of the height growth history of 10 mature *F. sylvatica* L. trees, which were collected from two distinct sites in the Italian Apennines, characterized by contrasting climates. Our sampling strategy involved measuring ring-level vessel diameters at multiple positions along the stem, from the pith to the bark. This meticulous approach allowed us to capture the full range of variation in vessel diameter within each tree, resulting in a robust and detailed dataset.

Materials and methods

Study sites and plant material

Plant material came from two selected sites along the Italian peninsula characterized by different climatic conditions**.** The first site is an even-aged *F. sylvatica* stand in the Abetone forest (Northern Apennines, Italy, 44.10 N, 10.70 E; 1300 m a.s.l.; hereafter Site1) characterized by a perhumid climate with 2046 mm of annual precipitation and a mean annual temperature of 7.8 ◦C (1981–2010 period). The second even-aged *F. sylvatica* site is in Southern Italy (Serra San Bruno, Southern Apennines; 38.56 N, 16.29 E; 1050 m a.s.l.; hereafter Site2), where the climate is typically Mediterranean with hot dry summers and predominantly winter rainfall; the annual precipitation is 1706 mm and the mean annual temperature is 11.1 ◦C (1981–2010 period).

Five dominant *F. sylvatica* trees for each site were felled from the study sites and measured for their total height (Table S1 available as Supplementary data at *Tree Physiology* Online). Part of the selected plant material come from previous work (Rita et al. 2020). We reconstructed the apexto-tree base axial trend within each tree ring to provide an annual resolution the variability of xylem anatomical traits along the stem axis. For each tree, eight transversal stem discs were taken at eight different heights along the main stem i.e. 1.3, 4.0, 8.0, 12.0, 16.0, 20.0, 28.0 and 32.0 m from the base (Fig. 1). In the lab, the width of tree-rings

Figure 1. Scheme of the sampling protocol used to reconstruct the axial trend from apex to tree base within each ring. Eight stem discs were harvested at intervals of 1.30, 4, 8, 12, 16, 20, 28 and 32 m along the heights of eight trees (see Table S1 available as Supplementary Data at Tree Physiology Online). The annual stem elongation rate was calculated by linearly interpolating the distance between adjacent discs and dividing it by the difference in age between neighboring discs. For each ring, the distance (L) from the stem apex was calculated as the difference between the reconstructed tree height and the distance from the ground. On the left the conceptualized hypotheses concerning the vessel diameter-stem length relationship: (A) uniform tip-to-base vessel widening (*β*) occurs with height growth, while the allometric constant (*α*) increases proportionally with tree height; (B) tip-to-base vessel widening is not consistently stable over ontogeny, yet the scaling exponent (*β*) shows an increase with tree height, leading to a slight adjustment in vessel diameter for a given tree height; (C) both tip-to-base vessel widening (*β*) and the allometric constant (*α*) remain relatively consistent as tree height increases.The background tree design is by Freepik.

was first measured to the nearest 0.01 mm using a treering measuring system (LINTAB 6 coupled with TSAP-Win Scientific software, Rinntech, Heidelberg, Germany) and then analyzed using COFECHA software (Holmes 1983) to check for cross-dating errors. Further details of stand characteristics, sampling and cross-dating are described in Gentilesca et al. (2018). The retrospective reconstruction of annually resolved height growth rates was generated by stem analysis by means of ring-width data sampled from successive sections along the stems with the '*IncrementR*' package (Kašpar et al. 2019) in the R statistical environment (R Core Team 2021). The annual height growth rate between two successive stem discs was based on a linear interpolation of their axial distance by the difference in tree-ring counts (Prendin et al. 2018). Accordingly, the total tree height in each year was reconstructed as the height of the last stem disc plus the annual linear growth rate toward the next disc multiplied by the number of years since the last stem disc was reached (i.e. cambial age of the outermost tree ring from the last stem disc). For each site, mean annual temperatures and precipitations were obtained from the 30-arcsecond resolution data developed by Brunetti et al. (2014) and Crespi et al. (2018), respectively, as extensively detailed in Gentilesca et al. (2018).

Preparation of woody samples

For each collected disc, one radial strip (1 cm wide) running from the bark to the pith was selected to analyze the total radial growth. Samples were first checked to exclude reaction wood, eccentricity or wounding (Jourez et al. 2001, Kašpar et al. 2019) and then radial woody samples were split into 3–4 cm length pieces for processing. After that, anatomical micro cross-sections (15–20 μ m of thickness) were cut with a sliding microtome (Microm HM 400, Thermo Sci., Walldorf, Germany), stained with Safranin (1%) and Astra Blue (2%), dehydrated with ethanol and fixed on permanent slides with Eukitt mounting solution (Kindler GmbH, Freiburg, Germany). Digital images were captured with an integrated digital camera (DCM300, ScopeTek) installed on a light microscope (Zeiss Axiophot, Carl Zeiss Microscopy, Jena, Germany). Overlapping images of the whole micro-section were stitched together with Image Composite Editor software (ICE 2.0.3, Microsoft Corp., Redmond, WA, USA).

Quantitative wood anatomy

The images were processed using image-analysis software ImageJ (National Institutes of Health, Bethesda, MD, USA) which allowed measurement of tree-ring lengths and the anatomical features of vessels inside all tree rings. A semiautomated procedure of image analysis was utilized to ascertain the x and y coordinates of the centers of each vessel, as well as their lumen areas. A manual correction was necessary to exclude non-vascular elements or include vessels that had not been selected automatically. Briefly, digital images were converted from 24-bit color in a threshold binary image (mask) in which only the vessel lumens were kept. Before any measurement, the images of all micro-sections were calibrated from a scale bar of known length. For each tree ring, a chosen surface (Area $=$ *TRW* $*$ *l*, where *TRW* is the tree-ring width and $l = 2$ mm the tangential length) was analyzed for the xylem vessel lumen cross-sectional area (*A*). Vessel area was converted to diameter (*D*) assuming circularity of vessels, where

$$
D = \sqrt{\frac{4 \cdot A}{\pi}}
$$

This transformation allows partially excluding the influence of possible large true tracheids that may have been measured. Based on the vessel contribution to hydraulic conductance, we calculated the hydraulically weighted mean diameter (D_h) for each ring (Tyree and Zimmermann 2002):

$$
D_h = \left(\frac{1}{n}\sum_{1}^{n} D^4\right)^{\frac{1}{4}}
$$

In total, more than one and a half million xylem vessels were measured. Then, for each tree ring, we calculated the average vessel size (A_{av}) , the vessel density (D_v) , the theoretical hydraulic conductivity (*Ktheo*) and vessel composition (*S*). The vessel density (D_v) was estimated as the number of vessels per unit of surface area, while the vessel composition (*S*) was calculated as the ratio between A_{av} and D_v , according to Zanne et al. (2010)*.* The xylem-specific specific hydraulic conductivity (K_s) was estimated according to the modified Hagen– Poiseuille equation reported by Tyree and Ewers (1991, their Eq. (4) :

$$
K_s = \frac{\pi \cdot \rho}{128 \cdot \eta \cdot Si} \sum_{i=1}^{n} D^4
$$

where *ρ* and *η* are the density and dynamic viscosity of water at 20 \degree C, respectively, S_i is the ring surface (i.e. tree-ring width [∗] 2 mm), *n* is the number of vessels per unit of surface area and *D* is the vessel diameter.

Statistical analyses

To examine multivariate associations between variables, including categorical and continuous variables, we carried out ordinary principal component analysis (PCA) using the *PCAmix* function from the R package *PCAmixdata* (Chavent et al. 2017) in R environment. Entered variables were the tree-ring width (*TRW*), the hydraulically weighted vessel diameter (D_h) , the vessel density (D_v) , the specific hydraulic conductivity (K_s) , the distance from the apex (D_{apex}) , the vessel composition (*S*) and the cambial age (*Age*). The study site was set as factor variable into the analysis.

The allometric relationships between the measured traits were analyzed using power scaling model $D_h = \alpha \cdot L^\beta$, where α is the allometric constant and β the scaling exponent. Model

parameters were estimated using the standardized major axes (SMA*s*) with the *smartR* package (Warton et al. 2012). All variables underwent log10 transformation before analysis to satisfy the assumptions of normality and homoscedasticity and to better capture the functional implications of conduit diameter variation (e.g. Kerkhoff and Enquist 2009). This analysis was performed on a maximum common time span and based on data from a moving window of three neighboring tree rings shifted by one to increase the number of axial points and increase the explained deviance of each iteration.

The variability of the α and β coefficients with tree growth was examined using a dual approach, which involved the application of two distinct models for each of the selected predictors. Specifically, the influence of cambial age (*Age*) and annual height increment (*Hincr*) on the variability of both α and β was initially assessed through a linear mixedeffects approach, incorporating tree ID as a random term. The estimation of parameters in linear mixed-effects models utilizing restricted maximum likelihood (REML) was accomplished using the *lmer* function within the *lme4* package (Bates et al. 2015) in the R environment. We then optimized the random-effect structure of the model testing if including extra random-effects terms for tree ID improved the fit of the model; different random structures were then compared through a Likelihood Ratio Test (LRT). Statistical significance (*P <* 0.05) and Satterthwaite approximation for degrees of freedom were obtained from the *lmertest* package and confidence intervals of the estimates were computed using the Wald estimation method.

Additionally, we fit linear quantile mixed models to further explore relationships between variables outside of the mean of the data. This model encapsulates a wider quantile (*τ*) range (from the fifth to the 95th percentile) of the outcome distribution, representing the conditional quantiles of a dependent variable as a linear function of the explanatory variables (R package *lqmm*, v.1.5.8, Geraci 2014). A 95% bootstrap confidence interval for the interquartile regression coefficients was computed with a bootstrap approach (100 replication).

Finally, we integrated all potentially relevant variables, including ontogenetic (i.e. tree age, height and stem diameter increment) and climatic factors (annual temperature and precipitation), into a comprehensive linear mixed model. This model aimed to elucidate the variation in both the allometric constant and the scaling exponent of $D_h \propto L$. The models incorporated tree cambial age (*Age*), annual height increment (*Hincr*), annual increment of stem diameter (*DBHincr*, at breast height), site-specific annual average temperature (*Temp*, ◦C) and cumulated annual precipitation (*Prec*, mm) as fixed effects. Following Zuur et al. (2009), the most parsimonious models were selected using the *lme4* R package (Bates et al. 2015) starting with a saturated model where the fixed component contained all explanatory variables with all possible interactions. Before analysis, all terms were squareroot transformed to achieve normality assumption; then all fixed terms were centered and scaled to improve parameter estimates and allow direct comparisons of the regression coefficients. Nested study site and tree ID were included as random variables. Subsequently, we fine-tuned the random-effect configuration of the model by assessing whether the incorporation of random slopes for site ID and tree ID enhanced the model's fit. Different random structures were compared using an LRT.When comparing saturated models that varied in their random structure but not fixed effects, the models were fit using REML to avoid biased estimators for the variance terms. The final models presented here contains only very limited collinearity (VIFs *<* 1.5). For all models, marginal and conditional \mathbb{R}^2 were calculated to examine the variation explained by fixed and fixed plus random factors, respectively, using the *r.squaredGLMM* function in the *MuMIn* package (Barton and Barton 2015). The residual diagnosis was performed to check the normality and homoscedasticity model assumptions.

To quantify the proportion of variance explained by fixed and random terms we partitioned the variance into four components, i.e. variance attributable to fixed terms, variance attributable to random terms and residual variance using the *r2mlm* R package (Shaw et al. 2023). Then, the standardized regression coefficients and the contribution of fixed effects to the variance of dependent variables was computed via *partR2* (R package *partR2* v. 0.9.1, Stoffel et al. 2021) using 1000 parametric bootstrap iterations to calculate 95% confidence intervals of estimates. The within-tree and between-tree variability of *α* and *β* coefficients as trees grow taller was tested by estimating the repeatability at both the tree (ID) and site levels. Adjusted repeatability estimates after controlling for fixed effects were obtained using the package *rptR* (Stoffel et al. 2017), all with a Gaussian distribution, using 1000 parametric bootstrap iterations and 100 permutations. The statistical significance of repeatability estimates was inferred from whether confidence intervals included zero.

Results

The first two axes of PCA exploring the interactions between the investigated traits explains 42.7% of the total variance in the data set (Fig. 2a). The first axis is strongly (i.e. PCA score *>*0.6) and positively associated with most wood anatomical traits (including *Dh*, *Ktheo* and *S*) and negatively with D_{ν} . The tree age correlates best with the second axis and negatively associated with the tree-ring width (*TRW*). Some common patterns of average vessel area (*A*av), average vessel density (*Dv*) and vessel lumen composition (*S*) appeared along the stem length in both study sites (Fig. 2b, c and d). Particularly, *A*av and *S* decrease with distance from the stem base while *Dv* increases. The average vessel area shows more than a two-fold reduction from stem base to top, despite an unusual hump-shaped pattern at breast height, particularly noticeable at Site1. In contrast vessel density there is a sharp increase in the number of vessels along the xylem pathway from 16 m height. Accordingly, the average vessel area shows an increasing pattern with cambial age across the study sites, while vessel density tends to decrease with cambial age (Fig. 3d and e).

An analysis of xylem traits scaling along the stem axis reveals that the power function ($y = a \cdot L^{\beta}$) provides the best fit to the data (Fig. 3). The hydraulic diameter (D_h) of pooled data per site scaled with the distance to apex reconstructed at annual resolution with scaling exponents *β* of 0.20 and 0.21 for Site1 and Site2, respectively (model statistics are reported in the Table S2 available as Supplementary data at *Tree Physiology* Online). The vessel density (D_{ν}) scaled with scaling exponents β of -0.24 for Site 1 ($R^2 = 0.24$, $P < 0.001$) and -0.26 for Site 2 ($R^2 = 0.38, P < 0.001$) (Fig. S4 available as Supplementary data at *Tree Physiology* Online).

Regarding the D_b –*L* relationship, the β values exhibit a range of variability spanning from 0.10 to 0.36, encompassing the 95% of the sample distribution. Meanwhile, 50% of the values fall within the range of 0.19 to 0.27, with a median of 0.24. Concerning the allometric constant α , the median value is -4.47 . The range extends from -5.51 to -3.71 for the 95% of the sample distribution and from −4.88 to −4.33 for the 50% of the distribution (Fig. S2 available as Supplementary data at *Tree Physiology* Online). This indicates a considerable range of variability in *β* and *α* values among individuals. Notably, for certain cases (i.e. specific annual growth rings), a significant portion of the sampled trees exhibit high values, namely outliers i.e. values of *β* greater than 0.4 and *α* less than -6 .

All the models we run agree that no significant trends of *β* and *α* exists with respect to *Age* and *Hincr* (Fig. 4, Table S3 available as Supplementary data at *Tree Physiology* Online). For both scaling parameters, their mean values and percentiles are not influenced by cambial age and annual height increment, with marginal R-squared values well below 1%. However, the conditional R-squared values indicate a pronounced contribution of individual variability in explaining the total variance, i.e. significant between-tree variation (Table S3 available as Supplementary data at *Tree Physiology* Online).

The models that included all potential predictors of the variability in β and α did not show any significant patterns (Fig. 5 and Tables S4 and S5 available as Supplementary data at *Tree Physiology* Online). Although the percentage of total variance explained by both models was quite high (for the *β* model, R-squared equals 0.70; for the *α* model, R-squared equals 0.61), the variance partitioning highlighted that only 13% of *β* variability and *<*10% of *α* variability can be attributed to fixed factors. This indicates that most of the variability is attributable to random factors, which pertain to differences between and among individuals and between sites (34% and 30% of the total explained variance for *β* and *α*, respectively). Upon detailed examination of the variance explained by the random component, we observed that the component most significantly influencing the variability of *β* and α is almost exclusively attributable to differences among individuals (LRT *P*-value *<*0.001). In contrast, the differences between sites did not show significant contribution (LRT *P*-value *>*0.05, Fig. S6 available as Supplementary data at *Tree Physiology* Online). The within-individual variability is an equally significant component of variance as the betweenindividual variability, accounting for over 56% and 63% of the total explained variance for β and α , respectively. We are aware that due to potential inter-annual variability of growth rates, the precision in the estimate of annual stem elongation rate likely decreased with tree age (due to lower inter-disc distance toward the stem base and likely faster growth commonly occurring during the early ontogeny) and we do not exclude the potential impact this may have had on biasing the patterns we observe, particularly in certain years.

Discussion

In this study, we aimed to characterize axial variation of vessel diameter within *F. sylvatica* trees to assess the consistency of predicted tip-to-base conduit widening values with distance from the tree apex. To do this, we analyzed an unparalleled dataset comprising records of long-term xylem anatomical series from different heights along the stems generated

Figure 2. Biplot of the first two axes of the PCA output (panel a). The pattern of average vessel size (A_{av}) , vessel density (D_v) and vessel lumen composition ($S = A_{av}/D_v$) for the common tree outermost rings along with stem cross-section height (panels b, c and d). Each data point represents the mean value of different sampling heights, while whiskers represent the mean \pm standard deviation of the mean. TRW, tree-ring width; S, vessel lumen composition; Age, cambial age; D_h , hydraulically weighted vessel diameter, D_v vessel density; K_s , specific hydraulic conductivity; L, distance from the stem apex.

through a retrospective reconstruction of the height growth history of 10 mature trees. The negative relationship between vessel density and vessel diameter, consistently observed in our sampled trees, is a well-established phenomenon across woody plants and angiosperms where vessels become smaller with height, leading to increased density (Petit et al. 2010; Lechthaler et al. 2019b). Accordingly, Olson et al. (2020) unveiled a remarkably strong correlation between vessel diameter and density, where vessel density explains a third of the total variation in vessel diameter. This phenomenon occurs because wider vessels occupy a greater proportion of crosssectional area of the stem than narrower vessels, geometrically resulting in the inverse relationship between vessel diameter and density along the stem. The distinctive pattern of vessels to narrow at the base of the stem, as observed for Site1 (Fig. 2*b*), has also been documented in tropical tree species (James et al. 2003; Dória et al. 2019; Li et al. 2019) and even in *Eucalyptus* spp. (Petit et al. 2010; Pfautsch et al. 2018). This trend deviates from the prevailing pattern of continuous

vessel widening from the tip to the base of the tree. While we lack substantial evidence to draw similar conclusions for our angiosperm species, the reasons for basal narrowing of vessels warrants further investigation. Some hypotheses put forth include considerations of xylem construction costs (e.g. Mencuccini et al. 2007) or rather the alleviation of mechanical stress (e.g. Gartner 1995) or invoke a trade-off between fluid dynamic resistance and carbon cost of the conducting system (Koçillari et al. 2021).

Predictability of vessel diameter change with tree height

The results from our long-term stem anatomical series strongly emphasize that vessel diameter follows a predictable pattern with distance from the stem apex. This finding robustly supports the hypothesis that, with height growth, trees do not alter their axial configuration (both *α* and *β* in the equation $D_h = \alpha \times L^{\beta}$ to compensate for hydraulic limitations imposed by tree height during growth. Our

Figure 3. Axial variation of hydraulic vessel diameter (D_h) versus distance from the tree top (L) (a). The thin lines are the predicted value of linear log-log models for each cambial age; the thick lines are the predicted value for Site1 ($\alpha = -4.44$, $\beta = 0.20$, $R^2 = 0.71$, $P < 0.001$) and Site2 ($\alpha = -4.42$, $\beta = 0.21$, $R^2 = 0.55$, $P < 0.001$), respectively. The dashed lines represent the fifth and 95th percentiles. Insets show log–log scaling between variables. Estimated univariate density distribution of *β* scaling exponents with highest density regions (HDR; red 50% and blue 95%) displayed; the vertical full line corresponds to the median value 0.24 (b). Box plots showing *β* scaling exponent variability for sampled trees. Each box represents the 75th to 25th percentiles and the line inside the median; the upper and lower marks are the largest to smallest observation values, which are less than or equal to the upper and lower quartile plus 1.5 the length of the interquartile range; the circles outside the lower-upper mark range are outliers (c). Radial variation of hydraulic vessel diameter (D_h) versus tree cambial age for Site1 (d) and Site2 (e). The thin lines are the predicted value of linear log–log models for collected stem disk.

finding that hydraulic vessel diameter increases with distance from the tree apex is consistent with existing knowledge and supports the hypothesis that the widening of xylem vessels is a result of natural selection, aimed at minimizing the negative impact of path length on resistance along the conductive pathway (Anfodillo et al. 2013). Whatever the mechanisms regulating the vessel diameter dimensions along the stem, i.e. turgor (Woodruff and Meinzer 2011; Cabon et al. 2020) or hormones (Anfodillo et al. 2012; Johnson et al. 2018), such tip-to-base widening profiles are observed in over a hundred species regardless of phylogenetic affinity (including ferns, clubmosses, horsetails and spikemosses), plant heights, growth habit or climate (James et al. 2003; Anfodillo et al. 2006; Mencuccini et al. 2007; Petit et al. 2010, 2023; Williams et al. 2019; Olson et al. 2014; Yang et al. 2021, but refer to Olson et al. 2021 for a comprehensive review). The observed axial widening pattern of *Dh* within the *F. sylvatica* stem appears to be a result of natural selection to favor a mechanism that mitigates the decline in hydraulic conductance with increasing path length. This concept is

Figure 4. Average values (depicted as points) along with 95% confidence intervals (illustrated as error bars) of the scaling exponent *β* (indicated by empty dots) and the allometric constant *α* (on a log scale, represented by full dots). The average values encompass pooled data from all individuals categorized by age (a) and annual height increment rate (d). Individual data can be found in Fig. S5 available as Supplementary data at Tree Physiology Online. The background dots represent the input data and the dashed horizontal lines represent *β* = 0.2. The slope coefficient (shown as dashed lines) and its 95% confidence interval (shaded area) of the estimated linear mixed model regression of the scaling exponent *β* (b, e) and the allometric constant α (c, f). These regressions are presented in relation to the tree cambial age (Age, upper panels) and annual height increment (H_{incr}, lower panels). The full lines and points indicate the slope of the estimated linear mixed quantile regression with respect to the *τ*th quantile *τ* (represented by full circles, each corresponding to the fifth quantile). The shaded area represents the 95% confidence interval (obtained through 1000 replicate bootstrap iterations) for the quantile estimates.

Figure 5. Estimated coefficients (points) and their 95% confidence intervals of the linear mixed model for *β* (upper panels) and *α* (lower panels) (a, d). Cambial age (Age), annual height increment (H_{incr}), annual stem diameter increment (DBH_{incr}), annual average temperature (Temp.) and cumulated annual precipitation (Prec.), * denotes variable interaction. Inclusive iR^2 defined as the variance explained by a predictor irrespective of covariances with other predictors (b, e). Variance components of the linear mixed effects models (c, f) explained by fixed variables (fix.), random variation among trees and between sites (mean var. and slope var.) and the within-tree residual variability (resid.).

consistent with the original theories proposed by the WBE model and the widened pipe model (West et al. 1999; Enquist et al. 2000; Anfodillo et al. 2006; Koçillari et al. 2021) and fully consistent with the patterns found in Petit et al. (2023) for the same species, highlighting that the observed widening pattern minimizes, but does not fully eliminate, the path length resistance.

In our dataset, the power scaling exponent between the hydraulically weighted vessel diameter (*Dh*) and the distance from the stem apex (*L*) mostly felled inside the range 0.15 to 0.33 values reported for various tree species (Anfodillo et al. 2006). Moreover, *β* values reported for both our sites remarkably match with the predictions of hydraulic optimality models, specifically, the expectation of a vessel diameterstem length exponent of 0.2 as supported by empirical measurements by Anfodillo et al. (2006). When compared with other studies on *F. sylvatica* species (i.e. $\beta = 0.23$ in Petit et al. 2023), we detected no significant difference in the *β* scaling exponent, as indicated by the overlapping 95% confidence intervals.

In some cases, we observed widening values significantly exceeding the predicted threshold (i.e. $\beta = 0.31$). This phenomenon, referred to as overwidening ("overtapering" sensu Anfodillo et al. 2006), in vessel structure, has also been documented in other species (e.g. *Fraxinus excelsior* in Anfodillo et al. 2006 and in *Eucalyptus regnans* in Petit et al. 2010) and partly attributed to trees with elevated height increment However, our analyses did not support the hypothesis that high degrees of vessel widening are frequently linked with trees displaying fast height growth in their early ontogeny.

Nevertheless, despite the variability in the *β* exponent within species and even within the same individual (see discussion below), the optimality predictions with respect to widening rates, as supported by empirical evidence, converge on the idea of the "just right" exponent (as per Olson et al. 2021), corresponding to values close to 0.2. Values of *β* close to this threshold, resulting in hydraulic resistance remaining relatively constant with increasing stem length, are likely to be favored by natural selection over other patterns of axial variation of vessel traits as vessels offer an efficient conductivity rate without excessively increasing the risk of embolism or metabolic cost.

The D_{ν} –*L* relationship (Fig. S4 available as Supplementary data at *Tree Physiology* Online) closely mirrored the *Dh*–*L* relationship in that it was well described by the similar power trajectories for all trees. While longer stem length corresponds to higher D_h , the negative relationship between D_v and L was expected given the well-documented geometrical relationship between D_h and D_v (Petit et al. 2010, Olson et al. 2020).

Tip-to-base vessel widening remains stable with height growth

Our results, based on detailed anatomical measurements at the individual level, explicitly demonstrate that vessel widening remains consistent throughout each year of growth, from the ring apex to the stem base. The fact that allometric parameters *α* and *β* remain unchanged as an individual tree grows taller has limited but valuable supporting evidence to date, with only a few studies, including Weitz et al. (2006) on a specimen of *Fraxinus americana*, Prendin et al. (2018) on eight specimens of *Larix decidua* and Petit et al. (2023) on a specimen of *F. sylvatica* and *Picea abies*, providing confirmation. Despite the expected changes in xylem structure

as trees grow, the slope of the relationship D_h –*L* remains essentially constant, suggesting strong developmental control over axial design. This suggests that the vascular system does not adjust to higher cambial age and tree height by modifying its axial scaling components. This lack of increase in either stem apex vessel diameter or tip-to-base widening rate means that the observed widening pattern minimizes, but does not fully eliminate, the path resistance length. However, Petit et al. (2023) have recently shown that while species-specific differences would alter the effects of these residual path length influences at the tree-ring level, the contribution of inner sapwood rings to the total xylem conductance becomes crucial in effectively mitigating the hydraulic limitations imposed by tree height on leaf-specific conductance.

Given the negative relationship between *β* and *α* (Fig. S7 available as Supplementary data at *Tree Physiology* Online), we expected that α of the within-individual tip-to-base widening profile would remain constant as an individual grows taller, indicating an absence of pattern of *α* with height growth. This is consistent with other intraspecific investigations that report only negligible variation with ontogeny, such as Prendin et al. (2018); Williams et al. (2019) and Petit et al. (2023). However, these results are not in line with interspecific outcomes that estimate mean vessel diameter at the distalmost twig from the base becomes predictably wider across angiosperms (Echeverría et al. 2019. Olson et al. 2014, 2020, see below discussion on leaf size effect).

We observed that the widening vessel rate of *F. sylvatica* remains consistent between sites with contrasting climate. This means that the basal vessel size–tree height relationship is chiefly influenced by tree height rather than by climate. This finding is consistent with studies conducted on other tree species across precipitation gradients, which observed no significant changes in the slope and intercept of tip-to-base vessel widening profiles (e.g. seven *Acacia* spp. in Lechthaler et al. 2019b; two *Cedrela* spp. in Chambers-Ostler et al. 2023; *Nothofagus antarctica* in Fajardo et al. 2020, although the latter did not measure axial profiles of xylem vessels). While there were minor differences reported in *Fraxinus* trees growing in soils with varying moisture retention (Kiorapostolou and Petit 2019), temperate conifers have consistently exhibited a constant tip-to-base vessel widening regardless of nutrient availability, temperature (Coomes et al. 2007) and even experimental manipulation of $CO₂$ concentration and soil temperature (Prendin et al. 2018). This suggests that tree size is a more reliable predictor of basal vessel size than climate across various tree species and sizes, as demonstrated by Olson et al. (2014), where stem length explained 63% of the total variation in vessel diameter. However, it is important to note that climate-related variables can still affect intra- and interannual xylem vessel patterns, but these effects should be considered in the context of the causal relationship between plant size and vessel diameter, as suggested by Rosell et al. (2017).

Another piece of evidence supporting the consistency of the widening coefficients between sites emerged when we tested the temporal correlation among individuals from the same site. In accordance with a key assumption of dendrochronology, if the local climate had a predominant effect, we would have expected trees growing in the same site and climate to exhibit similar patterns in the examined traits (Fritts 1976). However, the temporal correlation tests conducted among individuals from the same site revealed only partial or

contrasting significant Pearson correlation coefficient, indicating the absence of common patterns among individuals (Table S6 available as Supplementary data at *Tree Physiology* Online). This further corroborates the legacy of developmental history of individual tree in the stand (Olson et al. 2021). Indeed, individual development is intricately linked to both common site-specific factors (e.g. microclimate, soil, nutrients) and factors driving forest dynamics (competition for resources among trees) that influence phenotypic variation in plant traits. Accordingly, we are aware that certain sitespecific factors, such as nutrient availability (Borghetti et al. 2017) and water storage capacity (Tokumoto et al. 2014), possible variability randomly introduced by reconstructing tree height at annual resolution, as well as genetic background (Eilmann et al. 2014), which were not considered in our analysis, may partially affect the interpretation of our results.

We demonstrate that within-tree variation is a substantial, rather than a minor, contributor to the overall variance of the widening exponent. Numerous studies have highlighted that certain species exhibit intra-specific variations in xylem traits in response to site-specific or environmental conditions, while other research has shown limited intra-specific differences (Herbette et al. 2010; Schreiber et al. 2015; Hajek et al. 2016; Baer et al. 2021; Weithmann et al. 2022), although a large body of it does not denote standardized sampling approaches (see Petit et al. 2022 for a critique on this matter). Yet, significant variability in xylem traits can exist within an organism, both along its main plant axis and between its roots and stems (Jacobsen et al. 2018; Prendin et al. 2018; Rodriguez-Zaccaro et al. 2019; Baer et al. 2021). For instance, Baer et al. (2021) investigated intra-organismal variability in xylem traits such as density, biomechanics and water storage within a model angiosperm species, revealing that these variations arise in response to different biomechanical stresses on shoots and roots, as well as varying root-leaf water potential gradients within the plant. This leads to trade-offs within the xylem of a single individual, particularly between biomechanical and water-storage functions. Hence, in response to selective pressures from the surrounding environment, trees can exhibit significant intra-specific and intraindividual differences in xylem traits, potentially leading to systematic variations in the vessel diameter–plant height relationship within the boundaries of developmental possibilities (Olson et al. 2021). Within certain limits, authors concur that the vessel diameter-stem length exponent is not constant, suggesting the presence of systematic developmental variations that allow different slopes and intercepts (Rosell et al. 2017; Olson et al. 2021). Currently, there are emergent hypotheses indicating that features such as leaf phenology, wood density, porosity type and perforation plate type are linked to significant variations in vessel scaling slope and intercept that have adaptive implications (see Olson et al. 2020 for an overview). For example, there is every reason to believe that leaf size could play a crucial role in this context, particularly in determining the diameter of the conductive vessels in the distal parts of the branches (Lechthaler et al. 2019a; Cao et al. 2022), likely influencing the tip-to-base conduit widening profile in the stem. Additionally, some evidence suggests that taller individuals within a species may exhibit slightly lower tip-to-base scaling exponents (Mencuccini et al. 2007; Petit et al. 2008) to balance the need for efficient hydraulic transport with the constraints of carbon cost. Collectively, these studies indicate that there exists room for selection to influence not only narrower but also wider vessels than

typically observed for a given stem length (Olson et al. 2021). Within the framework of vessel scaling, individuals within a population that deviate significantly from the typical scaling pattern, either above or below, are expected to exhibit reduced performance or fitness compared with those individuals that closely adhere to the common scaling pattern (Anfodillo et al. 2016; Anfodillo and Olson 2021). Considering the potential functional significance of this variation, gaining a comprehensive understanding of factors contributing to intraindividual variation in widening exponent is a priority in comprehending how plants adapt and acclimate to climaterelated disturbances.

Conclusion

In our study, we explored axial variation in xylem vessel characteristics in a diffuse porous tree species to assess the consistency of theoretical predictions regarding the power scaling widening exponent of the relationships $D_h \propto L$. We examined 10 *F. sylvatica* L. trees from two contrasting climate sites. Although we identified a steady scaling exponent with height growth and between study sites there was still inherent variability in tip-to-base vessel widening among and within individual trees. Our findings emphasize that the primary source of variance in the relationship between tree height and hydraulic vessel diameter is the variation within and between trees. These results highlight our current lack of understanding regarding the origins of variation in the scaling exponent or the y-intercepts, indicating these as critical areas for future research. Given the potential functional importance of this variation, it is crucial to deepen the understanding of factors that contribute to the variation in intraindividual widening exponent in relation to plants' adaptation and acclimation to climate-related disturbances.

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Authors' contributions

A.R. designed and conducted the study and wrote the manuscript; O.P. and T.G. carried out the measurements; A.R. and J.T. performed the analysis. All authors contributed to discussing and interpreting the data throughout the study.

Supplementary data

Supplementary data are available at *Tree Physiology* Online.

Conflict of interest

The authors declare no conflict of interest.

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Data availability

The data and materials that support the findings of this study are available from the corresponding author upon reasonable request.

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