# 1 Water availability dynamics have long-term effects on mature

# 2 stem structure in Vitis vinifera

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- 14 **Running title:** Anatomical structure is affected by early season water availability.
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## 16 Highlights:

- 17 Water availability early in the season determines vegetative growth and stem anatomical structure
- 18 in mature *Vitis vinifera* vines.

19

## 20 Abstract:

*Vitis vinifera* is a climbing vine with wide vessels and high hydraulic conductivity. There is a
lack of data on the anatomical structure of the mature vine stem, and most current knowledge is
based on first-year shoots. Moreover, the effect of drought stress on anatomical structure has
been partly reported in shoots of *Vitis vinifera* but not in stems.
In current study two irrigation approaches were applied on *Vitis vinifera* Merlot vines: constant
(low, medium and high irrigation) and dynamic (early/late season water deficit). The following

- 27 parameters were measured: trunk diameter, annual ring width and area, vessel diameter, specific
- 28 hydraulic conductivity and stem water potential.

High water availability early in the season (high irrigation and late deficit) resulted in vigorous
vegetative growth (greater trunk diameter, ring width and area), wider vessels and increased

- 31 specific hydraulic conductivity. The distribution of large xylem vessels was altered by drought
- 32 stress, where high water availability early in the season caused a shift of the vessel population
- 33 towards the wider frequency classes. Interestingly, the early deficit vines showed more negative
- 34 water potential values late in the season compared to the low irrigation vines. This may imply an
- 35 effect of anatomical structure on vine water status.
- 36 Key words: Drought stress, specific hydraulic conductivity, vessel diameter, vessel distribution,
- 37 *Vitis vinifera,* water availability.

#### 38 Introduction

39 The genus *Vitis* has been distinguished for its wide (Adkinson 1913; Pratt 1974) 40 and long xylem vessels (Zimmermann and Jeje 1981; Ewers et al. 1990; 41 Jacobsen et al. 2012), which characterize lianas. Such hydraulic architecture 42 makes *Vitis vinifera* an excellent model of water flow in plants according to the 43 "unit pipe model" as described by Tyree & Ewers (1991), since Vitis vessels are 44 relatively optimal pipes. Since vessel length and diameter are correlated with 45 stem diameter, mature vine trunks tend to have wider and longer vessels 46 compared to young stems/shoots (Ewers and Fisher 1989; Jacobsen et al. 2012; 47 Hacke 2015). Moreover, since vessels in Vitis are completely inactivated after 4 to 7 years (Tibbetts and Ewers 2000), in mature vine trunks (older than 7 years) 48 49 only the secondary xylem is functional, while the primary xylem created in its 50 first year is nonfunctional. This fact is crucial, since in contrast to the scalariform 51 arrangement of intervessel bordered pits of secondary xylem vessel elements, the 52 primary xylem vessel elements have partial secondary wall thickenings, making 53 them much more vulnerable to cavitation (Pratt 1974; Choat et al. 2005; Sun et 54 al. 2006; Brodersen et al. 2011; Craig R Brodersen et al. 2013; Rolland et al. 55 2015; Hochberg, Herrera, et al. 2016). Indeed, visualization techniques (microCT 56 / MRI) have shown that in one-year-old stems of Vitis the spread of embolism 57 proceeds from the pith towards the cambium through the primary xylem (Choat 58 et al. 2010; Craig R Brodersen et al. 2013; Craig Robert Brodersen et al. 2013; 59 Knipfer et al. 2015; Hochberg, Albuquerque, et al. 2016). An additional 60 important anatomical feature of mature vine trunks is the ratio between the area 61 of the pith and the xylem, which decreases with stem maturation (Sun et al. 62 2006). All of the above suggest that the xylem architecture of the mature trunk differs from that of a young shoot, in a way that makes it less vulnerable tocavitation.

65 Despite those significant differences, there is a scarcity of information about the 66 anatomical structure of mature vine trunks, while current reported anatomical 67 information on vines is based mainly on analysis of one-year-old stems (Schultz 68 and Matthews 1993; Lovisolo et al. 1998; Schubert et al. 1999; Sun et al. 2006; 69 Brodersen et al. 2011; Chatelet et al. 2011; Santarosa et al. 2016). In one-year-70 old stems, hydraulic structure is reported to vary among Vitis cultivars and 71 rootstocks (Chouzouri and Schultz 2005; Chatelet et al. 2011; Gerzon et al. 2015; 72 Hochberg et al. 2015; Santarosa et al. 2016; Shtein et al. 2016), and to be 73 affected by environmental parameters (Schubert et al. 1999). Only a few studies analyzing anatomical features of xylem in mature Vitis trunks have been 74 75 published (Zimmermann and Jeje 1981; Ewers et al. 1990; Tibbetts and Ewers 76 2000; Shtein et al. 2016).

77 Another interesting feature of Vitis xylem is the bimodal distribution pattern of 78 vessel diameters, meaning two distinct vessel size groups - wide and narrow 79 (Carlquist 1985; Ewers et al. 1990; Wheeler and LaPasha 1994; Shtein et al. 80 2016). Wide diameter vessels are considered to be more hydraulically efficient, 81 and tend to be more vulnerable to embolism within the same species (Sperry and 82 Tyree, 1988; Lo Gullo and Salleo, 1991; Hargrave et al., 1994; Cai and Tyree, 83 2010; Christman et al., 2012; Scoffoni et al., 2016). The accepted "air-seeding" 84 theory suggests that the increased vulnerability to embolism of wide vessels is 85 linked to their enlarged total area of intervessel pits. A wide pit area raises the 86 average size of the "rare" largest pore, consequently increasing the risk of air 87 seeding (Choat et al. 2003; Wheeler et al. 2005; Jansen et al. 2009; Cai and 88 Tyree 2010).

89 Most cultivated vineyards worldwide are located in semi-arid and arid regions 90 where drought stress is prevalent (Chaves et al. 2007). Nevertheless, compared to 91 other woody plants, grapevines are often described as relatively vulnerable to 92 drought stress (Choat et al. 2010; Zufferey et al. 2011; Jacobsen and Pratt 2012; 93 Hacke 2015). It has been reported that drought stress induces embolism and a 94 loss of hydraulic function (Schultz and Matthews 1988; Hargrave et al. 1994; 95 Lovisolo et al. 1998; Choat et al. 2010; Brodersen et al. 2014). Drought stress 96 also negatively affects vegetative growth and pruning weight of vines (Matthews

97 1987; Intrigliolo and Castel 2010; Shellie and Bowen 2014; Munitz et al. 2016).

98 There is a lack of available information on the effects of drought stress on

99 hydraulic conductivity resulting from modifications to Vitis xylem anatomy, as

100 only a few studies have examined this subject (Lovisolo et al. 1998; Hochberg et

al. 2015). The current study focuses on the long-term anatomical acclimation to

- 102 drought stress of mature *Vitis vinifera* Merlot vines.
- 103

## 104 Materials and Methods

105 Plant material and experimental design

106 This study was carried out in a 100-ha commercial vineyard located in the Judean Plain, Israel (31<sup>0</sup>49'N, 34<sup>0</sup>53'E, elevation 124 m). This region has a semi-107 arid climate with predominantly winter rainfall (average 463 mm year<sup>-1</sup>) and high 108 evapotranspiration (average 1512 mm year<sup>-1</sup>). The vineyard was planted in 1998 109 110 with Vitis vinifera L cv. 'Merlot' grafted to 140 Ruggeri, and trained onto a two-111 wire vertical trellis. Row direction was North/South with a slight tendency to the 112 West, and vine and row spacing were 1.5 m and 3 m respectively (2222 vines ha 113 <sup>1</sup>). The soil was loam (48% sand, 29% silt and 23% clay, field capacity 28 % 114 vol., wilting point 14 % vol.). Pest management and fertilization in the vineyard 115 were applied according to standard local agricultural practice.

The experimental design was a complete randomized block design with five irrigation treatments each replicated four times. Each block comprised three rows (one data and two border rows). Each plot comprised 16 vines per line, with the outer two vines at each end being buffer vines and the inner 12 vines being measurement vines (a total of 240 measurement vines, i.e. 12 vines  $\times$  5 treatments  $\times$  4 replicates).

122 Irrigation treatments

During 2009-2012, five irrigation treatments representing different levels of 123 124 deficit irrigation were applied as percentages of crop evapotranspiration (ET<sub>c</sub>). 125 Crop evapotranspiration was calculated by multiplying reference 126 evapotranspiration (ET<sub>o</sub>) by the crop coefficient (K<sub>c</sub>), i.e.  $ET_c = ET_o \times K_c$ .  $ET_o$ 127 was calculated using data obtained from the adjacent meteorological station, and 128 K<sub>c</sub> was calculated according to Netzer et al (2009) following nondestructive 129 measurements of leaf area index. For more details about the irrigation method 130 see Munitz and Netzer (2016). Irrigation treatments followed two different

131 strategies: static irrigation and dynamic (seasonally changing) irrigation. 132 Dynamic treatments involved alternation of the percentage of ET<sub>c</sub> along the 133 growing season according to phenological stages (stage I, stage II, stage III) as 134 defined by Kennedy (2002): stage I - from bloom to bunch closure, stage II -135 from bunch closure to veraison (color change to red) and stage III - from 136 veraison to harvest. Dynamic irrigation treatments were: early deficit (0, 20, 50%)137 of ET<sub>c</sub>) and late deficit (50, 20, 20% of ET<sub>c</sub>). Static irrigation treatments were: 138 low irrigation (20% of  $ET_c$ ), medium irrigation (35% of  $ET_c$ ) and high irrigation 139 (50% of ET<sub>c</sub>).

140 Stem water potential ( $\Psi_s$ )

141 Stem water potential ( $\Psi_s$ ) was measured using a pressure chamber (Arimad 2, 142 Kfar Charuv, Israel). Three sunlit, mature, fully-expanded leaves from each plot 143 (12 leaves per treatment) were bagged 2 h prior to measurement in plastic bags 144 covered with aluminum foil. The time elapsing between leaf excision and 145 chamber pressurization was less than 15 s. The measurements were conducted 146 one day before irrigation was applied.

147 Trunk diameter

148 Measurements were performed monthly with a digital caliper (075430, Signet, 149 Taiwan) on 48 vines per treatment (12 vines per plot  $\times$  4 replicates). In order to 150 obtain consistent data, all measured vines (240) were marked 30 cm above 151 ground with colored tape, and all measurements were taken at this point.

152 Anatomical sampling

153 At the end of the experiment (December 2012) xylem cores from representative 154 vines were sampled 50 cm above ground with an increment borer (5.15 mm Core 155 3-Thread 8", Haglof, Sweden). Twelve cores were sampled from each treatment 156 (3 cores per plot  $\times$  4 replicates, 60 cores total). Trunk diameter (D; mm) at the 157 drilling location was recorded. Cores were placed in sterilized water and stored at 158 4<sup>°</sup>C until cross sectioned with a sliding microtome (NR17800, Reichert, Austria) 159 at a thickness of 90  $\mu$ m. In order to increase visual contrast, cross sections were 160 stained for 60 s in Reactif Genevois solution (FAHN 1954), then flushed with 161 distilled water. Photographs of stained cross sections were obtained using a 162 stereo microscope (Olympus SZ2-ILST) coupled with a digital camera (Olympus 163 LC20) equipped with image acquisition software (LCmicro 5.1, Olympus, 164 Tokyo, Japan) at  $\times 20$  magnification.

## 165 *Image analysis*

166 Analysis of cross sections was performed by separately quantifying various 167 parameters in the visible field for each of four recent growth rings (2009-2012) using ImageJ software (Rasband, W.S., ImageJ, U.S. National Institutes of 168 Health, Bethesda, Maryland, USA, http://imagej.nih.gov/ij/, 1997-2016). The 169 170 abbreviations for structural parameters were taken from Scholz et al. (2013). The 171 following anatomical parameters were measured (Fig. 1): annual ring width ( $W_r$ ;  $\mu$ m), bark width ( $W_b$ ;  $\mu$ m), xylem radius ( $r_x$ ;  $\mu$ m) and inner xylem radius ( $r_i$ ; 172 173  $\mu$ m). Vessel lumen area ( $A_{\nu}$ ;  $\mu$ m) and number of vessels (n) were measured using 174 the 'analyze particles' tool (see Fig. 2 for explanation); analyzed area (A;  $mm^2$ ) 175 was also measured. A total of 12,177 vessels were measured and used for 176 subsequent hydraulic conductivity calculations. The detailed calculations for 177 trunk and vessel parameters are presented in Table 1.

178 Theoretical specific hydraulic conductivity  $(K_s)$  calculations

179 The theoretical specific hydraulic conductivity ( $K_s$ ; kg m<sup>-1</sup> MPa<sup>-1</sup> s<sup>-1</sup>) was 180 calculated using the modified Hagen–Poisseuille's equation (Tyree and Ewers 181 1991):

$$K_s = (\pi \rho/128\eta A_w) \sum_{i=1}^n (d_i^4)$$

where  $K_s$  is the specific hydraulic conductivity, p is the density of the fluid in kg m<sup>-3</sup> (assumed to be 1000 kg m<sup>-3</sup>),  $\eta$  is the dynamic viscosity of the fluid in MPa s<sup>-1</sup> (assumed to be  $1 \times 10^{-9}$  MPa s<sup>-1</sup>),  $A_w$  is the area (m<sup>2</sup>) of the xylem cross section analyzed, d is the diameter (m) of the i<sup>th</sup> vessel and n is the total number of vessels in the measured area.

187 Hydraulic conductivity per annual ring ( $K_{ar}$ , kg m<sup>1</sup> MPa<sup>-1</sup> s<sup>-1</sup>) was calculated by 188 multiplying the theoretical xylem specific hydraulic conductivity ( $K_s$ ) by annual 189 growth ring area ( $A_r$ , m<sup>2</sup>).

190 Frequency classes for calculation of total vessel number and total conductivity 191 were established at intervals of 20  $\mu$ m (Fig. 3). For calculation of vessel density 192 and vessel average diameter, vessels were separated into two size categories 193 (>100  $\mu$ m,  $\leq$  100  $\mu$ m), since *Vitis* has a bimodal distribution of xylem vessels 194 (Fig. 3A).

195 Statistical analysis

196 The software program JMP7 (SAS Institute, Cary, NC) was used for all 197 statistical procedures. Data were analyzed via analysis of variance (ANOVA), 198 and means were separated according to the least significant difference (LSD) at p 199  $\leq 0.05$  using the Tukey-Kramer test.

200

## 201 **Results**

# 202 Stem vessel distribution

The distribution of stem xylem vessels in all irrigation treatments showed a classic bi-modal pattern. The small vessels (< 100  $\mu$ m) constituted the majority (61.3%) of total vessel number, while the large vessels comprised only 38.7% of total vessels (Fig. 3A). In contrast, the theoretical hydraulic conductivity showed a reverse distribution, where the large vessels contributed 97.2% of total conductivity (Fig. 3B), whereas the contribution of the more abundant small vessels was negligible.

210 Seasonal changes in trunk diameter

As an integrative indicator of vegetative growth, seasonal changes in trunk 211 212 diameter were monitored monthly during 2011-2012 (following two years of 213 differential irrigation application after 11 years of identical irrigation). Seasonal 214 trends of trunk diameter development were similar in both years in all irrigation 215 treatments (Fig. 4); an increase in trunk diameter began two weeks after bud 216 break and continued until stage II (mid-June), then remained stable until the next 217 season. In 2011, a decrease in trunk diameter was apparent during stage III, in all 218 irrigation treatments. Vines in different irrigation treatments showed significant 219 differences in trunk diameter throughout the entire experimental period (Fig. 4). 220 Among static irrigation vines, trunk diameter was positively correlated with 221 applied water amounts, even though the trunk of the medium irrigation vines was 222 slightly narrower than expected (Fig. 4A). In the dynamic irrigation treatments, 223 the early deficit vines exhibited the narrowest trunk diameter of all vines in all 224 irrigation treatments during the entire measuring period. The late deficit vines 225 had an intermediate trunk diameter throughout the measuring period (Fig. 4B).

226 Structural parameters

Vines subjected to the static irrigation treatments differed significantly in their
annual ring width, with a positive effect of applied water amounts on ring width
(Table 2). Within the dynamic irrigation treatments, the late deficit vines had the

230 widest annual ring width (901.5 µm), very similar to the width of the high 231 irrigation vines (Table 2). Surprisingly, the early deficit vines had the narrowest 232 annual ring width (686  $\mu$ m) even compared to the low irrigation vines (719  $\mu$ m). 233 The general trend of annual ring area resembled the trend of annual ring width. 234 Ring area was positively affected by increasing water amounts in static treatment 235 vines. In dynamic irrigation treatments, early deficit vines had the smallest 236 annual ring area of the five treatments, while the late deficit vines had a 237 relatively large ring area, intermediate between that of the high and medium 238 irrigation vines (Table 2).

239 The density of 'small' ( $\leq 100 \mu m$ ) and 'large' (>100 \mu m) vessels was not 240 significantly affected by irrigation treatments, although a reduction in the density 241 of 'small' vessels in high irrigation vines was observed. The 'large' (>100µm) 242 vessel diameter of the high irrigation vines was significantly wider than that of 243 the low and medium irrigation vines. In the dynamic irrigation treatments, early 244 deficit vines had significantly narrower 'large' vessels than all vines in all other 245 irrigation treatments, while the late deficit vines had the wideset (Table 1). The 246 trend for 'small' (≤100µm) vessel diameter was less clear, with early deficit and 247 medium irrigation vines exhibiting significantly wider vessels than all other 248 vines. The trend in specific hydraulic conductivity was similar to that of 'large' 249 (>100µm) vessel diameter, where the high irrigation and late deficit vines had 250 significantly higher hydraulic conductivity (Table 1) than vines in other 251 treatments. Hydraulic conductivity per annual ring increased significantly with 252 increasing water amounts in the static irrigation treatments, whereas in dynamic 253 treatments the early deficit vines had the lowest conductivity, while the late 254 deficit vines had high conductivity (slightly lower than the high irrigation vines). 255 The relationship between hydraulic conductivity per annual ring and seasonal water amount (Fig. 6a) was weak and non-significant ( $R^2 = 0.21$ ), while the 256 257 relationship between hydraulic conductivity per annual ring and water amounts 258 applied during stage I (bloom to bunch closure, Fig. 5b) was stronger and significant ( $R^2 = 0.60$ , p < 0.001). 259

260 Stem hydraulic conductivity distribution among vessel size classes

261 Specific stem hydraulic conductivity was calculated separately for each vessel 262 size class (Fig 6). Among static irrigation treatments, the conductivity 263 distribution was similar, but several differences among treatments were observed. In low irrigation vines a higher percentage of the calculated hydraulic conductivity was derived from narrow vessel classes, in the medium irrigation vines - from wide vessel classes, and in the high irrigation vines - from the widest vessel classes (Fig 6a). In the dynamic irrigation treatments, the early deficit vines had a higher amount of hydraulic conductivity derived from narrower frequency classes, while the hydraulic conductivity in late deficit vines shifted towards the wide frequency classes (Fig 6b).

271 Stem water potential

272 Significant differences in water potential ( $\Psi_s$ ) between vines in different 273 irrigation treatments were observed during stages II and III (Fig. 7). The daily 274 trend of  $\Psi_s$  was similar among treatments, with a steep decrease in  $\Psi_s$  values 275 being recorded throughout the morning, followed by stabilization and 276 improvement in  $\Psi_s$  during the afternoon (Fig. 7). In stage II an improvement in 277  $\Psi_s$  values during the afternoon was apparent only in the high and medium 278 irrigation vines (Fig. 7B). On both measuring days, the high irrigation vines had 279 the highest  $\Psi_s$  values during the course of the day, the medium irrigation vines 280 were at an intermediate level and the low irrigation vines had the lowest  $\Psi_s$ . In 281 the dynamic irrigation treatments, the early deficit vines had low  $\Psi_s$  values 282 during stage II (lower than the low irrigation vines) and intermediate values 283 during stage III (resembling those of the medium irrigation vines). The late 284 deficit vines had low  $\Psi_s$  values on both dates, even compared to the values of the 285 low irrigation vines (Fig. 7).

286

# 287 Discussion

288 *Vegetative growth* 

289 Vegetative growth can serve as a good indicator of water availability (Tyree and 290 Ewers 1991; Munitz et al. 2016). In the present study we continuously monitored 291 trunk growth. In all irrigation treatments, an increase in trunk diameter occurred 292 mainly early in the season (from two weeks before blooming until bunch closure) 293 (Fig. 4). Early season trunk growth has been previously reported in Merlot (Ton 294 and Kopyt 2004) and in other Vitis vinifera cultivars (Myburgh 1996; Ton and 295 Kopyt 2004; Intrigliolo and Castel 2007; Montoro et al. 2011; Papi and Storchi 296 2012; Edwards and Clingeleffer 2013).

297 The high irrigation and late deficit vines, which received higher water amounts 298 early in the season, exhibited the widest trunk diameter of all vines. Similarly, 299 annual ring width and area, which represent annual vegetative growth, were also 300 positively correlated with high irrigation early in the season (Stage I). The high 301 irrigation and late deficit vines had significantly wider ring width and area in 302 comparison to the low irrigation and early deficit vines (Table 2). The dominance 303 of early season vegetative growth in Vitis vinifera can be explained by the fact 304 that cambial activity to produce new vascular elements takes place mainly during 305 the early stage of the growing season (until 20 days after bunch closure, 306 Bernstein and Fahn, 1960).

307 *Hydraulic structure* 

308 A significant increase (9 - 11%) in the large vessel diameter was recorded in the 309 high irrigation and late deficit vines (Table 2), even though no significant 310 difference in the density of the large vessels was found. This implies that water 311 availability early in the season affects large vessel diameter rather than vessel 312 density. The same effect of drought stress reducing average vessel diameter has 313 been reported in Vitis vinifera shoots (Lovisolo et al. 1998) and petioles 314 (Hochberg et al. 2015). Interestingly, in small vessels the opposite was found, 315 where the vessels of the early deficit vines, which received minimal water early 316 in the season, exhibited the widest diameter (Table 2). Although the small 317 vessels comprised the majority of the total vessels (61%), they contributed only a 318 negligible 3% of total hydraulic conductivity (Fig. 3). A similar phenomenon of 319 dominance of wide diameter vessels with respect to total hydraulic conductivity 320 has also been recorded in other species (Hargrave et al. 1994; Tibbetts and Ewers 321 2000).

322 From inspection of the hydraulic conductivity distribution, it can be deduced that 323 high water amounts early in the season result in a higher percentage of hydraulic 324 conductivity derived from wider frequency classes (Fig. 6). This phenomenon is 325 more pronounced in late deficit vines, possibly due to their dynamic irrigation 326 treatment. It is well known that in *Vitis*, as in many other lianas, wide vessels are 327 formed at the beginning of the growing period and narrow vessels are formed at 328 the end (extreme diffuse-porous), causing a bi-model distribution of vessels 329 (Pratt 1974; Kozlowski 1983; Ewers et al. 1990; Wheeler and LaPasha 1994). 330 Applying high irrigation amounts during the formation of wide vessels and

reducing water allocation during the formation of narrow vessels (late deficit), should result in larger wide vessels and smaller narrow vessels. This will lead to an increased proportion of hydraulic conductivity being derived from wide vessels. As a result, late deficit vines are expected to be more susceptible to embolism formation in comparison to low irrigation vines.

336 Specific hydraulic conductivity  $(K_s)$  represents the conducting efficiency of the 337 xylem tissue (Tyree and Ewers 1991). The high irrigation and late deficit vines 338 had significantly increased  $K_s$  (24 - 33%) compared to other vines (Table 2). This 339 means that limited water availability early in the growth season (bloom to bunch 340 closure) results in decreased  $K_s$ . Since there is a good correlation between 341 measured and calculated hydraulic conductivity (Salleo et al. 1985; Hargrave et 342 al. 1994; Lovisolo et al. 1998; Nolf et al. 2017) our theoretical  $K_s$  results may 343 indicate an actual change in the hydraulic conductivity of vines in response to 344 drought stress.

345 Hydraulic conductivity per annual growth ring  $(K_{ar})$  is an integrated parameter 346 combining both hydraulic conductivity  $(K_s)$  and vegetative growth (trunk 347 diameter and ring width). As with ring area and hydraulic conductivity, the high 348 irrigation and late deficit vines had significantly increased hydraulic conductivity 349 per annual ring ( $K_{ar}$ , Table 2). Interestingly, the high irrigation vines had 9% 350 higher  $K_{ar}$  (not significant) than the late deficit vines. This difference can be 351 attributed to the reduction of water amounts in late deficit vines during bunch 352 closure, while cambial activity in the stem persisted for a further 20 days 353 (Bernstein and Fahn 1960).

Reinforcement of the notion that most of the vegetative growth and xylem development occur early in the season can be found in the correlation between the amounts of applied water and hydraulic conductivity per annual ring,  $K_{ar}$ (Fig. 5). While annual water applied had a weak and non-significant correlation with  $K_{ar}$  (R<sup>2</sup> = 0.21), water amount applied during stage I was strongly and significantly correlated to  $K_{ar}$  (R<sup>2</sup> = 0.6, P < 0.001).

360 *Water potential* 

361 Stem water potential is known to be a sensitive indicator of vine water status 362 (Choné et al. 2001; Munitz et al. 2016). Indeed, on both measuring days static 363 irrigation vines differed significantly in their stem water potentials, according to 364 the water amounts applied, throughout the entire day. Those differences in stem 365 water potential were present from bunch closure until harvest (Munitz et al. 366 2016). Interestingly, late deficit vines were more drought stressed (more negative 367 stem water potential) than low irrigation vines on both measuring days, even 368 though they received on average 31% more water prior to the measuring days. 369 Those differences in stem water potential cannot be attributed to a broader 370 canopy, since the late deficit and low irrigation vines had similar leaf area on 371 measuring days (Munitz et al. 2016). Water potentials of -1.4 MPa, recorded on 372 both measuring days, have been reported to cause a 30 - 80 % decrease in 373 hydraulic conductivity of Vitis vinifera shoots(Alsina et al. 2007; Choat et al. 374 2010; Jacobsen and Pratt 2012).

375 The increased drought stress in late deficit vines can be explained by a greater 376 hydraulic conductivity loss. As stated before, wide vessels are more susceptible 377 to embolism formation. Late deficit vines, with a higher percentage contribution 378 of wide vessels to total hydraulic conductivity, are expected to experience greater 379 hydraulic loss at a similar water potential. Increased hydraulic loss will, in turn, 380 lead to increased drought stress e.g. more negative water potential (Tyree et al. 381 1991). Another explanation for the increased drought susceptibility of late deficit 382 vines compared with low irrigation vines, can be found by examining their 383 Carlquist s "vulnerability index" (1977). The index is calculated by dividing 384 average vessel diameter by vessel density, where a lower value is interpreted as 385 greater redundancy of vessels and improved capability of withstanding drought 386 stress. The large vessels of the late deficit vines have an index of 17.0 while low 387 irrigation vines have an index of 14.9.

388 Structural parameters as compared to values from literature

389 Generally, our results seem to be in agreement with values presented in the 390 literature. The annual increase in trunk diameter of mature vines measured in this 391 study (1.5 - 2.5 mm, Fig. 4), is similar to that reported for a number of Vitis 392 vinifera cultivars (0.5 - 3.5 mm) (Bernstein and Fahn 1960; Myburgh 1996; Ton 393 and Kopyt 2004). Similarly, the range of annual ring width measured in this 394 study (720 - 901 µm) is consistent with values reported for mature Vitis vinifera 395 cultivars (100 - 1300 µm) (Perold 1927; Bernstein and Fahn 1960), and the range of annual ring area found in this study (104 - 134 mm<sup>2</sup>) resembles that of 396 Cabernet Sauvignon vines (120 - 240 mm<sup>2</sup>, Shtein et al., 2016). 397

398 The average diameter of large vessels measured in this study (147 - 158 µm) is 399 considerably smaller than that reported for Vitis vinifera cv. Cabernet Sauvignon. 400 While in the current study water availability of Merlot vines in different 401 irrigation regimes triggered a maximal difference of 11 µm in vessel diameter 402 (Table 2), the difference between the two cultivars stands at more than 50  $\mu$ m. 403 This implies an inherent genetic distinction in stem xylem anatomy between Vitis 404 vinifera cultivars, as reported previously for petioles and shoots (Chouzouri and 405 Schultz 2005; Chatelet et al. 2011; Tombesi et al. 2014; Gerzon et al. 2015; 406 Hochberg et al. 2015; Santarosa et al. 2016). Typical values of specific hydraulic conductivity of liana stems (Milburn 1979) are 65 - 349 kg m<sup>-1</sup> MPa<sup>-1</sup> s<sup>-1</sup>; our  $K_{s}$ 407 results (142 - 188 kg m<sup>-1</sup> MPa<sup>-1</sup> s<sup>-1</sup>) are in the middle of this range. Values of  $K_{ar}$ 408 reported for *Vitis vinifera* cv. Cabernet Sauvignon (0.07 - 0.11, kg m<sup>1</sup> MPa<sup>-1</sup> s<sup>-1</sup>) 409 are considerably higher than those calculated in this study, due to the wider 410 411 vessels and ring area measured in Cabernet Sauvignon vines.

412 *Conclusions* 

413 In the current study, we conducted a comprehensive structural analysis of the 414 mature stem xylem of Vitis vinifera, combined with physiological measurements. 415 The stem comprises the perennial part of the deciduous vine and its anatomical 416 structure constitutes the long term "memory" of the vine, yet very little 417 information about the anatomical features of mature Vitis stems is available. One example of this "memory" is the fact that the current year's vine canopy develops 418 419 while consuming water conducted through vessels differentiated in previous 420 years. Most of the vine's canopy develops in the early stages of the growing 421 season, about 60 days from budbreak (Ben-Asher et al. 2006; Intrigliolo et al. 422 2008; Romero et al. 2010; Munitz et al. 2016). Cambial tissue begins its activity 423 about two weeks after budbreak (Bernstein and Fahn 1960). Vessels become 424 hydraulically active about four weeks after their initial differentiation is initiated 425 (Halis et al. 2012; Hacke 2015): two weeks for differentiation, expansion and 426 formation of secondary walls and at least two additional weeks for the creation of 427 perforation plates by autolysis of axial cell walls. Practically this means that 428 current year vessels are functional no less than 54 days after budbreak, when 429 canopy development has almost ceased.

There is a lack of information about the effects of drought stress on *Vitis* xylemstructure (Lovisolo et al. 1998), especially in mature stems. Our research can

- 432 contribute to the understanding of mature stem xylem structure and how it is
- 433 affected by drought stress in the long term.
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# References

Parameter	Abbreviation	Unit	Formula		
Vessel diameter	d	μm	$d = (4*A_v/\pi)^{0.5}$		
Vessel density	$V_D$	mm <sup>-2</sup>	$V_D = n/A$		
Trunk radius	R	μm	R = D/2		
Xylem radius	r <sub>x</sub>	μm	$r_x = R - W_b$		
Annual ring area	$A_r$	$mm^2$	$A_r = \pi^* (r_x)^2 - \pi^* (r_i)^2$		

Table 1. Calculations used for trunk and vessel parameters.

Irrigation treatment	Water amount (mm season <sup>-1</sup> )	Annual ring width (µm)	Annual ring area (mm <sup>2</sup> )	Large vessel density (mm <sup>-2</sup> )	Small vessel density (mm <sup>-2</sup> )	Larege vessel average diameter (µm)	Small vessel average diameter (µm)	Specific hydraulic conductivity (kg m <sup>-1</sup> MPa <sup>-1</sup> s <sup>-1</sup> )	Hydraulic conductivity per annual ring (kg m MPa <sup>-1</sup> s <sup>-1</sup> )
Low	105	719.9 <sup>bc</sup>	104.4 <sup>bc</sup>	9.9	16.9	147.9 <sup>h</sup>	37.5 <sup>b</sup>	142.5 <sup>b</sup>	0.0152 <sup>b</sup>
Medium	184	835.6 <sup>ab</sup>	124.2 <sup>ab</sup>	9.6	14.7	147.1 <sup>b</sup>	43.3 <sup>ª</sup>	144.0 <sup>b</sup>	0.0179 <sup>b</sup>
High	256	891.4 <sup>ª</sup>	141.5 <sup>a</sup>	10.3	12.6	154.9 <sup>a</sup>	40.3 <sup>b</sup>	188.6 <sup>n</sup>	0.0273 <sup>a</sup>
Early deficit	143	686.0 <sup>e</sup>	100.9 <sup>c</sup>	9.7	15.5	142.1°	45.5 <sup>a</sup>	127.2 <sup>b</sup>	0.0128 <sup>b</sup>
Late deficit	141	901.5 <sup>a</sup>	134.3ª	9.3	15.8	158.5 <sup>a</sup>	38.3 <sup>b</sup>	188.3 <sup>a</sup>	0.0257 <sup>a</sup>

Table 2. Water amounts, anatomic and hydraulic parameters. 'Hulda' Merlot vineyard, 2009 - 2012.

Values represent means (n = 12). Within each column, means followed by different letters are significantly different (P < 0.05) according to Tukey's test.

## Figure legends

*Figure 1.* Schematic cross section diagram with abbreviations used for trunk parameter calculations. *Figure 2.* Stem cross section of vitis vinifera cv. Merlot. (A) Cross section image as acquired by stereo microscope. (B) Image processing to 8-bit. (C) Image conversion to black and white (binarization) and selection of annual ring to be analyzed (yellow line). (D) Measurement of vessel area.

**Figure 3.** (A) Distribution of stem xylem vessels according to diameter classes ( $\mu$ m) in Merlot vines of all irrigation treatments. (B) Distribution of hydraulic conductivity according to diameter classes ( $\mu$ m) in Merlot vines of all irrigation treatments. Vessel classes were divided into two size categories:  $\leq 100 \mu$ m (black bars) and  $< 100 \mu$ m (gray bars). Data represent vessels from all irrigation treatments during 2009-2012, n = 12177 vessels.

*Figure 4.* (A) Biennial pattern of trunk diameter development of Merlot vines exposed to low irrigation (closed circles), medium irrigation (closed squares) and high irrigation (closed triangles), in 2011 and 2012. (B) Biennial pattern of trunk diameter development of Merlot vines exposed to early deficit (open circles) and late deficit (open squares), in 2011 and 2012. Each point is the mean of 48 vines (12 vines per replicate). The vertical bars denote one standard error.

*Figure 5.* (A) Relationship between annual water applied and hydraulic conductivity per annual growth ring in Merlot vines of all irrigation treatments. (B) Relationship between water amount applied during stage I (bloom to bunch closure) and hydraulic conductivity per annual growth ring in Merlot vines of all irrigation treatments. Each point is the mean of 12 vines (3 vines per replicate). The vertical bars denote one standard error. \*Significant at P < 0.001.

*Figure 6.* (A) Distribution of hydraulic conductivity according to diameter classes ( $\mu$ m) in Merlot vines exposed to low irrigation (solid line), medium irrigation (dotted line) and high irrigation (dashed line). (B) Distribution of hydraulic conductivity according to diameter classes ( $\mu$ m) in Merlot vines exposed to early deficit (solid line) and late deficit (dashed line). Data represent vessels from all analyzed years (2009-2012), n = 12177 vessels. *Figure 7.* (A) Daily pattern of stem water potential of Merlot vines exposed to low irrigation (closed circles), medium irrigation (closed squares), high irrigation (closed triangles), early deficit (open circles) and late deficit (open squares), on 10/08/11. (B) Daily pattern of stem water potential of Merlot vines deficit (closed triangles), early deficit (open circles), medium irrigation (closed squares), high irrigation (closed triangles), early deficit (open circles) and late deficit (open squares), on 26/07/12. Each point is the mean of 12 leaves (3 vines per replicate). The bars denote one standard error.

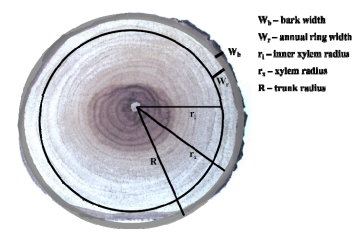
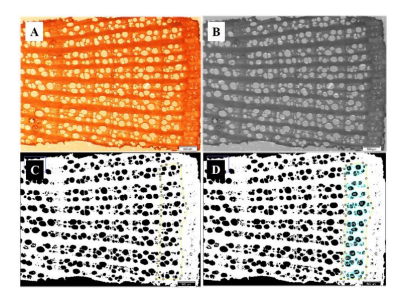


Fig.



*Fig.* 2

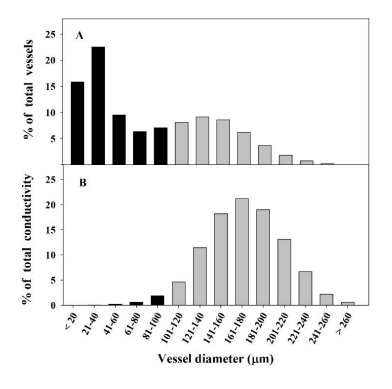


Fig. 3

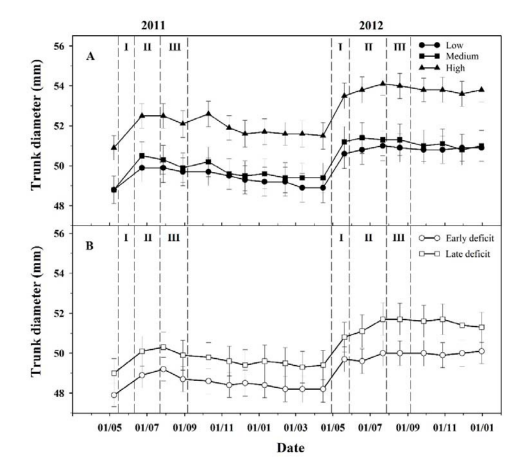
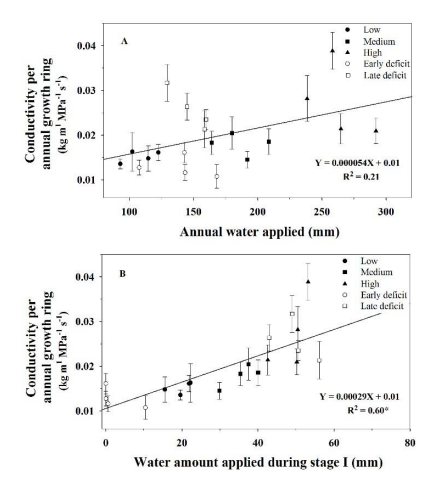


Fig. 4



*Fig.* 5

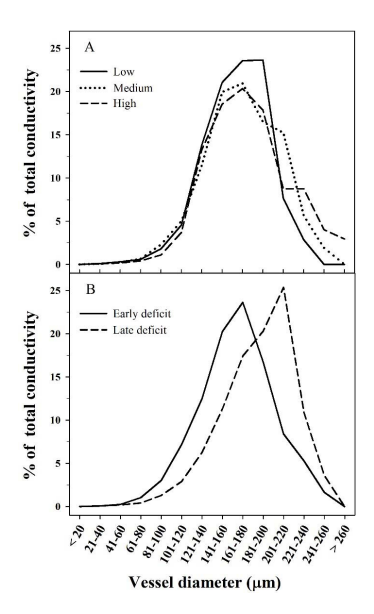
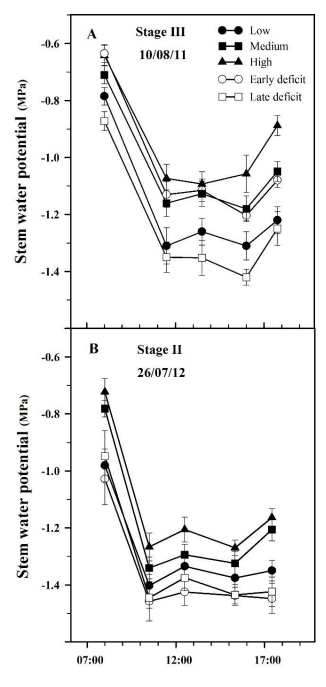


Fig. 6



Time of day (hours)

Fig. 7