

Hydraulic plasticity and water use regulation act to maintain the hydraulic safety margins of Mediterranean trees in rainfall exclusion experiments

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- 1 Hydraulic plasticity and water use regulation act to maintain the
- 2 hydraulic safety margins of Mediterranean trees in rainfall
- exclusion experiments

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- 6 Myriam Moreno^{1,2}, Jean-Marc Limousin³, Guillaume Simioni¹, Eric Badel⁴, Jesus Rodríguez-
- 7 Calcerrada⁶, Hervé Cochard⁴, José M. Torres-Ruiz⁴, Jean-Luc Dupuy¹, Julien Ruffault¹, Elena
- 8 Ormeno⁵, Sylvain Delzon⁷, Catherine Fernandez⁵, Jean-Marc Ourcival³, and Nicolas Martin-StPaul¹

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- 10 ¹INRAE, URFM, Avignon, France
- 11 ²French Environment and Energy Management Agency, Angers, France
- 12 ³CEFE, Univ Montpellier, CNRS, EPHE, IRD, Montpellier, France
- 13 ⁴ Université Clermont Auvergne, INRAE, PIAF, Clermont-Ferrand, France
- 14 ⁵ Aix Marseille Univ, Avignon Université, CNRS, IRD, IMBE, Marseille, France
- 15 ⁶Research Group Functioning of Forest Systems in a Changing Environment, Universidad Politécnica
- 16 de Madrid, Spain
- 17 ⁷INRAE, BioGeCo, Université de Bordeaux, Pessac, France.

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19 * Corresponding author: myriam.moreno@inrae.fr

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- 21 Key words: hydraulic failure, hydraulic adjustment, phenotypic plasticity, stomatal control, cuticular
- 22 conductance, turgor, throughfall exclusion, evergreen and deciduous trees.

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Abstract:

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Hydraulic failure due to xylem embolism has been identified as one of the main mechanisms involved 34 in drought-induced forest decline. The tree vulnerability to hydraulic failure depends on the hydraulic 35 safety margin (HSM), which can be computed as the difference between the highest water stress 36 experienced under drought or turgor loss point and the water stress causing hydraulic failure. While 37 it has been shown that HSM globally converge between tree species and biomes, there is still limited 38 knowledge regarding how HSM can adjust to locally varying drought conditions within species. In 39 this study, we relied on three long-term partial rainfall exclusion experiments in Southern France (15, 40 9 and 6 years of rainfall exclusion treatment respectively for Puéchabon, Font-blanche and O3HP 41 sites) to investigate the plasticity of hydraulic traits and HSM under intensified drought conditions 42 for three Mediterranean tree species (Quercus ilex L., Quercus pubescens Willd., and Pinus halepensis 43 Mill.). Our findings show, for all species, a similar HSM in trees submitted to intensified drought and 44 trees in the control treatments, despite reduced precipitation and, hence soil water availability. This 45 homeostasis of HSM in response to rainfall reduction is achieved through different mechanisms, 46 which appear to be related to the water use strategies of the three species. For Q. ilex, our results 47 indicate that the convergence in HSM is attributed to the adjustment of two plant hydraulic traits, 48 namely the turgor loss point (Ψ_{tlp}) and the water potential at which 50% of xylem conductivity is lost 49 due to embolism (P50). In contrast, both P. halepensis and Q. pubescens exhibited similar minimal 50 water potentials under control and drought treatments, due to isohydric behavior for the first and 51 probably other traits adjustment for the latter. Thus, the plasticity of the hydraulic vulnerability was not required to maintain their HSM in these two species. Our results suggest that trees exposed to 53 54 moderately drier conditions over several years may be able to maintain plastically their HSM. However, it remains to be seen whether this acclimatization of HSM to drier conditions will be 55 sufficient to withstand the more extreme droughts expected in the Mediterranean region. 56

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Introduction:

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Over the last decades, many studies have been conducted to better understand the mechanisms responsible for tree dieback following severe water stress (Barigah et al., 2013; Choat et al., 2018; Sperry et al., 2002; Tyree & Sperry, 1988). Among the different processes involved in plant mortality under drought (Allen et al., 2015; Anderegg et al., 2015), hydraulic failure has been identified as one of the key triggers of tree death (Adams et al., 2017; Anderegg et al., 2016; Arend et al., 2021;

McDowell et al., 2008). During drought, soil and atmospheric drying result in an increase in xylem water tension (i.e., a decrease in plant water potential). Excessive water tension promotes the occurrence of xylem cavitation, which cause the rupture of water columns between roots and leaves by embolism. Hydraulic failure occurs when the water transport in the plant is severely impaired by embolism (Tyree & Sperry, 1989), causing different damages to living tissues that may prevent the recovery of the tree when drought ends, and eventually kill the tree (Mantova et al., 2022).

71 The timing and likelihood of hydraulic failure during drought is influenced by various physiological characteristics that control plant desiccation resistance (Choat et al., 2018; Duursma et 72 al., 2019; Ruffault et al., 2022). The xylem vulnerability to cavitation is a crucial trait that define the 73 rate of spread of embolism with water potential decrease. It is derived from vulnerability curves 74 relating increased xylem embolism (e.g. the loss of hydraulic conductance) with decreasing xylem 75 water potentials (Cochard, 2002; Cochard et al., 2008, 2013). The main parameter of this vulnerability 76 curve is the water potential causing 50 % of hydraulic conductance loss (P50), which is negatively correlated with survival time during drought and the dryness of species habitat (Lens et al., 2016; 78 Martin-StPaul et al., 2017). The rate of water potential decline during drought (i.e. plant desiccation 79 dynamics resistance) depends, among other factors, on water loss regulation through stomatal control 80 (Martin-StPaul et al., 2017). Thus, the earlier the stomata close after the drought onset, the more the 81 plants are able to limit the decrease in their water potential, and preserve xylem conduits from 82 cavitation. The water potential at turgor loss point (Ψ_{tlp}) can be used as a surrogate of the water 83 potential at which stomata close (Brodribb & Holbrook, 2003; Martin-StPaul et al., 2017), given that 84 85 stomata close once guard cells lose turgidity. In addition, plant residual transpiration, occurring through incompletely closed stomata or the cuticle (Machado et al., 2021), also determine the rate of 86 the plant-drying process and the hydraulic risk (Duursma et al., 2019). 87

The hydraulic safety margin (HSM) is an integrative trait used to evaluate the risk of hydraulic 88 failure under drought, with a higher HSM indicating a lower risk. Two versions of HSM have been 89 proposed. The first one corresponds to the difference between the minimum xylem water potential 90 (e.g. $\Psi_{\text{midd min}}$) reached by a plant and the threshold water potential causing severe xylem embolism 91 92 (e.g. P50) (Choat et al., 2012). According to this definition, HSM represents the hydraulic risk associated to the water stress actually experienced by the plant. In the second definition, $\Psi_{\text{midd min}}$ is 93 replaced by the water potential causing stomatal closure (Martin-StPaul et al., 2017). This second 94 version of HSM integrates the degree to which stomatal control can prevent hydraulic risk. Since the 95 minimum potential achieved by a plant depends on stomatal control, the two definitions of HSM are 96 closely linked. The HSM has been shown to be conserved across forest tree species worldwide (Choat et al., 2012), as a result of the coordination of multiple traits optimizing the xylem safety-efficiency trade-off with species habitat (Franklin et al., 2022; Guillemot et al., 2022; Martin-StPaul et al., 2017; Pivovaroff et al., 2018). However, there has been little investigation into the intraspecific variations of the HSM in response to drier conditions. To our knowledge, the only publication focusing on this topic report narrower HSM for Mediterranean species occurring at the dry edge of their geographical distribution (Alon et al. 2023).

104 Understanding the impact of plant trait plasticity on plant desiccation dynamics is therefore crucial for predicting hydraulic risk in the current context of rapid climate change (Jinagool et al., 105 106 2018). Yet, limited research has thus far investigated the adjustment of multiple traits after long periods of increased drought. Some studies suggested that leaf traits, such as Ψ_{tlp} change more in 107 108 response to sustained drought than the xylem vulnerability to cavitation (Bartlett et al., 2012; Torres-Ruiz et al., 2019), which is sometimes invariant (Lobo et al., 2018; Rosas et al., 2019) and sometimes 109 plastic (Bert et al., 2021; Herbette et al., 2021; Lemaire et al., 2021). For the Mediterranean species 110 Quercus ilex, past studies have reported limited plasticity of xylem vulnerability to cavitation 111 (Limousin et al., 2022; Limousin et al., 2010; Martin-StPaul et al., 2013). Additionally, it is important 112 to note that, until recently, the methods used to measure xylem vulnerability to cavitation in long-113 vessel branches, such as those of *Q. ilex*, were affected by a bias due to open vessel methodological 114 artefact (Martin-StPaul et al., 2014; Torres-Ruiz et al., 2014). Similarly, the methods used to estimate 115 leaf hydraulic vulnerability curves based on leaf hydraulic conductance were also biased by the 116 integration of an extra-xylary water pathway (Li et al., 2020; Limousin et al., 2022; Trifiló et al., 117 2016). Hence, it is important to revisit the impact of plant trait plasticity on plant desiccation dynamics 118 119 with up-to-date and unbiased methods, such as Cavitron (Cochard et al. 2002) or optical vulnerability (Brodribb et al. 2016) techniques. 120

In this study, we aimed to assess whether, and by which mechanisms, the HSM of a given species can 121 change with increasing drought conditions. We focused on three of the most widespread tree species 122 in the Mediterranean basin, namely Pinus halepensis Mill., Quercus ilex L. and Quercus pubescens 123 Willd. Those species are being monitored in three long term partial rainfall exclusion experiments set 124 125 up in natural forests in southern France: the Puéchabon experimental site (a Q. ilex forest), the Oak Observatory at the Haute-Provence Observatory (a Q. pubescens forest), and the Font-blanche site (a 126 127 mixed forest of *P. halepensis* and *Q. ilex*). In those three sites, we measured and compared hydraulic 128 traits in adults of the three species under control and aggravated drought conditions.

Materials and Methods:

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Sites description

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134 The study was conducted on three experimental forest sites located in the French Mediterranean region (Figure 1A) that stand out by their vegetation types: the Font-blanche (FB hereafter) forest is 135 composed of P. halepensis in the overstory and Q. ilex in the understory, the oak observatory at the 136 Haute-Provence Observatory (O₃HP hereafter) is dominated by *O. pubescens*, and the Puéchabon site 137 138 (P hereafter) by O. ilex. The climate of the sites is Mediterranean, with wet, mild winters and dry, hot summers (Fig. S1). The subsoil of all sites is composed by a hard limestone bedrock. The soils of the 139 FB and P sites are similar and classified as silty clay loam, while the O3HP site is a clay-loam soil. 140 They have a high proportion of rocks and soil depth varies between 35 to 50 in FB and P and 30-40 141 cm in O₃HP. On all sites, a rainfall exclusion experiment has been implemented for for several years 142 (15, 9 and 6 years respectively for Puéchabon, Font-blanche and O₃HP sites at the time of this study 143 (more details provided in Table 1) excluding approximately 30 % of the precipitation reaching the 144 ground. At FB and P, the system excluding precipitation is passive, with PVC gutters hung under the 145 tree canopies. At O₃HP, rainfall exclusion is achieved with a mobile rainout-shelter deployed 146 manually only during some rainfall events from spring to autumn (Fig. 1). Table 1 provide more 147 details on sites characteristics and rainfall exclusion treatments. On each site, meteorological 148 variables are monitored (including half-hourly precipitation, radiation, air temperature and humidity).

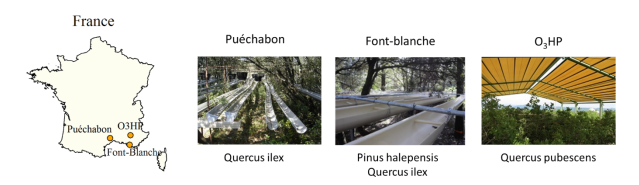


Figure 1. Locations of studied sites and pictures of the rainfall exclusion systems. PVC gutters placed under the canopy were used to collect around 30 % of precipitation at Puéchabon and Font-blanche sites, while in the O₃HP site, a mobile shelter placed above the canopy is deployed manually.

154 Table 1: Characteristics of studied sites.

	Font-Blanche	Puéchabon	ОзНР
Location	43°14'27"N, 5°40'45"E	43°44'29"N, 3°35'45"E	43°56′115″ N, 05°42′642″ E

Precipitation (mm) / temperature (°C) (annual mean value between 2016- 2019)	615 /14.2	1033 /14.0	871 /12.9	
Soil characteristics: type, depth (cm), available soil water (mm)	Silty clay loam, 20 to 50, 160 Silty clay loam, 30 70,140		Clay-loam, 30 to 40, NA	
Vegetation type	Evergreen P. halepensis and	Evergreen	Deciduous	
vegetation type	Q. ilex forest	Q. ilex forest	Q. pubescens forest	
Basal area (m²/ha) (mean 2018)	29.7	27.2	21	
Tree height (mean 2018)	13.5 for <i>P. halepensis</i> 6.5 for <i>Q. ilex</i>	5.5	5	
Rainfall exclusion device,				
treatments area (m²), year	Gutters,	Gutters,	mobile rainout-shelter,	
of the start of the	625,	140*4 replicates,	300,	
exclusion	2009	2003	2012	
Precipitation exclusion ratio	≈ 30 %	≈ 30 %	≈ 30 %	
References	(Moreno et al., 2021)	(Limousin et al., 2009)	(Genard-Zielinski et al., 2018)	

Water potentials in the field

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Water potentials data were taken from Moreno et al. (2021) for P. halepensis and Q. ilex at FB, from 157 158 Limousin et al. (2022) for Q. ilex at P, and from Genard-Zielinski et al. (2018) for Q. pubescens at O_3HP . Predawn (Ψ_{pd}) and midday (Ψ_{midd}) water potentials were measured from 2015 to 2019 in each 159 160 site (Table 2) during summer using Scholander pressure chambers (different brands depending on year and site). The samplings were performed at several dates from the onset to the end of summer 161 (between two to six measurements per year depending on year and site). At each site and date, 162 measurements were made on twigs (FB and P sites) or leaves (O₃HP site) of at least three trees per 163 164 species and per treatment (control/rainfall exclusion). One to three samples per tree were measured depending on site and date during the two hours preceding sunrise for Ψ_{pd} , and between 2 to 4 pm for 165 Ψ_{midd} . Ψ_{midd} were measured on transpiring twigs/leaves. For Ψ_{pd} , we have selected the data from the date when the lowest values were recorded per species for each year to compute the extreme annual predawn water potentials ($\Psi_{pd \, min}$). For Ψ_{midd} , we have selected the data from the date when minimum values were observed, combining all years, to compute the extreme midday water potentials (Ψ_{midd} min).

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Vulnerability to cavitation

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174 Depending on the length of xylem conduits of each species, two techniques were used to measure the xylem vulnerability to cavitation. For *P. halepensis*, a gymnosperm species with only 175 176 tracheids, we used the cavitron technique (Cochard, 2002). In March 2018 and October 2019, south facing branches, approximatively 50 cm long, were sampled from the top of the canopy of a total of 177 178 17 control trees and 10 trees submitted to rainfall exclusion of the FB site (Table S1: no significant difference of P50 between sampling date, Fig. S2). Once cut, branches were placed under water and 179 180 the basal 10 cm recut to eliminate cavitated tracheids due to conduits left open during the first cut. Then, they were placed in humidified black bags in order to limit transpiration and prevent 181 182 dehydration. At the laboratory, branches were placed under water, the foliage was removed, and a straight 30-cm-long section of the branch was recut. The sections were rolled into humidified 183 184 absorbent paper, wrapped in plastic film, and placed in an icehouse for shipping them to the to the INRAE-PIAF laboratory at Clermont-Ferrand (central France). There, the bark was removed and the 185 vulnerability to cavitation of the samples measured with the cavitron (Cochard, 2002). Tension in the 186 xylem was gradually increased, with the rotation speed and the percent loss of conductivity (PLC) 187 being calculated using Cavisoft (v.4.0; University of Bordeaux) as: 188

189 PLC (%) =
$$100*(1-K/K_{max})$$
 (1)

- where K is the conductivity recorded at a given rotation speed and K_{max} the maximal conductivity recorded at the lowest rotation speed.
- 192 Xylem vulnerability curves were then constructed for each sample by plotting PLC as a 193 function of the xylem tension caused by rotation. The following sigmoid function was fitted to data:

194 PLC =
$$100/(1+\exp(s(P-P50)))$$
 (2)

- where s is the slope at the inflexion point and P50 is the water potential causing 50 % loss of hydraulic conductivity.
- Because xylem vessel length in the two oaks species can reach up to 1m (Martínez-Vilalta et al., 2002), we were unable to employ the cavitron technique with oaks, due to the bias for long-vessel

199 species when branch samples are shorter than the vessel length (Beikircher et al., 2010; Cochard et al., 2013; Martin-StPaul et al., 2014; Sergent et al., 2020). Instead, we used the optical technique 200 201 (Brodribb et al., 2016) to measure the xylem vulnerability to cavitation in the leaves. We opted for the leaf optical technique rather than the stem optical technique, as it does not require to remove tissue 202 203 limiting potential bias caused by endommaged xylem if not well executed. All the branches used to measure VC were not flushed to remove native embolism, as such procedure can bias vulnerability 204 curves (https://doi.org/10.1093/treephys/tpad088) due to open-vessels artifact. Drawing from the 205 findings of Li et al., (2020), and our own observation of similar P50 using the optical method and 206 respectively X-ray microtomography for Q. ilex stems (Sergent et al, 2020) and cavitron for Q. 207 pubescens branches (personal data given by Delzon) (Fig. S3), we assumed that hydraulic 208 segmentation did not occur in the two oak species. Anyway, despite the use of different hydraulic 209 techniques across species, it is important to note that the aim of this study is to evaluate intraspecific 210 plasticity in hydraulic traits, rather than interspecific drought tolerance. 211

212 During summer 2019, 4 to 5 branches more than 1 m long of Q. ilex and Q. pubescens sampled from different trees, were harvested in each control and rainfall exclusion treatments of the three 213 studied sites. Branches were placed in humidified bags and stored in a dark cold room with the cut 214 end in water for at least one night to rehydrate. Once fully rehydrated, branches were allowed to 215 dehydrate progressively on the lab bench (bench dehydratation technique). One or two mature leaves 216 per branch, still attached to the branch, were placed on a flatbed light transmission scanner (Epson 217 V850). The leaves were kept flat using microscope slides attached to the scanner glass with 218 transparent tape. Leaf scans of a resolution of 2400 dpi were recorded in light transmission mode 219 220 every 5 min while the branch was progressively dehydrating. Concomitantly, xylem water potential was measured on bagged leaves of the same branch using a Scholander pressure chamber (PMS 221 222 instrument), approximatively every 3 hours throughout branch desiccation (along 3 to 5 days, a minimum period of 3 days being enough to reach 100% emboli in the xylem, Fig. S4). 223

224 Once the branch fully desiccated (and the xylem water potential was no longer measurable with Scholender pressure chamber), all images were analyzed to detect cavitation events as changes 225 226 in light transmittance between successive scans (http://www.opensourceov.org/). Image J (FIJI, (Schindelin et al., 2012)) was used to reveal embolism events by overlapping successive leaf scans 227 228 and calculating the pixel area of embolized vessels. Vulnerability curves were constructed for each leaf by plotting the relative cumulative embolized area (corresponding to the cumulative embolized 229 area divided by the total embolized area at the end of leaf dehydration) as a function of leaf water 230 potential, and fitting the same sigmoid function as for *P. halepensis*. 231

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233 Foliar traits

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235 Pressure-volume curves (hereafter PV curves) were used to characterize different foliar traits related to the maintenance of leaf turgor and hydration (Bartlett et al., 2012; Tyree & Hammel, 1972). 236 237 For each species, two well-lit twigs per tree were collected from the outer crown of 4-6 trees in the control and exclusion treatments. The sampling took place in March 2018 for *P. halepensis* and *Q.* 238 ilex at FB and P sites and in September 2019 for Q. pubescens at O₃HP site. In all cases, once 239 collected, twigs were bagged and placed in a cooler at 4°C until reaching the laboratory. Once in the 240 241 lab, each twig end was re-cut under water with a razor blade, then put overnight in a cool chamber in distilled water to rehydrate. Half of the samples were used to estimate the leaf mass area and the other 242 243 half to perform the PV curves

PV curves were established using the bench drying method proposed by Hinckley et al., 244 (1980). Briefly, the weight and the water potential of the twigs were measured all along their 245 dehydration using a precision balance (FS-220, resolution 0.1 mg) and a Scholander pressure chamber 246 247 (PMS instrument), respectively. To overcome oversaturation of rehydrated twigs, the first measurement of water potential and weight was removed, and full turgor weight was extrapolated 248 249 from the regression between twig weight and water potential before the turgor loss point. The osmotic potential at full turgor (π_{100}) and at the turgor loss point (Ψ_{tlp}) , the relative water content at turgor loss 250 point (RWC_{tlp}), the symplastic water fraction at full turgor (Fs), and the bulk modulus of tissue 251 252 elasticity (ε) were obtained by plotting the inverse of water potential (-1/Ψ, in MPa) against twig water saturation deficit (1-Relative Water Content, in %). PV curves data used for Q. ilex are the same 253 as those reported in Limousin et al. (2022). 254

Minimum leaf conductance to water vapor (g_{min}) was obtained considering the average twig water loss (corresponding to sample transpiration) after stomatal closure. To this end, relative humidity and temperature were recorded every second with a probe (PT100) to compute VPD, and minimum leaf conductance was obtained as follows for each species:

$$g_{min} = \frac{E}{VPD \times DA} \times Patm, (3)$$

with E being the transpiration after turgor loss, Patm the atmospheric pressure (set at 101 kPa), and DA the developed leaf area. The developed leaf area (DA) was obtained from the projected leaf area by using the ratio between leaf dry mass and leaf mass per area. For both oaks, with flat leaf surface, DA was calculated as twice the projected leaf area. For *P. halepensis*, we calculated DA considering

that needles formed a perfect half-cylinder, which was confirmed by measurements of width and thickness at the top, middle and bottom of 15 needles (data not shown). We choose to express gmin in function of developed leaf area to report more realistic value. Indeed, gmin does not result only from water loss through not fully close stomata but also by cuticular leaks, that can happen from all the leaf surface.

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270 Hydraulic safety margins

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- 272 Two types of Hydraulic safety margins (HSM) were computed at each site and for both treatments by
- 273 subtracting either (i) the minimal midday water potentials ($\Psi_{\text{midd min}}$) to P50, following the approach
- 274 outlined by Choat et al. (2012), or (ii) the tugor loss point, as a surrogate as water potential causing
- 275 stomatal closure to P50 as in Martin-StPaul et al. (2017).
- 276 To avoid damaging trees, the samples collected to construct hydraulic vulnerability curves were taken
- 277 from different trees than those sampled for leaf water potential and turgor loss point measurements.
- 278 Therefore, it was not possible to calculate the HSM at the tree level. Instead, we determined both
- 279 HSM versions at the species and treatment levels using a nonparametric bootstrap approach. For each
- 280 species and treatment (control and rainfall exclusion), we randomly sampled, with replacements, 5
- values of P50 and $\Psi_{\text{midd min}}/\Psi_{\text{tlp}}$ and determined 5 HSM as the difference between these values. This
- 282 process was repeated 1,500 times to obtain a distribution of HSM values for each site and treatment
- 283 that was subsequently used for statistical comparisons.

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Native stem xylem embolism measured by X-ray microtomography

- Native stem xylem embolism was estimated for *Quercus ilex* and *Pinus halepensis* trees in both rainfall exclusion and control treatments at FB and P. The sampling took place in March 2018. Four to eight well-lit branches of more than 1m were collected depending on the sites, the treatments and the species considered. Once collected, branches were immediately recut under water to avoid
- 291 cutting artefacts.
- The cut surface of the branches was kept under water until reaching the laboratory. Then, short segments of branches (diameter < 0.7 cm and length < 4 cm) were cut under water, plunged in liquid paraffin, and stored in a cold chamber at 4°C until analysis. Measurements were performed following the protocol described in Cochard et al. (2015). Samples were inserted in an X-ray microtomograph (Nanotom 180 XS; GE, Wunstorf, Germany) and analyzed using a field of view of 5 × 5 × 5 mm³, X-

ray voltage of 60 kV, current of 240 μA, and a scan time of 21 min. The final spatial resolution of the 3D images was 2.5 μm after 3D-reconstruction. For each sample, one transversal 2D slice was extracted from the middle of the branch using VGStudio Max© software (Volume Graphics, Heidelberg, Germany). The surface area of embolized conduits was estimated from slices using the software ImageJ (Schneider et al., 2012).

As *Q. ilex* is a semi ring-porous species with log normal distribution of xylem conduit sizes, we measured the surface area of large embolized vessels (contributing the most to hydraulic conductivity) to estimate their mean diameter and their corresponding hydraulic conductivity. The maximal hydraulic conductivity was estimated thanks to a second scan we performed on the same branch sample once it was fully embolized. The level of embolism was calculated as the ratio between estimated native hydraulic conductivity and maximum hydraulic conductivity.

Table 2. Summary of measurement periods according to sites and species. FB and P are the acronyms for respectively Font-Blanche and Puéchabon sites.

Measurements	P. halepensis	Q. pubescens	Q. ilex FB	Q. ilex P
Trees water potentials $\Psi_{pd min}(MPa)$	From 2013 to 2018	2014, 2015, 2018	From 2013 to 2018	From 2015 to 2018 2017
$\Psi_{midd\;min}\left(MPa\right)$	2018	2015	2017	2017
Pressure volume curves: Ψ_{tlp} (MPa), $\pi 100$ (MPa)	March 2018	September 2019	March 2018	March 2018
g_{min} (mmol s ⁻¹ m ⁻²)	March 2018	September 2019	March 2018	March 2018
Xylem vulnerability to cavitation: P50 (MPa)	March 2018, October 2018	September 2019	Onset of summer 2019	Onset of summer 2019
Native stem xylem embolism (%)	March 2018	/	March 2018	March 2018

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312 Statistics 313

To assess whether plant and leaf traits were different between treatments we used t-test for all traits 314 measured. For the HSM, we made a Student t-test for each of the 1500 subsamples obtained via 315 bootstrapping. We considered that rainfall exclusion and control treatments were significantly 316 different if the probability of having a P value > 0.05 among the 1500 t-tests realized was lower than 317 0.05. Statistical analyses were performed with the R software (3,5,2, R Development Core Team 318 319 2018). Differences in parameters derived from xylem vulnerability curve (slope and P50; Table S2-3, Fig S5), $\pi 100$, Ψ_{tlp} or g_{min} between rainfall exclusion and control treatments were found to be 320 similar for O. ilex in both Font-blanche and Puéchabon sites (Limousin et al., 2022). Hence, to 321

322 increase the statistical power in testing for a rainfall exclusion effect on traits values, as in Limousin

323 et al. (2022) we decided to combine the data from both sites for this particular species. All Statistical

324 analyses were performed with R software (3,5,2, R Development Core Team 2018)

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326 Results

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328 Rainfall exclusion impacts on tree water status

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- 330 Variations in the response of $\Psi_{pd \, min}$ to rainfall exclusion treatments were observed among different
- 331 tree species and sites (Fig. 2). For *Q. pubescens* at O₃HP, Ψ_{pd min} tended to be more negative for trees
- 332 growing in the rainfall exclusion treatments (P value = 0.005) until 2015 (two years after the onset
- 333 of the rainfall exclusion). After that, $\Psi_{pd \, min}$ was similar to that of control trees (P_value > 0.1). For
- 334 *P. halepensis* at FB, $\Psi_{pd \, min}$ were similar across years, except in 2015 (P value = 0.05), when they
- 335 were slightly more negative in the rainfall exclusion treatments (Fig. 2). It should be noted, however,
- 336 that if we consider the whole water potential dynamic rather than the extreme values, significantly
- 337 more negative potentials in rainfall exclusions appear between treatments at lower levels of water
- 338 stress (Genard-Zielinski et al., 2018; Moreno et al., 2021). Moreover, soil water content taken at
- 339 shallow ground level (< 50 cm), tended to be more negative in the rainfall exclusion treatment than
- 340 in the control (Genard-Zielinski et al., 2018; Moreno et al., 2021). All of these findings attest to the
- 341 effectiveness of the exclusion treatments for these two species.
- 342 Concerning Q. ilex at both FB and P sites, $\Psi_{pd min}$ tended to be more negative in the rainfall exclusion
- 343 treatments (Fig. 2, P_value < 0.05 in FB site until 2015; P_value < 0.006 in P site until 2016, then
- 344 P value < 0.05).

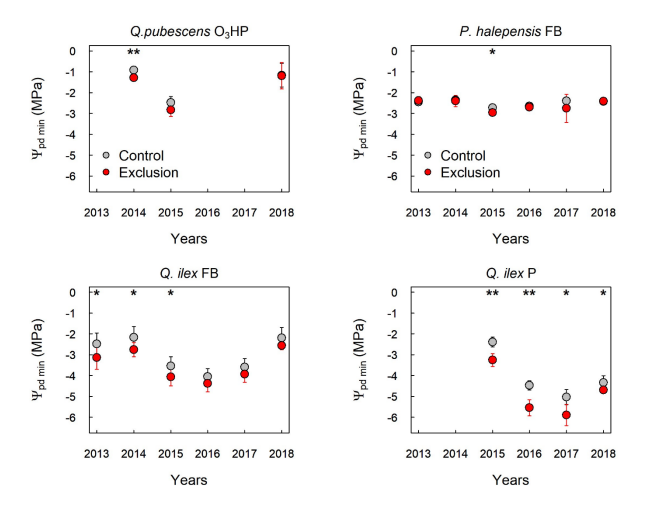


Figure 2. Average minimum predawn water potentials ($\Psi_{pd \, min}$) measured for each species in both control and rainfall exclusion treatments according to year and site. Bars represent standard deviations associated to the mean. Significant differences between treatments (t-test) are indicated by asterisks (*, $0.01 \le P_value < 0.05$; **, $0.001 \le P_value < 0.01$; ***, $P_value < 0.001$).

Effects of rainfall exclusion on hydraulic and leaf traits

No significant differences in P50, π_{100} , Ψ_{tlp} nor g_{min} , were found between the rainfall exclusion and control treatments for Q. pubescens and P. halepensis (Fig. 3, P_value > 0.2 for each of these traits). For Q. ilex, P50, π_{100} and Ψ_{tlp} were significantly more negative for trees growing in the rainfall exclusion (P_value = 0.04 for P50; P_value = 0.02 for π_{100} ; P_value = 0.03 for Ψ_{tlp}), whereas g_{min} remained unchanged (P_value > 0.05).

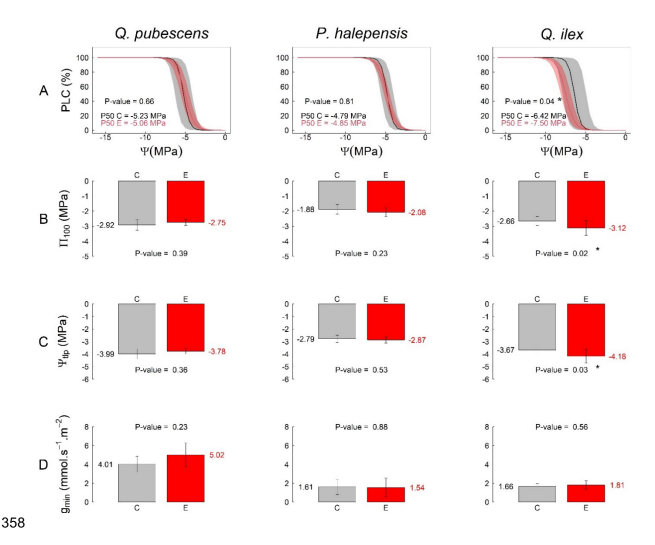


Figure 3. Effects of rainfall exclusion on hydraulic traits for the three studied species. A. Xylem vulnerability curves to cavitation, represented by the percent loss of conductivity (PLC) as a function of water potential (Ψ). Averages and standard deviations are respectively in bold lines and grey (control) or red (rainfall exclusion) areas. B. Osmotic water potential at full turgor (π_{100}), C. water potential at turgor loss point (Ψ_{tlp}), D. minimum leaf conductance (g_{min}) in the control (C, in grey) and rainfall exclusion (E, in red) treatments, for the three studied species. In the barplots, bars represent mean values; error bars represent standard deviation. P_values of treatment effect are presented; asterisks indicate a significant difference at P_value < 0.05.

Effect of rainfall exclusion on species hydraulic safety margins and xylem embolism.

For all species, we did not observe any statistical difference in HSM between rainfall exclusion and control trees, based on either $\Psi_{\text{midd min}}$ (Fig. 4A, P_value > 0.8 for each species) or Ψ_{tlp} (Fig. 4B, P_value > 0.9 for each species). Levels of native stem xylem embolism were also similar in both treatments for *P. halepensis* or *Q. ilex*, the only species analyzed (Fig. 5).



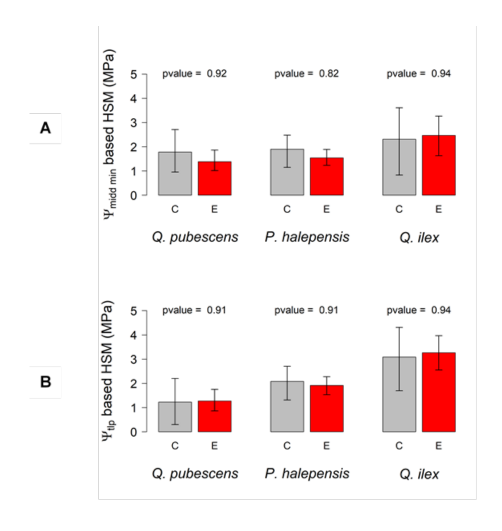


Figure 4. A. Hydraulic safety margins (HSM) in relation to species and treatment (C: control; E: rainfall exclusion). A. HSM defined as the difference between minimum midday water potentials ($\Psi_{\text{midd min}}$) and water potential causing 50% loss of hydraulic conductivity (P50). $\Psi_{\text{midd min}}$ correspond to the minimal values measured in 2015 for *Q. pubescens*, 2016 for *Q. ilex* in P and FB sites and 2018 for *P. halepensis*. B. HSM defined as the difference between turgor loss point (Ψ_{tlp}) and P50. Bars and errors bars represent respectively the median and the first/third quartiles (confidence interval) of the nonparametric bootstrap analysis.

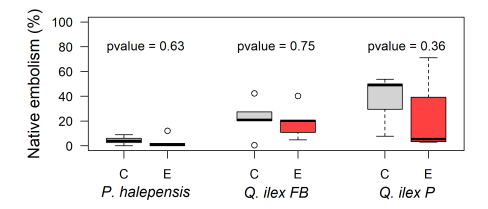


Figure 5. Native stem xylem embolism (%) estimated from X-ray microtomography on 4 years-old branches of *P. halepensis* and *Q. ilex* trees from the control (C) and rainfall exclusion (E) treatments at both Font-blanche (FB) and Puéchabon (P) sites. The measurements were performed in March 2018. P_value of Wilcoxon rank sum tests comparing native stem xylem embolism between treatments are represented for both sites.

Discussion

In the current context of rapid global change, tree capacity to acclimate to increasing drought appears crucial for the maintenance of forest structure, composition and function. Under drier conditions, hydraulic failure risk should increase for trees if they cannot adjust key hydraulic traits involved in the hydraulic safety. In the present study, we aimed to prospect the ability of three functionally different species (different drought response strategies, clades, phenology, etc.) to modify these traits under increasing drought conditions, and to assess the hydraulic risk ensued. Among the three studied species, only *Q. ilex* exhibited significant plasticity in the hydraulic traits prospected, but all three species maintained the hydraulic safety margins (HSM). In the following, we first discuss the effects of the hydraulic adjustments of *Q. ilex* for its drought tolerance capacity. Then, we argue about the likely causes leading to a lack of hydraulic adjustment under rainfall exclusion treatment for both *P. halepensis* and *Q. pubescens*. Finally, we discuss the consequences of HSM homeostasis in response to drier conditions across species for their hydraulic risk.

Homeostasis of HSM due to plasticity in *Q. ilex*

Q. ilex reached lower values of both $\Psi_{pd min}$ and $\Psi_{midd min}$ in the rainfall exclusion treatments 402 of both FB and P sites (Fig. 2, S6). Thus, the higher water stress experienced by this species in 403 response to increased drought could be the trigger for P50 and Ψ_{tlp} adjustments towards more negative 404 values (Fig. 3, S7). 405 Until now, studies focusing on long-term drought acclimation of Q. ilex had concluded that 406 no plasticity of hydraulic vulnerability occurred for adult trees undergoing rainfall exclusion, in either 407 branches (Limousin et al., 2010; Martin-StPaul et al., 2013) or in leaves (Limousin et al. 2022). Our results contradict those previous works. This could be explained by the facts that: 1) previous 408 409 estimates of P50 in the xylem of branches were possibly biased by the open-vessel artifact (Martin-StPaul et al., 2014; Torres-Ruiz et al., 2014); 2) estimations of xylem vulnerability to cavitation in 410 branches include several wood rings, which may hide xylem plasticity occurring after a specific 411 aggravated drought; and 3) the rehydration kinetics method used to calculate leaf hydraulic 412 conductance may not reflect only the xylem vulnerability to cavitation but also an extra-xylary 413 component in the leaf mesophyll (Li et al., 2020; Limousin et al., 2022; Trifiló et al., 2016). The 414 optical technique (Brodribb et al., 2016) used in this study for O. ilex and O. pubescens appears as a 415 good solution to overcome these limitations. It allows the estimation of hydraulic vulnerability in the 416 xylem only and in leaves, that are shorter lived, and therefore less affected than branches, by the 417 potential biases due to regrowth and cavitation fatigue, or by methodological issues related to the 418 419 flushing of native stem xylem embolism and vessels open at both ends (Gauthey et al., 2020).

420 In addition to the adjustment of xylem vulnerability to cavitation, our findings also highlighted that turgor-related traits also changed for this species under aggravated water stress conditions. 421 Indeed, for Q. ilex trees growing in the rainfall exclusion treatments, lower π_{100} values were reported 422 (Limousin et al., 2022), that translate into a significant reduction of Ψ_{tlp} (Bartlett et al., 2012). As 423 stomatal closure is realised through the loss of stomata guard cells turgidity, Ψ_{tlp} could be used as a 424 425 surrogate of the point of stomatal closure (Martin-StPaul et al., 2017). This suggests that Q. ilex trees 426 from the rainfall exclusion treatments could close their stomata at lower water potentials than trees from the control treatments thus allowing them to maintain more photosynthetic activity or root 427 428 growth under similar drought conditions. This adjustment may lead to an increased risk of xylem embolism if the vulnerability to cavitation, P50, is not also reduced. The apparent coordinated 429 430 plasticities of Ψ_{tlp} , that is lowered by 0.5 MPa, and of P50 that is lowered by 1 MPa, could be beneficial 431 to leaf gas exchange, without affecting the hydraulic safety margin (Fig. 4B). This maintenance of 432 the HSM in spite of drier conditions seems confirmed by the similar levels of native stem xylem embolism observed in the Q. ilex trees growing inside the rainfall exclusion treatments and those 433 434 growing under control conditions (Fig. 5). Limousin et al., (2022) showed that Fs increased significantly in the rainfall exclusion treatments, suggesting that lower Ψ_{tlp} do not result from internal 435 cell water loss but rather from the accumulation of osmolytes. Hence, we can assume that, besides its 436 action on stomatal control, osmotic adjustment in the exclusion treatments could also play a role in 437

limiting cell dehydration and hence drought-induced damages on leaves, which is crucial for this evergreen species with a leaf lifespan of two to three years (La Mantia et al., 2003).

Regarding the minimal leaf conductance (g_{min}), which is implied in water loss after stomatal closure (Duursma et al., 2019), we observed no plasticity in response to long-term increased drought for Q. ilex or any of the other species (Fig. 3). This suggests either that g_{min} is not or little plastic, or that more precise methods, such as the drought box (Billon et al., 2020), should be used to detect plasticity in this trait.

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7 Homeostasis of HSM despite lack of plasticity in P. halepensis and Q. pubescens

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The most likely explanation for the lack of adjustments in hydraulic traits for P. halepensis and Q. 449 pubescens is that these species experienced similar levels of extreme water stress under both control 450 and rainfall exclusion treatments, and that that was no advantage in increasing HSM. This hypothesis 451 appears plausible for P. halepensis, for which $\Psi_{pd min}$ were similar between rainfall exclusion and 452 control trees (Fig. 2). Such result is consistent with its behavior in response to drought. Indeed, P. 453 454 halepensis is an isohydric species that strongly regulates its transpiration at the onset of drought, allowing it to limit soil water consumption and maintain both Ψ_{pd} and Ψ_{midd} (Figs. 2, 3) over a safe 455 threshold of -3 MPa (Moreno et al., 2021). Recently, (Moreno et al., 2024) proposed that such 456 isohydric behaviour is the results of two meachnisms: (i) its stomatal closure that happened at 457 relatively high level of water stress and (ii) its ability to isolates itself from the ground and the 458 atmosphere. Hence, its water use strategy seems sufficient in itself to limit hydraulic damages caused 459 460 by rainfall reduction, without requiring xylem adjustments, at least under our experimental conditions. The fact that native stem xylem embolism were similar between rainfall exclusion and 461 462 control treatments (Figs. 5, 6) further supports this idea.

For *Q. pubescens*, given its anisohydric behaviour like for *Q. ilex* (Damesin & Rambal, 1995; Poyatos et al., 2008), we expect trees from the rainfall exclusion to be exposed to more negative water potentials. In 2014, two years after the onset of the rainfall exclusion experiment, a significant difference between Ψ_{pd} of control and rainfall exclusion trees was observed. This result attests the effectiveness of the rainfall exclusion, which is in line with the significant reduction in soil moisture measured in the rainfall exclusion treatments (Genard-Zielinski et al., 2015, 2018). Nevertheless, for the following studied years, $\Psi_{pd \, min}$ was similar between treatments. This result, which seems at first

470 sight contradictory, probably underpins other adjustments in unprospected traits. The figure S8 shows 471 that at the onset of the experiment, the amount of leaf loss by the trees of the rainfall exclusion is 1.5 higher than the one of the control treatment, suggesting highest leaf area index in the rainfall exclusion 472 Throughout the experiment, the gap between the litterfalls of the two treatments narrows to 473 become similar in 2018. This result suggests that trees in the exclusion plot have reduced leaf 474 production and therefore their overall leaf area, limiting overall transpiration and hence water 475 476 potential decrease. Other adjustment could also occured as, for example, an increase of rooting depth could be induced after aggravated water stress, as proposed by Martin-StPaul et al. (2013), enabling 477 trees to access deeper soil water and thus limiting their drought exposure. Likewise, earlier leaf senescence could also occur in trees facing drier conditions (Wu et al., 2022). These additional 479 adjustments occurring at the whole tree scale could be sufficient to dampen the rainfall exclusion 480 effect on extreme tree water stress and prevent hydraulic damages (Fig. 5). Further investigations are 481 482 however needed to identify the other traits implied in the homeostasis of hydraulic risk for this species. 483

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485 The risk of maintaining the HSM for species vulnerability under future climatic 486 conditions.

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Although achieved through different means, our data report a maintenance of HSM under 488 experimental 30 % rainfall exclusion (Fig. 4) for the three studied species. This suggest that no matter 489 490 the drought response strategies adopting by these species, trees exposed to drier conditions adjusted functionally to preserve this trait. One might have expected that, in order to reduce hydraulic risk, 491 trees could have adjusted to increase HSM values, through the development of a more cavitation 493 resistant xylem. A reasonable explanation that could be advanced to explain this lack is the need for the plant to optimize fluxes relative to the hydraulic risk and that can be achieve through a myriad of 494 495 strategies. The maintenance of HSM in response to plasticity that we reported in this study is in line with metanalysis from Choat et al. (2012), which reports a worldwide convergence of forest 496 497 vulnerability to drought (estimated through tree HSM) at the interspecific level independently of the level of drought. The fact that trees seem to maintain HSM despite several years of increasing drought 498 499 calls into question the natural capacity of forests to withstand accelerating climate change. To what extent HSM can be maintained in the future and at what cost, is still an open question. It remains to 500 501 be known how much future extreme drought or heat waves may affect acclimated and non-acclimated trees. The next step is to use traits based mechanistic process-based models (Martin-StPaul et al., 502

2017; Blackman et al., 2019, Ruffault et al. 2023), in order to characterize hydraulic risk in various conditions. In particular, they would enable to assess how the combination of several traits could

translate into different levels of risk in a warmer, drier future.

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