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1 Intra-specific variability in deep water extraction between trees growing on a Mediterranean karst

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14

15 **Abstract**

16 Plant transpiration is a major component of water fluxes in the critical zone, which needs to be
17 better characterized to improve our ability to understand and model the hydrological cycle. In water-
18 limited ecosystems such as those encountered on karst environments, climate-induced changes in
19 transpiration are expected to be strongly influenced by the ability of the vegetation cover to resist or
20 adapt to drought. However, because of the high heterogeneity of karst environments, the amount of
21 water available for trees can change within a stand, which may lead to significant differences in
22 drought vulnerability resistance between trees of the same species. So far it is not known if soil and
23 subsoil environment influence the magnitude of deep water extraction, at the intra-specific scale.
24 Here, we investigate the variability in deep water extraction for six individual *Quercus ilex* trees
25 growing on a karst substrate in a Mediterranean forest. We combined three approaches: (i) electrical
26 resistivity tomography to determine the variability of soil/subsoil characteristics, (ii) isotope tracing
27 to determine the origin of water transpired by plants, and (iii) predawn and midday leaf water
28 potential (Ψ) to assess the trees' water stress and transpiration regulation. Along the summer
29 season, deep water extraction increased with drought intensity. Deep water use varies between
30 individuals and according to drought intensity. At moderate water stress levels, we found no

31 significant relationship between the origin of xylem water and soil/subsoil characteristics or
32 individual stress level. However, at the peak of the drought (average predawn $\Psi < -2$ MPa),
33 individuals that had the least total available water in soil/subsoil (0-2m) relied more on deep water
34 and were also subject to less water stress. These results suggest that trees with less favorable
35 soil/subsoil conditions (i.e. low water retention capacity) in the near surface (0-2m) adapt their root
36 systems to exploit deep water reserves more intensively so as to enhance their drought tolerance,
37 while trees with more favorable surface conditions exhibit greater water stress and may be more
38 vulnerable to extreme droughts because of a lower root development in deeper horizons.

39

40 **Introduction**

41 On one side, a large part of continental precipitation is transpired at the world scale (e.g. Oki et
42 Canae 2006; Maréchal et al. 2009; Fisher et al. 2017). On the other side, the water availability is one
43 of the most important factors driving transpiration, biomass productivity, and plant species
44 distribution in water-limited ecosystems (Rambal et al. 2003; Mathys et al. 2014). Understanding
45 forest response to droughts is a crucial issue under climate change, because of their multiple impacts
46 on ecosystems and society (Kirilenko and Sedjo 2007; Bonan 2008; Taylor et al. 2013). Indeed,
47 ongoing climate change has a strong impact on vegetation (Allen et al. 2010; Anderegg 2015) and
48 changes in vegetation cover have a strong impact on hydrological processes (Scanlon et al. 2005;
49 Noretto et al. 2012). Deciphering the complex interactions between vegetation cover and
50 hydrological processes is crucial to improve the predictability of climate changes impacts on the
51 hydrological cycle.

52 The capacity to withdraw water in deep water is a key feature of drought tolerance (Ehleringer et
53 Dawson 1992; Querejeta et al. 2007; Pivavorroff et al. 2016; Brum et al. 2019) that is known to vary
54 among species and contributes, together with other traits, to define the hydrological niches (Brum et
55 al. 2019).

56 Drought tolerance variability at the intra-specific level, and within a same population, is expected to
57 have a key influence on the evolutionary potential of forests under climate change, and could
58 thereby constitutes an important source of resilience to drought (Bontemps et al. 2017). It could
59 result from of adaptation or plasticity in response to micro local environmental variations.

60 However there are less data available at intra-specific level, and in particular, the role of deep water
61 exploration for drought tolerance is less clear. Voltas et al. (2015) showed that genetic factors could
62 explain differences in water use on Aleppo pines of various origins grown in common garden test.
63 Barbeta et al. (2015) showed that plasticity could lead to differences in water exploitation for
64 individuals located in a rain exclusion zone and those located in the control zone. Some studies have
65 suggested that micro local variability in soil/subsoil conditions (i.e. water retention capacity,
66 stoniness, soil depth) also explains intra-specific variations of both leaf and tree scale traits (e.g. leaf
67 mass area, foliar $\delta^{13}\text{C}$, leaf area index, predawn leaf water potential), (Love et al. 2019; Preisler et al.
68 2019; Carrière et al. 2020a). More specifically, they suggest that trees with less favorable soil
69 environments have greater acclimation to drought and experience lower water stress and dieback
70 during extreme drought (than trees with higher total available water (TAW)) within soil. But this
71 theory implies two related hypotheses that so far have not been rigorously tested: i) a relationship
72 exists between near-surface soil/subsoil conditions and the ability of individuals to exploit deep
73 water, and ii) this relationship may affect the resistance of individuals to drought.

74 In this study we aim to assess the relationship between tree water stress, soil/subsoil characteristics,
75 and tree capacity to withdraw deep water, on 6 individual *Quercus ilex* stems. In particular we
76 explore if soil/subsoil conditions consistently affect tree level water stress and trees' ability to draw
77 water at several depths.

78 For these purposes, we chose to work in a karst settings where the soil/subsoil conditions are
79 expected to be strongly contrasted between individuals due to the intrinsic heterogeneity of this
80 media (Hartmann et al. 2014). In this type of environment, strong contrasts in soil thickness,

81 stoniness and water retention capacity can induce a strong variability of soil/subsoil conditions in a
82 few meters, which in turn might influence the develop of root systems for vegetation. We combine
83 three distinct approaches: geophysical, isotopic, and eco-physiological. The first approach, Electrical
84 Resistivity Tomography (ERT), is a non-destructive near-surface geophysical technique used to
85 characterize variability in soil/subsoil conditions (Dahlin 2001; Chalikakis et al. 2011). Unlike classical
86 techniques (e.g. pedologic pits, neutron probe, or Time Domain Reflectometry (TDR) sensors for
87 water), ERT provides an integrated and spatialized (horizontally and vertically) characterization of the
88 near surface at the plot scale. The second approach is water isotope tracing ($\delta^{18}\text{O}$), which is used to
89 determine the source of water extracted by plants (Dawson 1993; Querejeta et al. 2007; Nie et al.
90 2012; Ding et al. 2018). The third approach uses shoot water potential (Ψ), which is widely used to
91 assess plant water stress (Turner 1981). When measured at predawn, transpiration is close to zero,
92 leaf water potential provides an estimate of soil water deficit in the rooting zone. However, when
93 measured during daytime, water potential measurements results from both, soil water deficit and
94 plant hydraulics and transpiration. All datasets used in this study have been described and
95 investigated in previous papers. Carrière et al. (2020a) used ERT to study the variability of foliar traits
96 in response to drought. Carrière et al. (2020b; c) used water isotope combined with predawn and
97 midday leaf water potential to study the variability of deep water use among three species. In this
98 paper, we combine geophysical, isotopic, predawn and midday leaf water potential dataset in an
99 original way to explore the variability of deep water use between individuals of the same species.

100

101 **Material and Methods**

102 **Experimental site**

103 The experimental forest site is located in southern France within the Fontaine-de-Vaucluse
104 observatory (43°56'12"N; 5°27'58"E; 530 m a.s.l.), which is part of the French critical zone
105 observation network (OZCAR - <http://www.ozcar-ri.org/>; Jourde et al. 2018). The plot, which

106 measures 150*50 m, is dominated by *Quercus Ilex* L. (85% of the basal area) and an understory (15%
107 of the basal area) dominated by shrubs including *Buxus sempervirens* L., *Juniperus communis* L. and
108 *Juniperus phoenicea* L. (more details can be found in Carrière et al. 2017). The forest was managed as
109 a coppice for charcoal production for centuries before the last clear-cut 90 years ago.

110 The climate is Mediterranean with dry and hot summers; most rainfall occurs during spring and
111 autumn. Between 2003 and 2015, average annual rainfall was 909 mm and ranged between 407 and
112 1405 mm. Average annual temperature over the same period was 12.9°C. The soil is a stony rendzina
113 whose thickness is highly variable over the study site. Soil pits excavated along the profile revealed
114 soil depths ranging from 0 to 70 cm. This soil has developed on a karstified calcareous bedrock of
115 Urganian facies.

116 Under the experimental plot, a former military bunker has been converted into a scientific laboratory
117 (<http://lsbb.eu/presentation/>). This underground facility provides an opportunity to sample water
118 flows in the vadose zone at depths of 33 and 256m. Surface-based geophysical exploration showed,
119 with ERT, that the first two meters of the soil/subsoil can store considerable amounts of water
120 (several tens of millimeters) but with high spatial variability at the plot scale (Carrière et al. 2015).
121 Deeper (2 to 90 m) magnetic resonance sounding (MRS) and gravimetry showed seasonal relative
122 variations in water storage that can reach 50% within the vadose zone (Carrière et al. 2016).

123

124 **Electrical Resistivity Tomography**

125 The ERT dataset used in this study was described by Carrière et al (2015) and then re-used by
126 Carrière et al (2020a) to assess spatial variability of soil/subsoil conditions (i.e. TAW) at the plot level.
127 Additional methodological details about ERT measurements can be found in those two studies. ERT
128 measurements were done using an ABEM Terrameter SAS 4000 device (Dahlin, 2001) with 64
129 stainless steel electrodes and 2 m spacing along a 126 m transect. The measurement protocol that
130 was used recorded 1140 points. The transect was chosen to align almost perpendicularly to the slope

131 and geological structures (Carrière et al. 2013) to maximize the variability in soil/subsoil conditions.
132 The inversion quality of the ERT data was evaluated with RMS and Chi^2 , calculated using the raw data
133 and the inverted model (lower-left corner of each cross-section; Figure 1). The resistivity model had a
134 2 m lateral resolution and a vertical resolution ranging gradually from 0.5 m near the surface to 1.5 m
135 at the base of the cross-section (10 m). A temperature correction was applied on the final model,
136 following Keller et Frischknecht (1966):

$$137 \quad \rho = \rho_T [1 + \alpha (T - 25)] \quad (1)$$

138 where ρ ($\Omega\cdot\text{m}$) is electrical resistivity at the reference temperature of 25 °C, ρ_T ($\Omega\cdot\text{m}$) is electrical
139 resistivity measured at temperature T (°C), and α is 0.0202. T is underground temperature,
140 logarithmically interpolated between daily mean air temperature aboveground and groundwater
141 temperature at a depth of 30 m.

142 The Percent Variation in Resistivity (*PVR*) was used to describe the spatial variability of soil/subsoil
143 conditions under each *Quercus Ilex* L. individual. *PVR* was calculated for each tree as a function of the
144 dry (ρ_{dry}) and wet (ρ_{wet}) resistivity cross-section, as in Clément et al. (2010) and Robert et al. (2012):

$$145 \quad PVR(\%) = \frac{\rho_{\text{dry}} - \rho_{\text{wet}}}{\rho_{\text{dry}}} * 100 \quad (2)$$

146 Two ERT measurements were successively carried out in autumn 2011 at an interval of a few days
147 ranging from an extremely dry (i.e. 3 months of drought) to an extremely wet (i.e. after an episode of
148 230 mm rainfall) status (Figure 1a). The temporal proximity of these measurements and similar
149 meteorological conditions between these two days resulted in very limited variations in air and
150 groundwater temperatures. In Carrière et al. (2020a), we showed that *PVR* between these two dates
151 can be used as a proxy for the total available water (TAW) for vegetation (Figure 1b). The near
152 surface *PVR* was calculated for each individual by a lateral average within a distance of 2.5 m
153 (corresponding to crown radius) to the stool center and vertically between 0 and 2 m. This thickness
154 corresponds on average to the depth of the massive limestones previously detected by ground

155 penetrating radar (Carrière et al. 2013). We consider that the TAW (0-2m) of each tree is fix
156 parameter in time for a 90 years old stand and that it is its degree of filling that varies over the
157 seasons according to drought level. Therefore, the PVR, which is a proxy of TAW, was measured only
158 once in 2011.

159

160 **Isotopic tracing**

161 **Sampling**

162 The isotopic dataset was published in Carrière et al. (2020c). Field work was conducted on 6 trees at
163 monthly time step between June and August 2015 (06/11/2015 ; 07/06/2015 ; 08/11/2015). On each
164 date, three to four sunny branches with a diameter of 4 to 6 mm were collected from each tree at
165 midday. Phloem and bark were removed to prevent any contamination from phloem sap. The
166 samples were immediately packed in parafilm and placed in sealed vials. These three to four
167 branches were pooled before extraction. A portable cooler was used to transfer the samples to the
168 laboratory where they were stored frozen until water extraction and analyses. Liquid samples of
169 precipitation and drainage water were collected every 1 to 3 weeks. Precipitation was collected
170 through a pluviometer and stored in containers installed in a pit so as to limit temperature variations.
171 Drainage water was collected at 20 cm below the surface using a mini-lysimeter and this water was
172 stored in a separate container (see figure in SI1). These containers were maintained at atmospheric
173 pressure through a capillary to limit exchange between the atmosphere and collected water (IAEA
174 2014), (see figure SI1). The samples were collected on a monthly time step basis and were therefore
175 a mixture of all the water flowed into the cans during the previous month. At this experimental site,
176 the rocky karst soil prevents soil water sampling using auger drilling or porous cups.

177 **Water extraction and isotopic analyses**

178 Xylem water was extracted from wood by cryogenic vacuum distillation following the protocol of
179 West et al. (2006). The twigs were cut into small pieces and placed in an electrothermal heating and
180 stirring mantle at 90 to 100 °C for 1 hour. Two successive liquid nitrogen traps were used to collect 3
181 to 5 ml of xylem water. This water was stored in small vials until analyzed.

182 Deep water and shallow water samples were analyzed using a Los Gatos Isotope Ratio Infrared
183 Spectrometer (IRIS) at Avignon University (LGR DLT-100 liquid water stable analyzer accuracy $\pm 0.2\text{‰}$
184 vs V-SMOW for $\delta^{18}\text{O}$). The isotopic ratios were expressed as:

185

$$186 \quad \delta^{18}\text{O} = \left[\left(R_{\text{sample}} / R_{\text{standard}} \right) - 1 \right] \times 1000\text{‰} \quad (3)$$

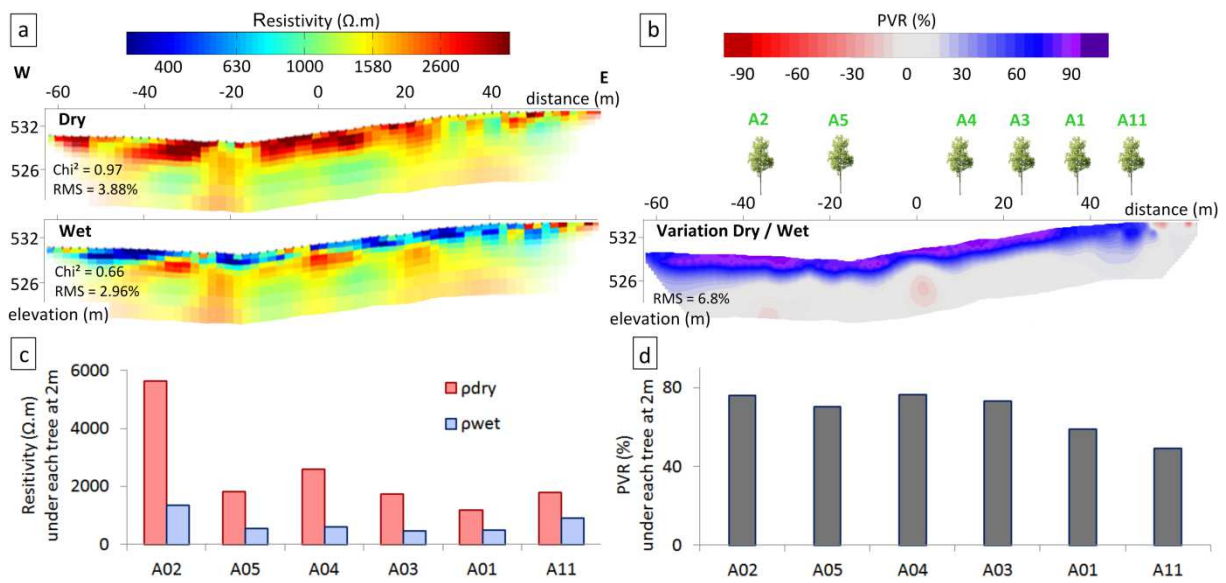
187

188 where R_{standard} and R_{sample} are the light/heavy isotope ratios ($^{16}\text{O}/^{18}\text{O}$) of the sample and the standard
189 (Vienna Standard Mean Ocean Water (VSMOW)), respectively. Possible spectral perturbations of the
190 IRIS measurements due to organic contaminants in xylem and drainage samples (Martín-Gómez et al.
191 2015) motivated us to use an Isotope Ratio Mass Spectrometer (IRMS). Xylem and drainage samples
192 were analyzed using the Isoprime IRMS at the LAMA laboratory of HydroSciences Montpellier, using
193 the CO_2 equilibration technique in dual inlet mode, yielding $\delta^{18}\text{O}$ results with a $\pm 0.06 \text{‰}$ precision.
194 This value was obtained by calculating the standard deviation related to repeated measurements of a
195 standard sample during the analysis of our samples. This precision is reported in Figure 2b and c in
196 the $\delta^{18}\text{O}$ error bars.

197 **Leaf water potential**

198 The predawn (Ψ_P) and midday (Ψ_M) leaf water potential dataset was previously published in Carrière
199 et al. (2020a). Both metrics were measured on the 6 individuals chosen for isotope sampling
200 throughout the 2015 summer season using a Scholander pressure bomb. Samples for Ψ_P were
201 collected in the morning, before sunrise. For Ψ_M , samples were collected from sun-exposed
202 branches around midday (2PM GMT), when the sky was not cloudy. For each tree, at least four leaves

203 were sampled and immediately placed in a plastic bag saturated with water vapor and stored in a
 204 portable cooler until measurement (few minutes later). Between two and four leaves were measured
 205 to check the consistency between measurements. These multiple measurements were used to
 206 determine the error bars shown in Figure 2 b and c. In the results we divided our interpretation into
 207 two periods: moderate stress ($\Psi_p > -2\text{MPa}$) and severe stress ($\Psi_p < -2\text{MPa}$) following the work of
 208 Lempereur et al (2015).



209

210 Figure 1: Electrical Resistivity Tomography (ERT) results. (a) ERT models for the “dry” and “wet” dates
 211 (respectively before and after a rain event in November 2011), involved in *PVR* computation
 212 (equation 2); Gradient arrays, 64 electrodes; ERT model uncertainty, based on current line density, is
 213 represented by the color attenuations at the edges and the bottom of the resistivity cross-section. (b)
 214 Percent variation in resistivity (*PVR*) between the two resistivity models (dry/wet). (c) Resistivity
 215 averaged over 2 m under each tree in dry (ρ_d) and wet period (ρ_w). (d) *PVR* averaged over 2 m under
 216 each tree.

217 Data analysis

218 We assumed that the isotopic dataset (Carrière et al. 2020c) is affected by an isotopic fractionation in
 219 ^2H (D-fractionation), therefore we use here only ^{18}O data. D-fractionation is visible in many works

220 (e.g. Brooks et al. 2010; Evaristo et al. 2016; Bowling et al. 2017; Evaristo et al. 2017; Geris et al.
221 2017; Vargas et al. 2017). The causes of this fractionation are still poorly understood (Barbeta et al.
222 2019). Early studies have assumed that D-fractionation is related to the mode of water absorption by
223 roots (Sternberg and Swart 1987; Lin and Sternberg 1993; Ellsworth and Williams 2007). Barbeta et
224 al. (2020) suggest that D-fractionation is rather related to the heterogeneity of the water signal in soil
225 pores and plant tissues. Ellsworth et al. (2007) and Barbeta et al. (2020) agree that fractionation is
226 much stronger in ^2H than in ^{18}O and that ^{18}O provides more satisfactory analysis.

227 Therefore, considering only one valid tracer (^{18}O), we had to simplify our analysis into only two
228 potential pools of plant supply: i) the shallow pool or ii) the deep pool (Fig. SI2). Rain and drainage
229 water were analyzed separately, but the two datasets are similar during summer. We arithmetically
230 averaged the two values to represent the "shallow water" signal in Figure 2b and c. Deep water was
231 collected at two seepage points (named "Point C" and "Point D") within the LSBB tunnel
232 (<http://lsbb.eu/presentation/>) under the experimental site. These two points were arithmetically
233 averaged to represent the "deep water" signal of the karst vadose zone in Figure 2b and c. We
234 assume that this "deep water" signal corresponds to water contained within rock, as described by
235 Bowling et al. (2017) and Geris et al. (2017).

236 The analysis presented in this paper is based on two previously published dataset (Carrière et al.
237 2020a, b). Combining both datasets, only six trees were prospected by geophysics and were
238 monitored in $\delta^{18}\text{O}$ and Ψ along the summer period 2015. The logistical difficulty for data acquisition
239 explains this small sample. We therefore evaluated the correlation between the parameters using
240 the Pearson correlation, but also the Spearman correlation which is recommended for the analysis of
241 small samples.

242

243 **Results and interpretation**

244 **Spatial variability in soil/subsoil properties at tree scale assessed with ERT**

245 We observed wide variations in resistivity both within and between the driest and wettest profiles
246 measured in 2011 (Figure 1a), from less than 300 $\Omega\cdot\text{m}$ in wet conditions to more than 2500 $\Omega\cdot\text{m}$ in
247 dry conditions. We note that the resistivity signal was more stable between the two profiles in deep
248 (> 2 m) than in shallow (<2m) layers (Fig. 1a), which caused *PVR* to be smaller in deeper layers (Figure
249 1b).

250 *PVR* patterns showed consistent signals with field soil observations and total available water (TAW)
251 measurements. For instance, the cross section reveals a low *PVR* to the right of tree A11, which is
252 consistent with field observations showing limestone rock outcrops in this area (see SI3). We also
253 observed a consistent trend between *PVR* and TAW estimated through pedologic pits (see SI4). This
254 was discussed in Carrière et al. (2020a) and provides further evidence that *PVR* is well suited to
255 quantifying TAW relatively.

256 In the following, *PVR* (0 - 2 m) is used as a proxy for TAW. As an example, *PVR* values indicate that
257 tree A4 has a higher TAW than A11. We will examine in the following results whether these trees
258 have contrasting $\delta^{18}\text{O}$ and Ψ_{M} signals. Note that the *PVR* is a constant value for each tree, unlike $\delta^{18}\text{O}$
259 and Ψ_{M} , which vary throughout the season.

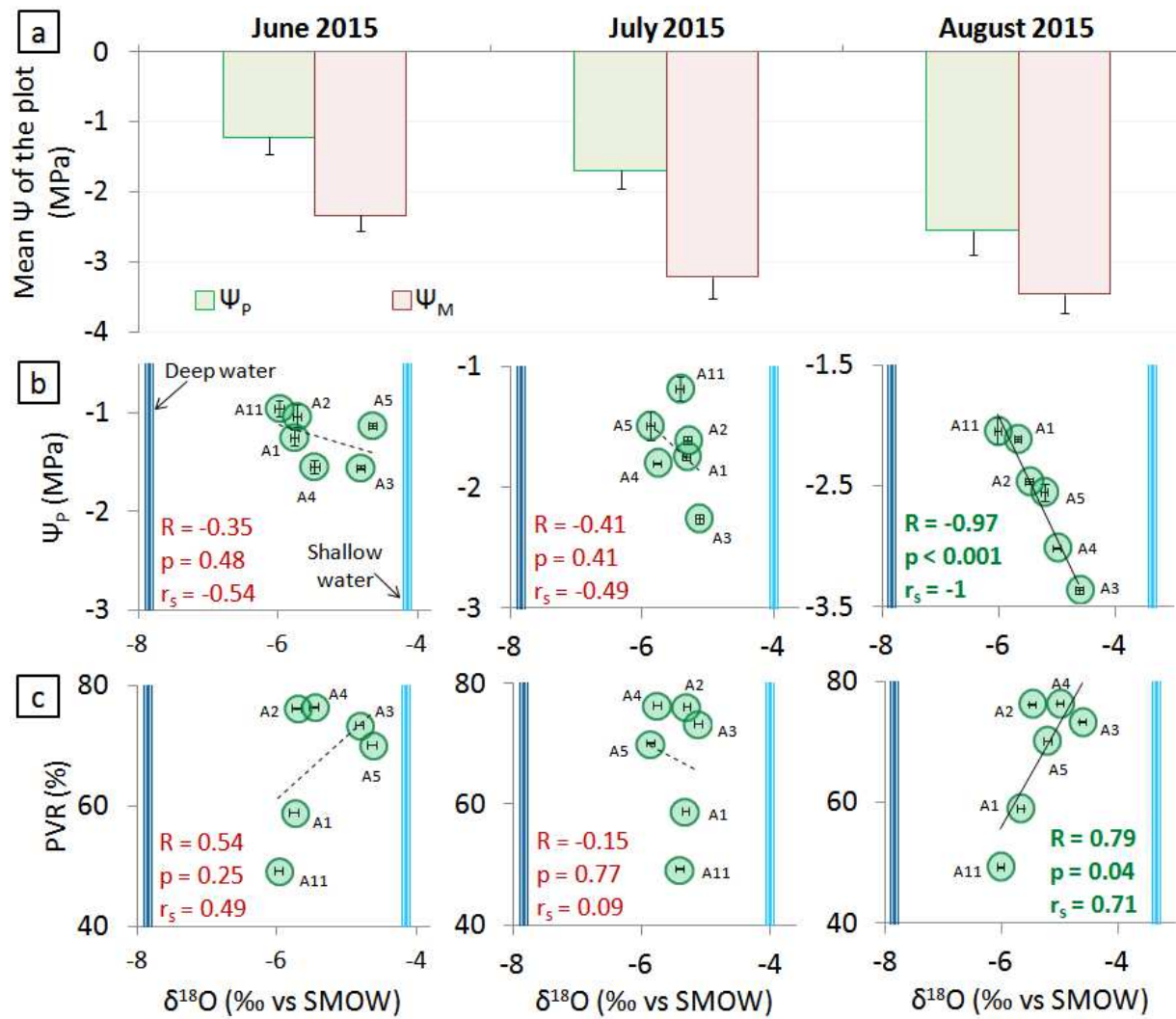
260 **Individual drought adaptation strategies**

261 Tree water deficits increased during the 2015 summer season, as shown by the clear decrease in
262 average Ψ_{P} and Ψ_{M} leaf water potentials from June to August 2015 (Figure 2a). Concurrently, the
263 difference between Ψ_{P} and Ψ_{M} tended to decrease during the summer season, suggesting that trees
264 regulate their transpiration as drought increases. At the individual scale (Figure 2b), we observed
265 that tree A11 was systematically less stressed than tree A4. Moreover, the range of Ψ_{P} among
266 individuals increased with drought.

267 During moderate water deficit ($\Psi_{\text{P}} > -2$ MPa, June and July), there was no clear relationship between
268 Ψ_{P} and $\delta^{18}\text{O}$ of the xylem (Figure 2b ; p-value > 0.4). By contrast, at the peak of the drought (August
269 2015), we observed a significant negative relationship between Ψ_{P} and $\delta^{18}\text{O}$ of the xylem (p-value

270 < 0.001). This implies that the least stressed trees (with the highest water potentials) had an isotopic
271 signal closer to the isotopic signal of groundwater while the more stressed trees (with the lowest
272 water potential) had an isotopic signal closer to shallow water. There was no relationship between
273 Ψ_M and $\delta^{18}O$ at any period in 2015 (Fig. S15). This is probably due to the fact that Ψ_M is dependent on
274 soil water status but also on additional environmental (light and vapor pressure deficit) and biological
275 factors including plant hydraulic conductance, tree transpiration) that blurred its relations with soil
276 related metrics.

277 Similarly, no significant relationship was observed between the isotopic signal and *PVR* among
278 individuals when stress is moderate, but a significant relationship (p -value < 0.05) was observed in
279 drier conditions (Fig. 2c). Trees with a low *PVR* between 0 and 2 m (i.e. low TAW) had a $\delta^{18}O$ signal
280 closer to groundwater while trees with a high *PVR* (0-2 m) had an isotopic signal closer to shallow
281 water.



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290 Discussion and perspectives

291

292

Figure 2 : a) Mean and standard deviation values for leaf water potential of the stand at predawn (Ψ_P) and midday (Ψ_M) for the months of June, July, and August 2015; b) Relationship between Ψ_P and the xylem isotope signal ($\delta^{18}O$) for each individual (each point is a tree), the dark blue and light blue vertical bars represent the groundwater and shallow water isotope signals, respectively. R is Pearson correlation, p is p-value, and r_s is Spearman correlation; c) Relationship between PVR and xylem $\delta^{18}O$ during the summer 2015.

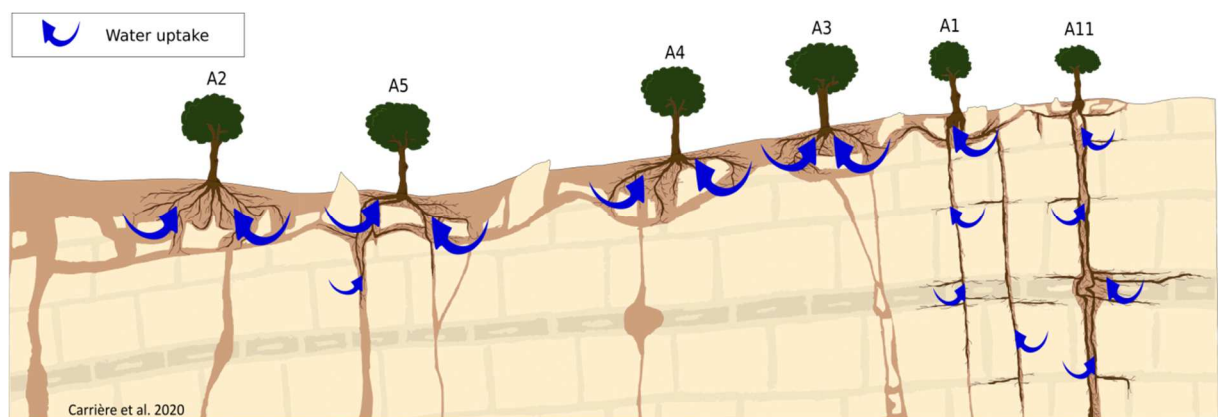
293 in such variability remain poorly known. In Carrière et al. (2020a) we showed that trees with low near
294 surface TAW (low PVR) experienced higher stomatal control and lower water stress during drought
295 peak compared to trees with higher TAW, which experienced higher water stress and a higher
296 defoliation rates following extreme drought (see SI6). In this study, for the first time, we analyzed
297 together ERT, water potential measurements, and isotopic data to further show that there is a link
298 between near-surface TAW, water status, and tree ability to exploit deep water.

299 We found a negative correlation between tree $\delta^{18}\text{O}$ and Ψ_p of individuals at the drought peak in
300 August 2015, indicating that the individuals experiencing higher water stress relied proportionately
301 less on deep water sources than the others. This drought dependence of the xylem isotope signal had
302 already been demonstrated across different species (Flanagan et al. 1992; Jackson et al. 1995; Brum
303 et al. 2019) or in the course of drought for a single species (Barbetta et al. 2015; Carrière et al.
304 2020b). However, to our knowledge, this is the first time that such mechanism is reported at the
305 intra-specific level. Furthermore, we show that trees with low TAW (0-2 m) rely proportionally more
306 on deep water during drought peaks. A likely explanation for this pattern is that trees with low TAW
307 would have developed deeper root into the karst vadoze zone on which they could rely when
308 drought becomes critical (Figure 3). This hypothesis would explain why we observed an isotopic
309 signal closer to deep water and a lower vulnerability to intense droughts for trees with higher TAW. A
310 schematic interpretation of our results is presented in Figure 3.

311 Drought resistance variability among individuals of the same species is rarely considered in ecological
312 studies despite its role in forest structure and outcomes under ongoing climate change. Here we
313 show strong variability between individuals that can be partially explained by underground factors
314 (TAW, ability of trees to extract deep water). This implies that stand resistance to drought may be
315 more heterogeneous than is currently predicted by current ecosystem models. For instance, our
316 conclusions may explain the results of several studies showing that tree dieback following drought

317 occurs more frequently in areas where trees benefit from well-developed soils (Nourtier et al 2014;
318 Preisler et al. 2019; Carrière et al 2020a).

319 Further experiments should be carried out on a larger scale and on multiple species to generalize our
320 results, and to evaluate if inter and intra-specific differences overlap. We already obtained
321 encouraging results by enriching the Ψ_p and $\delta^{18}O$ relationship (Figure 2b) with data observed in
322 August 2015 on beech and silver fir trees at a nearby site located at slightly higher altitude (Figure
323 S17). It would also be interesting to assess if deep water extraction scale with root biomass in
324 factures, that can be estimated with non-invasive method (Mary et al. 2019).



325
326 Figure 3: Schematic interpretation of tree root implantation as a function of their soil/subsoil
327 condition identified by geophysics (Figure 1b), and their response to drought (Ψ_p and $\delta^{18}O$). The
328 representation of the crown size of each tree is proportional to its individual leaf area, which is also
329 related to soil/subsoil conditions (Carrière et al. 2020a).

330

331 Conclusions

332 The combination of geophysics, isotopes, and foliar water potentials open fruitful research avenues
333 to clarify plant-water relationships and tree response to drought. Significant differences in water
334 stress were observed between the trees of the same species growing on a Mediterranean karst. Our
335 analysis clearly suggest that such differences are due to karst heterogeneity, which imposes

336 contrasted soil/subsoil conditions (i.e. TAW) between individual. We have shown that trees with the
337 least favorable near-surface (0-2m) soil/subsoil conditions proportionately extract more deep water
338 in periods of severe stress. We hypothesis that deep water extraction can be considered as an
339 adaptation mechanism to recurrent water deficit. It is crucial to better understand these adaptation
340 mechanisms in order to better anticipate the impact of changes in vegetation cover on hydrological
341 processes.

342

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