Listening to ultrasound from plants reveals xylem vessel anatomy

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Article

Keywords: xylem vessel anatomy, ultrasound,

DOI: https://doi.org/10.21203/rs.3.rs-452046/v1

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Listening to ultrasound from plants reveals xylem vessel anatomy

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Plants emit ultrasound pulses under drought stress, which originate in their water-carrying xylem vessels, and can be recorded externally. We demonstrate that these ultrasound pulses consist of superposed damped oscillations at plant-specific frequencies in the range of 10 – 150 kHz, that are correlated to xylem dimensions. We present a method to relate geometrical and viscoelastic properties of xylem vessels with the time- and frequency-domain characteristics of the observed oscillations. We apply the method to ultrasound pulses from drying shoots of three vascular dicot plant species. The extracted parameters are validated with destructive measurements of xylem vessel radii, wall thickness, length of xylem vessel elements, and the elastic modulus of the vascular bundle by optical and scanning cryo-electron microscopy and tensile loading. Our method demonstrates the potential for non-invasive and continuous monitoring of plant vascular anatomy. We foresee applications in high-throughput phenotyping and early detection of vascular wilt diseases.

Plant hydraulics, the study of water transport in plants, is vital to our understanding of plant function and stress resilience1-2. In vascular plants, the xylem is responsible for water and nutrient transport from the roots to the leaves3. Transpiration through leaves results in a tensile force on the water-column, which, combined with the strong cohesion of water molecules, results in ascent of water from the roots to the leaves4. During drought or strong transpiration rate, the tension in the water column increases rapidly. Beyond a critical tension, the stress is released by the formation of vapour or gas-bubbles5,6 in the xylem. The bubble formation results in a sudden release of the elastic energy stored in the water column, a fraction of which is converted to a sound pulse7. The rate at which such pulses are emitted has been used as a marker of a plant’s response and vulnerability to drought-stress8-11. The time- and frequency-domain features of these ultrasound pulses, measured directly from plant shoots, were shown in a recent study12. Yet, the physical origin and relevance of the observed damped oscillations in these acoustic pulses13-16 has remained elusive.

Xylem vessels resemble cylindrical tubes with fused ends3. These tubes consist of several xylem vessel elements that are separated by perforation plates. The diameters of these vessels range from ~ 1 μm in small herbs to ~100 μm in woody trees, and their lengths range from ~100 μm to ~10 cm17-19 across the plant kingdom. The viscoelastic walls of xylem cells are composed of an interwoven matrix of cellulose, hemicellulose, pectin and lignin fibres, which can have a wide range of elastic moduli depending on their relative composition20 and the water content11,22. The elasticity of macroscopic segments of plant stems can be measured via various mechanical loading techniques23,24, which are invasive. Existing techniques to measure xylem dimensions, such as paint-injection, X-ray micro-CT, and optical and electron microscopy25-28, are also destructive and time-consuming.

Here we present a physical model that links the dimensions and (visco-)elasticity of xylem vessels, to measured ultrasound pulses. We compare information about the radius, length, and viscoelasticity of xylem vessels obtained by analysing ultrasound pulses, to that gained by independent destructive techniques on Hydrangea quercifolia. In addition, the correlation between viscous damping in the ultrasound pulses and the vessel radius distribution is further elucidated by experiments on Hydrangea macrophylla, and Solanum lycopersicum. Lastly, via pulsed ultrasound spectroscopy with an external sound source, we show that acoustic resonances in the vascular tissue can be artificially excited, the characteristics of which agree with those excited naturally during drought-stress.

Results

Ultrasound waveforms. We first examined the waveforms of ultrasound pulses that were emitted by drying plant shoots. A total of three specimens, samples A, B, and C, were taken from three H. quercifolia plants (Methods). Ultrasound pulses were recorded with a broadband ultrasound microphone placed along the axial and radial direction of the stem (Fig. 1a). The microphone recorded the time series of the ultrasound emissions starting ~ 5 minutes into the drying process (Fig. 1b; Methods), where time t = 0 s corresponds to the start of the recording. The pulses occurred sporadically and with varying
amplitudes. We observed that the time-domain waveforms of these pulses resembled damped oscillations, both when recorded along the axial and in the radial directions (Figs. 1c and 1d). The pulse amplitude in time-domain decayed exponentially with a 1/e time constant \( \tau \); the settling time (Methods). For stem sample A, we extracted \( \tau = 28.8 \pm 6.4 \mu s \) (mean \( \pm \)s.d.), for the pulses recorded in the axial direction. The corresponding value of \( \tau \) for the radially recorded pulses was 41.7 \( \pm \) 12.4 \( \mu s \), which was statistically similar. The \( \tau \) for the many individually measured axial and radial sound pulses of all the three stem samples A, B, and C are shown in Supplementary Figs. 1 – 3, respectively. The determined settling times of samples B and C agreed with those of sample A. All pulses died out within \( \sim 0.3 \) ms, in agreement with reported work. Based on this observation, we hypothesize that the damped oscillations are generated by resonant vibrations within the xylem vessels. In the following paragraphs, the settling times and characteristic frequencies of ultrasound pulse waveforms were interpreted to estimate xylem vessel dimensions and elasticity (Fig. 1e).

Xylem vessel radius. In order to explain the origin of the observed ultrasound waveforms and to use them to extract information about the plant’s microstructure, we develop a model relating the micromechanics of the xylem to the waveform of the generated ultrasound. We hypothesize that the damped oscillations are identical to those of an organ pipe filled with water. The bubble formation excites axial standing waves in the sap (water), whose resonance frequencies depend on the longitudinal speed of sound in the pipe \( c_{\text{pipe}} \), and the xylem vessel element length \( L \) (Methods). We modelled the xylem vessel as a resonant cylindrical pipe containing a series network of vessel elements of length \( L \), which are bounded by scalariform perforation plates (Fig. 2a). The perforation plates serve as non-ideal (leaky) reflecting surfaces at the termination of a vessel element for the pressure waves. The sound waves propagate along the length of the xylem vessel, and are likely to dominate the recorded ultrasound. These waves undergo damping, primarily due to the dynamic viscosity of sap (water) \( \eta \) in the xylem, which dominates the settling time \( \tau \). The resonating element can be described using a linear second order resonator model consisting of lumped acoustic inductance, capacitance and resistance (Methods). Using this acoustic model, we have established that the effective xylem radius \( R \) in terms of the measured settling time using equation (8) (Methods), where \( \rho \) is the mass density of the sap (water). In this model, \( R \) can be calculated independently of \( L \), from the settling time of the measured time-domain waveform. The histograms of xylem radii \( R \) were extracted from the axially recorded ultrasound pulses, and also from optical micrographs of the stem samples (Figs. 2d, Supplementary Fig. 4). The mean (\( \pm \)s.d.) acoustic radius \( R \) for sample A was 9.93 \( \pm \) 1.6 \( \mu m \). Similar values were obtained for stem samples B and C (Supplementary Fig. 4). Using optical micrographs of latex-paint stained transverse cross-sections of the stem sample A (Fig. 2b), we observed the vessel radius \( R = 11.9 \pm 2.6 \mu m \) (Fig. 2d and Table 1). These values were confirmed with scanning electron cryo-microscopy (Fig. 2c). Thus, the calculated \( R \), using the ultrasound analysis, agrees with that observed by optical and scanning electron microscopy.

We further validated our method using other plant species, namely H. macrophylla and S. lycopersicum (Figs. 2e – 2j). The mean \( \tau \) for H. Macrophylla and S. lycopersicum were 26.4 \( \pm \) 7.0 \mu s and 116 \( \pm \) 85.0 \mu s, respectively. Histograms and mean \( R \) derived from direct measurements (Figs. 2g, 2h), were in good agreement (Figs. 2i, 2j). The relatively larger \( \tau \) for S. lycopersicum was in agreement with its wider mean vessel radius (20.4 \( \pm \) 7.1 \mu m), compared to that of H. macrophylla (10.9 \( \pm \) 2.4 \mu m). The corresponding vessel radii, obtained with the acoustic model, were 9.6 \( \pm \) 1.2 \mu m and 20.5 \( \pm \) 8.6 \mu m for the two species, respectively. The ultrasound methodology was thus validated for multiple plant species, showing the link between the vessel radii, and the settling time of the ultrasound pulses.

Xylem vessel (element) length and Young’s modulus. To estimate the length \( L \) of the xylem vessel element, we analysed the frequencies in the ultrasound pulses. The resonance frequencies \( f \) are integer multiples of the ratio \( v_{\text{eff}} / L \) (Methods). We found that the Fourier spectra of representative ultrasound pulses (recorded axially) exhibited characteristic peak frequencies (Fig. 3a). The peak frequency with the largest amplitude, \( f_{\text{peak}} \) for sample A was 34 \( \pm \) 5 kHz. In addition, peaks close to integer multiples of \( f_{\text{peak}} \) were observed (Supplementary Table 1). Analysis of pulses from samples B and C showed similar trends (Supplementary Fig. 5). Similar data were observed in the pulses recorded in the radial direction of the stem samples (Fig. 3b, Supplementary Fig. 5, Supplementary Table 2).

The resonance frequency \( f_{\text{peak}} \) was calculated from \( f_{\text{peak}} \) (equation (6); Methods). Note that the two values differ due to the high damping (small \( \tau \)) in the sound pulse. To extract \( L \) from the resonance frequencies, we need the vessel wall thickness \( h \) and the Young’s modulus of elasticity \( E \) (equation (9); Methods). We found \( h \) to be \( \sim 1 \mu m \) via scanning electron cryo-microscopy (Fig. 2c, Methods). We determined \( E \) of stem segments cut from the same plant, and from shoots similar in age and size. For this, we measured the stress-strain curves via uniaxial tensile loading (Fig. 3c, Methods). The mean mass density per stem segment was also estimated from the measured weights and dimensions. The linear slope of the stress-strain curve (Fig. 3c) at small values of strain (\( \approx 10^2 \)) yields the value of \( E \), which was extracted to be 0.2 \( \pm \) 0.1 GPa for fresh (hydrated) stem samples (Fig. 3d). For dry stem samples, \( E > 0.6 \) GPa were obtained. We observed an overall decline in \( E \) with increasing mass density. This indicates that the water-content dominates the variations in \( E \). This agrees with an earlier empirical model, where the dependence of \( E \) on the relative water content in the xylem is taken into account.

We calculated \( L \) using \( h \approx 1 \mu m \) and \( E = 0.2 \pm 0.1 \) GPa (equation (9); Methods). The histogram of \( L \) was extracted from the axially recorded ultrasound pulses for stem samples. For sample A, \( L = 0.99 \pm 0.08 \) mm under a unimodal Gaussian fit (Fig. 3e). Similar values were obtained for samples B and C (Supplementary Fig. 5). This highlights the reproducibility of our method and the similarity of the recorded ultrasound pulses in the axial direction.
We validated the assumption that \( L \) represents the actual length of xylem vessel element. First, we extracted the mean xylem vessel length (a vessel contains several vessel elements) using latex paint staining\(^{27} \), by counting the number of stained vessels on transverse cross-sections of the stem (Methods). These counts decrease exponentially with the distance\(^{50} \) from the lower end of the stem at which the paint was taken up (Supplementary Fig. 6). The mean xylem vessel lengths were found to be in the range ~ 12 – 17 mm for the three stem samples. The xylem vessel length is thus much larger than the \( L \) extracted from the ultrasound pulses (~ 1 mm, Fig. 3e). This is because the Latex paint cannot penetrate the fused ends, but can pass through the perforation plates between adjacent vessel elements\(^{50} \). Next, we observed individual vessel elements in longitudinal sections of stem samples using scanning electron cryo-microscopy (Fig. 3f). The observed length ranged from 0.5 to 0.9 mm for individual xylem vessel elements (Fig. 3g, Table 1). Thus \( L \), as obtained from our acoustic model, is a good estimate of the length of individual vessel elements.

Ultrasound pulsed transmission spectroscopy of xylem vessels. We further elucidated the observed link between the characteristics of drought-induced ultrasound pulses and the xylem vessel anatomy by artificially exciting ultrasonic resonances in the vascular tissue of a stem. A piezo-transducer transmits an acoustic pulse when excited electrically (Methods). This pulse was applied such that it propagated through a stem segment of \( H. \) macrophylla along either the axial or the radial direction, and was subsequently detected by the broad-band microphone (Fig. 4a). Figures 4b and 4e show the time-domain and frequency-domain waveform of the ultrasound pulse detected axially, while Figs. 4d and 4e show the same for the pulse detected radially. The ultrasound pulse exhibited an envelope settling time of 36.3 \( \mu \)s, which was in close agreement with that obtained from the drought-induced ultrasound pulses (26.4 ± 7.0 \( \mu \)s). In both axial and radial directions, characteristic frequencies were observed in the Fourier spectra, which match those observed in drought-induced pulses. This could enable the use of acoustic excitation as a technique for non-invasive monitoring of vascular geometry and moisture-dependent elasticity.

Discussion

Our results have shown how ultrasound emissions from drought-stressed plant stems can be used to extract and monitor the geometry and viscoelasticity of xylem vessels. In this section, we first interpret our results further and discuss the applicability of our method to monitor the vascular physiology of plants. We end the section by commenting on its potential in non-invasive plant health monitoring.

Xylem radius \( (R) \). We have shown that by modelling the xylem vessel as cylindrical acoustic resonator, the radius \( R \) can be extracted from the settling time of the ultrasound pulse, resulting in comparable values as those obtained from common microscopy techniques. Using \( H. \) macrophylla and \( S. \) as example plant species with relatively narrow and wide vessel radii respectively, we validated the dependency of \( R \) on \( L \). Optically determined xylem vessel radii were slightly bigger (~ 2 \( \mu \)m) than the acoustically determined radii (Figs. 2d, 2i, 2j, Supplementary Fig. 4). We attribute this to the assumption of a constant dynamic viscosity of xylem sap \( \eta \). In practice, \( \eta \) depends on ambient temperature, and concentration of dissolved nutrients\(^{31} \). Moreover, water close to the sap-wall interface is held with adhesive forces, and thus has a slightly higher dynamic viscosity\(^{32} \). As a corollary to our analysis, if the distribution of \( R \) is known directly from optical microscopy, one can evaluate the effective kinematic viscosity \( (\eta_l)/\rho \) of the xylem sap. Note that the solid walls of the xylem vessels also possess shear or extensional viscosity\(^{33} \). This means that elastic forces arise in them as a response to elongation, compression or shear stresses. Shear viscosity is a property of solids to resist a change in deformation (shear rate). This additional viscosity likely sets an upper bound on \( R \) and \( L \), whereas the agreement between optical and acoustic radii likely deteriorates.

Xylem vessel element length \( (L) \) and Young’s modulus \( (E) \). The xylem vessel element length \( L \), extracted from the ultrasound pulses (Fig. 3e) consistently exceeded the physical length (via SEM; Fig. 3g) by ~ 0.3 mm (~ 30 \%). We attribute this to two factors. Firstly, the perforation plates serve as non-rigid and leaky boundaries (not accounted for in the model), due to which the standing waves penetrate beyond the physical length of a single vessel element. Secondly, the uniaxial tensile loading measurements that we performed (Fig. 3c, 3d) on stems provide an overestimation of the xylem Young’s modulus. This is due to the presence of stiffer \( S. \) sclerenchyma and \( C. \) collenchyma tissue\(^{34} \), with Young’s moduli exceeding ~ 1 GPa\(^{35} \), close to the circumference of the stem. Hence, as a corollary to our analysis, instead of fixing the Young’s modulus, one can alternatively fix the xylem vessel element length via microscopy. Xylem cells differentiate very early during the growth of a plant\(^{36} \), subsequently growing to their maximum lengths before maturing (dying)\(^{37} \) to become hydraulically active vessel elements. Thus, once the vessel element length is determined via diagnostic techniques for a given plant, the Young’s modulus can then be continuously and non-invasively monitored to diagnose variations in water-content\(^{38} \), ageing, or even pathogen-induced occlusions within the xylem\(^{39,40} \). \n
Relationship between \( L \) and \( R \). Our method of analysing ultrasound emissions enabled us to generate a set of length versus radius data for xylem vessel elements within a given stem segment. We observed that in a single plant \( (H. \) quercifolia\), \( L \) scales as \( R^{0.74} \) (Fig. 5). Basic fluid and structural mechanics can help us in predicting an upper bound on \( L-R \) dependency. In plants of
height within ~1 m, transpiration pull is the governing force of ascent of water through xylem vessels, which creates a gradient in the hydrostatic pressure along the vascular column. With a constant volume flow rate of water through the series-connected vessel elements (continuity), the pressure-drop along a length $L$ can be obtained from the Darcy-Weisbach equation (Methods). Further, a vessel element can withstand a maximum pressure drop to avoid rupture (Methods). This critical pressure is also a function of both $L$ and $R$. Combining the two dependencies, we can derive that $L_{\text{crit}} \propto R^{1.25}$, where Young’s modulus and wall thickness are assumed to be constants. This reasoning gives us an upper-bound on the scaling exponent from a purely mechanical viewpoint.

**Application to intact plants.** Plants vary in their drought-resistance. It may take several days for the water potential in the leaves to fall below the reported threshold for cavitation based ultrasound emissions of at least one per minute. Therefore, we detached plant shoots to induce accelerated drought-stress. This enabled us to record a large set of ultrasound pulses in a relatively short time. We measured similar waveforms in both axial and radial directions of the stem. The latter direction avoids physical incision of the stem and is, therefore, preferred for non-invasive measurements on intact plants. Ultrasound does not propagate far and events occurring within a maximum distance of 20 – 30 mm are likely to be useful. This distance depends on the species, and the level of hydration in the stem, and thus adjusting the proximity of the microphone to the stem may be necessary during growth or movements of the plant. For large trees/shrubs, the radius and lengths of the xylem vessels exceed ~100 μm, and ~10 cm. This would require shifting the sensitive frequency band of the microphone down to the audible range (100 Hz – 10 kHz).

**Impact and scope.** Despite centuries of research into plant hydraulics, our insights into xylem vessel properties and their influence on abiotic and biotic stress resilience, are still constantly evolving. This has largely been possible due to advancement in non-destructive measurement techniques to determine xylem vessel properties. Using methods like latex paint staining and scanning electron microscopy to monitor xylem vessels is time-consuming, and is of limited applicability in the field. Recently, X-ray micro-tomography was recommended in non-destructive measurement techniques to determine xylem vessel properties. Using methods like latex paint staining and latex may be necessary during growth or movements of the plant. For large trees/shrubs, the radius and lengths of the xylem vessels exceed ~100 μm, and ~10 cm. This would require shifting the sensitive frequency band of the microphone down to the audible range (100 Hz – 10 kHz).

We foresee applications of our method to a multitude of plant species with varying vessel dimensions and viscoelasticity. This enables in-vivo studies to mechanical resonances of a plants’ vascular tissue via external acoustic transducers. In turn, this provides a non-invasive method for rapid phenotyping. Crops could be selected for breeding based on their xylem vessels and thus based on their response to drought and/or susceptibility to vascular wilt pathogens. Drought-stress directly impacts the viscoelasticity of the vascular tissue, which can be monitored with ultrasound. Correlation between vessel radius and drought-stress have been reported in poplar and apple trees. Pathogens within the xylem vessels have a parasitic effect on the sugar/nutrient concentration in the sap, which can in turn change the kinematic viscosity of the xylem sap.

Lastly, from the viewpoint of a complete sensor system, the presented methodology only uses Fourier transforms and envelope detection. These are standard signal processing functions, which can be implemented in commercial integrated chip (IC) technology. This will help with future development of low-cost and compact tools for monitoring plant stress. This will in turn boost climate-smart agriculture, and indoor farming by providing farmers with new tools for optimal irrigation strategies and early disease-detection.

**Conclusions.**

We showed for the first time that the radius, length, and viscoelasticity of xylem vessel elements can be co-determined non-destructively and rapidly. This was achieved using a lumped mechanical model of the water-carrying xylem vessel. We analysed the time- and frequency-domain characteristics of ultrasound emission from drought-stressed stems of *Hydrangea quercifolia*. The ultrasound pulses were recorded along the radial and axial direction of stems using a broadband ultrasound microphone. A consistent set of characteristic peak frequencies across a multitude of ultrasound pulses in the range 10-150 kHz was observed. These remotely detected ultrasound pulses were attributed to damped acoustic resonances triggered by bubble formation inside the xylem vessels. We validated the model with results from common destructive methods of optical microscopy, latex paint-staining, scanning electron microscopy and tensile stress testing of plant stems. In particular, we showed that the mean settling time of the sound pulses increases with increasing mean xylem vessel radii through experiments on *Hydrangea macrophylla* and *Solanum lycopersicum*. As a further validation of our method, ultrasound pulsed transmission spectroscopy was performed on stem segments of *H. macrophylla*. A good agreement of the extracted settling time and the characteristic frequencies was obtained with those extracted from drought-induced sound pulses. The presented methodology provides a new outlook on plants “talking” during drought-stress, and presents ultrasound sensing as an inexpensive technique for rapid, non-invasive and in-vivo characterization of plant vasculature.
Methods

Plant material. Three potted plants of *Hydrangea quercifolia* were obtained from a commercial garden center and moved to the laboratory within 1 hour. One shoot sample per plant was cut, keeping the leaves intact, and immediately placed in tap water (Supplementary Fig. 7) to prevent embolism in the xylem vessels at the cut-end. From each shoot sample, a 60-70 mm long and trimmed (i.e., without leaves and petioles) stem segment was cut under water to prevent air entry and blockage. The segments were roughly cylindrical, with a cross-section diameter of ~5-6 mm, and were used for vessel staining and optical microscopy. The rest of the sample was left intact to measure ultrasound emissions. Additionally, one plant each of *Hydrangea macrophylla* and *Solanum lycopersicum*, was also obtained for optical microscopy and ultrasound recording.

Recording ultrasound pulses and signal processing. The shoot samples were taken out of water, dried using tissue paper, and left on the bench for air-drying, resulting in accelerated drought stress. A M500-USB ultrasound microphone, with a reliable detection window between 10 kHz and 150 kHz, from Pettersson Elektronik AB (Uppsala, Sweden) was placed first in the axial (~2 mm from the cut-face of stem normal to the cross-section) and then in the radial (on the cylindrical surface of the stem) directions (Fig. 1a) to record the ultrasound bursts at a sampling rate of 500 kHz in continuous time windows of 120 seconds. The sensor consists of a piezoelectric material which produces an electrical voltage proportional to the pressure of the incident sound wave. From the time-domain waveforms, the pulse envelope was obtained with the built-in “envelope ()” function in MATLAB, which returns the upper and lower envelopes of the input sequence, as the magnitude of its analytic signal. The analytic signal of the input sequence was found using the Hilbert transform. The peak of the envelope curve was determined and the decreasing part of the envelope curve was stored, which was subsequently fitted with the exponential function $\exp(-t/\tau_s)$ using the Least-Squares method. This yielded the settling time $\tau_s$. The frequency spectra of the measured signals were obtained via a 250-point Discrete Fourier Transform, spanning a time frame of 1.5 ms. Due to the low intensity of the emitted sound, the spectra are shown until 150 kHz beyond which the signal merges with the noise floor of the sensor (~80 dB). The raw data was then post-processed and analysed in MATLAB R2018b (MathWorks, Massachusetts, USA).

Analytical model for longitudinal vibrations. We modelled the xylem vessel as a cylindrical pipe of radius $R$, and effective length $L$ sustaining longitudinal standing waves in the water of density $\rho$, whose resonance frequencies depend on the mode order $m$, and the longitudinal speed of sound in the pipe $v_{\text{eff}}$. The resonant frequency of the $m$th order ($m = 1,2,\ldots$) is given by

$$f_m = \frac{m \cdot v_1}{L}$$

where $v_1$ is the speed of sound in the liquid (~1482 m/s in bulk water at 20 °C). We denote the fundamental resonance frequency ($m = 1$) as $f_1$ in the rest of this section. In practice, equation (1) cannot be applied directly because in a real pipe with an elastic wall, sound propagates at a slower speed than that in the bulk liquid. If the walls of the pipe have a non-zero acoustic thickness $h$ and finite Young’s modulus $E$, then the effective speed of sound is given by

$$\left(\frac{1}{v_{\text{eff}}} \right)^2 = \frac{1}{v_1^2} + \rho_h \beta_{\text{xylem}}$$

where $\beta_{\text{xylem}}$ is known as the cross-sectional compressibility, and $\rho_h = 996 \text{ kg.m}^{-2}$ is the mass density of water. Thus, $v_1$ is replaced by $v_{\text{eff}}$ in equation (1).

These sound waves (expected to be dominant in the axially recorded ultrasound) undergo damping primarily due to the dynamic viscosity of water $\eta$ in the xylem. The resulting time-domain response of the resonating pipe can be described using a lumped circuit model consisting of acoustic inductance ($L_a$), capacitance ($C_a$) and resistance ($R_a$), analogous to an electrical L-C-R circuit, where voltage and current are replaced by pressure and flow rate respectively. $L_a$ is a consequence of the kinetic energy in the water, while $C_a$ arises due to the compressibility of water. $R_a$ leads to energy dissipation and can be obtained from Poiseuille’s law for capillary flow. The three lumped parameters can be expressed as:

$$L_a = \frac{L \cdot \rho_1 \cdot \pi \cdot R^2}{\eta_1 \cdot v_1}, \quad C_a = \frac{L \cdot \pi \cdot R^2}{\rho_1 \cdot v_1^2}, \quad R_a = \frac{2 \cdot \eta_1 \cdot L}{\pi \cdot R^4}$$

where $\eta_1 = 8.9 \times 10^{-4} \text{ Pa.s}$ is the dynamic viscosity of water. By describing the circuit as a linear 2nd order differential equation, we obtained the damping ratio $\zeta$, envelope settling time $\tau_s$ (the time needed for the amplitude to decrease by a factor of e), and the driving frequency $f_d$ as:

$$\zeta = \frac{R_a}{2 \cdot L_a} \cdot \sqrt{C_a} = \frac{4 \cdot \eta_1 \cdot L}{\rho_1 \cdot v_1 \cdot R^2}$$

$$\tau_s = \frac{1}{\zeta \cdot f_d} = \frac{\rho_1}{4 \cdot \eta_1} \cdot R^2$$

$$f_d = f_1 \sqrt{1 - \zeta^2}$$

The lumped model is valid as long as the dimensions $L$ and $R$ are smaller than the acoustic wavelength (~1-10 cm in water).

Noting that $f_d$ is the same as the observed $f_{\text{pulsu}}$ in the ultrasound pulses, $\zeta$ is obtained by combining equations (5) and (6) as:

$$\zeta = \frac{1}{\sqrt{1 + \left(f_{\text{pulsu}} \cdot \tau_s\right)^2}}$$
And the acoustic xylem radius was obtained by rearranging equation (5):

\[ R = \sqrt{\frac{4 \eta_0 \tau_x}{\rho}} \tag{8} \]

Combining equations (1) and (2), the effective xylem length \( L \) was obtained as:

\[ \frac{1}{L^2} = \frac{4 \mu^2}{m^2 v^2 \rho} + \frac{4 \lambda^2}{m^2 v^2 \rho} \left( \frac{2 \rho R}{h} \frac{1}{E} \right) \tag{9} \]

Scanning electron (cryo-) microscopy. Transverse sections from hydrangea stems were made using a razorblade. The cross-section was left on filter paper for 1-2 minutes to remove most of the adhering water. Thereafter, the section was fixed to a sample holder using Tissue-Tek. The sample was frozen by plunging the sample holder into liquid nitrogen. Subsequently, the sample was transferred to a cryo-preparation chamber (Leica Microsystems, Wetzlar, Germany) under vacuum where it was kept at -90°C for 3 minutes to remove ice from the surface (freeze etching to remove water vapor contamination). While still under vacuum the sample was coated with 12 nm of tungsten and transferred using a VCT100 shuttle (Leica) to a field emission scanning electron microscope (Magellan 400 from FEI, Oregon, USA). The samples were analysed at 2 kV, 13 pA at -120°C.

Longitudinal sections were made by carefully cutting through the region that contains the xylem vessels. The rest of the sample preparation was identical.

Uniaxial tensile loading for Young’s modulus determination. Multiple stem segments of lengths in the range of 4-7 cm were cut and mounted vertically between two clamps of a tensile testing machine (Z005; Zwick/Roell, Ulm, Germany; inset of Fig. 3c). The initial pre-strained length \( l_0 \) is equal to the vertical separation between the clamps and was kept as 20 mm. The uniaxial stress was calculated as the tensile force applied by the equipment divided by the average cross-section area of the stem segment. The longitudinal strain was calculated as the change in stem length per unit initial length \((\Delta \ell / \ell_0)\). The Young’s modulus \( E \) was then extracted as the slope of the linear part of the stress-strain curve (Fig. 3c) at small strains of strain \((< 10^6)\). The average mass density of each sample was also calculated from measured weight and volume just before tensile loading. The weights were measured with a Sartrex BSC 33 precision balance (Sartex Instruments GmbH, Göttingen, Germany), while the dimensions were measured with a standard Vernier Calliper with a resolution of 0.1 mm. Note that the measurement error for elastic moduli and mass density (\(\approx 20\%\)) is predominantly due to error propagation from length and diameter measurements.

Vessel staining and optical microscopy. An aqueous solution 1 % (v/v) suspension of red latex paint was left standing for at least 24 hours to allow large particles to settle at the bottom. The supernatant was subsequently transferred to a glass container and degassed. The stem segments were mounted vertically over the glass container, with one end immersed in the paint and the other end tightly inserted into a plastic tube connected to a suction pump (to allow large particles to settle at the bottom. The supernatant was subsequently transferred to a glass container and degassed. The stem segments were mounted vertically over the glass container, with one end immersed in the paint and the other end tightly inserted into a plastic tube connected to a suction pump (Supplementary Fig. 7) which applied a pressure difference of 400 mbar. The stem-tube junction was taped and smeared with Vaseline to prevent air leakage. As the solution was sucked through the stem for 12 hours, the paint remained confined in one xylem vessel (macromolecules in the paint cannot move through the bordering pits of xylem vessels) while the clear water was conducted through the entire stem. Subsequently, the stem samples were sliced with a blade at intervals of 5 mm. The number of painted vessel segments were mounted between the transducer and the microphone such that the longitudinal axis of the stem was perpendicular to the line of flight of the sound pulse (Fig. 4a). The transducer was excited with a voltage step of 5 V and an on-time of 500 ms. The Fourier transform of the detected ultrasound pulse was performed over a time span of the first 100 μs (Fig. 4d), to observe the frequency components present in the pulse that propagated only through the stem.

Darcy-Weisbach equation and critical pressure in xylem vessel. The Darcy-Weisbach equation is an empirical relation that relates the pressure drop \( p \) along a given length \( L \) of a viscous and incompressible fluid flowing through a conduit of radius \( R \) as:

\[ p = \frac{8 \mu L}{D^2} \]
The data that support the findings of this study are available from the corresponding author upon reasonable request.

**Data availability**

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

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Acknowledgments

This work has been carried out under the “Plantenna” research programme, a collaboration among the members (technical universities) of the 4TU federation in the Netherlands. The authors would like to thank Lars Pettersson for technical support.
concerning the ultrasound sensor equipment, dr. ing. Marcel Giesbers for support with scanning electron microscopy, and Patrick van Holst for technical support in measuring the stress-strain curves of the stem samples.

**Author contributions**

S.D. and G.V. conceived the idea. G.V., E.K., and P.G.S. supervised the work. S.D. performed the ultrasound measurements, elastic modulus measurements, and the analysis/modelling of data. E.K. obtained the plant specimens. P.M.M. and S.D. carried out the paint infusion method and analysed the microscopic images for determining vessel length and diameter. E.K. and P.M.M. provided necessary advice on plant physiology. S.D., and G.V. prepared the manuscript. E.K., P.G.S. and G.V. revised the manuscript. All authors read and approved the manuscript.

**Competing interests**

The authors declare no competing interests.

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**Additional information**

Supplementary information is available for this paper.
Listening to ultrasound from plants reveals xylem vessel anatomy (Figures)

Figure 1: Ultrasound measurement set-up and time-domain signal. (a) Schematic set-up for recording of ultrasound pulses from shoots of *Hydrangea quercifolia* along axial and radial directions. The zoom-in represents a schematic of the vascular bundle of the stem, showing the peripheral arrangement of tubular xylem vessels around the pith in the core. Bubbles seeded in the vessels trigger the emission of ultrasound pulses. (b) Example raw time-domain data for ultrasound recorded along the axial direction of a stem. Time $t = 0$ represents the start of the recording, which occurs after ~5 minutes of drying. (c), (d) Zoomed-in time-domain example ultrasound pulses recorded axially and radially, respectively. Black curves represent the amplitude envelope that decays exponentially (damping), and blue curves represent the exponential fit of the envelope decay. (e) Schematic flow-chart illustrating the steps in our analysis. The settling time and peak frequencies are obtained from the time-domain and frequency-domain waveforms (Figs. 2e, 2f, 3a, and 3b). The resonant frequency is obtained from peak frequency and settling time (Methods, equation (6)). Using these, xylem radius and xylem vessel element length are extracted (Table 1). Parameters of sap viscosity, vessel wall-thickness and young’s modulus are taken as input.
Figure 2: Xylem vessel radius extraction from damping in axial sound waves. (a) Schematic of water-conducting xylem vessels in vascular plants. Vessels are the dominant and more efficient cells to transport water in *H. quercifolia*. They consist of a series network of vessel members/elements, which are interconnected via perforation plates. Also shown is the simplified cylindrical acoustic-resonator model for a vessel element sustaining damped longitudinal standing waves in its sap, the damping factor being a function of sap viscosity and the radius of the vessel. (b) Optical micrograph of the transverse section of stem sample A, showing the xylem vessels filled with latex paint. (c) Cryo-SEM image of the transverse section of a stem sample showing the diameter (2R) and the wall thickness (h) of a xylem vessel. (d) Histogram showing the model-extracted xylem radii (in blue), and that of the observed xylem radii (in yellow) obtained via latex-staining and optical microscopy for stem sample A of *H. quercifolia*. The red curve represents a unimodal Gaussian fit. (e),(f),(g) Time-domain ultrasound waveform, cross-section optical micrograph (300X) of stem, and histograms of observed (in yellow) and acoustic (in blue) xylem vessel radii, respectively, in *H. macrophylla* stem sample recorded along the axial direction. (h), (i), (j) Time-domain ultrasound waveform, cross-section optical micrograph (200X) of stem, and histograms of observed (in yellow) and acoustic (in blue) xylem vessel radii, respectively, in *Solanum lycopersicum* stem sample recorded along the axial direction.
Figure 3: Ultrasound frequency spectra, Young’s moduli of stem, and extraction of xylem vessel element lengths. (a), (b) Observed characteristic peak-frequencies in the example Fourier transform of the ultrasound pulses recorded axially and radially, respectively, in sample A of *H. quercifolia*. The black curve represents the spectrum of a representative pulse with the indicated timestamp of the recording. Grey shaded regions indicate the variation in the peak frequencies among the individual pulses (means ± standard deviation). (c) Measured stress-strain curve for freshly cut stem segments of *H. quercifolia* with mass densities indicated in the legend. The blue line represents the linear fit of a stress-strain curve at low strain, the slope of which, yields the Young’s modulus in accordance with Hooke’s Law. The inset shows the photograph of the set-up for uniaxial tensile loading of the stem. (d) Extracted Young’s modulus versus mass density for freshly cut and dried stem segments, extracted from stress-strain measurements (solid circles) with indicated error bars (± 0.1 GPa). The red curve represents a fit based on the empirical model of Young’s moduli as a function of relative water-content in stems. (e) Histogram showing the extracted xylem vessel element lengths in stem sample A, extracted via the acoustic model. The black curve represents a unimodal Gaussian fit. (f) Example cryo-SEM image of the longitudinal section of a stem segment (*H. quercifolia*), showing the structure of individual vessel elements terminated by scalariform perforation plates (marked by red arrows). Observed length of each element is in the range 600 μm – 900 μm. (g) Scatter plot showing the observed length of xylem vessel elements via cryo-SEM technique. The grey bars indicate the upper and lower bounds due to the slant length of the perforation plates along the longitudinal axis (~ 100 μm).
Figure 4: Ultrasound pulsed transmission spectroscopy of vascular tissue. (a) Schematic measurement set-up showing an external piezo-ultrasound transducer (resonant frequency of 40 kHz) and the broad-band microphone placed along the axial and radial direction to a *H. macrophylla* stem with the indicated dimensions. The piezo-transducer is excited with a voltage step in order to emit a broad-band acoustic pulse. (b) Time-domain waveform (in blue) with the fitted envelope, and (c) Fourier spectra of the axially transmitted sound pulse (in blue). The spectrum of the sound pulse emitted by the transducer (source) with transmission through air is shown in grey as a reference. (d) Time-domain waveform and (e) Fourier spectra of the radially transmitted sound pulse.

| Parameter                      | Ultrasound method     | Destructive measurement                                      |
|--------------------------------|-----------------------|-------------------------------------------------------------|
| Xylem radii [μm]               | 9.89 ± 1.6            | 12.4 ± 2.6                                              |
| Mean Xylem vessel length [mm]  | ---                   | 16.9 (sample A), 12.8 (sample B), 14.4 (sample C).        |
| Xylem vessel element length [mm]| 0.98 ± 0.14           | 0.63 ± 0.14                                              |
| Young’s modulus of elasticity [GPa]| ---                   | 0.2 ± 0.1                                                |

Table 1: Summary of extracted parameters via the ultrasound analysis for the three stem samples of *H. quercifolia*.
Figure 5: Acoustic xylem vessel element length versus vessel radius. Black circles: scatter plot of model-extracted (acoustic) xylem vessel element length ($L$) and radius ($R$) corresponding to each analysed ultrasound pulse. Data from all three stem samples of *H. quercifolia* are merged here. The radii were obtained from the settling time of the ultrasound pulses, while the lengths were obtained from the resonance frequency of the sound pulses and by measuring the Young’s modulus (Fig. 1e). The data points are classified into bins of $L$ with intervals of 0.5 mm. In each bin, the median $L$ and $R$ are calculated (pink squares), and fitted with a power law function (blue line). The green dashed line indicates the predicted $L$-$R$ dependency in accordance with Darcy-Weisbach equation for fluid flow, combined with mechanical failure of the xylem vessel (Methods). Red circles: observed vessel element lengths and radius in *H. quercifolia* via cryo-SEM technique.
Figures

Figure 1

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Supplementary Files

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- SUPPL1.pdf