Grapevine petioles are more sensitive to drought induced embolism than stems: evidence from in vivo MRI and microCT observations of hydraulic vulnerability segmentation.

Running title: Hydraulic Vulnerability Segmentation in Grapevine

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Abstract

The “hydraulic vulnerability segmentation” hypothesis predicts that expendable distal organs are more susceptible to water-stress induced embolism than the main stem of the plant. In the current work we present the first in-vivo visualization of this phenomenon.

In two separate experiments, using Magnetic Resonance Imaging (MRI) or synchrotron-based micro computed tomography (microCT), grapevines (Vitis vinifera) were dehydrated while simultaneously scanning the main stems and petioles for the occurrence of emboli at different xylem pressures ($\Psi_x$). MRI imaging revealed that 50% of the conductive xylem area of the petioles was embolized at a $\Psi_x$ of -1.54 MPa, whereas the stems did not reach similar losses until -1.9 MPa. MicroCT confirmed these findings, showing that approximately half the vessels in the petioles were embolized at a $\Psi_x$ of -1.6 MPa, whereas only few were embolized in the stems. Petioles were shown to be more resistant to water-stress induced embolism than previously measured with invasive hydraulic methods.

The results provide the first direct evidence for the hydraulic vulnerability segmentation hypothesis and highlight its importance in grapevine responses to severe water stress. Additionally, these data suggest that air entry through the petiole into the stem is unlikely in grapevines during drought.

Key words: MRI, microCT, grapevine, cavitation, hydraulic conductance, Vitis vinifera, xylem, vulnerability curves
Introduction

Water flows through xylem conduits at negative pressures (i.e., tension) that can be well below vapor pressure. Under such tensions, water is in a metastable state (Dixon 1914) and at risk of cavitation that results in subsequent loss of function in the conduit (Tyree & Sperry 1989). The concept of ‘plant segmentation’ was introduced over 30 years ago (Zimmermann 1983) and proposes that hydraulic constraints enable the plant to sacrifice organs of lesser importance and investment, in order to save organs that are critical for long-term survival and propagation. Zimmerman (1983) suggested ‘hydraulic segmentation’ as a mechanism in which high resistance results in a high gradient of xylem pressure between the basal and distal plant parts. Eight years later, Tyree & Ewers (1991) coined the term ‘vulnerability segmentation’, suggesting a second mechanism in which expendable organs (e.g. leaves) in woody plants are more susceptible to embolism, compared to more permanent structures. Supporting this idea, cavitation vulnerability curves of petioles and stems in walnut trees (Juglans regia L.) showed that petioles are significantly more susceptible than stems; petioles lost 50% of their hydraulic conductivity at xylem pressures ($\Psi_{50\%}$) of -1.5 MPa, as compared to -2.2 MPa in stems (Tyree et al. 1993). Similarly, Tsuda & Tyree (1997) reported $\Psi_{50\%}$ of -2.0 and -0.5 MPa for stems and petioles, respectively. Choat et al. (2005) have even shown that in sugar maple trees (Acer saccharum Marsh.) the increase in air seeding threshold of different organs is a function of their distance from the trunk. While this pattern has been documented in some species, the opposite was found in others (i.e. stems more susceptible than leaves; Cochard et al. 1992), suggesting that the phenomenon is not universal across species.

A comparison of vulnerability curves collected by different research groups suggests that grapevines (Vitis vinifera L.) belong to the group of species that do exhibit vulnerability segmentation, with stems that are more resistant to water-stressed induced embolism than the petioles (Alsina et al. 2007, Choat et al. 2010, Zufferey et al. 2011, Tombesi et al. 2014). A comparison of acoustic emissions in detached stems and petioles of Syrah and Grenache vines revealed that most of the petiole vessels embolized at a xylem pressure ($\Psi_x$) of -1.5 MPa, whereas the formation of emboli was still occurring in the stem at $\Psi_x$ as low as -2MPa (Schultz 2003). A comparison performed by Lovisolo et al. (2008) of embolism in different organs of water stressed grapevines showed that at $\Psi_x$= -1.4 MPa 80% loss of conductance (PLC) was measured in the petioles, whereas the stems exhibited only 45% PLC. These lines of evidence suggest that there is a high probability for vulnerability segmentation in grapevines (Zufferey et al. 2011), although this hypothesis has yet to be explicitly tested. Furthermore, the debate about the validity of previous measurements of embolism (Cochard, Delzon & Badel 2015) calls for a direct measurement and observation of the phenomenon in vivo.

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Over the last five years, the validity of previous embolism measurements in long vessel species, such as grapevines, have been questioned. It appears that measurements of some species with a large population of long vessels (i.e. longer than the length of the sample) are prone to a bias when the centrifuge technique is used or when cutting stem samples under tension (even when under water; Choat et al. 2010, Wheeler et al. 2013, Torres-Ruiz et al. 2015), calling for non-invasive methods to measure or visualize the formation and spread of emboli in vivo. Magnetic Resonance Imaging (MRI) and X-ray computed micro-tomography (microCT) have been successfully used to visualize embolism formation in a number of studies (Holfbrook et al. 2001, Scheenen et al. 2007, Choat et al. 2010, Zwieniecki, Melcher & Ahrens 2013, Knipfer et al. 2014, Brodersen et al. 2010, Brodersen et al. 2013).

In the current study we used MRI and microCT as complementary diagnostic imaging tools to evaluate embolism formation in stems and petioles while they are still connected to one another. Both imaging techniques were used to test the vulnerability segmentation hypothesis independently and to validate previous indirect measurements of grapevine petiole vulnerability curves.

Materials and methods

This study consisted of two separate experiments. Both compared the occurrence of xylem emboli in the petioles and stems of drying grapevines. In the first experiment four plants were disconnected from their root system by cutting the rootstock and subjected to fast dehydration for a time period of between 12-40 hours, while continuously monitoring events of embolism formation by MRI. To acquire improved spatial resolution images of the phenomenon and to validate that results were not affected by cutting the rootstock, a second experiment utilized synchrotron-based microCT technology to observe the same phenomenon. Intact plants were dried to the Ψ50% that was observed in the MRI experiment and then imaged.

Plant material

The MRI experiment was conducted on one year old Syrah vines grafted on SO4 rootstocks, grown in commercial potting soil (Einheitserde Classic, type ED73) in 10 L pots. During the 65-85 days growth period, between budbreak and the imaging, the vines were grown in a growth chamber at 24°C and 80% relative humidity. Artificial lighting of 300 µmol m⁻² s⁻¹ photosynthetic photo flux density (PPFD) with 14/10 day/night cycles was used. The vines were pruned to yield a single long stem without side branches to fit the bore of the MRI imager, and were approximately 2.5 m long and had between 15 and 19 leaves at the time of imaging. Irrigation to soil saturation was applied
daily prior to imaging. Random sampling of the xylem water pressure with a pressure chamber was done to ascertain that vines were maintained at a xylem pressure higher than -0.5MPa.

The four plants utilized in the microCT experiment were own-rooted Syrah grapevines donated by Foundation Plant Services (Davis, CA, USA) and grown in 0.5L pots containing a potting mix (Berger BM-6- with a composition of mostly dolomite, perlite, peat moss and micro nutrients) in a UC Davis glasshouse facility. The plants were in their first year, eight weeks after budburst. All samples were maintained in a well-watered condition prior to use in the experiment and received 10% Hoagland solution twice a day. Watering for the drought-stressed plants was ceased approximately one week prior to the experiment and the pots dried down slowly over this time. Samples were transported to The Advanced Light Source at Lawrence Berkeley National Lab, Berkeley, CA, USA (ALS) on the same day of microCT scanning.

**Preparation and installation of the vines in the MRI scanner**

The MRI experiment was designed to image embolism formation in fast drying xylem with no open vessels in the imaged area. To study the maximal vessel length originating from the rootstock, we followed the procedure described by Ewers & Fisher (1989). Plants were cut in the middle of the rootstock and air was forced at 200 kPa into the open cut. The apical part of the stem was immersed in water and progressively cut back in two cm intervals until bubbles were observed escaping the cut surface. This method verified that no xylem vessel crossed the grafting union to more than 30 cm.

MRI measurements were conducted on four plants that were dark acclimated for ~2 hours in order to maximize leaf water content. The dehydration occurred under dark conditions inside the MRI bore and was similar to the bench top dehydration as described by Choat et al. (2010). The imaged area was approximately 60 cm above the cut end and 50 cm (3-6 internodes) above the grafting point, while the stem apex was still intact. The petiole and a reference tube with a 10mM Ni(NO$_3$)$_2$ solution were fixed parallel to the stem by means of parafilm and subsequently enclosed in a 5 mm openable RF coil (Fig. 1a,d). Four leaves were enclosed in plastic bags and later used to measure $\Psi_x$. The rootstock was cut under water and the vine transferred into the MRI setup, while at all times keeping the cut end of the rootstock submerged in a 50ml tube filled with water. The first image in the measurement series was acquired with the cut end still submerged under water and at xylem pressures no lower than -0.3 MPa, as was verified by measurements obtained immediately after imaging. After acquiring the first image the cut end of the vine was exposed to air by sucking the water out of the 50ml tube from outside the imager. In this way the vines were
allowed to commence drying, without having to remove the vine from the closed cylindrical bore of the magnet and without causing movement of the sample. All subsequent images were taken while the vines were drying inside the MRI under continuous darkness at a temperature of 19°C. Three of the four vines were dried for a period of between 12 and 16 hours, ultimately reaching an average Ψₓ of -1.6 MPa. The fourth vine was left to dry in the MRI for a period of 40 hours, reaching a Ψₓ of -1.94 MPa. In the latter case, Ψₓ was extrapolated from the relative petiole water content (as described below).

**MRI imaging**

MRI imaging was done using a superconducting, vertical bore 4.7T Varian VNMRS MRI system (Varian, Palo Alto, USA) fitted with a 300 mT/m gradient set with an inner diameter of 205 cm and a custom built openable four turn solenoidal RF coil of 5 mm in diameter. The RF coil was mounted at an angle of 45° with respect to the vertical main magnetic field (B0). CPMG (Carr–Purcell–Meiboom–Gill) amplitude-T₂ measurements were performed using the following settings: field of view, depending on object size, between 8 x 8 and 12x12mm; slice thickness: 2 mm; matrix size: 128 x 128; number of averages: 2; echo time: 4 ms; number of echos: 32; repetition time: 5s; total scan time per image: 25m; and spectral width: 50 kHz. The acquired MRI datasets were processed using fitting routines written in IDL software (Research Systems Inc., Boulder, Colorado, USA). The datasets were fitted on a per pixel basis using a mono exponential decay function (van der Weerd et al. 2000), which yielded quantitative maps of amplitude and T₂ (Donker et al. 1997, Edzes, Van Dusschoten & Van As 1998). In the current study only the resulting amplitude maps are shown.

All imaged petioles were later fixed in 70% ethanol and were embedded and sectioned following the methods of (David-Schwartz et al. 2013). Xylem vessel diameter were analyzed using imageJ (Abramoff et al., 2004).

**Xylem pressure (Ψₓ)**

Measuring Ψₓ continuously is never trivial, but even more so inside an MRI scanner where the strong magnetic field prohibits the use of magnetic materials. To overcome the problem we combined the pressure bomb method with the MRI capability to accurately measure water content, in order to create a model linking the two. In four leaves per plant Ψₓ was measured using a pressure bomb (Soil Moisture Equipment Corp., USA) according to the procedure described by Scholander et al. (1964). The measurements were taken before emptying the 50ml tube (~0.3MPa), 25 minutes after drying (~0.65MPa), 100 minutes after drying (~1.2MPa) and 12-16 hours after drying (~-
Measurements were performed on fully expanded leaves that were bagged (enclosed in plastic bags) for at least 30 minutes. Each leaf was excised from the stem using a sharp blade and then placed into the pressure chamber with the petiole protruding from the chamber lid. The chamber was pressurized using a pressurized nitrogen tank, and Ψₓ was recorded when the initial xylem sap was observed emerging from the cut end of the petiole.

The Ψₓ for all other MRI samples, apart from the four measurement points that were done as described above, were derived from a modification of the pressure-volume model (Turner 1988) applied to the petiole (Fig. 2). This model describes the relation between the cell’s water content and water potential allowing the conversion of one to the other. We assumed that such a model would be more powerful applied to the petiole (rather than the stem) due to its composition of larger portion of parenchyma cells (Fig. 1C). The water content of each petiole was quantified using ImageJ (Abramoff et al., 2004) as the sum of the amplitudes of all its pixels normalized to the sum of all the pixels of the H₂O reference tube. Assuming that dark acclimated well-watered plants are close to 100% relative water content (RWC), the petiole’s water content of each image was normalized to that of the first image. The 3rd and 4th pressure bomb Ψₓ measurements (at ~ -1.2 and -1.6MPa) were taken after wilting had occurred; their RWC was subsequently linearly regressed with 1/Ψₓ to model the osmotic modulus (Fig. 2; R²=0.92).

**Quantification of the degree of embolism formation**

To be able to fit the stem together with an intact petiole and a reference tube in the field of view (FOV) of the MRI scanner, a large FOV by necessity was chosen. This limited the spatial resolution that could be obtained to a pixel size of 40x40 µm, which is larger than most vessels of the petioles (Hochberg et al. 2015). This precluded a straightforward embolism evaluation based on vessel count as demonstrated before (Choat et al. 2010, Holbrook et al. 2001). To overcome this problem we had to determine embolism based on a per pixel basis, with the assumption that a pixel within the xylem area that changed from water-filled to air-filled had embolized and was non-functional. We assessed the degree of embolism through binary transformation, using ImageJ (Abramoff, Magalhães & Ram 2004). Binary transformation was used to assure that parenchyma cells would not be counted as embolism. Since the values of the parenchyma cells were much closer to the values of conductive xylem vessels than to those of empty xylem vessels, binary transformation always tagged them as conductive tissue. The organs shrinkage during dehydration was accounted for by modifying the analyzed area to match the boundaries of the petiole or stem.
Zero value pixels (marked in white in Fig. 1b,e) were counted and assumed to be embolized (pix\textsubscript{emb}). The percent of embolism formation (\%\textsubscript{emb}) was calculated as:

\begin{equation}
\text{(1) } \%\text{emb} = 100 \times \frac{\text{pix}_{\text{emb}} - \text{pix}_{\text{initial}}}{\text{pix}_{\text{max}} - \text{pix}_{\text{initial}}}
\end{equation}

\text{pix\textsubscript{initial}} is the number of zero value pixels in the first image, and \text{pix\textsubscript{max}} is the number of zero value pixels at 100% embolism inflicted by cutting the vessels open (Fig. 1d,e). The normalization to \text{pix\textsubscript{initial}} means that \%\text{emb} represents the appearance of new emboli, but does not account for emboli that were already present before the start of the experiment (this, in contrast to the microCT embolism quantification method – described below). The two are probably similar, but not identical since even under well-watered condition (as the plants were grown) a certain degree of embolism (<10\% of the total vessel population) is expected (microCT results in Figure 5). The normalization to \text{pix\textsubscript{max}} suggests that any effect that was not accounted for by the binary transformation was normalized, as it was probably also present in the last image.

To quantify \text{pix\textsubscript{max}} both the stem and petioles were cut directly under the coil, less than 1 cm beneath the imaged tissue, at the end of the dry down experiment. To verify that indeed the majority of vessels were longer than 1 cm and would therefore be embolized when cut under tension, we performed the following test. Short stem and petiole segments (5 cm) were flushed of all native embolism with membrane filtered distilled water (1 minutes, 150 kPa). The same segments, now free of emboli, were perfused with air at high pressure (200 kPa) in order to embolize all the open xylem vessels. The segments were divided under water with a sharp razor blade into 2 smaller (1 cm and 4 cm) segments, 1 cm away from the pressurized end. The percent loss of conductance (PLC) of both segments was evaluated using the Xylem apparatus (Xylem Embolism meter, Instructec, Montigny-les-Cormeilles, France) as described by Zufferey et al. (2011). In the 4 cm segment, 93.4\% and 100\% of the conductance of petioles and stems, respectively, was lost indicating similar percentage of conducting area is open 1 cm away from a random cut.

**MicroCT imaging**

The xylem pressure in the stem was measured on bagged leaves with a Scholander Pressure Chamber (Soil Moisture Co., Santa Barbara, CA, USA), ~10-40 min before scanning each sample. Intact plants were prepared for scanning by carefully placing the axis of a petiole against the stem internode that it subtended, and these were held together with Kapton\textsuperscript{\textregistered} tape (DuPont, Wilmington, DE, USA). The petiole and stem were scanned simultaneously, and were located towards the base of each plant.
Imaging was conducted following the methods described by McElrone et al. (2012). The scans took approximately 6 minutes to complete using the continuous tomography setting at 20 keV at the Lawrence Berkeley National Laboratory Advanced Light Source, beamline 8.3.2. A 2x lens was utilized to produce images with a voxel resolution of 3.19µm. Images were then reconstructed using Octopus 8.3 software (Institute for Nuclear Sciences, University of Ghent, Belgium) and analyzed with ImageJ software (Abramoff et al., 2004) to count water-filled and embolized vessels. The high spatial resolution allowed the identification of individual vessels. The degree of embolism was quantified by dividing the number of embolized vessels by the total number of vessels.

**Statistical analysis**

In the MRI experiment, apart from the first five images, where dehydration was very fast, all images of the same vine in 0.1MPa intervals were averaged, both in their Ψx and %emb. Values were averaged between the four replicated vines (n=4) and compared in a T-test with the matching value of the other organ (petioles vs. stems). To evaluate Ψ 50% sigmoidal curve fitting was performed using SigmaPlot 11 by adjusting %emb versus Ψx as follows:

\[
(2) \%_{emb} = \frac{100}{1 + e^{a(Ψ_x - Ψ_{50})}}
\]

The slope (s) at the inflection point was calculated as:

\[
(3) s = a \times 25
\]

The averages acquired in the microCT experiment were compared with a T-Test.

**Results**

MRI was used to continuously and non-invasively measure the appearance of emboli in the stem and petiole of plants that were allowed to dry down (Fig. 3; also see supplementary time-lapse animation S1). Embolism formation was almost negligible for both organs in the first 100 minutes of dehydration, as the Ψx of the vines dropped from -0.3 to -1.2MPa. Significant embolism formation was not noticed in either organ until values lower than -1.2MPa were reached. Significant differences (P<0.01) in the %emb between stems and petioles were measured at all pressures less than and equal to -1.3MPa. The quantification of %emb resulted in two significant (p<0.01 for ‘a’ and Ψ 50%) sigmoidal vulnerability curves for the stems and petioles (Fig. 4). Ψ 50% was 0.37MPa lower
(more negative) in the stems (-1.9MPa) than in the petioles (-1.54MPa). The slope at the inflection point of the petiole trace (159%/MPa, p<0.001) was nearly twice as large as that of the stems (87%/MPa, p<0.001) once embolism started to occur, further confirming the larger vulnerability of the petioles. In the stem, the larger vessels allowed the continuous visualization of the spatial pattern of embolism formation.

The microCT imaging experiment confirmed the existence of the aforementioned vulnerability differences. The petioles exhibited a much larger degree of embolism than the stems at a similar $\Psi_x$ (Fig. 5). In fact, the MicroCT results suggest an even larger difference between the degree of embolism of petioles and stems when compared to MRI results. The high spatial resolution of microCT makes it possible to spatially resolve and count individual vessels. It could thus be determined that at a $\Psi_x$ of -0.2MPa both organs had very little embolized vessels (3-9% of vessels embolized). At -1.6MPa in the petioles 45% of the vessels were embolized, whereas in the stems on average only 12% of the vessels were embolized.

**Discussion**

Our results provide the first non-invasive observations of the hydraulic segmentation phenomenon *in vivo*, and indicate that in intact Syrah plants under water deficit petioles lose conductivity at less negative $\Psi_x$ than stems (Fig. 3; supplementary time-lapse animation S1). These observations of hydraulic segmentation in *V. vinifera* agree with previous measurements in deciduous plants (Tyree et al. 1993, Tsuda & Tyree 1997, Choat et al. 2005, Lovisolo et al. 2008), giving further support to the hypothesis proposed by Tyree & Ewers (1991) that this strategy may be common in woody, deciduous plants. Our methods also provided a unique opportunity to simultaneously study the functional status of two hydraulically linked organs, and to then generate vulnerability curves to predict their respective loss of hydraulic conductivity.

Greater embolism resistance is often associated with smaller vessel diameters, as they are likely to have a lower surface area and fewer intervessel pits compared to larger diameter vessels (Wheeler et al. 2005). The average vessel diameter of the stem (25.44 ± 1.09 µm) is significantly larger than that of the petiole (9.95 ± 0.53 µm), and the stem’s larger diameter vessels should, theoretically, be more vulnerable to embolism formation and spread. Our data suggest a different origin for the differences in embolism resistance. As the primary xylem is more sensitive to embolism than the secondary, the larger primary/secondary xylem ratio in petioles (Choat et al. 2005) could, in part, lead to the differences in embolism vulnerability reported here. Additionally,
there may be other xylem network properties (e.g. pit morphology, pit membrane thickness, vessel ending frequency, xylem/tracheid proportion, conduit redundancy, etc. Jansen, Choat & Pletsers 2009, Guet et al. 2015) that were not considered in this study and may play an important role in the differences in cavitation resistance between the two tissue types.

Measurements of Ψₓ were conducted through leaf bagging, thereby stopping leaf transpiration and allowing the xylem pressure of the leaf to equilibrate with that of the stem. The Ψₓ values reported here thus represent an average of the Ψₓ of both the petiole and the stem. Because of the equilibration between the two organs, we were not able to determine the Ψₓ gradient between the stem and petiole and thus could not determine whether the differences in cavitation resistance between the two organs are a product of vulnerability to embolism (Tyree & Ewers 1991) or hydraulic segmentation (where Ψₓ is expected to be more negative in distal organs; Zimmermann 1983). However, the contribution of hydraulic segmentation to these observations is questioned, as it will require high transpiration rates, which are not likely to occur in the dark conditions under which the MRI experiment took place. The phenomenon is more likely to occur when plants were illuminated, such as during the microCT experiment, which might explain the larger differences that were observed as compared with MRI. Under field conditions, pressure differences of 0.2 MPa between grapevine leaves and stems are commonly found (Williams & Araujo 2002), and thus hydraulic segmentation may increase the differences in embolism between the two organs.

Zwieniecki et al. (2013) showed that reliable vulnerability curves can be generated using MRI, even in cases where the spatial resolution does not allow the individual vessels to be fully resolved. As Zwieniecki et al. (2013) used an air injection method to induce embolism, water loss from non-vessel cells was minor, allowing the quantification of PLC on the basis of pixel brightness. This was not the case during the long dehydration we performed in the current study, which will have resulted in significant water loss from parenchyma and fiber cells and thus in an equally significant loss in MRI signal amplitude. We circumvent this problem by means of the binary image transformation approach, which enabled us to detect the occurrence of emboli, irrespective of the water status of the surrounding tissue. The vulnerability curves obtained by means of MRI and microCT were consistent with each other and in good agreement with the data in the literature. The stem vulnerability curves were similar to the ones published by Choat et al. (2010), who employed high resolution MRI as well as traditional bench dehydration of long stems to acquire their PLC values. In addition, Brodersen et al. (2013) reported a vulnerability curve for small V. vinifera (cv Chardonnay) stems with microCT that agree well with the MRI curves presented here (Fig. 4).
At \( \Psi_x \) of down to -1.1 MPa we observed the formation of only a negligible number of emboli (\( \%_{emb} \)) close to 0, fig. 4) in both stem and petiole. These findings, which are the first in vivo visualizations of embolism in grapevine petiole, strongly contrast with previous invasive measurements. Zufferey et al. (2011) and Lovisolo et al. (2008) already found a substantial degree of PLC (40%) in petioles of grapevines at \( \Psi_x \) above -1 MPa. Tombesi et al. (2014) measured \( \Psi_{50} \) values that were slightly lower (-1.08 or -1.25 MPa for Montepulciano and Sangiovese cultivars, respectively), but also already measured about 30% PLC at a high \( \Psi_x \) of -0.5 MPa. These differences suggest the existence of a discrepancy between invasive and non-invasive methods of embolism evaluation. The difference cannot be attributed to the methodical differences in calculating PLC and \( \%_{emb} \) (percentage loss of conductivity vs. % of vessels embolized), as such a distinction cannot be claimed when \( \%_{emb} \) is zero. A more likely explanation was provided by Wheeler et al. (2013). These authors showed that cutting the xylem under tension may artificially introduce emboli into the xylem and result in apparent higher degree of PLC. Evidence suggesting that the studies by Lovisolo et al. (2008) and Zufferey et al. (2011) may have suffered from this artefact is provided by the observation that most PLC disappeared as soon as \( \Psi_x \) was increased when evening approached. Nonetheless, since the Wheeler et al. (2013) findings cannot be generalized to be valid for all species (Wheeler et al. 2013; Venturas et al., 2015), and since concerns about potential fast xylem refilling were raised (Trifilo et al., 2014), there still is a need to more closely test for the existence of a cutting artefact in grapevine petioles. Alternatively, the high PLC found at \( \Psi_x \) could also have resulted from emboli that already had been introduced before, for example during episodes when \( \Psi_x \) was more negative, and were never completely removed from the system. Our data thus suggest that in grapevine, the formation of emboli in the stem and petiole at high \( \Psi_x \) is more rare an occurrence than previously presumed.

It has been hypothesized that emboli could play a role in the regulation of transpiration. Zufferey et al. (2011) suggested that vulnerability segmentation could be an efficient mechanism for grapevines to regulate stomatal opening. The xylem vessels in the petioles could act as “hydraulic fuses” and their embolization would thus limit leaf transpiration. However, it seems that embolism formation in stem and petioles of grapevines does not occur until \( \Psi_x \) reaches values that are much more negative than the ones inducing stomatal closure. Even more so, the hypothesis that the induction of embolism could act as a signal for stomata closure (Nardini & Salleo 2000) is not supported by our findings. For any kind of regulation to be associated to the appearance of emboli, the emboli should precede the proposed effect. Stomata conductance is normally reduced by half (compared with well-watered conditions) when water potentials are reduced to -0.75 MPa (Zufferey et al. 2011, Tombesi et al. 2014, Hochberg et al. 2013). In the current study, at such \( \Psi_{so} \), we were not
able to detect any embolism in the petiole or in the stem (fig. 4). Similarly, the reduction of leaf and vine hydraulic conductance normally appear shortly after dehydration starts (Zufferey et al. 2011), therefore the involvement of stem or petiole embolism is not likely. Rather, it reinforces the view that vulnerability segmentation serves as a second line of defense in the event that leaf conductance declines and stomata regulation fails. Only under severe stress the petioles would experience significant embolism induction, thus limiting further water loss and most importantly, assuring the preservation of the functional integrity of the xylem in the stem. This view is corroborated by the observation that grapevines tend to shed their leaves under severe water stress, yet often are able to recover rapidly upon irrigation. It should be noted that this does not necessarily mean that embolism did not play a role at all. Embolism formation in the leaf blade may precede embolism formation in the petioles and thus reduce leaf conductance and limit transpiration (Johnson et al. 2012, Nolf et al. 2015).

In conclusion, in the current study we present the first in vivo observations of vulnerability segmentation in intact plants. Under water deficit, xylem vessels in the petioles of grapevines on average embolize before vessels in the stems do. Our vulnerability curves suggest that embolism induced xylem failure in grapevine may be a much rarer occurrence than is commonly assumed.

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References


Figure 1. Assessment of embolism degree. Binary transformation was applied to all MRI images to dichotomously distinguish between conductive (black) or non-conductive (white) pixels. The percent of embolism of each image was calculated after normalization to the number of pixels at full conductivity (A, Bpixel) and the number of pixels at full embolism (pixmax). Pixmax was validated by comparison to xylem area (in red) derived from microscopic cross section images of petioles (C) and stem (F). The lines at the bottom left of all images represent 1 mm.
Figure 2. Relation between the petiole relative water content (RWC) and the xylem water potential ($\Psi_x$). To determine the $\Psi_x$, the linear regression below turgor loss point was used as suggested by Turner (1988).
Figure 3. Five representative MRI images acquired during the vines dehydration and their correspondent xylem water potential. The petiole was paralleled to the stem and to a reference water tube.
Figure 4. Vulnerability curves of Stems (closed symbols) and petioles (open symbols) - sigmoidal fitting between the percent of embolized area (%emb) and the shoot water potential (Ψx). Data represent averages ± S.E (n=4).
Figure 5. Representative transverse microCT images of grapevine shoots and petioles (A,B,C,D) and the average degree of embolism (E). Images A and B were taken from the same plant (shoots and petiole, respectively), and images C and D were taken on another plant (shoot and petiole, respectively). MicroCT images A and B were taken on a well-watered vine ($\Psi_x = -0.3$ MPa), while C&D were taken on a drought-stressed vine ($\Psi_x = -1.5$ MPa). % embolized vessels for the four samples were: A= 5.7, B=3.8, C=8.4, D=56.3. Scale bars located to the bottom right of each image represent 500 microns. The mean % of embolized vessels for stems and petioles of well-watered plants and drought-stressed plants were compared (E). Bars represent the mean S.E. n=4.
In this work we examined the “hydraulic vulnerability segmentation” hypothesis which predicts that expendable distal organs are more susceptible to water-stress induced cavitation than the main stem of the plant. We explored the differences in embolism formation of intact petioles and stems of grapevines by means of MRI and microCT. The paper presents the first direct evidence and the first imaging of hydraulic vulnerability segmentation in living plants and highlight its importance in grapevine responses to severe water stress.