- 1 Combined drought resistance strategies and the hydraulic limit in co-
- 2 existing Mediterranean woody species
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21 Summary

22	•	Woody species employ various strategies to cope with drought stress. We
23		investigated similarities and differences in response to chronic drought to
24		understand resistance strategies in co-occurring Mediterranean species.
25	•	We studied five predominant Mediterranean species; Quercus calliprinos,
26		Pistacia palaestina, Pistacia lentiscus, Rhamnus lycioides, and Phillyrea
27		latifolia over two summers at three sites with different aridities. We measured
28		key hydraulic and osmotic traits related to drought resistance, including
29		resistance to embolism (Ψ_{50}), carbon isotope signature (δ^{13} C), pre-dawn (Ψ_{PD})
30		and mid-day (Ψ_{MD}) water potentials, and native (Ψ_s) and full turgor (Π_0)
31		osmotic potentials.
32	•	Significant differences among species appeared in resistance to embolism. The
33		species also showed differences in the water potential plastic response over the
34		dry season. This interspecific variation increased at the end of the dry season
35		and resulted in very narrow hydraulic safety margins (HSM). Consequently,
36		predicted loss of hydraulic conductivity revealed species with significant
37		native embolism. Two of the species also had seasonal changes in osmotic
38		adjustment.
39	•	Our detailed analysis indicates that co-existing Mediterranean woody species
40		combine various drought resistance strategies to minimize mortality risk.
41		However, all of them risk mortality as they approach their hydraulic limit near
42		the dry margin of their distribution.
43		
44		

Key words: Climate change, Drought resistance, Hydraulic failure, Hydraulic safetymargins, Osmotic adjustment, Tree hydraulics.

47 Introduction

Drought is projected to increase in intensity and duration in many regions 48 worldwide, including the Mediterranean (Spinoni et al., 2018; Xu et al., 2019) a hot spot 49 for biodiversity (Myers et al., 2000). The current massive tree mortality in parts of the 50 region is a major cause of concern for the extinction of native species (García de la 51 Serrana et al., 2015; Cramer et al., 2018). Deciphering drought resistance strategies and 52 their limitations in Mediterranean woody species is crucial for understanding changes in 53 structure and function of plant communities threatened by climate change, and will help 54 improve forests and woodlands' sustainable management programs (Trumbore et al., 55 2015). 56

The various strategies used by woody species to cope with drought stress can be 57 divided into three categories; escape, avoidance and tolerance (Delzon, 2015; Volaire, 58 2018). Escape is the temporary shedding of leaves and branches through which water is 59 60 lost. Avoidance is actively minimizing water loss by stomatal closure or increasing water uptake through deep roots. Tolerance is maintaining physiological functionality during 61 water loss, mainly by increased xylem resistance to embolism and osmotic adjustment to 62 prevent turgor loss at the cell level. Woody species vary in their strategies to cope with 63 drought, especially under natural conditions of prolonged and severe drought. Therefore, 64 field studies of natural populations for a wide range of species and traits, and along 65 aridity gradients, are necessary. 66

Drought can lead to embolism, a process that occurs via cavitation events in the 67 xylem and causes hydraulic dysfunction by disrupting water conduction in the xylem 68 (Tyree & Zimmermann, 1983). Embolism resistance is often expressed as the value of 69 xylem water potential (Ψ_x) corresponding to 50 or 88 percent loss of conductivity (PLC), 70 Ψ_{50} and Ψ_{88} , respectively (Tyree & Sperry, 1989). Embolism has been shown to be one 71 of the leading causes of tree mortality worldwide (Anderegg et al., 2016; Adams et al., 72 2017). The variation in embolism resistance between species is large, and it appears that 73 species habitat dryness plays a significant role in this variation (Maherali et al., 2004; 74 Delzon et al., 2010; Choat et al., 2012; Larter et al., 2017; Skelton et al., 2018). As 75 opposed to the large interspecific variation, it seems that intraspecific variation in 76 resistance to embolism is limited. However, an analysis of 46 species suggests that 77 significant intraspecific variation may occur (Anderegg, 2015). 78

Leaf water potential (Ψ_1) is an indicator of plant water status. Pre-dawn leaf Ψ 79 (Ψ_{PD}) is measured when the plant is in equilibrium with soil water, and is a measure of 80 the soil water availability as perceived by the plant. Ψ_{PD} is affected by drought severity 81 and root depth (Nardini *et al.*, 2016). The difference between either the Ψ_{50} or Ψ_{88} value 82 and the minimum water potential observed in field conditions (Ψ_{min}) is defined as the 83 hydraulic safety margins (HSM) (Meinzer et al., 2009; Martin-StPaul et al., 2017). A 84 narrow HSM means proximity to thresholds where there is a risk of hydraulic failure. A 85 meta-analysis showed that most plants that resist drought do so by stomatal closure at a 86 87 much higher water potential than that which causes hydraulic failure (Martin-StPaul et al., 2017). This suggests that species in nature avoid hydraulic failure by sacrificing 88 photosynthesis (Meinzer et al., 2009; Johnson et al., 2011; Martin-StPaul et al., 2017; 89 Creek et al., 2020). 90

Photosynthetic performance is usually measured in ecological studies through leaf carbon isotope discrimination, δ^{13} C (Cernusak *et al.*, 2013). Leaf δ^{13} C is used to assess leaf gas exchange characteristics in C3 terrestrial plants. It is commonly used as a proxy for intrinsic water use efficiency (WUEi), which is the ratio between carbon assimilation (A) and stomatal conductance (g). WUEi is considered an integrative, long-term evaluation of photosynthetic performance rather than an instantaneous measurement (Dawson *et al.*, 2002).

Stomatal closure has been shown to be correlated with the cell turgor-loss-point (TLP) (Mencuccini *et al.*, 2015; Bartlett *et al.*, 2016; Martin-StPaul *et al.*, 2017). To keep turgid cells, plants may invest energy to reduce the cell TLP during dehydration. The water potential for TLP (Ψ_{TLP}) is reduced through active accumulation of solutes, i.e. osmotic adjustment (Bartlett *et al.*, 2012). The seasonal course leaf osmotic potential reveals the plasticity of osmotic adjustment as related to environmental changes (Bartlett *et al.*, 2012).

105 Studies focusing on leaf traits of co-existing woody species in Mediterranean 106 climates reveal variation in response to drought during the long rainless summer. Leaf 107 defoliation in response to extreme drought was observed in *Juniperus phoenicea*, 108 *Rosmarinus officinalis*, and *Rhamnus lycioides* but not in four other co-existing species 109 (Gazol *et al.*, 2017). Dehydration avoidance via stomatal closure at the cost of low carbon 110 assimilation was demonstrated by *Pinus nigra*, while neighboring *Quercus ilex* and 111 *Quercus faginea* showed dehydration tolerance via osmotic adjustment. The observed

osmotic adjustment was more robust in the evergreen Q. ilex than in the semi-deciduous 112 Q. faginea, explaining its better resistance to drought (Forner et al., 2018a). However, Q. 113 faginea responded to intensified drought conditions by a more robust plastic stomatal 114 response than Q. ilex (Klein et al., 2013; Forner et al., 2018b). Independent studies that 115 measured δ^{13} C agreed that pine exhibited better WUEi than co-existing oak species under 116 prolonged drought or inter-annual precipitation differences. A significant variation in 117 stomatal regulation, detected through δ^{13} C and oxygen isotope composition in ten co-118 existing species in Spain, further emphasized the contrasting WUEi among species 119 (Moreno-Gutiérrez et al., 2012). Combined analysis of Ψ_{PD} and sap flow of five species 120 allowed designating Pinus halepensis, Pistacia lentiscus, and Erica multiflora as water 121 savers, versus Quercus coccifera and Stipa tenacissima as water spenders (Chirino et al., 122 2011). 123

The above studies imply interspecific differences in the plastic stomatal response 124 or osmotic adjustment; however, resistance to embolism, a key trait in species resistance 125 to drought, was missing from those studies. Large variation in resistance to embolism in 126 Mediterranean climates was found among 19 fynbos species in South Africa (Pratt et al., 127 2012) and nine chaparral species of the Rhamnaceae in California (Pratt et al., 2007) 128 Both studies concluded that resistance to embolism is linked to the species' post-fire 129 130 recruitment strategy. In a study conducting year-round measurements of resistance to embolism, water potential, and Ψ_{TLP} in three co-existing species (Väänänen *et al.*, 2019), 131 it was found that *Phillyrea latifolia* tolerates drought via xylem resistance to embolism 132 and osmotic adjustment, while co-existing Pistacia lentiscus and Quercus calliprinos 133 avoid drought via stomatal regulation. However, these results did not explain the high 134 mortality rate of *O. calliprinos* compared to the other species, suggesting that an 135 additional mechanism might occur in these co-existing species. This emphasizes the 136 importance of evaluating a range of traits, species, and aridity gradients to represent 137 better each species' total traits repertoire used for drought resistance. 138 Here, we tested the hypothesis that co-existing Mediterranean species differ in drought 139 resistance strategies, however, all can respond to drought intensification through plastic 140 traits. To this end, we studied five co-existing species at three sites with different aridity, 141 one of which is near the dry margin of the species distribution. We measured predawn 142 and midday water potentials during the dry season and $\delta^{13}C$ at the end of the dry season, 143 resistance to embolism at the end of the wet season, and native and full-turgor osmotic 144

potentials as related to midday water potential over the course of the dry season. The

resulting data set allowed us to evaluate the response of co-existing species to prolonged

- 147 drought.
- 148

149 Materials and methods

150 *Sites and species*

The steep climatic gradient in Israel is governed by Mediterranean weather patterns, 151 characterized by long, dry summers, and changes gradually from mesic-152 Mediterranean to arid from north to south (Tielbörger et al., 2014). Three sites were 153 154 chosen to represent Mesic-Mediterranean (MM), Mediterranean (M) and Semi-arid (SA) climate conditions along the natural rainfall gradient (Table 1). The three sites 155 were undisturbed for the last 40 years, and thus represent ecologically equilibrated 156 environments. Climate data was taken from stations of the Israel Meteorological 157 Service (IMS, ims.gov.il); station Michmanim for the MM site, station Ramat 158 Hanadiv for the M site, and station Netiv HaLamed-Heh for the SA site. All sites 159 were characterized by a prolonged summer-fall rainless dry period, from the end of 160 April to October. Aridity indexes (mean annual precipitation divided by potential 161 evapotranspiration) for the SA, M, and MM sites are 0.27, 0.39, and 0.46, 162 respectively. Annual precipitation (from September to August) in 2017-2018 was 337, 163 624, and 700 mm, and in 2018-2019 was 504, 651, and 1039 mm at the SA, M and 164 MM sites, respectively. Temperature also differed along the climate gradient and 165 average daily maximum temperature in the summer was 35°C, 32°C, and 29°C, 166 respectively (Fig. 1a). Maximum VPD in the summer ranged from 4 to 2.5 kPa from 167 SA to M, respectively (Fig. 1b). Soil types varied between the sites and the degree of 168 clayey soil decreased from north to south. Five predominant native woody species 169 were selected for the research (Table 2), which co-existed in 2,500 square meter plots 170 at each of the three research sites. The SA site is close to the dry southern limit of the 171 Mediterranean zone and the studied species (Supporting Information Fig. 1) (Danin & 172 173 Plitmann, 1987). For all trait measurements described below, samples were taken from specific five 174

175 labeled individuals per species at each site, unless otherwise stated.

176

177 Hydraulic vulnerability measurements

178 Vulnerability curves (VC) for percent loss of conductivity (PLC) of stem samples as a

- 179 function of water potential were measured in a Cavitron (Cochard, et al., 2005). This
- 180 was done for all species except QC, which was not measured due to its long vessels (>
- 181 30 cm, personal data). It was impossible to find QC branches 1 m long for
- measurements in a nonstandard Cavitron with a large rotor diameter. Thus, we
- measured the PLC values of *Quercus coccifera*, which is an evergreen oak that
- belongs to the subgenus *Quercus* section *Cerris* and is considered a subspecies of QC
- 185 (Toumi & Lumaret, 2010).
- 186 To avoid native embolism in branches that were tested for VC and PLC, samples were
- 187 collected from the three sites at the end of the rainy season (April 2018), when water

potentials were less negative than -1.5 MPa. Samples of RL at the MM site were not

included in this analysis, as its branches were too short for the Cavitron. Two terminal

branches (1 cm diameter and 100 cm in length) were harvested from the upper canopy

[1]

- 191 of 5-7 individuals of each species and were sent in overnight mail to France (to
- 192 Bordeaux and Clermont-Ferrand). VC curves were determined as previously
- described (Lamy et al., 2014). PLC was calculated every 1-2 MPa, following the
- 194 equation:

195
$$PLC = 100 * (1 - \frac{K}{K_{max}})$$

The sigmoidal curve was fitted to the following equation (Pammenter & Van derWilligen, 1998):

198 [2]

$$PLC = \frac{100}{[1 + \exp(\frac{S}{25} * (\Psi - \Psi_{50})]]}$$

Where
$$\Psi_{50}$$
 (MPa) is the xylem pressure inducing 50% loss of conductance and S (% MPa⁻¹) is the slope of the vulnerability curve at the inflection point. Predicted PLC (PLC_P) was calculated according to the actual leaf water potential and Equation 2.

204 *Field measurements of water potential*

Field campaigns were conducted monthly at all sites during the rainless period (May -205 September) in two consecutive years; 2018, where predawn and midday water 206 potentials were measured (Ψ_{PD} and Ψ_{MD} , respectively) and in 2019, where only Ψ_{MD} 207 was measured. The measurements were made in a Scholander-type pressure chamber 208 (PMS, Corvallis, OR, USA). The decline in Ψ_{MD} in relation to soil dehydration (as 209 reflected by Ψ_{PD}) was analysed according to Meinzer et al. (2016). Regression lines 210 were calculated for all species for each site. Slopes, Hydroscapes (which is a metric of 211 stomatal control based on the area between the 1:1 line and the regression slope), and 212 Ψ_{g_0} (extrapolated to find the value at which $\Psi_{PD} = \Psi_{MD}$) were calculated from the 213 above analysis both for different sites for each species and for each species separately 214 at all sites included (Supporting Information Table S8, S9). Ψ_{min} was taken as the 215 lowest value of measured midday water potential in the field. Hydraulic safety 216 margins (HSM) were calculated as Ψ_{min} - Ψ_x , where Ψ_x is the xylem pressure inducing 217 12, 50 or 88% loss of branch hydraulic conductivity. 218

219

Leaf $\delta^{13}C$ 220

Carbon isotope ratio (δ^{13} C) was measured in mature healthy sunlit leaves collected at 221 the end of the dry period (August 2018) with a ¹³C cavity ring down analyzer 222 (G2131i, Picarro, Santa Clara, CA, USA) as described by Nemera et al. (2020). Leaf 223

intrinsic water-use efficiency (WUEi) was calculated using the species mean based on 224 a leaf-scale model of C3 photosynthetic isotope discrimination (Farquhar et al., 1989):

226

225

227

[4]

228
$$WUI_{i} = \frac{C_{a}(b - \delta^{13}C)}{1.6(b - a)}$$

229

Where C_a is the atmospheric CO₂ concentration in PPM and a and b are fractionation 230

factors occurring during diffusion of CO₂ through stomata pores (4.4‰) and 231

enzymatic carbon fixation by Rubisco plus a small component accounting for 232

mesophyll conductance (27‰), respectively. 233

235 Osmotic potential

236	Leaf samples in which water potential was measured were frozen in liquid nitrogen
237	for native osmotic potential (Ψ_s) measurements. For full turgor osmotic potential (Π_0)
238	measurements, an additional shoot, harvested from each individual, was cut under
239	water, rehydrated for 2 hours, and measured in the pressure chamber to verify
240	rehydration. For both cases (Ψ_s and Π_0), samples were packed into 250 µl tubes and
241	were frozen in liquid nitrogen. Upon thawing, holes were drilled in the bottom of the
242	frozen tubes, which were then put into other clean tubes that collected the liquid when
243	centrifuged at 15000 RCF (g) for 2 min. Ten microliters from each Osmolality
244	(mmol) of the samples was assessed with a vapor-pressure osmometer (VAPRO 5520
245	Wescor, Logan, UT). Conversion to pressure units was done by Van't Hoff equation:
246	
247	[5]
248	$\Psi_{\pi}(MPa) = nRT/10000$
249	in which n is the solute concentration in mol/L, R is the universal gas constant
250	(8.314472 L bar K ⁻¹ mol ⁻¹), and T is the temperature in °K. Temperature was taken as
251	25°; conversion ratio was 403.33 mmol/MPa.
252	Ψ_s and Π_0 were analyzed by correlation with the Ψ_{MD} values, and by covariance
253	analyses which were performed to test the influence of Ψ_{MD} , site, and the interaction
254	between Ψ_{MD} and site on Ψ_s and Π_0 . Slope regression analysis of Ψ_s Vs. Ψ_{MD} is a
255	proxy for the osmotic potential due to both cell shrinkage and osmotic adjustment
256	(OA, i.e., active solute accumulation). Slope regression analysis of Π_0 Vs. Ψ_{MD} is a
257	proxy for OA only.
258	
259	Species characterization by drought-resistance strategies
260	Tolerance and Avoidance were quantified as numbers between 0 and 100 (less to
261	most, respectively) for each species. Each strategy was evaluated from the below
262	measured parameters which were converted to normalized values (NV) as follows:
263	NV=(x-min)/(max-min) [6]
264	Where x is the measured value, and min and max are the minimum and maximum
265	thresholds of each parameter.

"Tolerance" refers to "Xylem tolerance", and was taken to be a normalized value of Ψ_{50} . Values for normalization were from 0 to -18 MPa, the latter being the most negative Ψ_{50} reported (Larter et al., 2015).

269 "Avoidance" was calculated as the average of "Water access", "Stringency of

stomatal control", and "Osmoregulation". "Water access" was normalized from the

- 271 minimum seasonal Ψ_{PD} at the SA site. Values for normalization were from 0 to -10
- 272 MPa. Normalized values were subtracted from 1. "Stringency of stomatal control"
- was normalized from Hydroscapes (HS), calculated from Ψ_{PD} vs. Ψ_{MD} according to
- 274 Meinzer et al. (2016), where the full range of values was from 0 to 10 MPa.
- Normalized values were subtracted from 1. "Osmoregulation" was normalized from

the slope of Ψ_{MD} vs. Π_0 . Values for normalization were from 0 to 1.

277 "Escape" was the rank for "Drought-deciduous". "Drought-deciduous" was evaluated

from the literature (1, 0.5, and 0, refer to full-, partial-, and non- deciduous,

respectively). Among the studied species, only RL is known to be partially drought-

deciduous (Gazol et al., 2017).

281

282 Statistical Analysis

Analysis of variance (ANOVA) was used (Python software, Python Software

Foundation; JMP 14 software, SAS Institute Inc., Cary, NC, USA) to identify

significant differences between species and sites. The Tukey-Kramer post hoc test

was used to compare the results. Data fitting was carried out using Python software.

- 287 Bartlett's test for homogeneity of variances, using JMP 14 software was used to
- compare interspecific variation among sites and along the season within sites.
- Analysis of co-variance was used to test influence of both site and Ψ_{MD} on Π_0 and Ψ_s ,
- and also used to test influence Ψ_{PD} and site on Ψ_{MD} .
- 291

292 **Results**

293 Effect of environmental drought on hydraulic traits

Resistance to embolism – Large differences in resistance to embolism were found

between species, with the highest Ψ_{50} for PP (~ -5 MPa) and lowest for PHL (< -10

MPa). All parameters of vulnerability curves for resistance to embolism per species,

including slope, Ψ_{12} , Ψ_{50} , and Ψ_{88} , were similar at the three sites (Fig. 2, Table 3,

Supporting Information Tables S1-S6), and were not influenced by the site, as shown

by a two-factorial ANOVA (Table 3).

Leaf water potential - Minimum Ψ_{MD} was found at the end of the season at the SA 300 site. For QC and PHL minimum values were close to, but did not decline below Ψ_{12} , 301 while for the other species minimum Ψ_{MD} values were significantly lower than Ψ_{12} 302 (Fig. 3). Significant differences in Ψ_{PD} between sites for each of the species were 303 found in each sampling along the dry season (Fig. 3, Supporting Information Table 304 S7). A strong influence of site on Ψ_{PD} was found at the beginning and end of the dry 305 season (Table 3). Ψ_{PD} was more negative at the SA than at the MM and M sites (Fig. 306 **3a-e**, Supporting Information Table S7). Ψ_{min} was significantly influenced by site in 307

308 both 2018 and 2019 (Table **3**, Fig. **3f-g**).

HSM and PLCp – The HSM in 2018 was narrower at the SA site than at the M and 309 MM sites for three of the species, while the other two species had narrower HSM's at 310 the SA and M site in comparison to the MM site (Fig. 4a,b, Supporting Information 311 Tables S1, S3, S5). These differences were less prominent in 2019 (Fig. 4d.e, 312 Supporting Information Tables S2, S4, S6). Values of HSM₅₀ less than 1 MPa were 313 found at the SA site in 2018 (for PP and RL, Supporting Information Table S1), and the 314 predicted PLC (PLCp) for those species reached values of 30% or more (Fig. 4). In both 315 2018 and 2019 HSM₁₂ declined to negative values at the SA site (Fig. 4a,e, Supporting 316 Information Tables S3, S4). The calculated HSM of Ψ_{12} , Ψ_{50} , and PLCp were affected 317 by site in 2018, but not in 2019, which was a wetter year (Table 3, Fig. 1). 318

Carbon-water balance - Leaf δ^{13} C and its derivative WUEi was higher at the drier site than at the wetter sites. Differences were significant for two of the species (Fig. 5). The site influence on leaf δ^{13} C were significant in the true fortenial ANOVA (Table

5). The site influence on leaf δ^{13} C was significant in the two-factorial ANOVA (Table 3).

Osmotic potential - Changes in osmotic potential were significant for all the species along the dry season, in relation to the decline in Ψ_{MD} (Fig. 6 f-j, Table S14).

325 Covariance analysis for Ψ_s revealed a significant site effect only for PL, and a

significant Ψ_{MD} effect for all species (Supporting Information Table S14). Osmotic

- adjustment differed substantially among species, being large in PHL and QC and
- minor in RL (which was expressed in significant Ψ_{MD} effect) negligible in PL, and

nonexistent in PP (Fig. 6f-j, Supporting Information Table S10). Covariance analysis

for Π_0 revealed a significant site effect only for PL and PHL (Supporting Information

331 Table S15).

Accordingly, most of the reduction in Ψ_s during the dry season in QC is from active

osmolyte accumulation, while in PL, PP and RL most of the reduction is due to cellshrinkage. PHL seems to retain both mechanisms.

335

336 Species comparison by trait

Resistance to embolism - Resistance to embolism, expressed by Ψ_{50} (Fig. 2), Ψ_{12} ,

and Ψ_{88} , showed interspecific variation at the SA and M sites (Table 3, Supporting

Information Tables S1-S6). PHL demonstrated the most negative values followed by

RL and PL, while PP showed the least negative value.

Water potential - Interspecific variations were evidenced at each site, and at each measurement event (Supporting Information Tables S11, S12). PHL had the most negative Ψ_{PD} , followed by RL and PL, while QC and PP had the highest Ψ_{PD} values (Fig. **3a-e**). Two-factorial ANOVA showed that Ψ_{min} , which is the lowest Ψ_{MD} value at the end of the dry season, was affected by the species and site. However, it was affected by the interaction of species-on-site only in 2018 (Table 3).

A comparison of the interspecific variation among sites using Bartlett's test for

homogeneity of variances, resulted in a significant difference (P = 0.0259) between

349 sites at the beginning of the dry season. Interspecific variation was evidenced at the

SA site more than at the M and MM sites (Table 4). The rest of the sampling dates did

not show a significant difference in interspecific variation between sites; however, the

352 SA site had higher values than the M and MM sites (Table 4).

Species showed different evolution of Ψ_{PD} along the dry season, and reached different

values at the end of the season (Fig. **3a-e**). In each site, the interspecific variation

increased during the dry season (Table 4), however, a comparison of the variations

along the season at the different sites (using Bartlett's test) found a significant (P =

0.0258) increase in interspecific variation only at the M site (Table 5). Slopes of Ψ_{PD}

along the dry season were different between species in each site, while the higher

value was that of RL and PHL, followed by PL, PP, and QC (Supporting InformationTable S18).

HSM – Similar to Ψ_{PD} , the HSM of Ψ_{50} , Ψ_{12} , and Ψ_{88} showed interspecific variation

362 (Tables 3, Supporting Information Tables S1-S6). PP (at SA and M sites) and RL (at

SA site) had a very narrow HSM, i.e. less than 1MPa (Fig. 4). However, in 2019,

- which was a wetter year (Fig. 1), the HSM of all species was wider than in 2018. In
- 2018, the HSM based on Ψ_{12} at the SA site reached negative values for all species,
- except for QC, which had 0.02±0.21 MPa (Fig. **4a**, Supporting Information Table S3).
- In 2019, negative HSM values for Ψ_{12} were recorded only for PP and RL at all sites
- 368 (Fig. **4d**, Supporting Information Table S4).
- 369 Carbon-water balance One-way ANOVA suggested differences between species in

leaf δ^{13} C and WUEi per site (*P* = 0.0682, Table 3, Supporting Information Table

- S13). A comparison of the interspecific variation among sites using the Bartlett's test
- suggested a strong tendency to significance (P = 0.0693), with the MM site showing
- the highest interspecific variation, while SA and M sites showed reduced variation
- (Table 4). These results were attributed to PHL and QC, which had significant

differences in leaf δ^{13} C between sites (Fig. 5a).

Osmotic potential - Osmotic potential differed significantly between species in Ψ_s and Π_0 (Table 3). In addition, significant differences between species were revealed by covariance analysis for the different species at different sites (Supporting Information Tables S16, S17).

380

381 *Correlation between traits*

The relationships between Ψ_{PD} and Ψ_{MD} - Analysis of the various sites for each 382 species showed that the slopes at the drier sites (M and SA) were steeper than the 383 slope at the M site, except for QC. However, covariance analysis did not show 384 significant differences between the slopes, except for PL. Hydroscape values for all 385 species were always larger at the SA site than at the two wetter sites (Fig. 3f-j, 386 Supporting Information Table S8, S19). Analysis which includes the data from the 387 various sites for each species showed that slopes ranged from 0.83 for QC to 0.68 for 388 RL (Supporting Information Table S9). For QC and PP, the extrapolated values were 389

several MPa lower than the lowest data points, -8.9 and -8.7 MPa, respectively.

- However, for PHL, RL and PL the lowest points were close to the regression values at
- equality, -8.7, -7.6, and -7.1 MPa, respectively (Supporting Information Table S9).
- ³⁹³ Hydroscape values were 10.7, 9.15, 7.31, 6.31, and 3.52 MPa², for RL, PHL, PL, PP,
- and QC, respectively (Table S9).
- 395

396 The relationships between osmotic parameters and Ψ_{MD}

For all species significant linear correlations were found between osmotic potential, 397 Ψ_s , and Ψ_{MD} along the season (Fig. **6a-e**). The effect of Ψ_{MD} on Ψ_s was evident for all 398 species, while a site effect was found only for PL (Supporting Information Table 399 S14). A linear correlation was also found between Π_0 and Ψ_{MD} (Fig. 6f-j, Supporting 400 Information Table S10), while only QC, PHL, and RL had a Ψ_{MD} effect on Π_0 401 (Supporting Information Tables S15). There was evidence for a site effect on Π_0 for 402 PHL and PL, while no effect for the interaction of Ψ_{MD} with site was revealed for any 403 of the species (Supporting Information Table S15). The largest osmotic adjustment 404 405 was observed for PHL, and ranged from -2 to -5 MPa along the dry season, with similar responses at all sites (Fig. 6j). Osmotic adjustment for QC ranged from -2 to -406 3 MPa during the dry season, mostly at the M and SA sites (Fig. 6f). A weak, but 407 significant, osmotic adjustment was observed for PL and RL at the SA site (Fig. 6g-i, 408

- 409 Supporting Information Table S10).
- 410

411 Species characterization by drought-resistance strategies

The results described above allowed the assessment of drought resistance strategies

413 for each of the studied species (Fig. 7). That was achieved by quantifying tolerance,

414 avoidance, and escape strategies, as described in the Material and Methods

415 (Supporting Information Table S19). Thus, we found that extreme resistance to

- 416 embolism confers tolerance in PHL, which also uses osmotic adjustment to resist
- 417 drought. Osmotic adjustment is a major trait in QC, which probably has deep roots to
- 418 access water, supporting its stomatal opening during drought. PL and RL do not use
- 419 osmotic regulation but have intermediate values of resistance to embolism and almost
- 420 complete stomatal closure. PP has low resistance to embolism and no osmotic
- regulation, but its continued water uptake along the dry season suggests it has deep
- roots. RL is known to escape drought by partial leaf defoliation (Gazol et al., 20017).

423 Discussion

The resulting data set of the current study, in addition to known species-specific 424 characteristics, demonstrated that co-existing Mediterranean species minimize 425 mortality risk by combining drought resistance strategies. The results support the 426 hypothesis that regulation of water potential is a result of a robust plastic response, 427 while resistance to embolism is a fixed rigid trait that is not affected by site aridity. 428 These characteristics led to a decline in the HSM with increasing drought intensity 429 leading to negative HSM in several cases and suggesting hydraulic failure at the dry 430 SA site. 431

432

433 *Relating environmental factors to phenotype*

The large data set provided by this study, encompassed three aspects of environmental 434 drought, including site aridity, seasonal drought, and inter-annual climate differences. 435 While all site characteristics were less favorable at the SA site than at the two wetter 436 sites, precipitation and VPD differed the most, and actually played major roles in 437 determining the drought intensity of the site. The rainless summer, together with the 438 inter-annual precipitation differences, further emphasized the stress intensity that the 439 studied species confronted. The SA site differed significantly from the other two sites 440 for most of the traits, suggesting that species were closer to their physiological limits 441 at the dry edge. The latter is in agreement with Feng, et al. (2019) and Guo, et al. 442 443 (2020) who emphasized that temporal and spatial variability in the environment is important in determining plant response to drought, as opposed to characterization of 444 445 species without considering environmental influences.

Our results show interspecific variation in resistance to embolism (Fig. 2). However, 446 no intraspecific variation was evident for this trait. A lack of intraspecific variation is 447 in agreement with previous studies that showed similar resistance to embolism in 448 distantly separated populations (Martínez-Vilalta et al., 2009; Lamy et al., 2014; 449 González-Muñoz et al., 2018; Lobo et al., 2018; Li et al., 2019; Bittencourt et al., 450 2020). However, although the three tested sites differ in climate characteristics, the 451 lack of intraspecific variation might also be due to the continuous geographical 452 distribution of the tested species (Figure S1), which prevented differentiation between 453 populations due to continuous gene flow. In addition, it is possible that other remote 454

populations, which were not included in the current study, do possess intraspecific 455 variation in resistance to embolism. Several studies have reported intraspecific 456 variation in resistance to embolism in angiosperms. Examples are *Cordia alliodora*, 457 Artemisia tridentate, Fagus sylvatica, Populus trichocarpa, and in Mediterranean and 458 chaparral shrubs (Kolb & Sperry, 1999; Sparks & Black, 1999; Choat et al., 2007; 459 Wortemann et al., 2011; Pratt et al., 2012; Jacobsen et al., 2014; Stojnić et al., 2017). 460 As opposed to the stability of resistance to embolism, the stomatal response was very 461 plastic as reflected in changes in leaf water potential (Ψ_{PD} and Ψ_{MD}) in response to 462 drought in all species in relation to site aridity, seasonality, and inter-annual climate 463 differences (Fig. 3, Table 3). The difference between species in the Ψ_{PD} slope along 464 the season, especially at the SA site (Fig.3, Table S18), suggests species 465 differentiation in the degree of plasticity, where RL and PHL showed the strongest, 466

- and QC showed the least plastic response. The interspecific variation in Ψ_{PD} seemed
- to increase with drought and along the dry season, i.e., it was inversely related toaridity (Tables 4, 5).
- 470 The hydroscapes (HS, Supporting Information Tables S8, S9), which showed
- 471 differences between species, can be used as proxies for stringency of stomatal
- regulation (Meinzer et al., 2016). They gave a nearly mirror image of plasticity, where
- 473 QC had the most stringency of stomatal regulation, while PHL and RL had the least
- 474 stringency (Fig. **3**, Supporting Information Tables S8, S9).
- 475 Variations in Ψ_{PD} and its plasticity between species along the dry season may reflect
- 476 differences in root depth, where Ψ_{PD} of more deeply rooted species, such as QC,
- 477 appears high with weak plasticity along the season (Crombie et al., 1988). Roots of
- five meter depth have been reported for PP (Jakoby et al., 2020), and QC (Canadell et
- al., 1996). Co-existing species with different root depths sustain niche segregation to
- 480 share soil water resources (Palacio *et al.*, 2017). Niche segregation has been shown in
- 481 Mediterranean-type ecosystems through the leaf life span and Ψ_{min} as "anchor traits"
- among different morphological traits used to distinguish contrasting strategies of
- drought tolerance vs. avoidance(Ackerly, 2004). Our study, which focused on
- embolism resistance and leaf water potential, also suggests niche segregation in
- 485 Mediterranean species. Another approach to exploring niche segregation divides
- 486 species by different water use patterns (Redtfeldt & Davis, 1996), which has been

recently shown to be important in increasing forest productivity and the carbon sink in 487 semi-arid regions (Rog et al. 2021). Our study suggests that niche segregation is 488 sustained under different drought conditions. 489

490

515

The dearth of hydraulic safety margins near the dry edge of species distribution 491

The value of Ψ_L for complete stomatal closure (Ψ_{g_0}) for QC and PP was more than 1 492

MPa lower than Ψ_{\min} , while for PL, RL and PHL Ψ_{\min} was close to Ψ_{g_0} (Fig. **3f-j**). 493

The proximity of the minimum Ψ_{PD} to Ψ_{g_0} at the SA site indicates that these plants 494

approached the point of null activity, which corresponds with the large reduction in 495 Ψ_{PD} across sites, as shown in Fig. 3. 496

The Ψ_{50} HSM seemed to change according to annual precipitation (MAP, Fig. 4), 497

especially at the SA site, suggesting that a drought year would have more impact on 498

species vulnerability. This is in agreement with Ziegler et al. (2019), who showed that 499

for tropical trees the Ψ_{50} HSM became narrower, but still positive, in dry years. In the 500

current study, all species (except for QC) crossed the Ψ_{12} threshold (Fig. **3f-j**) and 501

502 some degree of difference between Ψ_{PD} and Ψ_{MD} remained, suggesting that species

maintain some stomatal opening when embolism is low. Interestingly, PP had a 503

504 negative Ψ_{12} HSM in all sites in both years (Fig. 4), emphasizing the trade-off

between hydraulic safety and carbon assimilation in this winter-deciduous species. In 505

addition, results of predicted PLC (PLCp, Fig. 4c,f) suggest that all species 506

experienced embolism, which increased with drought intensity. These results suggest 507

that species approach the limit of hydraulic capacity at the site near the dry margins of 508 their distribution. 509

Evidence for embolism of stem xylem in nature is rare. Johnson et al. (2018) recently 510 reported measurements implying that negative HSM's in Quercus fusiformis and 511

Prosopis glandulosa occurred during the most severe drought in recorded history in 512

central Texas. Fontes et al. (2018) found negative HSMs in Eschweilera cyathiformis

513

- and Pouteria anomala that experienced extreme drought during the strong El Nin^o o 514 that occurred across Amazonia in 2015–2016. The two above reports resulted from
- extreme climate events, while the species in our study seem to confront severe
- 516

drought every year. Recent studies on Prunus ramonensis and Pyrus syriaca also 517

reported potential embolism in nature (Paudel et al., 2019a; Paudel et al., 2019b). 518

Taking a modeling approach, Benito et al. (2018) use minimum soil water potential

data and HSMs of 44 European woody species and found that negative HSMs explain

521 the mortality of 15 species at the driest margins of their distribution.

522

523 Interspecific variation in response to drought

524 The interspecific variation in Ψ_{PD} increased with aridity and along the dry season

525 (Tables 4, 5). This difference was supported by the significant effect of the species-

by-site interaction on Ψ_{PD} (Table 3). However, this effect appeared only at the

527 beginning of the dry season, where species operate at their relatively maximal

528 physiological activity. As drought progressed, species reached minimum water

529 potential, at which time the effect of site on differences was not significant.

530 The Ψ_{PD} results were supported by the leaf $\delta^{13}C$ values that increased with site aridity,

suggesting an increase in WUE_i (Fig. 5) (Farquhar et al., 1989). Similar results were

obtained by Rumman et al. (2018), who found a negative correlation between MAP

and WUE_i for precipitation up to 1000 mm/year, above which the trend flattened,

indicating that isotopic discrimination in wet environments remained nearly constant.

535 The current study shows that the interspecific variation in leaf δ^{13} C tends to be larger

at the MM site than at the M and SA sites (Table 4). This result suggests that genetic

variation in carbon assimilation rate is more pronounced in environmental conditions

favoring high stomatal conductance, as compared to the M and SA sites. It also

suggests that species under severe environmental drought, that demonstrate different

540 plasticity, reach similar minimum rates of carbon assimilation (Fig. 5). This is in

agreement with Forner et al. (2018b), who showed that the interspecific variation in

leaf δ^{13} C in three woody Mediterranean species in two consecutive wet years was

reduced after an extremely dry year. Together, these results encourage the

544 measurement of interspecific variation in carbon assimilation rates in wet rather than

545 dry environments. Furthermore, high versus low interspecific variation in leaf δ^{13} C

546 could be a proxy for evaluating water stress in multi-species ecological niches.

547 As all the species in our study suffered from severe drought at the SA site, as

548 manifested in a significant reduction in HSM's, not all showed osmotic adjustment,

549 suggesting that this mechanism is species dependent. Osmotic adjustment in drought

has been reported for PHL (Serrano *et al.*, 2005), and is also well known in Olive

species, which are related to PHL (*Oleaceae* family) (Lo Gullo & Salleo, 1988; Sofo

et al., 2008). It has also been found in *Quercus* species (Deligö & Bayar, 2018;

Aranda et al., 2020) but not in QC (as far as we know), and in PL (Álvarez et al.,

554 2018) only in response to salinity.

555 The low degree of osmotic adjustment in RL and nonexistent osmotic adjustment in

556 PP may also be related to their deciduous nature, and may result from a strategy of

allocating fewer resources to the leaf, similar to the findings of Liu *et al.* (2011) who

showed higher capacities of osmotic adjustment in evergreen shurbs than in decidous.

559

560 Conclusions

As illustrated in Figure 7, each of the co-occurring species in our study combines

drought-resistance strategies to minimize the risk of mortality. However, all

⁵⁶³ approached the limit of their hydraulic capacity at the site near the dry margins of

their distribution. The hydraulic limit was more pronounced in the drier year,

suggesting that a slight reduction in precipitation is more likely to put species at the

⁵⁶⁶ dry margins of their distribution at the risk of mortality.

567

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577 Author Contribution

AA, RDS, SC, SD, and HC designed experiments and interpreted data. AA, RB, and

579 VL collected and measured field samples. AA and RB performed vulnerability curves

580 measurements. AA and IR performed carbon isotope measurements. AA analyzed all

data. AA, SC, and RDS co-wrote the manuscript with contributions from SD, HC,

582 UH, and TK.

Table 1: Three selected sites with main geographical, edaphic, and climatic

584 characteristics.

Code Site	*AI	Latitude	Longitude	Elevation (m)	MAP * (mm)	SMT *(°C)	WMT *(°C)	Soil type**
MM	0.46	32°54'N	35°20'E	430	700	30	10	TR
М	0.39	32°33'N	34°56'E	80	550	32	16	TR and R
SA	0.27	31°40'N	34°56'E	300	400	31	17	Bright and Brown R

* AI, Aridity Index is the ratio of the annual precipitation and potential evapotranspiration.
MAP, mean annual precipitation; SMT, Average maximum daily temperature in July; WMT,
Average maximum daily temperature in January. *Data for 1981–2000, provided by the Israel
Meteorological Service. ** Data taken from GIS site of Israel Agriculture office
(https://moag.maps.arcgis.com/), TR and R correspond to Terra Rossa and Rendzinza,
respectively.

592

Table 2: Characteristics of species selected for the research.

Species	Code	Family	Life form	Leaf phenology
Quercus calliprinos	QC	Fagaceae	Tree	Evergreen
Pistacia palaestina	PP	Anacardiaceae	Shrub/Tree	Winter-deciduous
Pistacia lentiscus	PL	Anacardiaceae	Shrub	Evergreen
Rhamnus lycioides	RL	Rhamnaceae	Shrub	Partially drought-deciduous
Phillyrea latifolia	PHL	Oleaceae	Shrub	Evergreen

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595

Table 3: Two-factorial ANOVA of the effect of species, site and the interaction between site and species on all measured traits. For all embolism resistance parameters the 2 factors analysis was done on four species (PL, PP, PHL, and RL) for two sites (M and SA). For the other parameters, analysis was done on four species (QC, PL, PP, and PHL) for three sites. Bold P values indicate statistically significant result.

Trait	Species	Site	Species*Site
Ψ ₅₀	F(3,19)=84.5056	F(1,19)=0.4501	F(3,19)=1.4025
- 50	.P=<0.0001	,P=0.508	,P=0.2636
Ψ ₁₂	F(3,19)=17.4657	F(1,19)=0.0105	F(3,19)=0.6733
- 12	,P=<0.0001	,P=0.919	,P=0.5759
Ψ ₈₈	F(3,19)=27.4226	F(1,19)=0.719	F(3,19)=0.1714
- 88	,P=<0.0001	,P=0.4039	,P=0.9148
Slope	F(3,15)=0.6424	F(1,15)=0.0095	F(3,15)=0.9251
	,P=0.5944	,P=0.9229	,P=0.442
Ψ _{min} (2018)	F(3,75)=84.7131	F(2,75)=68.5311	F(6,75)=4.9622
	,P=<0.0001	,P=<0.0001	, P=0.0003
Ψ _{min} (2019)	F(3,39)=92.7639	F(2,39)=19.4777	F(6,39)=2.0137
	,P=<0.0001	,P=<0.0001	,P=0.087
Leaf $\delta^{13}C$ / WUEi	F(3,33)=2.4531	F(2,33)=22.3469	F(6,33)=1.3517
	,P=0.0806	,P=<0.0001	,P=0.2629
PLCp (2018)	F(3,75)=61.1677	F(2,75)=13.0761	F(6,75)=8.4556
	,P=<0.0001	,P=<0.0001	,P=<0.0001
PLCp (2019)	F(3,39)=62.9183	F(2,39)=9.6769	F(6,39)=6.3173
	,P=<0.0001	, P=0.0004	, P=0.0001
HSM ₁₂ (2018)	F(3,75)=109.954	F(2,75)=57.7828	F(6,75)=10.5175
	,P=<0.0001	,P=<0.0001	,P=<0.0001
HSM ₁₂ (2019)	F(3,75)=110.3402	F(2,75)=48.928	F(6,75)=10.6456
	,P=<0.0001	,P=<0.0001	,P=<0.0001
HSM ₅₀ (2018)	F(3,75)=111.7792	F(2,75)=40.8695	F(6,75)=11.3523
	,P=<0.0001	,P=<0.0001	,P=<0.0001
HSM ₅₀ (2019)	F(3,39)=87.4207	F(2,39)=17.8401	F(6,39)=5.0945
	,P=<0.0001	,P=<0.0001	, P=0.0006
HSM ₈₈ (2018)	F(3,39)=88.1662	F(2,39)=14.9253	F(6,39)=5.0271
	,P=<0.0001	,P=<0.0001	, P=0.0007
HSM ₈₈ (2019)	F(3,39)=89.8414	F(2,39)=12.7135	F(6,39)=5.5257
	,P=<0.0001	,P=<0.0001	, P=0.0003
$\Psi_{PD}(\text{Sep}_{2018})$	F(3,74)=73.7787	F(2,74)=39.6804	F(6,74)=2.3523
	,P=<0.0001	,P=<0.0001	,P=0.0391
$\Psi_{PD}(May_{2018})$	F(3,32)=17.1726	F(2,32)=85.8451	F(6,32)=21.302
	,P=<0.0001	,P=<0.0001	,P=<0.0001
Ψs (June 2019)	F(3,11)=10.7141	F(2,11)=0.3355	F(6,11)=1.8519
	,P=<0.0001	,P=0.7179	,P=0.1263
Π_0 (June 2019)	F(3,11)=21.5185	F(2,11)=7.5033	F(6,11)=4.9563
	, P=<0.0001	,P=0.0033	, P=0.0024

597

	Measurement	Standard deviation				Bartlett's
Parameter	Date	MM	М	SA	χ^2	value
$\delta^{13}C$	Summer 2018	1.153	0.328	0.464	χ2(2)=2.7	0.069
Ψ_{PD}	May 2018	0.374	0.201	0.922	χ2(2)=3.655	0.026
Ψ_{PD}	July 2018	0.860	0.894	1.474	χ2(2)=0.389	0.687
Ψ_{PD}	August 2018	0.998	1.273	1.726	χ2(2)=1.308	0.309
Ψ_{PD}	September 2018	1.308	1.284	1.731	χ2(2)=0.2	0.819

Table 4. Bartlett's test for homogeneity of variances for all species per date at the different sites.Bold indicates significant difference.

599

600

Table 5. Bartlett's test for homogeneity of variances for all species per site at the different dates. Bold indicates significant difference.

	_		Standard	_	Bartlett's		
Parameter	Site	May	July	August	September	χ^2	value
Ψ_{PD}	SA	0.922	1.474	1.726	1.731	χ2(3)=0.532	0.660
Ψ_{PD}	М	0.201	0.894	1.273	1.284	χ2(3)=3.092	0.026
Ψ_{PD}	MM	0.374	0.860	0.998	1.308	χ2(3)=1.139	0.332

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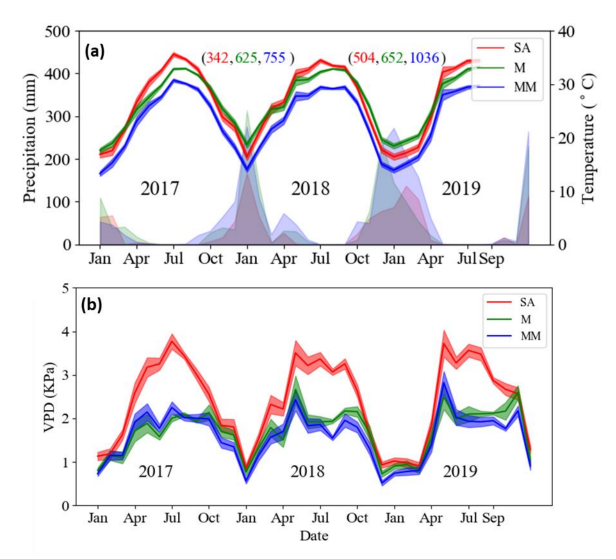
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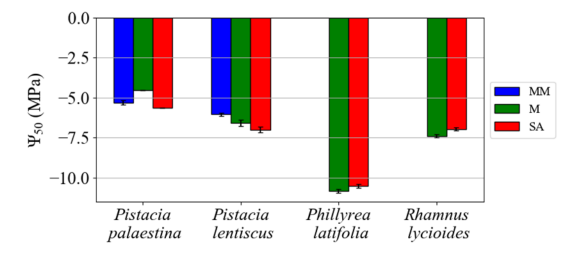
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Fig. 1: Meteorological data from the three study sites. (a) Monthly average daily
maximum temperature and monthly precipitation. Numbers in brackets represent the
annual precipitation for 2 consecutive winters (2017-2018, 2018-2019, calculated
from September to September) for the SA (red), M (green), and MM (blue) sites,
respectively. (b) Monthly average daily maximum vapor pressure deficit (VPD). Line
shadow represents standard error.

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Fig. 2: Ψ_{50} at the three sites (MM, Mesic-Mediterranean; M, Mediterranean; SA,

873 Semi-arid). *Quercus calliprinos* was not measured due to excessive vessel length.

874 Differences between species were significant, but no significant differences were

observed between sites. Error bars indicate standard error (n = 5-7).

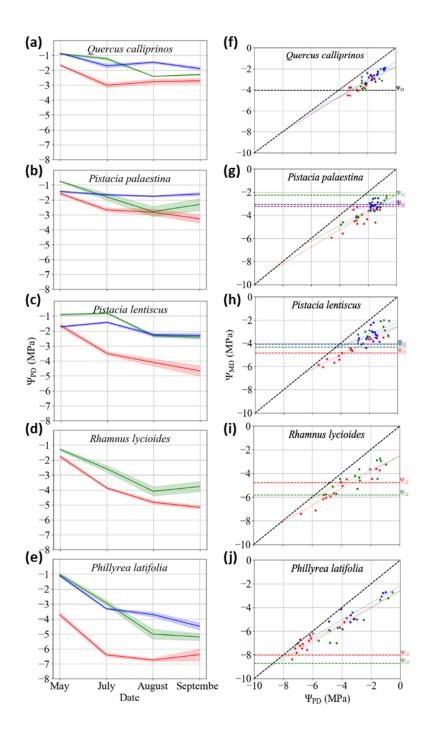


Fig. 3: Leaf water potential values during the dry season of 2018 at the Mesic-878 Mediterranean (blue), Mediterranean (green), and Semi-arid (red) study sites. a-e; 879 Courses of predawn leaf water potential for each species. Line shadow represents 880 standard error. **f-j**, Midday water potential (Ψ_{MD}) vs. predawn water potential (Ψ_{PD}) 881 for all species during the dry season. Each point on a plot is the average of two twigs. 882 Regression parameters can be found in Table S3. Thick dashed black line represents 883 1:1 ratio. The thin colored lines are the regressions of all points of each site. Vertical 884 dashed lines indicate the point of incipient embolism (Ψ_{12}) values for the MM (blue), 885 M (green), and SA (red) sites. Vertical dashed black line in f indicate point of 886 incipient embolism based on Q. coccifera data (see Material and Methods section). 887



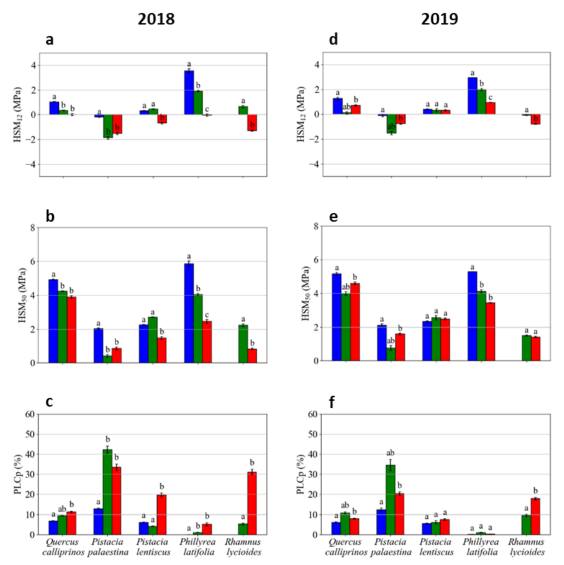
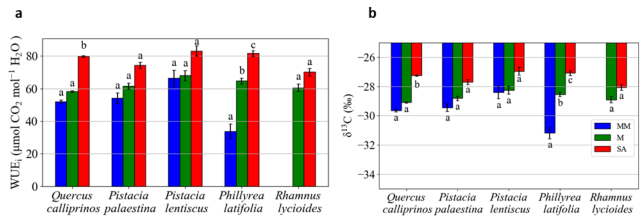


Fig. 4: HSM- Hydraulic safety margins (HSM12 and HSM50) and PLCp (% of
predicted embolism) for all species at the different sites, a-c represent 2018 data, d-f
represent 2019 data. MM, Mesic-Mediterranean; M, Mediterranean; SA, Semi-arid.
Different lowercase letters denote significant differences among sites. HSMs of QC

are based on the PLC curve of *Q. coccifera* (See Material and Methods).

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Fig. 5: Leaf δ^{13} C (a) and the derived WUEi (b) of the leaves of five species at the

- study sites at the end of the dry season. MM, Mesic-Mediterranean; M,
- 899 Mediterranean; SA, Semi-arid. Different lowercase letters denote significant
- 900 differences among sites.

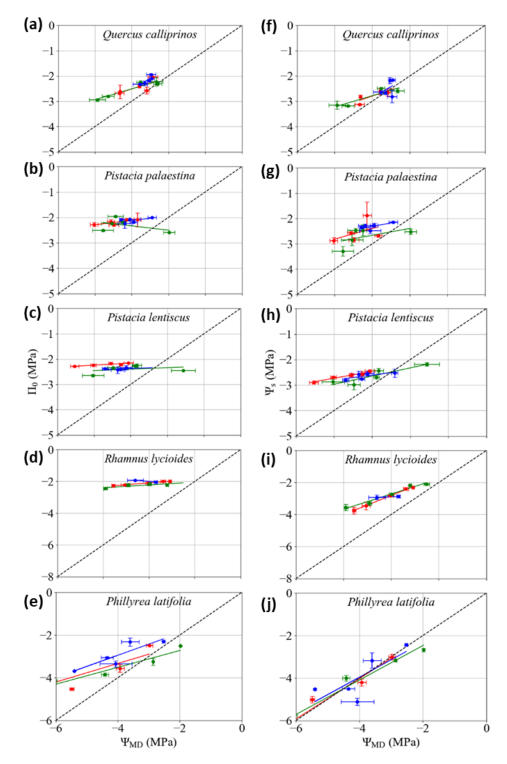
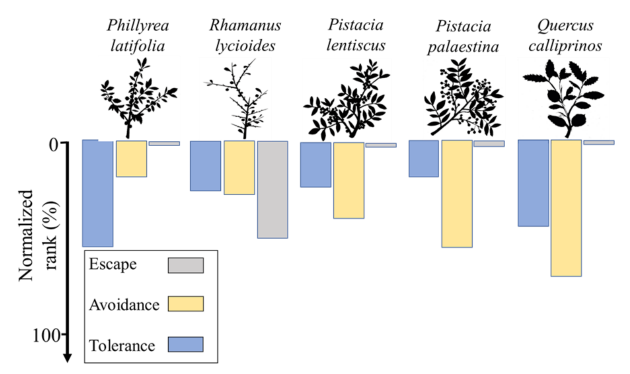




Fig. 6: Osmotic potential and leaf water potential for all species measured in summer 2019 at the Mesic-Mediterranean (blue), Mediterranean (green), and Semi-arid (red) sites. **a-e**: Leaf water potential at midday (Ψ_{MD}) vs. osmotic potential at full turgor (Π_0). f-j, Leaf water potential at midday (Ψ_{MD}) vs. native osmotic potential (Ψ_s). Regression parameters can be found in Supporting Information Table S10.



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910 Fig. 7: A superposition for the three different drought-resistance strategies (Tolerance,

Avoidance, and Escape) for a species, evaluated as normal parameters (scaled from 0

to 100%) derived from measured parameters. Detailed evaluations are presented in

Supporting Information Table S19. Drawing by Ilana Stein.

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916 Supporting Information

- Table S1: Ψ_{50} , Ψ_{min} and safety margins measured for the different species at the different sites in summer 2018.
- Table S2: Ψ_{50} , Ψ_{min} and safety margins measured for the different species at the different sites in summer 2019.
- Table S3: Ψ_{12} , Ψ_{min} and safety margins measured for the different species at the different sites in summer 2018.
- Table S4: Ψ_{12} , Ψ_{min} and safety margins measured for the different species at the different sites in summer 2019.
- Table S5: Ψ_{88} , Ψ_{min} and safety margins measured for the different species at the different sites in summer 2018.
- Table S6: Ψ_{88} , Ψ_{min} and safety margins measured for the different species at the different sites in summer 2019.
- Table S7: Ψ_{PD} ANOVA analysis between sites per species per measurement date.
- Table **S8**: Parameters for the linear regression of Ψ_{MD} vs. Ψ_{PD} for different species at different sites.
- Table **S9**: Parameters for the linear regression of Ψ_{MD} vs. Ψ_{PD} for different species.
- Table **S10**: Parameters correspond to Ψ_{MD} vs. Π_0 , in all species at all sites.
- Table **S11**: Ψ_{PD} ANOVA analysis between species per site per measurement date.
- Table S12: Ψ_{MD} ANOVA analysis between species per site per measurement date.
- Table **S13**: δ^{13} C ANOVA analysis between species per site per measurement date.
- Table **S14**: Summary of Covariance analysis testing influence of Ψ_{MD} , site and Ψ_{MD} X site on Ψ_{S}
- Table **S15**: Summary of Covariance analysis testing influence of Ψ_{MD} , site, and Ψ_{MD} 840 X site, on Π_0
- Table **S16**: Summary of Covariance analysis testing influence of Ψ_{MD} , species and their interaction on Π_0
- Table S17: Summary of Covariance analysis testing influence of Ψ_{MD} , species and their interaction on Ψ_{S} .
- Table S18: Parameters correspond to Ψ_{PD} slopes analysis in all species at all sites.
- Table **S19**: Measured and normalized parameters correspond to Fig. 7.