

Disentangling drought-responsive traits with focus on Arabidopsis

Thonglim, A.

Citation

Thonglim, A. (2023, November 9). *Disentangling drought-responsive traits with focus on Arabidopsis*. Retrieved from https://hdl.handle.net/1887/3656528

Version: Publisher's Version

Licence agreement concerning inclusion of doctoral

License: thesis in the Institutional Repository of the University

of Leiden

Downloaded from: https://hdl.handle.net/1887/3656528

Note: To cite this publication please use the final published version (if applicable).

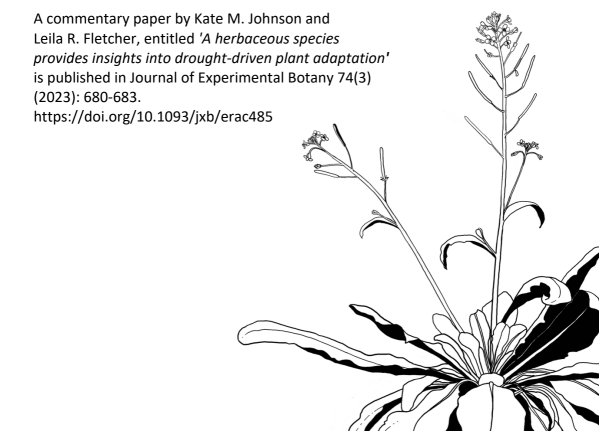
Chapter 3

Drought response in Arabidopsis displays synergistic coordination between stems and leaves

Ajaree Thonglim^{1,*}, Giovanni Bortolami¹, Sylvain Delzon², Maximilian Larter², Remko Offringa³, Joost J.B. Keurentjes⁴, Erik Smets^{1,5}, Salma Balazadeh^{6,†} and Frederic Lens^{1,5,†,*}

Adapted from

Journal of Experimental Botany 74(3) (2023, first published online in 2022): 1004-1021. https://doi.org/10.1093/jxb/erac446



 $^{^{1}}$ Naturalis Biodiversity Center, Research Group Functional Traits, PO Box 9517, 2300 RA Leiden, The Netherlands

 ² BIOGECO INRA, Université Bordeaux, 33615 Pessac, France
 ³ Leiden University, Institute of Biology Leiden, Plant Developmental Genetics, Sylviusweg 72, 2333 BE Leiden, The Netherlands

⁴ Laboratory of Genetics, Wageningen University, Droevendaalsesteeg 1, 6708 PB Wageningen, The Netherlands

⁵ Leiden University, Institute of Biology Leiden, Plant Sciences, Sylviusweg 72, 2333 BE Leiden, The Netherlands

⁶ Leiden University, Institute of Biology Leiden, Molecular Plant Stress Biology, Sylviusweg 72, 2333 BE Leiden, The Netherlands

^{*}For correspondence. E-mail ajaree.thonglim@naturalis.nl and Frederic.lens@naturalis.nl

†Shared last author

Abstract

The synergy between drought-responsive traits across different organs is crucial in the whole-plant mechanism influencing drought resilience. These organ interactions, however, are poorly understood, limiting our understanding of drought response strategies at the wholeplant level. Therefore, we need more integrative studies, especially on herbaceous species that represent many important food crops but remain underexplored in their drought response. We investigated inflorescence stems and rosette leaves of six Arabidopsis thaliana genotypes with contrasting drought tolerance and combined anatomical observations with hydraulic measurements and gene expression studies to assess differences in drought response. The soc1ful double mutant was the most droughttolerant genotype based on its synergistic combination of low stomatal conductance, largest stomatal safety margin, more stable leaf water potential during non-watering, reduced transcript levels of drought stress marker genes, and reduced loss of chlorophyll content in leaves, in combination with stems showing the highest embolism resistance, most pronounced lignification, and thickest intervessel pit membranes. In contrast, the most sensitive Cvi ecotype shows the opposite extreme of the same set of traits. The remaining four genotypes show variations in this drought syndrome. Our results reveal that anatomical, ecophysiological, and molecular adaptations across organs are intertwined, and multiple (differentially combined) strategies can be applied to acquire a certain level of drought tolerance.

Keywords: *Arabidopsis thaliana*, chlorophyll content, drought response, embolism resistance, gene expression, intervessel pit membrane thickness, stem anatomy, stomatal control.

Introduction

The increasing intensity and frequency of drought episodes are becoming major threats to current and future agricultural productivity around the globe. Even the countries that had not experienced drought stress during the last decades are now impacted by drought (Corso et al., 2020; Gleason et al., 2022). One of the major problems that plants experience when they are facing severe drought is that detrimental levels of drought-induced gas bubbles (embolisms) in the xylem sap generate massive obstruction of the root to shoot water transport (Sperry and Tyree, 1988; Tyree and Zimmermann, 2002; Cochard, 2006; Choat et al., 2012; Venturas et al., 2017; Johnson et al., 2022), which happens after stomata are closed (Martin-StPaul et al., 2017). Stomatal closure may result in reduced photosynthetic productivity, growth rate, and reproduction, and under conditions of intense and prolonged drought may eventually cause desiccation and dieback of tissues (Mantova et al., 2022), organs, and entire plants (Davis et al., 2002; Venturas et al., 2016; Pratt et al., 2020; Brodribb et al., 2021). Lethal levels of embolism, from which plants are unable to recover, are thought to be reached when the hydraulic conductivity is reduced to ~88% of its maximum conductance (P_{88}) (Urli et al., 2013; Li et al., 2015; but see Hammond et al., 2019; Johnson et al., 2021), although there are probably more accurate thresholds to drought-induced mortality than P₈₈ (Mantova et al., 2021, 2022). Due to the implications of dramatic levels of drought-induced embolism on productivity, tissue death, and longterm survival, there is increasing evidence that natural selection has shaped the hydraulic systems of plants to minimize embolism occurrence and water potential loss during periods of water shortage (Lens et al., 2022). This can be made possible when many drought-related traits from different organs act in concert (Dayer et al., 2022).

As an example, angiosperms can build more resistant xylem by modifying a whole array of xylem anatomical adaptations to prevent the spread of embolisms, such as fine-scale modifications of pits in vessel walls allowing lateral transport of water and gas between adjacent vessels (Lens et al., 2011; Li et al., 2016; Kaack et al., 2019, 2021; Levionnois et al., 2021), or increased levels of lignification (Lens et al., 2013, 2016; Thonglim et al., 2021). In addition, plants can also delay xylem sap pressures from reaching critical embolism thresholds throughout the whole-plant body by producing

the stress hormone abscisic acid (ABA) that induces stomatal closure in the leaves very rapidly at the onset of drought, well before embolism events start to exponentially increase (Brodribb et al., 2017; Martin-StPaul et al., 2017; Buckley, 2019; Creek et al., 2020). Consequently, stomatal closure is one of the primary responses that helps restrict water loss, which safeguards the water potential in the leaves and buffers the negative pressure in xylem sap (Brodribb et al., 2017; Martínez-Vilalta and Garcia-Forner, 2017; Martin-StPaul et al., 2017; Knipfer et al., 2020). The regulation of water potential in leaves during drought is crucial because it influences plant metabolic processes. However, declining transpiration rates reduce not only water loss but also carbon uptake, leading to decreased photosynthetic activity, which ultimately may lead to carbon starvation when stomata remain closed for a long time (McDowell et al., 2008). In other words, the interplay between embolism resistance inside the plant's xylem and the onset and duration of stomatal closure at the level of leaves will determine how long leaves can remain metabolically active without risk of detrimental levels of droughtinduced embolism (Allen et al., 2010; Choat et al., 2012; Mitchell et al., 2013; Brodribb et al., 2017; Martínez-Vilalta and Garcia-Forner, 2017; Buckley, 2019; Creek et al., 2020; Limousin et al., 2022). Accordingly, the stomatal safety margin (SSM), which can be defined as the difference between the water potential at stomatal closure (Ψ_{gs90}) and the pressure inducing 50% loss of hydraulic conductance (P_{50}) is physiologically more important to estimate a plant's ability to cope with massive levels of drought-induced embolism than only P₅₀ (Sperry and Tyree, 1988; Meinzer et al., 2009; Anderegg et al., 2016; Martin-StPaul et al., 2017; Creek et al., 2020; Dayer et al., 2020; Skelton et al., 2021). It is widely accepted that species with a narrower safety margin are operating more closely to their hydraulic threshold, while species that have a wider safety margin have a lower risk of facing a detrimental level of drought-induced embolism (Choat et al., 2012; Anderegg et al., 2016; Martin-StPaul et al., 2017; Eller et al., 2018; Creek et al., 2020; Oliveira et al., 2021; Skelton et al., 2021).

It is clear that anatomical and physiological traits need to be intertwined within and among organs, but the molecular mechanisms cross-linking different pathways remain elusive. For instance, there is increasing evidence from gene expression studies confirming the positive correlation between lignification and drought resilience in a whole range of species (Tu et al., 2020; Xu et al., 2020; Wen et al., 2021; Yan et al., 2021; Hou et al.,

2022; Li et al., 2022). Regarding drought responses in plants, the ABAmediated signalling pathway is probably the best-known pathway at the molecular level. ABA regulates the expression of stress-responsive genes via transcription factors (Bauerle et al., 2004; Cutler et al., 2010; Bauer et al., 2013; Dodd, 2013; Mehrotra et al., 2014; Chen et al., 2020). Once ABA is accumulated, it regulates ABA-responsive genes via the cis-element called ABRE (ABA-responsive element) in their promoter regions using AREB (ABRE binding) transcription factors (Choi et al., 2000; Uno et al., 2000; Yoshida et al., 2015; Chen et al., 2020). In Arabidopsis, AREB1 is mainly expressed in vegetative tissues and up-regulated during drought (Yoshida et al., 2010; Fujita et al., 2011, 2013; Singh and Laxmi, 2015; Chen et al., 2020). Other drought-responsive genes are regulated by dehydration-responsive element-binding (DREB) proteins through an ABA-independent pathway (Bartels and Sunkar, 2005; Sakuma et al., 2006; Song et al., 2018). For example, DREB2 transcription factors are induced by dehydration and are involved in gene transcription under water shortage (Agarwal et al., 2006; Song et al., 2018). Interestingly, many stress-inducible genes contain both ABREs and DREs in their promoter regions, such as Responsive to Desiccation 29 (RD29) (Shinozaki and Yamaguchi-Shinozaki, 2007). Hence, gene expression of drought-responsive genes occurs via ABA-dependent and/or ABA-independent signal transduction pathways (Umezawa et al., 2010; Rushton et al., 2012; Song et al., 2016), and allows us to evaluate the expression of drought-responsive genes during a drought experiment with a simultaneous assessment of physiological and anatomical traits involved in drought tolerance.

Most studies investigating drought-induced embolism in plants have been focusing on trees, while herbaceous plants have been largely ignored despite their importance as crops and food sources for humans and animals (Brodribb and Hill, 1999; Stiller and Sperry, 2002; Holloway-Phillips and Brodribb, 2011; Choat *et al.*, 2012; Ahmad, 2016; Lens *et al.*, 2016; Volaire *et al.*, 2018). In our previous study on the herbaceous model species, *Arabidopsis thaliana*, including genotypes with contrasting levels of embolism resistance and lignification in the inflorescence stems (Thonglim *et al.*, 2021), we found that the more lignified genotypes are more resistant to embolism and have thicker intervessel pit membranes. Surprisingly, in most structure—function studies published so far, the drought response is only partly observed due to methodological and time constraints. For

instance, resistance to embolism in branches/twigs is often recorded in xylem physiological studies (e.g. Choat et al., 2012; Anderegg et al., 2016), and less frequently integrated with leaf P₅₀ data (e.g. Cochard et al., 2004; Klepsch et al., 2018; Skelton et al., 2019; Levionnois et al., 2021) and/or root P₅₀ data (e.g. Rodriguez-Dominguez et al., 2018), and sometimes linked with other leaf physiological traits such as stomatal conductance (q_s) and water potential (e.g. Brodribb et al., 2003; Li et al., 2015; Cardoso et al., 2018; Charrier et al., 2018; Creek et al., 2020; Chen et al., 2021). Only occasionally are detailed hydraulic measurements in stems, leaves, and/or roots complemented with detailed anatomical traits on intervessel pits (Guan et al., 2022). Other papers only focus on the molecular pathway and gene regulation during drought (e.g. Bhargava and Sawant, 2013; Pandey et al., 2013; Janiak et al., 2016; Ebrahimian-Motlagh et al., 2017; Thirumalaikumar et al., 2018; Roca-Paixão et al., 2019; Zhang et al., 2019), while publications that integrate gene function with xylem physiology are scarce (e.g. Kitin et al., 2010; Lamarque et al., 2020). Integration of drought-related traits across organs in structure-function studies and intensive collaboration among plant anatomists, xylem physiologists, and molecular biologists will help us to make considerable progress in a holistic understanding of drought response at the whole-plant level. To contribute to that whole-plant approach, we measured hydraulic traits in stems and leaves during a experiment, combined with detailed stem anatomical measurements and an assessment of transcript levels of drought stress marker genes across Arabidopsis genotypes (two transgenic lines and four natural accessions).

In this study, we investigate the following two questions. (i) Is there a coupling between drought-related stem (anatomy, P_{50}) and leaf traits (stomatal regulation, leaf water potential, expression of drought marker genes) among Arabidopsis genotypes? (ii) Can these genotypes use different combinations of drought-response traits to reach a certain level of drought tolerance? To answer these questions, we investigated six genotypes with marked differences in embolism resistance and lignification of the inflorescence stems. We examined the detailed stem anatomical traits and hydraulic traits (stem P_{50}) of each genotype and quantified the drought response for all six genotypes using a drought experiment, during which we measured g_{5} and leaf water potential (ψ_{1}), allowing us to calculate the SSM (as defined by ψ_{g590} minus P_{50}). In addition, we compared the expression of

four drought-responsive genes from the ABA-(in)dependent (ABI2, AREB1, RD29A, and DREB2A) pathways from the rosette leaves at the end of the drought experiment to validate the level of drought stress among the six genotypes. By integrating all traits mentioned above, we want to assess how anatomical and ecophysiological traits across organs are intertwined to acquire a certain level of drought tolerance, and how these traits relate to the drought stress level at the end of the drought experiment based on a limited number of drought stress marker genes.

Materials and methods

Plant material

In addition to the four A. thaliana genotypes with contrasting levels of stem P₅₀ and stem lignification, we studied before the ecotypes Columbia-0 (Col-0; wild type with intermediate stem lignification), Shadarah (Sha; wild type with a higher level of stem lignification), Cape Verde Islands (Cvi; least lignified wild type), and the double loss-of-function mutant SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1 and FRUITFULL (soc1ful; most lignified genotype) (see Thonglim et al., 2021); we added one more wild type [Kelsterbach-4 Kel-4)] and a p35S:AHL15 line (AHL15 overexpression) in the Col-0 background (Rahimi et al., 2022). The two additional genotypes were selected based on their inflorescence length (at least 27 cm required for the centrifuge method used to estimate embolism resistance measurements) and their increased lignification in the basal parts of the inflorescence stem, respectively (Supplementary Figures S1A, B, G, H). Indeed, Kel-4, an early flowering ecotype from Germany, shows a relatively high proportion of lignification at the base of the inflorescence stem (Ak, 2020), and has been reported to be more drought tolerant compared with many other wild-type accessions (Bac-Molenaar et al., 2016; Kooke et al., 2016). The AT-HOOK MOTIF CONTAINING NUCLEAR LOCALIZED 15 (AHL15) gene has been found to suppress axillary meristem maturation, and its overexpression extends plant longevity (Karami et al., 2021), and promotes secondary growth in the inflorescence stem to a similar extent as the soc1ful mutant (Rahimi et al., 2022).

Growing conditions

The plants were grown at the Institute of Biology Leiden (Leiden University, The Netherlands) under the same controlled conditions as in Thonglim *et al.* (2020) to ensure comparable datasets. Briefly, we germinated the two additional genotypes from seeds directly into a mixture of soil and sand (4.5:1). After 10 d of germination, the healthy seedlings were transferred into pots. Plants were grown in a controlled growth chamber with the following parameters: 20 °C temperature during the day and 17 °C temperature at night, 70% relative humidity, and 16 h photoperiod condition with 100 μ mol m⁻² s⁻¹ light intensity. Sampling was synchronized based on differences in flowering time and subsequent inflorescence development. To synchronize flowering, *p35S:AHL15* plants were planted earlier (harvesting inflorescence stems 85 d after sowing). The Kel-4 individuals were planted slightly later (harvesting inflorescence stems 65 d after sowing) (Supplementary Figure S1A, B).

Drought experiment

A drought experiment was performed to assess the link between the anatomical and hydraulic traits and investigate the differences in drought tolerance across the six A. thaliana genotypes studied. The six genotypes were selected based on a previous screening of drought tolerance and the differences in stem lignification (Melzer et al., 2008; Bac-Molenaar et al., 2016; Thoen et al., 2017; Thonglim et al., 2021). The seeds of each genotype were directly sown in 6 cm pots (27 g) with the same amount of soil and sand mixture (4.5:1) at different times to synchronize flowering. The weight of the pot with dry and saturated soil was controlled (807 g and 1097 g, respectively). The pots were kept in a growth-controlled chamber under the same conditions as the individuals grown for stem P_{50} measurements. After germination, when seedings were 10 d old, they were thinned to one healthy seeding per pot and remained well watered. We equally divided 30 individuals of each genotype into a control and a drought batch during the experiment. The control plants were well irrigated every day to keep the soil constantly hydrated ($\Psi_{\rm I}$ was around -0.5 MPa to -0.6 MPa). The drought batch was subjected to water deficit by completely withholding watering for 3 weeks ($\Psi_{\rm I}$ values ranged between -1.85 MPa to -3.4 MPa among genotypes), starting 1 week before all the genotypes began to flower. When most genotypes started developing an inflorescence stem (7 d after watering was stopped), drought measurements were initiated. Rosette leaves were harvested on the last day of the drought experiment (depending on the water potential and phenotype), immediately frozen io liquid nitrogen, and stored in a –80 °C freezer for further gene expression and chlorophyll analyses. We initially intended to have three biological replicates per genotype. However, during sample preparation, some tubes containing ground leaf material popped open in the freezer. We assume that some liquid nitrogen used for grinding the samples was still left in the tubes, causing several closed tubes to burst open and potentially contaminate the other open tubes containing different genotypes. We opted to discard all the open tubes due to potential contamination and use only the closed tubes. We were able to still use three biological replicates for Cvi, Sha, and soc1ful, but only two for Col-0, Kel-4, and p35S:AHL15. For the latter genotypes, we included two biological and two technical replicates.

Chlorophyll content

Chlorophyll content was determined based on three biological replications for Cvi, Sha, and soc1ful, and four replicates (two biological and two technical) for Col-0, Kel-4, and p35S:AHL15, using the 80% acetone method (Porra et~al., 1989). Ground leaf samples of ~0.5 mg was transferred into 1.5 ml tubes containing 1 ml of 80% acetone. The mixtures were gently vibrated using a vortex to extract chlorophyll and centrifuged at 1000 g for 5 min to remove debris. The supernatants (800 μ l) were then transferred to UV-transparent microplates. The absorbance was measured at 647 nm (A_{647}), 664 nm (A_{664}), and 750 nm (A_{750}) using the DMF-chl conc._YU program. Chl g and g contents (g Chl mlg) in the extract were calculated with the following formulas:

Chl
$$a = (12.25 \times (A_{664} - A_{750}) - 2.85 \times (A_{647} - A_{750})) / 0.29$$

Chl $b = (20.31 \times (A_{647} - A_{750}) - 4.91 \times (A_{664} - A_{750})) / 0.29$
Total Chl $(a+b) = (17.76 \times (A_{647} - A_{750}) + 7.34 \times (A_{664} - A_{750})) / 0.2$

RNA isolation and qRT-PCR

Total RNA was extracted using the RNeasy Plant Mini kit (Qiagen, Hilden, Germany). Synthesis of cDNA, quantitative reverse transcription—PCR (qRT—PCR) using SYBR Green, and data analysis were performed as previously described (Balazadeh *et al.*, 2008). Gene expression was normalized with two reference genes (*ACTIN2* and *GADPH*). qRT—PCR primers were designed using QuantPrime (www.quantprime.de) (Arvidsson *et al.*, 2008). Primer sequences are given in Supplementary Table S1. Experiments were conducted in three biological replications for Cvi, Sha, and *soc1ful*, and two biological replicates with two technical replicates for Col-0, Kel-4, and *p35S:AHL15*.

Leaf water potential (Ψ_l) and stomatal conductance (g_s)

After 7 d of water deficit (i.e., the time required to dehydrate the moisturized soil in the pots of the drought batch), ψ_{l} was measured in both control and drought batches every day during the drought period until harvesting (15–17 d). The daily measurements were carried out using three mature leaves (one from control and two from drought treatment) for each method. Before the measurements, the leaves were covered with aluminium foil for 30 min. Subsequently, leaf discs were cut from the bagged leaves and placed in the PSYPRO leaf water potential system (Wescor, Inc., Logan, UT, USA) to measure the leaf water potential. At the same time, q_s (mmol H₂O m⁻² s⁻¹) was measured on single mature rosette leaves that were close to the leaves used for water potential measurements, using an SC-1 leaf porometer (METER Group, Pullman, WA, USA) that was calibrated every other day. The q_s was measured using Auto Mode configuration with desiccant. g_s , depending on leaf water potential, was fit according to the following sigmoid function for each genotype using the NLIN procedure in SAS:

$$g_{\rm s} = g_{\rm sm} \div \left[1 + \exp\left(S \times (\Psi - \Psi g s_{50})\right) \right]$$

 g_{sm} is the maximal stomatal conductance for Ψ_{I} =0, S is the slope of the curve, and Ψ_{gs90} the water potential inducing 50% stomatal closure. We then estimated the water potential inducing 90% of the stomatal closure (Ψ_{gs90}).

Stomatal safety margin (SSM)

The SSM was defined as the difference between the leaf water potential at 90% stomatal closure (Martin-StPaul *et al.*, 2017) calculated from the fitted curve ($\Psi_{\rm gs90}$) and the water potential at 50% loss of stem conductivity ($P_{\rm 50}$):

$$SSM = \Psi_{gs90} - P_{50}$$

Generating vulnerability curves (VCs) in stems

Sample preparation of inflorescence stems

All individuals (80 individuals per genotype) were harvested at the Institute of Biology Leiden with roots, leaves, and flowers still attached and immediately wrapped in wet tissue papers. They were then enclosed in plastic bags to avoid dehydration during the shipment to the PHENOBOIS platform (INRAE, University of Bordeaux, France), where the Cavitron centrifuge measurements were performed. Before the Cavitron measurements, the roots were cut off at the basal part of inflorescence stems and trimmed on both sides, obtaining a stem segment of 27 cm in length that matches a standard Cavitron rotor. The length of the stem segments exceeds by far the maximum vessel length of Col-0, reaching only 4 cm according to Tixier *et al.* (2013) to avoid potential open-vessel artefacts (Cochard *et al.*, 2013). Next, all siliques, leaves, and flowers were removed underwater immediately before placing the inflorescence stems in the Cavitron rotor (7–9 stem segments per VC).

Xylem vulnerability to embolism was evaluated using the Cavitron method, a custom-built centrifuge that allows measuring the water flow through the inflorescence stems while spinning them to create a negative pressure in the middle part of the stem segments (Cochard, 2002; Cochard et al., 2005, 2013). The negative pressure was gradually increased in each spinning step, as described in Thonglim et al. (2020). The degree of embolism in the xylem segment was quantified as the percentage loss of conductivity (PLC), calculated as follows:

$$PLC = 100 \times \left(1 - (K/K_{MAX})\right)$$

where K_{MAX} (m² MPa⁻¹ s⁻¹) is the maximum hydraulic conductivity which was calculated when stem segments were fully functioning (no embolism) at low spinning speed (near 0 MPa), and K is the decreased hydraulic conductivity due to embolisms. The extent of embolism formation at every rotation speed was measured using the Cavisoft software (Cavisoft v1.5, University of Bordeaux, France). We fitted the data points to reconstruct the VCs using a sigmoid function based on the NLIN procedure in SAS 9.4 (SAS Institute, Cary, NC, USA) (Pammenter and Van der Willigen, 1998):

$$PLC = 100 \div \left[1 + \exp\left(\frac{S}{25} \times (P - P_{50})\right) \right]$$

where P is the xylem pressure used at each rotation step, P_{50} is xylem pressure inducing 50% loss of hydraulic conductivity, and S (MPa⁻¹) is the slope of the VC at P_{50} . Due to the low hydraulic conductivity of Arabidopsis, we measured vulnerability to embolism of 7–9 inflorescence stems to generate one vulnerability curve. Eight VCs were constructed for each genotype.

Stem anatomy

Three stems from three representative VCs per genotype (nine stems per genotype) were randomly selected for light microscopy (LM) observations and one stem per VC from three VCs (three individuals per genotype) for TEM observations (Supplementary Figure S1C, D). Both basal and central parts of the 27 cm inflorescence stem segments were sectioned because they differ in the amount of lignification (Supplementary Figure S1E–H). We, however, invested more time in measuring trait data from the middle part than in the basal segment because that is the region where the negative pressures were applied during the Cavitron experiments, allowing us to accurately link the anatomical traits with embolism resistance (P_{50}). The anatomical traits are represented in Supplementary Table S2. ImageJ (National Institutes of Health, Bethesda, MD, USA) was used, and the guidance of Scholz et al. (2013a) was followed to measure the anatomical features in digital images from both LM and TEM observations.

Light microscopy (LM)

Inflorescence stems were cut into 1 cm long pieces and submerged in 70% ethanol. The samples were then gradually infiltrated in LR-white resin (Hamann *et al.*, 2011). After embedding in LR-white, specimens were sectioned with a rotary microtome (Leica RM 2265, Leica, Eisenmark, Wetzlar, Germany) with disposable tungsten carbon blades at 4 µm thickness. Next, the sections were heat-fixed onto the slides, stained with 1% (w/v) toluidine blue (VWR Chemicals BDH®, Radnor, PA, USA), and mounted with DPX new-100579 mounting medium (Merck Chemicals, Amsterdam, the Netherlands). Finally, various anatomical traits (Supplementary Table S2) were observed using a Leica DM2500 light microscope equipped with a Leica DFC-425 digital camera (Leica microscopes, Wetzlar, Germany).

Transmission electron microscopy (TEM)

The middle parts of inflorescence stem segments were collected immediately after Cavitron measurements and fixed in Karnovsky's fixative (Karnovsky, 1965). Subsequently, the samples were washed in 0.1 M cacodylate buffer and post-fixed with 1% buffered osmium tetroxide. The samples were then prepared for semi-thin and ultra-thin sectioning according to the protocol described in Thonglim *et al.* (2020) and were observed with a JEM-1400 Plus TEM (JEOL, Tokyo, Japan) with an 11 megapixel digital camera (Quemesa, Olympus). TEM observations were conducted to measure the intervessel pit membrane thickness and the pit chamber depth (Supplementary Table S2).

Statistical analysis

R version 3.6.3 in R Studio version 1.2.5033 was used for the statistical analyses of all traits studied, of which all the differences were considered significant when the P-value was <0.05. First, general linear models with a Newman-Keuls post-hoc test were used to check the differences in embolism resistance (P_{50} , P_{12} , and P_{88}), anatomical features, leaf physiological traits, chlorophyll content, and gene expression among Arabidopsis genotypes studied. Then, multiple linear regression was applied to assess the anatomical traits (predictive variables) that explain the differences in embolism resistance (responsive variables, including P_{50} , P_{12} , and P_{88}). The collinearity between variables was firstly checked to select the predictors. Then, the 'step' function (stats package; R Core Team, 2016) was applied to achieve the most parsimonious linear regression model based on the least Akaike information criterion (AIC). Subsequently, the model's residuals, heteroscedasticity, skewness and kurtosis, and variance inflation factor (VIF) were checked. Once we obtained the best model, the relative importance of each explanatory variable was analysed to assess the variable that explains the best P_{50} . Pearson's correlation was applied to plot the relationship between P₅₀ and predictive variables and leaf physiological traits, and among the variables. Lastly, we investigated whether the different Arabidopsis genotypes presented different q_s in well-watered control conditions using a generalized linear mixed model with the accession as a fixed effect, with the GLIMMIX procedure in SAS software (SAS 9.4; SAS Institute).

Gene codes

Arabidopsis gene codes are: *ACTIN2*, *AT3G18780*; *GAPDH*, *AT1G13440*; *RD29A*, *AT5G52310*; *ABI2*, *AT5G57050*; *AREB1*, *AT1G45249*; and *DREB2A*, *AT5G05410*.

Results

Drought-response phenotyping, chlorophyll content, and expression of drought-responsive genes in the basal rosette leaves

After 3 weeks of non-watering, we found differences in phenotypes of the drought-treated batch compared with the well-watered controls. The soc1ful mutant and the p35S:AHL15 overexpression line were least affected by drought based on the rosette phenotype (less wilting of leaves, less reduction of rosette size) and the small reduction of chlorophyll content when compared with the control individuals. The droughted individuals of Sha showed intermediate phenotypic drought stress-related signs compared with the control batch, such as a minor reduction in leaf rosette size, more wilting of leaves, and a slightly higher decrease of chlorophyll content (Figure 1A, B). In contrast, the rosette leaves were more reduced in size in the droughted individuals of Col-0, Kel-4, and Cvi compared with the well-watered control plants (Figure 1A); likewise, leaves and inflorescence stems in the droughted batch of these three genotypes were considerably more wilted compared with the control plants (Figure 1A), along with the stronger chlorophyll reduction in the rosette leaves (Figure 1B). With regards to Chl b reduction during the drought experiment, two significantly different genotype groups could be defined: one group comprising Col-0, Cvi, and Kel-4 (62%, 67% and 46% reduction, respectively) and the other comprising Sha, soc1ful, and p35S:AHL15 (31, 13, and 27% reduction, respectively) (F=15.83, P=0.00212). For Chl α reduction, significant differences were detected among the genotypes (F=181.6, P=1.84e⁻⁰⁶), except for soc1ful and p35S:AHL15 that presented a similar reduced value (10% and 12% reduction). This is also the case for total chlorophyll (Chl a+b) reduction (F=168.1, $P=2.32e^{-06}$) (Figure 1B).

In order to estimate how each Arabidopsis genotype senses drought stress at the molecular level, we measured the expression of four selected drought marker genes at the end of the 15–17 d drought treatment. In the ecotypes with an intermediate level of stem lignification (Col-0 and Kel-4) and the one with the least lignified stems (Cvi), all four drought-responsive genes were up-regulated under drought compared with well-watered conditions (Figure 1C). In contrast, of the four drought-response genes in the more lignified genotypes Sha, overexpression p35S:AHL15, and soc1ful

were significantly less induced under drought treatment. Interestingly, p35S:AHL15 showed no difference in ABI2 and AREB1 expression level between drought and control conditions (-0.45 and -1.37 log2 fold change, respectively). Regarding the changes in the expression of each gene between drought and control conditions among genotypes studied, we found that the change of RD29A expression was similar between Col-0 and Cvi (~6.9 log2 old change). Still, these two genotypes were significantly different from the rest (2.8–4.7 log2 fold change) (F=10.2, P=0.00021). For DREB2A, two significantly different groups were defined: one comprising Col-0, Cvi, and Kel-4 (4.55, 5.6, and 5.57, respectively) and the other comprising Sha, soc1ful, and p35S:AHL15 (3.37, 2.75, and 2.87, respectively) (F=21.05, P=2.71e⁻⁰⁶). The changes of AREB1 were significantly different among genotypes (F=13.28, P=4.63e⁻⁰⁵), except for Col-0, Cvi, and Kel-4 (3.48, 3.22 and 3.19 log2 fold change, respectively). Likewise, for ABI2, there was a significant difference among genotypes (F =40.95, $P=3.2e^{-08}$), except for Col-0 and Kel-4 (6.22 and 5.93), and Sha and soc1ful (4.57 and 3.58 log2 fold change) (Figure 1C).

Leaf water potential (Ψ_i) and stomatal conductance (g_s) dynamics during drought

 $\Psi_{\rm I}$ under well-watered conditions was similar in every genotype, ranging between -0.5 MPa and -0.6 MPa (Figure 2A). However, g_s of control plants was significantly different among the genotypes studied (F=236.12, P<0.0001, Figure 2B). Cvi (least lignified wild type) had the highest q_s (384) mmol m⁻² s⁻¹), followed by Col-0, Sha, and Kel-4, while the more lignified soc1ful and p35S:AHL15 genotypes presented the lowest q_s value (up to 216 mmol m⁻² s⁻¹); only q_s values of Sha and Kel-4 were not statistically different from each other (Figure 2B). In addition, we noticed that Col-0 closed its stomata at a less negative leaf water potential compared with the other genotypes. It reached 90% of stomatal closure (q_{s90}) at -0.9 MPa, followed by Kel-4 (-1.13 MPa), and the more lignified Sha (-1.27 MPa), soc1ful (-1.43 MPa), and p35S:AHL15 (-1.6 MPa). The least lignified Cvi reached more negative $\Psi_{\rm L}$, even before closing its stomata (-1.75 MPa; Figure 2A). When following stomatal conductance and leaf water potential decline during the drought experiment, we found that the lignified *soc1ful* and Sha genotypes never reached critical water potential values (i.e., the P_{50}) even after 17 d of drought, while other genotypes reached their respective P_{50} between 10 d and 14 d (Supplementary Figure S2A, B).

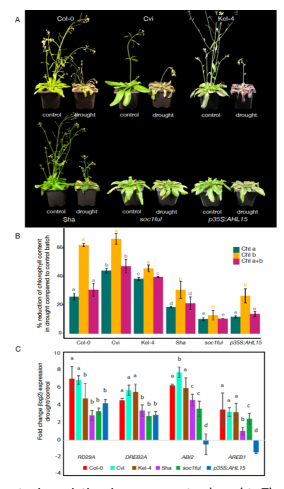


Figure 1 (A) Phenotypic variation in response to drought. The phenotype of six *Arabidopsis* genotypes subjected to drought, by water withholding, after 3 weeks and their untreated counterparts. Nine plants per genotype and condition were analyzed, and representative images are shown. (B) The variation in chlorophyll contents (chlorophyll a, chlorophyll b, and chlorophyll a+b) among genotypes studied. The Y-axis represents the percent reduction of chlorophyll content in drought compared to the control batch. A student t-test was performed, showing the differences between each accession compared to Col-0. * p-value < 0.05. (C) qRT-PCR analysis of the expression of selected drought-responsive genes (*RD29A*, *DREB2A*, *ABI2* and *AREB1*) in six Arabidopsis genotypes. The Y-axis represents the log2 fold change of the gene expression between drought and control conditions. A student t-test was performed, showing the differences between each accession

compared to Col-0. * *p*-value < 0.05. Note: the genes are significantly less upregulated by drought in Sha, *p35S:AHL15*, and *soc1ful* plants.

Stem vulnerability to embolism

When comparing all six genotypes, the most lignified soc1ful was the most embolism resistant, with P_{50} of -3.07 MPa (Figure 3; Table 1), whereas the least lignified Cvi remained the most vulnerable ($P_{50}=-1.58$ MPa). For the two added genotypes, Kel-4 (wild type with intermediate lignified stems) was among the most vulnerable genotypes with $P_{50}=-1.69$ MPa, whereas p35S:AHL15 (overexpression line) was intermediate, almost identical to the common wild-type Col-0 with $P_{50}=-2.13$ MPa. The P_{12} (stem water potential at onset of embolism) values of most of the genotypes studied were different from each other (F=420.6; $P<2e^{-16}$), but Cvi and Kel-4 presented similar P_{12} (P=0.5424). For P_{88} , p35S:AHL15 and Kel-4 were different from other genotypes (F=75.09; $P<2e^{-16}$) (Supplementary Figure S3). The slope of the vulnerability curve was similar across the genotypes, except Col-0, which had a lower slope (see Figure 3).

Water potential and SSM during drought

Assuming that leaf water potential values are similar to stem water potential values in the tiny Arabidopsis herbs, we calculated the SSM as the difference between $\Psi_{\rm gs90}$ and P_{50} . The SSMs of all genotypes studied were positive (from +0.53 MPa to +1.64 MPa), except for the least lignified Cvi with a narrow and negative SSM (-0.17 MPa) (Figure 4). Accordingly, Cvi also closed its stomata and reached a leaf water potential equivalent to P_{50} the soonest (10 d; Table 1). SSM was the widest in the most lignified soc1ful (+1.64 MPa), followed by Col-0 and Sha (+1.24 MPa and +1.22 MPa, respectively; Table 1; Figure 4). Kel-4 and p35S:AHL15 had intermediate SSMs (+0.56 MPa and +0.53 MPa, respectively).

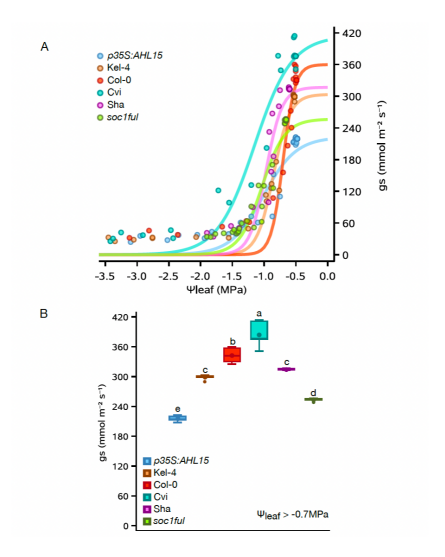


Figure 2 Drought response traits for the six *A. thaliana* genotypes studied. (A) The relationship between leaf water potential (Ψ_l) and stomatal conductance (g_s). (B) Stomatal conductance (g_s , mmols⁻¹ m⁻²) in control well-watered plants for the different Arabidopsis accession ($\Psi_l > -0.7$ MPa). Larger symbols within boxes correspond to means and smaller symbols outside boxes to outlier values. Colours refer to the genotype studied: Col-0, red; Cvi, turquoise; Sha, purple; *soc1ful*, green; *p35S:AHL15*, blue; Kel-4 brown.

The differences in anatomical features among genotypes studied

When comparing the anatomical dataset across the six genotypes, we found that the lignified soc1ful and Sha genotypes had the thickest intervessel pit membranes (T_{PM}), followed by an intermediate pit membrane thickness of p35S:AHL15 and Col-0 (F=3.857; P=0.0672), and thinner pit membranes in Kel-4 and the least lignified Cvi (F=4.467; P=0.0506) (Supplementary Figure S4A). Results of vessel wall thickness (T_V) showed the same pattern as that described for intervessel pit membrane thickness (F=2.546; P=0.13 and F=0.554; P=0.468, respectively) (Supplementary Figure S4B). Vessel grouping index (V_G) was markedly higher in the p35S:AHL15 overexpression line than in all the other genotypes (F=27.38; $P=5.46e^{-13}$) (Supplementary Figure S4C), which was also the case for the proportion of lignified area per total stem area (P_{LIG}; F=28.8; P=2.25e⁻¹³) (Supplementary Figure S4D). The lignified p35S:AHL15 overexpression line also had a higher proportion of fiber wall area per fiber cell area (PF_WF_A) than Kel-4, Col-0, and Cvi, but the fibers were less thick walled compared with the lignified genotypes soc1ful and Sha (F=49.05; P<2e⁻¹⁶) (Supplementary Figure S4E). Surprisingly, p35S:AHL15 showed no wood formation at the stem segment investigated (Supplementary Figure S1E) and was less lignified than soc1ful, although AHL15-SOC1-FUL belong to the same pathway. The vessel diameter (D) of Kel-4 was significantly narrower than that of the other genotypes. Among the remaining genotypes, Cvi (least lignified wild type) had the widest mean D, which was significantly different from the p35S:AHL15 overexpression line, but there was no statistical difference in D with Col-0, Cvi, Sha, and soc1ful (F= 9.46; P=2.52e⁻⁰⁶) (Supplementary Figure S4F). For theoretical vessel implosion resistance $(T_{VW}/D_{MAX})^2$, the lignified soc1ful and Sha showed the highest values as well, while there was no difference among p35S:AHL15, Kel-4, Col-0, and Cvi (F=3.955; P=0.0166). Finally, vessel density (V_D) of p35S:AHL15, Col-0, Cvi, Sha, and soc1ful was similar (F=1.899; P=0.13) and significantly higher than that of Kel-4.

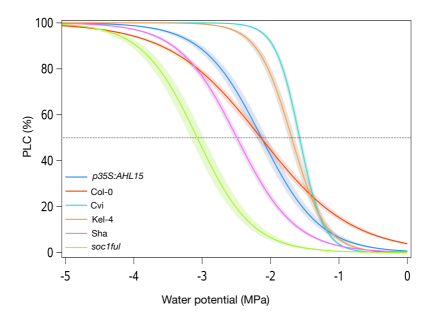


Figure 3 Mean vulnerability curves present the percentage loss of conductivity (PLC) as a function of xylem pressure (MPa) of each genotype studied. Shaded bands represent standard errors based on five to ten vulnerability curves per genotype. Colours refer to the genotype studied: Col-0, red; Cvi, turquoise; Sha, purple; *soc1ful*, green; *p35S:AHL15*, blue; Kel-4 brown.

Stem anatomical traits explaining variation in embolism resistance

According to the most parsimonious model derived from multiple linear regression (AIC= -194.59), the stem anatomical predictors that explain the embolism resistance variation were T_{PM} , T_V , V_G , and maximum vessel lumen diameter (D_{MAX}) (R^2 =0.924; P<2.2 e^{-16}) (Supplementary Table S3). T_{PM} was the anatomical feature explaining P_{50} variation best, with relative importance of 44%, followed by T_V (38%), V_G (9%), and D_{MAX} (2%) (Figure 5A). Likewise, T_{PM} and T_V together also explained most of the variation in P_{12} , with 41% relative importance (R^2 =0.795; P=1.135 e^{-14}) (Supplementary Table S4; Supplementary Figure S5A). P_{88} variation, on the other hand, was mostly explained by PF_WF_A (25% relative importance) (R^2 =0.516; P=1.07 e^{-07}) (Supplementary Table S5; Supplementary Figure S5B).

The relationship among embolism resistance, anatomical traits, and hydraulic traits

Based on a Pearson's correlation test, T_{PM} was strongly positively correlated with other anatomical traits, such as T_V, $(T_{VW}/D_{MAX})^2$, PF_WF_A, and V_D (r=0.77 and P=1.108e⁻¹¹; r=0.74 and P=1.956e⁻¹⁰; r=0.61 and P=8.96e⁻⁰⁷, r=0.58, $P=4.472e^{-06}$, respectively) (Supplementary Figure S6). Lastly, T_V and PF_WF_A were correlated as well (r=0.71, $P=2.3e^{-09}$) (Supplementary Figure S6). When also taking P_{50} into account, we saw that P_{50} was strongly correlated with T_{PM} , $(T_{VW}/D_{MAX})^2$, T_V , and PF_WF_A (r = -0.91, -0.87, -0.86, and -0.70; P<2.2e⁻¹⁶, respectively) (Figure 5B–E; Supplementary Figure S6). Similarly, P_{12} had strong relationships to T_{PM} , $(T_{VW}/D_{MAX})^2$, and T_V (r = -0.77 and $P=6.41e^{-12}$; r=0.84 and $P=3.93e^{-15}$; r=0.68 and $P=1.38e^{-08}$, respectively) (Supplementary Figure S6). P_{88} only showed a correlation with PF_WF_A (r=-0.54; $P=2.762e^{-0.5}$) and T_V (r=-0.44; P=0.0008146) (Supplementary Figure S6). We also found a strong correlation between P_{50} and the leaf water potential at the harvesting day (Ψ_{lh}), the number of days until reaching 90% stomatal closure (Day₉₀), and the SSM ($r = -0.9, -0.85, \text{ and } -0.84; P < 2.2e^{-16},$ respectively), but not between P_{50} and Ψ_{gs90} . Subsequently, the anatomical traits that were strongly correlated to P₅₀, such as T_{PM}, T_V, and V_G, were also significantly correlated to Ψ_{lh} , Day₉₀, and SSM (Supplementary Figure S6).

Discussion

We performed a drought experiment including six Arabidopsis genotypes, during which we compiled a detailed xylem anatomical—hydraulic dataset of inflorescence stems (among others intervessel pit membrane thickness, proportion of lignification, and P_{50}) and leaves (rate of stomatal conductance, leaf water potential, and chlorophyll content), and validated the drought response of the genotypes with the transcript abundance of four known drought marker genes at the end of a 15–17 d treatment without watering. Based on anatomical, hydraulic, and gene expression results, it is clear that the most lignified mutant soc1ful (Melzer $et\ al.$, 2008; Lens $et\ al.$, 2012, 2013) is the most drought-tolerant genotype, closely followed by the lignified ecotype Sha and the p35S:AHL15 overexpression line, while the lesser lignified Col-0, Kel-4, and especially Cvi ecotypes are much more sensitive. Interestingly, each genotype applies a unique combination of anatomical stem traits and hydraulic traits in stems

and leaves to acquire a certain level of drought tolerance, as will be discussed in the following sections.

Table 1 The hydraulic data of *Arabidopsis* genotypes studied measured during the drought experiment.

Genotypes	<i>P</i> ₅₀ (MPa)	ψ _{gs90} (MPa)	SSM (MPa)	Ψ _{lh} (MPa)	Days until 90% stomatal closure	Days until P ₅₀	PLC after 3 weeks of non- watering
Cvi	-1.58	-1.75	-0.17	-3.4	10	10	100%
Kel-4	-1.69	-1.13	0.56	-3.4	11	11	100%
Col-0	-2.14	-0.9	1.24	-2.97	10-11	12	75%
p35S:AHL15	-2.13	-1.6	0.53	-3.03	13	14	88%
Sha	-2.49	-1.27	1.22	-1.85	12	does not reach P_{50}	14%
soc1ful	-3.07	-1.43	1.64	-1.87	14	does not reach P ₁₂	10%

 P_{50} = stem water potential at 50% loss of hydraulic conductivity, $\Psi_{\rm gs90}$ = leaf water potential at 90% stomatal closure, SSM= stomatal safety margin, $\Psi_{\rm lh}$ = leaf water potential at the harvesting day, PLC = percentage loss of hydraulic conductivity

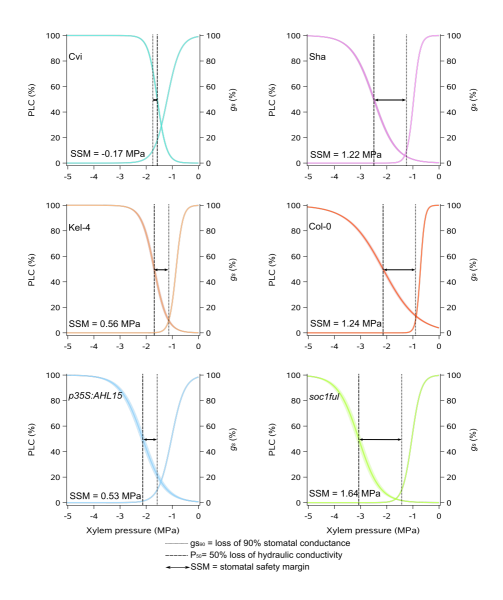


Figure 4 Stomatal safety margin (SSM) of each genotype studied. The graphs show the percent loss of hydraulic conductivity (PLC) and the percent of stomatal conductance (g_s) as a function of xylem pressure (MPa). The dotted lines represent water potential at 90% loss of stomatal conductance. The dashed lines show P_{50} . The difference between the dashed and the dotted line refers to the SSM.

Comparing extremes in drought response: most lignified soc1ful versus least lignified Cvi

Both the most drought-tolerant soc1ful and the most droughtsensitive Cvi use a similar set of traits with contrasting trait values to reach the two extremes of the drought tolerance spectrum among the genotypes studied. The drought-tolerant strategy of soc1ful (Figure 1A) is determined by a unique combination of traits, as exemplified by the most negative stem P₅₀ (Figure 3; cf. Choat et al., 2012; Lens et al., 2016; Thonglim et al., 2021), coupled with a low initial g_s that gradually slowed down during drought, allowing a more stable leaf water potential (Supplementary Figure S2A) (Li et al., 2017; Dayer et al., 2020; Lemaire et al., 2021). In addition to its low q_s, soc1ful started closing its stomata rapidly at the onset of drought (at high water potential) to further reduce water loss, but at the same time it reached full stomatal closure later than in the other genotypes (Ψ_{gs90} was reached after 14 d of non-watering, Table 1). Although we had not quantified carbon uptake during drought, we observed that stomatal closure in soc1ful occurred gradually over a longer period during drought, probably extending photosynthetic activities without risking a detrimental level of drought-induced embolism (Figure 2A; Supplementary Figure S2). This is further supported by a low reduction of chlorophyll content in rosette leaves of droughted soc1ful individuals compared with the well-watered control batch (Figure 1B), Moreover, this mutant line had the widest positive SSM (Figure 4), which is essential in estimating a plant's drought response (Choat et al., 2012; Delzon and Cochard, 2014; Anderegg et al., 2016; Eller et al., 2018; Oliveira et al., 2021; Skelton et al., 2021). Finally, as reported in Thonglim et al. (2020), this mutant also produced the thickest intervessel pit membranes and the largest wood cylinder at the base of the inflorescence stem. Both traits are thought to play an important role in preventing embolism spread (Lens et al., 2022). In contrast, the least lignified Cvi was the most vulnerable genotype as it showed the least negative stem P_{50} combined with a rapid drop in leaf water potential during drought, leading to rapid wilting (Figure 1A) and a strong decrease of chlorophyll content (Figure 1B). In addition, Cvi had the highest initial q_s , and it closed its stomata at low water potential, which led to more water loss due to transpiration (Figure 2A; Supplementary Figure S2). Although it reached Ψ_{gs90} earlier than the more tolerant genotypes (Table 1), it seemed like Cvi could not close its stomata in time because all the water was already consumed, giving rise to a rapid water potential drop during drought (Supplementary Figure S2A). Due to its less negative stem P_{50} , the Ψ_{gs90} exceeded stem P_{50} , leading to the only negative SSM among the six genotypes studied (Figure 4). This implies that Cvi experiences a considerable decrease in stem hydraulic conductivity right after or even before stomatal closure. In addition to all these physiological parameters pointing to the most sensitive drought response among the genotypes studied, Cvi also had the least lignified inflorescence stems with the thinnest intervessel pit membranes (Thonglim *et al.*, 2021).

The role of embolism resistance and stomatal regulation in drought tolerance and its impact on the stomatal safety margin

The previous section highlights the importance of embolism resistance as well as SSMs in determining drought tolerance, as has been demonstrated across many other lineages of plants (Meinzer et al., 2009; McDowell, 2011; Choat et al., 2012; Johnson et al., 2012; Cochard et al., 2013; Lens et al., 2013; Skelton et al., 2015, 2021; Martin-StPaul et al., 2017; Creek et al., 2020; Dayer et al., 2020). However, our dataset suggests that stem P₅₀—which is probably a good proxy for whole-plant P₅₀ based on our few leaf P₅₀ measurements in the p35S:AHL15 overexpression line and based on other herbaceous species showing no difference in P_{50} across organs (e.g., Skelton et al., 2017)—outperforms SSM in explaining the responses to drought among the genotypes studied. This is because stomatal regulation in Arabidopsis genotypes that were equally drought tolerant could be substantially different, while P₅₀ showed a more consistent pattern with whole-plant drought tolerance. However, it seems that the rate of g_s in Arabidopsis under well-watered conditions is more critical than the speed of stomatal closure, as shown by Cvi, Col-0, and Kel-4 (Table 1; Supplementary Figure S2B). Indeed, $\Psi_{\rm gs90}$ is not the driving force behind drought tolerance since the more drought-tolerant genotypes closed their stomata slightly later than the sensitive ones. In other words, Cvi, Col-0, and Kel-4 lost more water because of a higher transpiration rate, but they closed their stomata sooner than the more drought-tolerant genotypes (Table 1). These results align with previous studies stating that stomatal behavior only shows how each species respond to drought stress, but not how much they tolerate drought (Roman et al., 2015; Combe et al., 2016; Martínez-Vilalta and Garcia-Forner, 2017). Bearing this in mind, our observation shows that the two mutant genotypes studied in the Col-O background (soc1ful and

p35S:AHL15)—both belonging to the same regulatory SOC1-FUL-AHL15cytokinin pathway that induces wood formation in stems (Rahimi et al., 2022)—also have by far the lowest initial q_s values across all six genotypes studied, including the Col-O ecotype (Figure 2B). This makes it a promising gene regulatory pathway to discover how drought-responsive traits in stems (increased lignification or woodiness) and leaves (reduced q_s) are linked to each other at the genetic level. Our dataset aligns with earlier studies showing that safety margins across (mainly woody) angiosperms are overall positive, and considerable levels of embolisms only happen under remarkable, intense drought events (Choat et al., 2012; Delzon and Cochard, 2014; Martin-StPaul et al., 2017; Creek et al., 2020; Dayer et al., 2020; Skelton et al., 2021; Guan et al., 2022; Lens et al., 2022). The positive SSMs in five out of six genotypes indicate that stomatal closure typically occurs before embolism in order to prevent water loss and delay hydraulic dysfunction (Martin-StPaul et al., 2017; Creek et al., 2020). In contrast, Cvi the only genotype with a negative SSM—closed its stomata at 70% loss of maximum conductance, highlighting its high sensibility to drought.

Multiple strategies to acquire drought tolerance

In addition to the drought-responsive traits discussed in soc1ful and Cvi, different combinations among these traits were observed in the remaining genotypes. This shows that even in a species with a short life cycle, multiple strategies can be applied to acquire a certain level of drought tolerance. For instance, Sha and p35S:AHL15 had a similarly high level of drought tolerance based on their phenotype after 3 weeks of water shortage (Figure 1A), but their drought-responsive traits were different. Sha had high embolism resistance in stems combined with a relatively high initial transpiration rate in leaves that rapidly declines during drought, allowing a relatively stable leaf water potential (also confirmed by Bouchabke et al., 2008) and a large SSM. On the other hand, p35S:AHL15 had the lowest q_s of all the genotypes studied (Figure 2A), which means it can keep its leaf water potential relatively high during drought, whereas its stem P_{50} was intermediate and led to a smaller SSM compared with Sha (Figures 2-4). Another example is given by p35S:AHL15 (overexpression line) and Col-0 common wild type, which both had a similar stem P_{50} (-2.1 MPa; Figure 3). However, Col-0 was more drought sensitive than p35S:AHL15, even though the former closed its stomata earlier during drought, resulting in a wider

SSM (Figure 4). The reason for Col-0 being more drought sensitive is that stomatal conductance is much higher, leading to more water loss and consequently a more rapid decline in leaf water potential during the drought experiment, while the leaf water potential during drought in p35S:AHL15 drops more slowly (Supplementary Figure S2). Thus, a wider SSM does not always lead to prolonged survival during drought since the rate of g_s is not accounted for in the SSM. In other words, the width of the safety margin does not necessarily match all aspects of stomatal regulation and the resulting leaf water potential dynamics during drought (Martin-StPaul *et al.*, 2017; Martínez-Vilalta and Garcia-Forner, 2017; Knipfer *et al.*, 2020).

Expression levels of drought-responsive genes agree with drought-response traits

To assess the level of drought stress and compare it among the genotypes, we assessed the expression of selected drought-responsive genes on the final day of the drought treatment (15–17 d). As expected, the four drought-responsive genes *RD29A*, *DREB2A*, *ABI2*, and *AREB1* were most up-regulated in the more sensitive genotypes Col-0, Kel-4, and Cvi, and less up-regulated in the more tolerant genotypes Sha, *p35S:AHL15*, and *soc1ful* (Figure 1C). To study the casual relationship between physiological responses (e.g., stomatal closure) and gene activity (e.g., ABA biosynthesis genes), future work should focus on conducting a high-resolution time-course gene expression analysis, which is beyond the scope of this study.

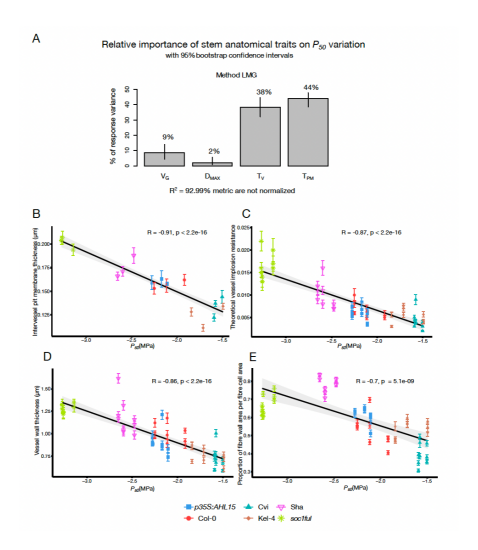


Figure 5 (A) Relative importance of stem anatomical traits on P_{50} variation. The P_{50} variation is mainly explained by intervessel pit membrane thickness (T_{PM}) and vessel wall thickness (T_{V}) based on R^{2} contribution averaged over orderings among regressors (based on LMG method). (B) Negative correlation between T_{PM} and P_{50} (C) Negative correlation between (T_{VW}/D_{MAX})² and P_{50} (D) Negative correlation between T_{V} and P_{50} . Colours and styles refer to the genotype studied: Col-0, red circles; Cvi, turquoise triangles point up; Sha, purple triangles point down; soc1ful, green stars; p35S:AHL15, blue squares; Kel-4, brown diamonds.

Intervessel pit membrane thickness as an important anatomical driver of embolism resistance, and the potential effect of stem lignification on P₅₀

Our extended database confirms our previous results that intervessel pit membrane thickness is the anatomical trait that explains best the variation in P_{50} across all six genotypes studied (Figure 5A). These results are in line with several other angiosperm studies showing a strong positive correlation between embolism resistance and T_{PM}, both at the interspecies level (Jansen et al., 2009; Lens et al., 2011, 2022; Plavcová and Hacke, 2012; Plavcová et al., 2013; Scholz et al., 2013b; Li et al., 2016; Dória et al., 2018; Trueba et al., 2019; Guan et al., 2022) and within species (Schuldt et al., 2016). The functional explanation for this relationship was intensively discussed in our previous paper (Thonglim et al., 2021). In brief, there is convincing evidence based on microCT and/or optical technique observations in stems (Brodersen et al., 2013; Knipfer et al., 2015; Choat et al., 2016; Skelton et al., 2017; Torres-Ruiz et al., 2017) and leaves (Brodribb et al., 2016a; Skelton et al., 2017, 2018; Klepsch et al., 2018; Lamarque et al., 2018) that embolism spread between adjacent vessels predominantly happens via porous pit membranes located inside the bordered pits between adjacent vessels. Although this explains why the thickness of intervessel pit membrane plays an important role in embolism propagation and, by extension, also whole-plant drought tolerance, the detailed mechanisms behind this embolism spread remain poorly known due to the complex 3D structure/composition of pit membranes and the enigmatic behavior of gas-liquid-solid-surfactant interfaces at the nano-scale (Kaack et al., 2019, 2021; Yang et al., 2020; Zhang et al., 2020; Lens et al., 2022).

It has also been shown in previous studies that intervessel pit membrane thickness is strongly linked not only with P_{50} , but also with other anatomical traits assumed to be involved in drought-induced embolism resistance, such as vessel wall thickness (Jansen *et al.*, 2009; Li *et al.*, 2016), and the amount of stem lignification or woodiness (Li *et al.*, 2016; Dória *et al.*, 2018; Thonglim *et al.*, 2021). How exactly lignification would impact embolism spread in stems is the subject of ongoing research. One hypothesis is that the amount of lignification in secondary cell walls may determine gas diffusion kinetics across xylem cell walls and, therefore, could reduce the speed of embolism propagation in species with increased levels of lignification or woodiness (Li *et al.*, 2016; Dória *et al.*, 2018; Pereira *et al.*,

2018; Thonglim *et al.*, 2021; Lens *et al.*, 2022). This may imply that older stems from herbaceous species could lead to increased embolism resistance, resulting from a possible increase in stem lignification and/or the amount of wood. In our study, this may especially apply to the *p35S:AHL15* overexpression line, which has the ability to develop as much wood as the *soc1ful* double knockout genotype (Rahimi *et al.*, 2022). However, this study shows that wood development is delayed in *p35S:AHL15* (Supplementary Figure S1E, G) compared with *soc1ful* in 80-day-old plants, despite the fact that SOC1, FUL, and AHL15 belong to the same wood pathway (Rahimi *et al.*, 2022). Older individuals of *p35S:AHL15* will therefore develop more wood and probably also thicker intervessel pit membranes in their inflorescence stems, most probably resulting in both higher embolism resistance and higher SSM, which synergistically may increase total plant tolerance of the overexpression line to the level of *soc1ful*.

In conclusion, there is a considerable difference in drought response among the six Arabidopsis genotypes studied. The genotypes soc1ful, Sha, and p35S:AHL15 synergistically increase their drought tolerance by building lignified inflorescence stems with thick intervessel pit membranes, developing the largest SSMs, keeping the water potential in their leaves pretty stable during periods of water shortage as a result of low stomatal conductance, maintaining relatively high chlorophyll content in rosette leaves, and by showing the lowest expression levels of drought-response genes compared with the control batch. In contrast, the most sensitive genotypes to drought (Cvi, Kel-4, and Col-0) are more susceptible to drought due to the opposite extreme of the same set of drought-responsive traits. This shows that stem anatomical traits and hydraulic stem and leaf traits are intertwined to acquire a certain level of drought tolerance. To further disentangle gene regulatory networks underlying drought-responsive traits across organs and to find out how they are linked with each other and synergistically strengthen the whole-plant drought response, future studies should combine a time series of gene expression data in roots, stems, and leaves during a drought experiment followed by rewatering. During such an experiment, a range of drought-responsive (anatomical and physiological) traits in all organs should be investigated. Only with this integrative approach, will we be able to make considerable progress in securing our food production by developing breeding tools that can make crops more

drought tolerant and propose solutions on how to protect our herbs and forests under the current global change scenario.

Acknowledgements

We would like to thank Alex Bos (IBL) for support with the gene expression analysis, Omid Karami and Arezoo Rahimi (IBL) for providing seeds and their knowledge in taking care of Arabidopsis plants, and Gaëlle Capdeville (BIOGECO INRA) for technical support. This work is funded by a PhD scholarship awarded to AT from the Institute for the Promotion of Teaching Science and Technology (IPST), Thailand, and by the Dutch Research Council NWO (grant number ALWOP.488).

Supplementary data

Supplementary Table S1 Oligonucleotide sequences.

Primers for qRT-PCR					
Gene name	Gene ID	Primer Fwd sequence (5´-3´)	Primer Rev sequence (5´-3´)		
ACTIN2	AT3G18780	TCCCTCAGCACATTCCAGCAGAT	AACGATTCCTGGACCTGCCTCATC		
GAPDH	At1G13440	TTGGTGACAACAGGTCAAGCA	AAACTTGTCGCTCAATGCAATC		
RD29A	AT5G52310	TGGACAAAGCAATGAGCATGAGC	AGGTTTACCTGTTACGCCTGGTG		
ABI2	ATG557050	CTCGCAATGTCAAGATCCATTGGC	TTACTCGCCGCACTGAAGTCAC		
AREB1	AT1G45249	AGTTACAACGAAAGCAGGCAAGG	CCTCCTTGCAGAAGATTCCTCATC		
DREB2A	AT5G05410	CAGTGTTGCCAACGGTTCAT	AAACGGAGGTATTCCGTAGTTGAG		

Supplementary Table S2 The anatomical characters and hydraulic values measured with acronyms, definitions, calculations, units, and techniques.

Acronyms	Definition	Calculation	Number of measurements	Unit	Technique
A _F	Fiber cell	Area of single xylem	Min. 30 fibers	μm²	LM
	area	fiber in cross-section			
A _{FL}	Fiber lumen area	Area of single xylem fiber lumen in cross-	Min. 30 fibers	μm^2	LM
	arca	section			
A_{FW}	Fiber wall	A _F - A _{FL} for the same	Min. 30 fibers	μm^2	LM
	area	fiber			
A _{LIG}	Lignified	Total xylem area +	9 stems per	mm^2	LM
	stem area	fiber caps area + lignified pith cell area in cross-section	accession		
Apith	Pith area	Total pith area in	9 stems per	mm²	LM
		cross-section	accession		
As	Total stem	Total stem area in	9 stems per	mm^2	LM
	area	cross-section	accession		
Day ₉₀	Days until	-	-	days	-
	reaching				
	90% of				
	stomatal closure				
D _{MAX}	Maximum	Diameter of single	Min. 30 vessels	μm	LM
DIVIAX	vessel lumen	vessel	141111. 30 VC33C13	μ…	LIVI
	diameter				
D_PC	Pit chamber	Distance from the	Min. 25 pits	μm	TEM
	depth	relaxed pit			
		membrane to the			
a	Stomatal	inner pit aperture	1 control	mmol	Porometer
g s	conductance	-	sample and 2	m ⁻² s ⁻¹	Porometer
	conductance		drought	3	
			samples each		
			measurement		
SSM	Stomatal	$\Psi g s_{90} - P_{50}$	1 SSM per	MPa	-
	safety margin		accession		
P ₁₂	Stem water		8 values per	MPa	Cavitron
, 12	potential at	-	each accession	ivii u	centrifuge
	12% loss of				J
	hydraulic				
	conductivity				
P ₅₀	Stem water	-	8 values per	MPa	Cavitron
	potential at		each accession		centrifuge
	50% loss of hydraulic				
	conductivity				
	•				
P ₈₈	Stem water	-	8 values per	MPa	Cavitron
	potential at		each accession		centrifuge

P _{FW} F _A	88% loss of hydraulic conductivity Proportion of fiber wall area per fiber cell area	A _{FW} /A _F for the same fiber; a measure of xylem fiber wall thickness	Min. 30 fibers	-	LM
ψ_{gs90}	Leaf water potential at 90% loss of stomatal conductance	-	1 control sample and 2 drought samples each measurement	MPa	PSYPRO meter
ψ_{lh}	Leaf water potential at the harvesting day	-	1 control sample and 2 drought samples each measurement	MPa	PSYPRO meter
P _{LIG}	Proportion of lignified area per total stem area	Alig/As	9 stems per accession	-	LM
Трм	Intervessel pit membrane thickness	Thickness of intervessel pit membrane measured at its thickest point	Min. 25 pit membranes	μт	TEM
Tv	Vessel wall thickness	Thickness of a single vessel wall	Min. 30 vessels	μm	LM
T _{VW} /D _{MAX}	Thickness-to- span ratio of vessels	Double intervessel wall thickness divided by the maximum diameter of the largest vessel	Min. 30 measurements	μт	LM
(T _{VW} /D _{MAX})	Theoretical vessel implosion resistance	(Tvw/D _{MAX}) ²	Min. 30 measurements	-	LM
V_D	Vessel density	Number of vessels per mm ²	Min. 5 measurements	No. of vessel per mm ²	LM
V _G	Vessel grouping index	Ratio of total number of vessels to total number of vessel groupings (incl. solitary and grouped vessels)	Min. 50 vessel groups	-	LM

Supplementary Table S3 The most parsimonious multiple linear regression model (based on AIC scores) of anatomical traits, explaining stem P_{50} variation of the six *Arabidopsis thaliana* accessions studied.

Predictors	Estimate	Std. Error	t value	Pr (> t)
(Intercept)	0.435	0.239	1.825	0.074
ТРМ	-11.096	1.329	-8.347	5.67e ⁻¹¹ ***
D _{MAX}	0.028	0.007	4.149	0.000132***
V _G	-0.242	0.073	-3.331	0.001651**
Tv	-1.074	0.146	-7.360	1.84 e ⁻⁰⁹ **

 T_{PM} = intervessel pit membrane thickness; D_{MAX} = maximum vessel lumen diameter; V_G = vessel grouping index; T_V = vessel wall thickness. *** *p-value* < 0.001; ** *p-value* < 0.01

Supplementary Table S4 The most parsimonious multiple linear regression model (based on AIC scores) of anatomical traits explaining stem P_{12} variation of the six *Arabidopsis thaliana* accessions studied.

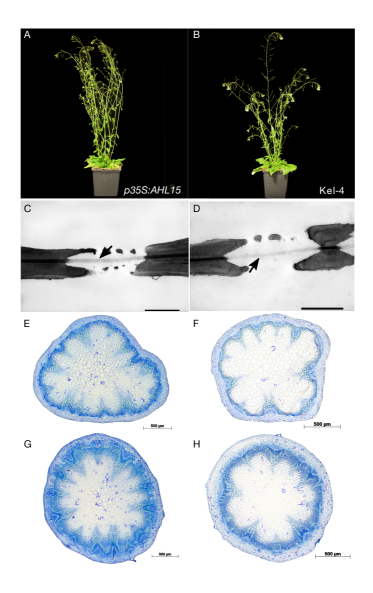
Predictors	Estimate	Std. Error	t value	Pr (> t)
(Intercept)	-0.599	0.687	-0.872	0.388
P _{LIG}	-3.365	1.793	-1.877	0.067
ТРМ	-12.657	3.660	-3.458	0.001**
D _{PC}	1.926	0.756	2.546	0.014*
D_{MAX}	0.086	0.016	5.492	1.76e ⁻⁰⁶ ***
PF_WF_A	1.093	0.660	1.656	0.105
\mathbf{V}_{D}	0.006	0.003	2.443	0.019*
V_{G}	-0.511	0.261	-1.959	0.056
T _V	-1.416	0.404	-3.502	0.001**

 P_{LIG} = proportion of lignified area per total stem area; T_{PM} = intervessel pit membrane thickness; D_{PC} = pit chamber depth; D_{MAX} = maximum vessel lumen diameter; PF_WF_A = proportion of fiber wall area per fiber cell area; V_D = vessel density V_G = vessel grouping index; T_V = vessel wall thickness. *** p-value < 0.001; ** p-value < 0.01; * p-value < 0.05

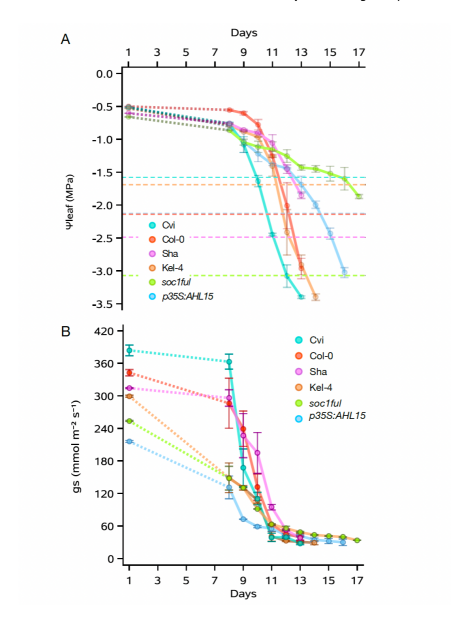
Supplementary Table S5 The most parsimonious multiple linear regression model (based on AIC scores) of anatomical traits explaining stem P_{88} variation of the six *Arabidopsis thaliana* accessions studied.

Predictors	Estimate	Std. Error	t value	Pr (> t)
(Intercept)	1.414	0.701	2.018	0.049*
P _{LIG}	3.967	1.556	2.550	0.014*
D _{PC}	-1.932	0.726	-2.662	0.011*
PF_WF_A	-2.208	0.469	-4.710	2.148e ⁻⁰⁵ ***
V_D	-0.010	0.003	-3.575	0.000810***
D _{MAX}	-0.072	0.018	-4.054	0.000184***

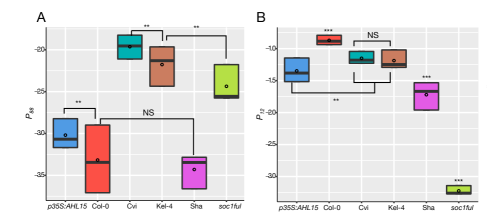
 P_{LIG} = proportion of lignified area per total stem area; D_{PC} = pit chamber depth; PF_WF_A = Proportion of fiber wall area per fiber cell area; V_D = vessel density; D_{MAX} = maximum vessel lumen diameter. *** p-value < 0.001; ** p-value < 0.05



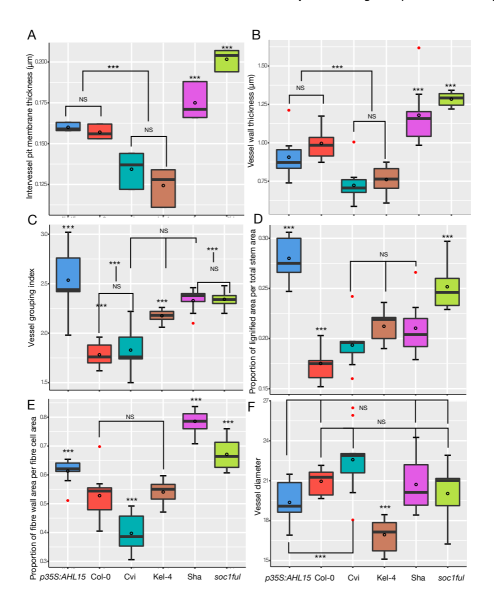
Supplementary Figure S1 Growth form and cross-sections of inflorescence stems of *p35S:AHL15* (left, 85d after sowing) and Kel-4 (right, 65d after sowing). (A, B) Growth form. (C, D) TEM images of intervessel pit membranes (arrows). Scale bars = $2 \mu m$. (E, F) LM images of cross-sections at the middle part of inflorescence stems. Scale bars = $500 \mu m$. (G, H) LM images of cross-sections at the basal part of inflorescence stems show more pronounced lignification. Scale bars = $500 \mu m$.



Supplementary Figure S2 Leaf water potential and stomatal conductance during drought experiment. (A) Leaf water potential (Ψ_l) over time. Dotted lines represent P_{50} value of each genotype. (B) Stomatal conductance (g_s) over time. Colours refer to the genotype studied: Col-0, red; Cvi, turquoise; Sha, purple; soc1ful, green; p35S:AHL15, blue; Kel-4, brown.

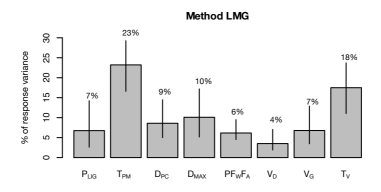


Supplementary Figure S3 Boxplots showing P_{88} and P_{12} variation within and between genotypes. (A) Boxplot showing P_{88} of every genotype studied. (B) Boxplot showing P_{12} of every genotype studied; ns = p-value > 0.05; *** p-value < 0.05.



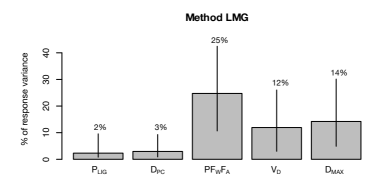
Supplementary Figure S4 Boxplots showing anatomical variation within and between all genotypes. (A) Boxplot of intervessel pit membrane thickness (T_{PM}). (B) Boxplot of vessel wall thickness (T_{V}). (C) Boxplot of vessel grouping index (V_{G}). (D) Boxplot of the proportion of lignified area per total stem area (P_{LIG}). (E) Boxplot of the proportion of fiber wall area per fiber cell area ($P_{FW}F_{A}$). (F) Boxplot of vessel diameter (D); $P_{FW}F_{A}$ 0. (F) Boxplot of vessel diameter (D); $P_{FW}F_{A}$ 1.

A Relative importance of stem anatomical traits on P₁₂ variation with 95% bootstrap confidence intervals



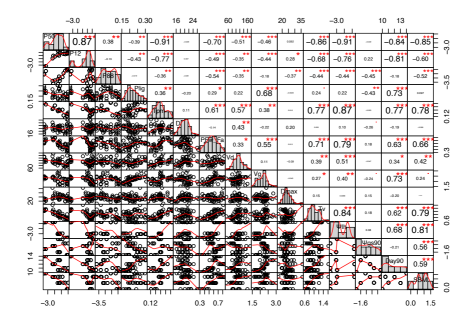
 R^2 = 82.62%, metrics are not normalized.

Relative importance of stem anatomical traits on P₈₈variation with 95% bootstrap confidence intervals



R² = 56.16%, metrics are not normalized.

Supplementary Figure S5 The relative importance of P_{12} and P_{88} evaluated. (A) The relative importance of P_{12} variation is mainly explained by intervessel pit membrane thickness (T_{PM}) and vessel wall thickness (T_{V}). (B) The relative importance of P_{88} variation is mainly explained by proportion of fiber wall area per fiber cell area ($PF_{W}F_{A}$).



Supplementary Figure S6 The pairwise scatter plots based on Pearson's correlation analysis show the correlations of P_{50} , P_{12} and P_{88} (response variables) and each stem anatomical and hydraulic traits studied (predictive variables) and between all the predictive variables. *** p-value < 0.01; ** p-value < 0.05.