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# The role of deep vadose zone water in tree transpiration during drought periods in karst settings – Insights from isotopic tracing and leaf water potential



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#### HIGHLIGHTS

- Mediterranean forest in karst environment exploits deep water in vadose zone.
- Beech, Silver Fir, and Holm Oak exhibit different strategies in groundwater use.
- Importance of including deep water resources in ecohydrological models

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## G R A P H I C A L A B S T R A C T



# ABSTRACT

Karst environments are unusual because their dry, stony and shallow soils seem to be unfavorable to vegetation, and yet they are often covered with forests. How can trees survive in these environments? Where do they find the water that allows them to survive? This study uses midday and predawn water potentials and xylem water isotopes of branches to assess tree water status and the origin of transpired water. Monitoring was conducted during the summers of 2014 and 2015 in two dissimilar plots of Mediterranean forest located in karst environments. The results show that the three monitored tree species (*Abies alba* Mill, *Fagus sylvatica* L, and *Quercus ilex* L.) use deep water resources present in the karst vadose zone (unsaturated zone) more intensively during drier years. *Quercus ilex*, a species well- adapted to water stress, which grows at the drier site, uses the deep water resource very early in the summer season. Conversely, the two other species exploit the deep water resource only during severe drought. These results open up new perspectives to a better understanding of ecohydrological equilibrium and to improved water balance modeling in karst forest settings.

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## 1. Introduction

Changes in temperature and rainfall patterns across many forest ecosystems are expected to increase the risk of droughtinduced tree mortality (Allen et al., 2010). Tree resistance to drought is therefore of concern to scientists (Chaves et al., 2002; Bréda et al., 2006) and stakeholders (Spies et al., 2010; Keenan, 2015). To cope with drought, trees have developed various strategies at different time scales.

Stomata closure is a universal process designed to limit transpiration (Limousin et al., 2010), avoid water potential drop, and prevent irreversible damage caused by embolism to the plant hydraulic system (Martin-StPaul et al., 2017). Other mechanisms that take place at longer time scales involve changes in leaf area to decrease water consumption (Limousin et al., 2009; Van Hees, 1997; Martin-StPaul et al., 2013). Instead of regulating water loss, trees can also take up water from groundwater when the water table is accessible (e.g. Murray et al., 2003; Naumburg et al., 2005; Querejeta et al., 2007). This process helps to maintain transpiration and carbon assimilation rates and limits water potential drop and cavitation risk during drought (Rambal, 1984; Bréda et al., 2006: Abrams, 1990: McElrone et al., 2004: Johnson et al., 2014). However, several examples in the literature show that despite extremely dry shallow resource conditions ( $\theta_{soil} < 0.5\%$  $\Psi_{soil} < -7$  MPa, respectively, Rambal, 1984; Plaut et al., 2012) and a water table that is inaccessible (>100 m) to root prospection (Canadell et al., 1996), trees can continue to transpire. Increased biomass allocation to roots is also observed (Van Hees, 1997). This makes it possible to tap water from deeper resources (Fig. 1b) during the dry season and thus to sustain water fluxes. How does this deep water reservoir contribute to transpiration throughout drought periods?

To explain summer tree survival in karst environments, it seems necessary to invoke deep water exploitation. Karst environments (Martel, 1902; Cvijic, 1960; Bakalowicz, 2005) are rocky and often have steep landscapes resulting from the weathering of carbonate or evaporitic rocks. These landscapes are common world-wide (Chen et al., 2017), especially around the Mediterranean Sea (Bakalowicz and Dörfliger, 2005). Soils in karst environments are generally poorly developed and rocky or even absent (Fig. 1) and karst's high permeability causes these environments to be generally dry and apparently not well-suited for vegetation development. Due to their low productivity, karst landscapes are rarely cultivated and are often covered with forest. These ecosystems are crucial because they constitute the recharge areas of karst aquifers that supply water to a large part of the world's population (e.g. 50% in Austria, 30% in Belgium, (Hartmann et al., 2014)). Karst hydrosystems have high water storage capacity in the saturated zone (Ford and Williams, 2007) and in the vadose zone (Fig. 1a). In the vadose zone, water storage can occur in: *i*) superficial horizons called epikarst, composed of soil and weathered rock (e.g. Perrin et al., 2003; Aquilina et al., 2006); or *ii*) deeper horizons (e.g. Emblanch et al., 2003; Carrière et al., 2016). However, the total soil available water (TSAW) generally measured via pedologic pits in these environments is low (a few tens of mm) and does not explain vegetation transpiration during drought periods.

In the three last decades, stable isotope ratios of hydrogen ( $\delta^2$ H) and oxygen ( $\delta^{18}$ O) have been used to determine the source of water uptake by plants from soil, streams, and groundwater (e.g. Ehleringer and Dawson, 1992; Bariac et al., 1994; Walker and Brunel, 1990; Ogle et al., 2014; Barbeta et al., 2015; Evaristo et al., 2015). It has been hypothesized that water that enters roots is not isotopically fractionated with respect to soil water (Dawson and Ehleringer, 1991; Zimmermann et al., 1966). This idea has led many authors to use deconvolution to estimate the mixing ratio of xylem water between two pools (e.g. Dawson, 1993; Phillips and Ehleringer, 1995) or three pools (e.g. Parnell et al., 2010; Barbeta et al., 2015). However, earlier studies show that in halophyte and xerophyte plants (Sternberg and Swart, 1987; Lin and da Sternberg, 1993; Ellsworth and Williams, 2007) and avocado (Per*sea americana*) trees (Vargas et al., 2017),  $\delta^2$ H isotopic fractionation can occur during root water uptake. This isotopic fractionation of  $\delta^2$ H is noted in many studies (e.g. Brooks et al., 2010; Bertrand et al., 2014: Barbeta et al., 2015: Evaristo et al., 2016: Bowling



Fig. 1. Conceptual diagram of (a) a karst hydrosystem, (b) water reservoirs available for trees in a karst setting: (i) shallow water resource in soil and weathered rock; (ii) deep water resource in the karst vadose zone.

et al., 2017; Evaristo et al., 2017; Geris et al., 2017). Fortunately, this fractionation effect has a disproportionately greater effect on  $\delta^2$ H than on  $\delta^{18}$ O (Ellsworth and Williams, 2007; Vargas et al., 2017). Moreover, Barbeta et al., (*submitted*) have shown that mixing proportion calculations performed with  $\delta^{18}$ O data provide consistent results when the  $\delta^2$ H estimation is distorted.

Leaf or stem water potential is a widely used proxy for plant water stress (Turner, 1981). The more negative the leaf water potential, the greater the stress on the plant. Leaf water potential is generally measured with a Sholander pressure bomb (Scholander et al., 1965) but can also be measured with psychrometers to conduct continuous potential measurements (Taylor-Laine et al., 2016). Leaf water potential measured at predawn  $(\Psi_{\rm P})$  is generally considered representative of soil water potential in the rhizosphere (Tardieu et al., 1990; Brisson et al., 1993). Leaf water potential at midday ( $\Psi_M$ ) depends not only on the soil water potential but also on plant hydraulic conductance and the evapotranspiration (ET) demand based on the Ohm's law analogy (Brisson et al., 1993; Olioso et al., 1996; Cruiziat et al., 2002; Martínez-Vilalta et al., 2014). Hence, the difference between predawn water potential and midday water potential ( $\Delta \Psi$ ) provides information about the sap flow driving force and indirectly the transpiration level (Brisson et al., 1993; Cruiziat et al., 2002). The greater the  $\Delta \Psi$ , the more the trees transpire; the smaller the  $\Delta \Psi$ , the less the trees transpire.

Used independently, the two approaches presented above make it possible to determine the origin of water and the tree stress level, respectively. In this study we combine isotopic tracing and leaf water potential to determine if it is possible to relate extraction depth to stress level. For this reason, our measurements were done during two summer periods (2014–2015) in two forest types located on Mediterranean karst. One is a mountain forest (1340 m. a.s.l.) dominated by *Abies alba* and *Fagus sylvatica* trees, and the other, at lower elevation (530 m.a.s.l.), is a typical Mediterranean ecosystem covered by *Quercus ilex*. We sampled xylem water from three tree species, as well as shallow and deep water in the karst vadose zone for  $\delta^{18}$ O analysis. The main goal of this experiment was to answer the following three questions:

(i) Do trees use deep karst water from the vadose zone during the dry season?

(ii) How does deep water use by trees vary according to the level of water deficit?

(iii) Does deep water uptake by trees vary among species?

In this project, we used tools from ecology to improve our understanding of how karst systems function. We hypothesize that vegetation may use water from the "deep" vadose zone (between a few meters and few tens of meters) to increase its resilience to drought. This article aims to provide a functional view of the interactions between vegetation and groundwater in the deep vadose zone.

## 2. Material and methods

## 2.1. Experimental site location

The experimental sites are located within the Fontaine de Vaucluse (FdV - France) karst hydrosystem, which hosts one of the largest karst springs in Europe (Fig. 2b; Cognard-Plancq et al., 2006; Jourde et al., 2018). This recharge area covers 1162 km<sup>2</sup> (Ollivier et al., 2019a, 2019b) and has a broad elevation gradient, making



**Fig. 2.** (a) The Fontaine de Vaucluse recharge area located in southeastern France; (b) The experimental sites, Mont Ventoux at 1340 m and Rustrel at 530 m are located in the Fontaine de Vaucluse recharge area. They cover areas of 100 m × 100 m and 150 m × 50 m, respectively.

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Name	Coordinates	Holm oak basal aRea (%)	Silver fir basal aREa (%)	Beech basal aREa (%)	Plot size (m)	Coarse fraction in soil (%)	Soil depth (cm)	MEAN ANNUAL RAINFALL (mm)	Number of pits	Ground- water
Rustrel	43°56′12.15″N 5°27′58.18″E 530 m.a.s.l.	85	0	0	$150\times 50$	50-60	0–70	909	3	Point D Point C
Mont Ventoux	44°10′43.93″N 5°14′36.85″E 1340 m.a.s.l.	0	86.3	13.7	$100 \times 100$	48-63	50-60	1258	4	MtS and Ctr springs

Main characteristics of the experimental sites. MtS and Ctr are Mont Serein and Contrat springs, respectively, and point D and point C are two seepages monitored within the Rustrel underground laboratory (http://lsbb.eu/presentation/).

it possible to find a variety of tree species with differing drought resistance. The FdV observatory is part of OZCAR (http://www.ozcar-ri.org/), the French network of critical zone observatories. We selected two experimental sites: i) one in the southern part of the FdV recharge area near Rustrel village at an elevation of 530 m, and ii) one in the northern part of the FdV recharge area, on the northern slope of Mont Ventoux at an elevation of 1340 m. Table 1 summarizes the characteristics of each site.

#### 2.2. Environmental and groundwater settings

The Rustrel forest is dominated by a sparse evergreen overstory of holm oak (*Quercus ilex L.*), which represents >85% of the basal area (Carrière et al., 2017). The dominant tree height is about 4 m. The understory is a sparse shrubby layer which represents 15% of the basal area and includes *Buxus sempervirens L.*, *Juniperus communis L.*, and *Juniperus phoenicea L.* (Carrière et al., 2017). In this paper, we study only the holm oak on a plot of  $150 \times 50$  m at the Rustrel site.

The stand located on Mont Ventoux's northern slope is dominated by silver fir (*Abies alba* Mill.) (86.3% of the basal area) and beech (*Fagus sylvatica* L.) (13.7% of the basal area). The dominant tree height is about 17.7 m. Our study was conducted on a plot of approximately  $100 \times 100$  m. This plot was previously studied in detail by Nourtier et al. (2014). They used ground-based geophysics (electrical resistivity tomography (ERT)) to characterize soil conditions and to relate silver fir mortality to drought.

The two sites are located on similar karstified Cretaceous limestone. The subsurface karst at the Rustrel site is complex and heterogeneous due to numerous faults and karst features described in detail by Carrière et al. (2013). Carrière et al. (2016) used multiple geophysical techniques (ERT, Magnetic Resonance Sounding (MRS), Gravimetry) to show that mobile water is stored in the karst vadose zone down to 90 m, as demonstrated by significant seasonal water content variations. At both sites, the saturated water zone of the Fontaine de Vaucluse aquifer is inaccessible to trees; the water table is approximately 400 m below surface at Rustrel and 1200 m deep at Mont Ventoux.

At both sites, the karst vadose zone isotopic signal is provided by underground seepage and springs. At Rustrel, two seepage zones named "point D" and "point C" (at 33 m and 256 m below the surface, respectively) have been monitored since 2003 (Garry et al., 2008; Blondel et al., 2010) in the Low Noise Underground Laboratory tunnel (http://lsbb.eu/presentation/). At the Mont Ventoux site, two springs are present: Mont Serein spring (MtS), located at a distance of 1.6 km from the experimental plot (44°10′51.47″ N; 5°15′48.25″ E; 1446 m.a.s.l.), and Contrat spring (Ctr) located at a distance of 3 km (44°11′03.83″ N; 5°16′51.24″ E; 1395 m.a.s.l.).

Soils at both sites are shallow and rocky rendzina has developed due to the karst settings and slope. At Rustrel, the soil contains 50 to 60% coarse elements and its thickness is relatively variable (from 0 to 70 cm). The total soil available water (TSAW), estimated according to the Baize and Jabiol (2011) protocol, ranges between 0 and 65 mm. At Mont Ventoux, the soil contains 48 to 63% coarse elements and its thickness ranges from 50 to 60 cm; TSAW ranges between 45 and 50 mm (Table 1).

The climate at both sites is Mediterranean, characterized by cool and wet winters, hot and dry summers, and a high interannual variability (see SI1). The two sites differ in terms of temperature and precipitation due to the elevation difference. Between 2003 and 2015, annual rainfall at Rustrel ranged from 407 to 1405 mm with a mean value of 909 mm, while the mean annual temperature was 12.9 °C. At Mont Ventoux, annual rainfall ranged from 818 to 1842 mm during the same period and its mean value was 1258 mm, while the mean annual temperature was 6.53 °C.

## 2.3. Isotopic sampling

Field work was conducted between July 2014 and August 2015, to sample two successive summer periods. The xylem isotopic signal was sampled on 3 to 5 trees every 1 to 2 months. Three to four sunny branches were collected from each tree at midday. Bark and phloem were removed to prevent interference with enriched water from the leaves. The twigs were immediately packed in parafilm and placed individually in sealed vials and then transferred in a portable cooler to prevent evaporation. All samples were stored frozen at the laboratory until water was extracted and analyzed. A punch was used to sample xylem from the trunk on one date (June 2015) at each site (Mont Ventoux and Rustrel). Sample conservation was similar to that of the stems. The shallow water isotopic signal was collected from precipitation and drainage water every 1 to 3 weeks. Precipitation was collected by pluviometer and stored in containers installed in a pit sheltered from light to limit temperature variations. In addition, these containers were equipped with an atmospheric pressure capillary in accordance with the IAEA protocol (IAEA, 2014) to limit exchanges between collected water and the atmosphere (see Fig. SI2). At each site, drainage water was collected at 20 cm below the surface through a mini-lysimeter directed to a container similar to those used for precipitation (see Fig. in SI2). We chose to collect drainage water because the subsurface is too rocky for auger drilling or porous candle sampling.

Initially, rain and drainage water were analyzed separately, and because the two trends were similar, the two values were merged (arithmetical average) to represent the "shallow water" signal (Fig. 1) in Figs. 3 and 4. Deep water was collected at two points for each site: i) seepage points D and C for Rustrel, ii) the Contrat spring (Ctr) and the Mont Serein spring (MtS) at the Mont Ventoux site. For each site, these two points were merged to represent the "deep water" signal (Fig. 1) of the karst vadose zone in Figs. 3

and 4. We assume that this deep water represents the rock moisture described by Bowling et al. (2017) and Geris et al. (2017).

### 2.4. Leaf water potential

The leaf water potential was measured at predawn  $(\Psi_P)$  and at midday ( $\Psi_M$  or minimum) for 10 individuals of each species on different dates throughout the summer seasons (June to September) of 2014 and 2015, using a Scholander pressure bomb. Samples for minimum water potentials were collected from sunny branches when the weather was not cloudy. On each date and for each tree, at least three leaves were sampled, immediately placed in a plastic bag saturated with water vapor (by blowing into the bag), and stored in a cooler until measurement. Measurements were made less than one hour after sampling. Two of the three leaves were measured and the third was measured only if a difference >0.2 MPa was found between the two first measurements.

## 2.5. Water extraction and isotopic analyses

Water from xylem samples was extracted by cryogenic vacuum distillation (Ehleringer and Osmond, 1989; West et al., 2006). The stems were quickly cut into small pieces and placed in a flask heated to 90 to 100 °C for 1 h. The water was collected in two successive liquid nitrogen traps. Generally, 3 to 5 ml of xylem water was extracted. These liquid samples were stored in small vials until analyzed.

Precipitation and deep water samples were analyzed on a Los Gatos Isotope Ratio Infrared Spectrometer (IRIS) at the University of Avignon (LGR DLT-100 liquid water stable analyzer accuracy  $\pm 0.2\%$  vs V-SMOW for  $\delta^{18}$ O). However, because of possible spectral perturbations of IRIS measurements due to organic contaminants in xylem and drainage samples (Martín-Gómez et al., 2015),  $\delta^{18}$ O of xylem and drainage samples were also analyzed using the Isoprime Isotope Ratio Mass Spectrometer (IRMS) at the LAMA laboratory of HydroSciences Montpellier, using the CO<sub>2</sub> equilibration technique in dual inlet mode, yielding  $\delta^{18}$ O results with a  $\pm 0.6\%$  precision. The isotopic ratios in this study are expressed as:

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

where  $R_{sample}$  and  $R_{standard}$  are the heavy/light isotope ratios (<sup>18</sup>O/<sup>16</sup>O) of the sample and the standard (Vienna Standard Mean Ocean Water (VSMOW)), respectively.

#### 2.6. Mixing model theory and uncertainty

We assume that xylem water presents a mixed signal between deep and shallow water. If the origin of xylem water is linked to two sources without isotopic fractionation, the proportional contribution of each source may be resolved using a single isotope in a two-source system mass balance equation (Dawson, 1993; Phillips and Ehleringer, 1995):

$$\delta_{xyl} = f_{sw} \cdot \delta_{sw} + f_{dw} \cdot \delta_{dw} \tag{2}$$

where  $\delta_{xyl}$  is the plant xylem water  $\delta^{18}$ O, and the proportions shallow water  $(f_{sw})$  and deep water  $(f_{dw})$  of the two sources have isotopic signatures  $\delta_{sw}$  and  $\delta_{dw}$ , respectively. We performed a single isotope ratio two-source mass balance approach and calculated the fraction of deep water  $(f_{dw})$  contribution to xylem from Eq. (2) as:

$$f_{dw} = \frac{\delta_{xyl} - \delta_{sw}}{\delta_{dw} - \delta_{sw}} \tag{3}$$

We show the results of the ratio (Fig. 4) with uncertainties  $Wf_{sw}$  quantified at each date following Genereux (1998):

$$Wf_{sw} = \left\{ \left[ \frac{\delta_{dw} - \delta_{xyl}}{(\delta_{dw} - \delta_{sw})^2} . W \delta_{sw} \right]^2 + \left[ \frac{\delta_{xyl} - \delta_{sw}}{(\delta_{dw} - \delta_{sw})^2} . W \delta_{gw} \right]^2 + \left[ \frac{-1}{\delta_{dw} - \delta_{sw}} . W \delta_{xyl} \right]^2 \right\}$$

$$(4)$$

where  $W\delta_{sw}$ ,  $W\delta_{dw}$ ,  $W\delta_{xyl}$  are the standard deviation for shallow water (precipitation and drainage), deep water (point D and point C or MtS and Ctr) and xylem water (value of the sampled tree at each site), respectively.

Determining the origin of xylem water (shallow or deep water) based on water isotopes is possible only when these two mixing pools are significantly distinct. Because the two pools showed little difference in autumn and spring (Fig. 3), it was not possible to unravel the origin of the xylem signal for these two seasons.

#### 3. Results and interpretations

#### 3.1. Environmental data

The two study years differed in terms of weather (Fig. 3), as the summer of 2014 ranks in the 30% wettest years (return



Fig. 3. Precipitation and evolution of isotopic signals of shallow water, deep water, and xylem water in 2014 and 2015, at a) Rustrel (Holm oak) and b) Mont Ventoux (Beech and Holm oak).



**Fig. 4.** For each species, (top) the proportion of shallow water and deep water in the xylem, (center) leaf water potential at predawn ( $\Psi_P$ ) and at midday ( $\Psi_M$ ), and (bottom) cumulative precipitation during the 20 days preceding sampling, a) Beech, b) Silver fir, c) Holm oak.

period = 3.4 years), whereas 2015 was drier than average summer conditions and is classified among the 40% driest years (return period = 2.2 years) (Fig. SI1). At Rustrel, during the 2014 growing season (May to September) the mean temperature was 19.9 °C, cumulative precipitation was 271 mm, and mean vapor pressure deficit (VPD) was 857 hPa. By contrast, in 2015 the mean temperature was 21.2 °C, cumulative precipitation was only 101 mm, and the VPD reached 1059 hPa. At Mont Ventoux the trend was similar, the mean temperature was 11.9 °C, cumulative precipitation was 432 hPa during the 2014 growing season, whereas in 2015 the mean temperature was 14 °C, cumulative precipitation was 370 mm, and the VPD reached 632 hPa.

## 3.2. Raw isotopic results: $\delta^{18}$ O vs time

The temporal variation of isotopic signals for the years 2014-2015 is presented in Fig. 3. The  $\delta^{18}$ O signal of the shallow resource is highly variable throughout the year because it is strongly affected by rainfall, which has a marked seasonal signal (Celle-Jeanton et al., 2001). The isotopic composition of rain is affected by several factors, including air temperature, the origin of atmospheric moisture, and rainfall intensity (Rozanski et al., 1992; Celle-Jeanton et al., 2004), which explains the high variability of the precipitation signal at both seasonal and rain event time scales. The isotopic composition of deep water has a more stable signal corresponding to the average precipitation signal during the recharge period. These results are consistent with those of Emblanch et al. (2003) and Lastennet (1994) on the same karst hydrosystem. Shallow water and deep water at Mont Ventoux are notably more depleted in <sup>18</sup>O than the water at Rustrel. This effect is consistent with the well-known altitudinal precipitation gradient (Rozanski et al., 1993; Lastennet, 1994; Celle, 2000).

The tree xylem water  $\delta^{18}$ O signal followed the shallow water signal during the summer of 2014. By contrast, in 2015, the xylem signal deviated from the shallow water  $\delta^{18}$ O signal and approached the deep water signal as drought increased. This means that during the summer of 2015, xylem water was a mixture of shallow and

deep water, and that trees therefore exploited the deep water reservoir.

#### 3.3. Determination of plant water sources and water stress

Significant temporal and inter-specific variations in the origin of xvlem water were observed (Fig. 4). All species of trees used comparatively deeper water in the drier summer of 2015 than in 2014. On average for all species, 9% of xylem water came from deep water in 2014 as compared to 38% in 2015. We also noticed a broad variation among species, but this observation should be interpreted with caution because it is difficult to unravel the effects due to the site and that due to species. The two species on Mont Ventoux used less ground water compared to those at Rustrel. Beech used 7% deep water on average during the 2014 summer, with a maximum of up to 13% at the end of summer and 46% in 2015 (summer average), with a maximum up to 63% at the end of summer. On average, silver fir consumed 0% of deep water in 2014 compared to 27% in 2015, and a maximum of 54% at the end of the summer of 2015. At Rustrel, holm oak consumed on average 21% deep water in 2014 and 40% in 2015 (up to 42% at the end of summer).

Leaf water potentials and cumulative precipitation corroborated the isotopic results and confirmed that the drought was considerably more severe in 2015 than in 2014. Leaf water potentials varied among species. Holm oak had the most highly variable water potentials ( $\Psi_P$  and  $\Psi_M$ ) over time, with values always lower than the other two species; this species reached  $\Psi_P < -2.5$  MPa in July 2015 due to very low rainfall. Holm oak water potentials gradually decreased during the 2015 summer period. The difference between  $\Psi_P$  and  $\Psi_M$  ( $\Delta \Psi$ ) also decreased during the same period. For beech,  $\Psi_P$  and  $\Psi_M$  potentials were relatively constant throughout the summer periods ( $-0.3 < \Psi_P < -0.5$  MPa for  $\Psi_P$  and  $-1.5 < \Psi_M < -2$  MPa until July 2015 for  $\Psi_M$ , respectively). However, during the severe drought of August 2015,  $\Psi_P$  almost doubled while  $\Psi_M$  increased. As a consequence, the  $\Delta \Psi$  amplitude dropped sharply from 1.5 to 0.8 MPa, indicating decreased transpi-

ration. The silver fir exhibited behavior distinct from the other species. Its water potentials ( $\Psi_P$  and  $\Psi_M$ ) showed a more limited variation range. When drought came,  $\Delta \Psi$  decreased regularly down to <0.5 MPa, because of increased  $\Psi_M$  and a concomitant decrease of  $\Psi_P$ .

#### 4. Discussion

The results of this paper show that when tree water stress increases, the fraction of deep water in xylem water increases. These results are consistent with those of other authors who have shown that trees take up water from deeper horizons during drought periods in a diverse range of ecosystems (e.g. Rambal (1984) and Barbeta et al. (2015) in Mediterranean forests; Le Roux et al. (1995) in savannas; Huang et al. (2011) and Liu et al. (2011) in tropical forest). Barbeta et al. (2015) showed that the groundwater proportion varies between 25 and 50%. However, they did not identify any difference between the studied species. Our study shows that the three species monitored in this study exploit the deep water pool in different ways. Silver fir exploits deep water only under severe drought conditions. Beech exploits more deep water and earlier in the summer period. Holm oak exploits deep water for most of the summer period and the deep water contribution increases regularly during drought. Such differences between species can be associated with both (i) site characteristics, and (ii) species physiology.

i) The characteristics of the study sites are similar from a geological and pedological point of view, but the climatic conditions are significantly different. The Rustrel site, where holm oak grows, receives less rain and has higher potential evapotranspiration. This climatic context makes the Rustrel site more arid (Table 1; Fig. 3), which would constrain oak trees to rely more on deep water to sustain their water demand than trees at the other site. The Ventoux site is wetter and drought occurs later in the summer period. This is probably why beech and silver fir exploit mainly shallow water until the beginning of severe drought (August 2015).

ii) Physiology can also explain the differences between species. Beech and fir can be easily compared because they have the same geological, pedological, and climatic conditions at the Ventoux site. We identify differences in how these two cohabiting species use the deep water pool. Our findings contrast with previous isotopic results on a different system (Barbeta et al., 2015) but are consistent with other studies (Evaristo et al., 2017). The difference (pvalue =  $3.10^{-4}$ ) in  $\Psi_{\rm P}$  between silver fir (-0.88 Mpa on average) and beech (-0.64 Mpa) indicate that they do not have access to the same total available water (TAW). Considering that the two species are present at the same site, we assume that these differences in TAW are due to differences in rooting depth. The systematically lower  $\Psi_P$  of silver fir means that its global rooting depth is lower. This hypothesis is consistent with isotopic results that show that silver fir takes almost no water from the deep water reserve except during severe drought (August 2015). Our results confirm the study conducted by Nourtier et al. (2014) at the same site, which attributed the high mortality of silver fir to drought. These authors highlighted with geophysics (ERT) that mortality was higher for individuals located on developed soil areas. They hypothesize that in areas where the shallow water reserve is large, trees live and grow without exploiting the deep water pool. These trees do not develop enough deep roots to exploit the deep water resource within karst vadose zone. As a result, they are more vulnerable to severe droughts that can cause massive tree mortality (Cailleret et al., 2014).

From a methodological point of view, we have shown that the combination of isotopic tracing and leaf water potential monitoring is relevant. Isotope tracing alone helps to determine the origin of water. Leaf water potential monitoring by itself makes it possible to evaluate the stress level related to water. By combining these two approaches, we have succeeded in showing that trees adapt their water uptake according to drought level. The greater the drought, the deeper the water that the trees exploit.

This study has major implications for water balance modeling in karst settings. These models generally use TSAW measured in the field in pedologic pits. In karst areas, the measured TSAW does not represent the TAW reserve for trees. In fact, the TSAW represents only the superficial part of the TAW within the soil because the depth of pedologic pits is usually limited due to high rock content. Thus, the TSAW measured in the field cannot explain tree survival (Davi and Cailleret, 2017) or the hydrodynamics of the studied karst system (Ollivier, 2019). Modelers therefore increase the TAW to simulate a deep reservoir that presents a calibration difficulty. For example, they often multiply by a factor of 2 the field-measured TSAW (via pedologic pits) (Davi and Cailleret, 2017). Ollivier et al. (2019a, 2019b) also increased TAW to better simulate the outflow simulated by a hydrogeological model of the Fontaine de Vaucluse system. These results show that trees extract water not only from the soil reservoir but also from a deeper reservoir. It remains difficult to quantify the exact contribution of this deep reservoir, but our results make it possible to qualitatively validate these assumptions. The goal of further studies will be to quantify this contribution by adding sap flow measurements or ecophysiological modeling to the current isotopic monitoring. In this way we could convert the water proportions as water volumes from each pool. These quantities would be easier to interpret and would be useful for ecophysiological and hydrogeological modeling. Another way to provide a better estimate of TAW, complementary to isotopic tracing, is to use geophysics. Geophysical tools such as magnetic resonance sounding (MRS) or electrical resistivity tomography (ERT) are promising tools for obtaining better estimates of TAW and its spatial variability at the plot scale (Carrière et al., 2019).

This paper shows that the interaction among climate, soil, and species determines deep water exploitation patterns. This observation needs to be repositioned in the climate change perspective because the impact of repeated and prolonged droughts on interactions between vegetation and groundwater resources is still poorly understood (Brolsma et al., 2010; Anderegg et al., 2013; Schäfer et al., 2014). Two extreme and opposite evolutions can be envisioned: i) droughts will cause massive dieback and defoliation of vegetation, which will limit the transpiration of vegetation cover, which can in turn increase infiltration and groundwater recharge. Or, ii) forests will adapt to drought, trees will consume less water quantitatively, but they will intercept and use a larger proportion of rainfall. In the latter case, with lower precipitation (P) and a higher ratio ET/P, groundwater recharge could decrease drastically. It is difficult to predict which scenario will occur, but it is likely that the future reality will follow an intermediate path between the two. Experiments on forest plots with rain exclusion seem to validate the second scenario (Barbeta et al., 2015). However, other studies using complementary approaches (e.g. surface based geophysics, remote sensing) need to be conducted to confirm these hypotheses at the broadest scales and in different ecological contexts. Importantly, our results suggest that deep water could constitute a much larger buffer than soil water alone. It is also essential to integrate human activities (e.g. planting new species, tree felling) to make these scenarios realistic.

# 5. Conclusions

The combination of isotope tracing and leaf water potentials in a monitoring study conducted over two summer periods has provi-

ded useful results. The two methods provide complementary results that furnish new insights into water regulation during drought conditions in karst environments. We observed a concomitant decrease in  $\Psi_{\rm P}$  and  $\Delta\Psi$  as drought increased. This indicates the presence of transpiration regulation, a primary mechanism of drought response (Maseda and Fernández, 2006). The results of this paper also show that when tree water stress increases due to drought, the fraction of deep water in the xylem increases in comparison to the shallow water resource. This increased contribution of deep water was observed during the drought period (summer) for all three species. However, each species exploits this deep water reserve in a different way. This reflects different adaptation to drought among species. We show that it is essential to take into account the deep water resource in the calculation of total available water for transpiration. Taking this deep water into account will improve water balance modeling for ecological purposes (i.e. tree resistance to drought) and hydrogeological purposes (i.e. characterization of karst recharge). The scientific community must continue to study these interactions between vegetation cover and groundwater to better predict the evolution of these two interdependent entities.

## **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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#### Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.scitotenv.2019.134332.

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