

Sap flow, leaf-level gas exchange and spectral responses to drought in *Pinus sylvestris*, *Pinus pinea* and *Pinus halepensis*

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In a climate change scenario, Mediterranean forest species such as pines may be endangered by rising temperatures and reduced precipitation, thus calling for studies on the transpiration and water balance in pines. In this paper, the response of young plants of *Pinus sylvestris* L., *Pinus pinea* L. and *Pinus halepensis* Mill. to different irrigation treatments has been studied. Significant differences were found in water potential, sap flow, leaf-level gas exchange and spectral variables. *P. sylvestris* had higher pre-dawn and midday water potentials, sap flow rates and leaf-level gas exchange rates compared to the other two species in well-watered conditions. Vapor pressure gradient correlated with stomatal conductance, net assimilation and transpiration, but the association between stomatal conductance and sap flow was weak. The environmental variables more strongly associated with sap flow were solar radiation and reference evapo-transpiration, especially in the well-watered plants, but those associations were weaker in the stressed plants. All three pine species showed the isohydric, drought-avoiding strategy common in the genus *Pinus*, maintaining relatively high water potentials in dry conditions. Nevertheless, *P. halepensis* showed a water-saving strategy, with a stomatal closure behavior under drought. Stomatal regulation was less strict in *P. sylvestris*, closer to a water-spending pattern, while *P. pinea* showed an intermediate behavior. Significant differences were recorded among species in spectral reflectance in the visible and infra-red regions. Photochemical Reflectance Index, Normalized Difference Vegetation Index and combinations of other ratios permitted the discrimination among the three pine species. These spectral variables showed association with sap flow rate, water potential and leaf-level gas exchange variables. Both cluster analysis and k-means classification discriminated Scots pine and Aleppo pine in two different groups. On the other hand, Stone pine showed differences in spectral behavior depending on the hydric status of the plants. Well-watered Stone pine plants had the same spectral behavior as Scots pine, while the plants subjected to drought stress were closer to Aleppo pine plants in spectral response. These findings may help to quantify the impacts of early and mid-summer water deficit on Mediterranean pines in future climate regimes.

Keywords: Carbon Assimilation, Aleppo Pine, Hydric Relations, Reflectance, Scots Pine, Stone Pine, Transpiration

Introduction

In a global warming scenario, many Mediterranean forest species may be endangered by a temperature rise and a decrease in precipitation. This trend has already been described and assessed for some pine

forests in North-Eastern Spain (Martínez-Vialta & Piñol 2002), where severe drought-induced mortality has occurred during recent dry episodes. Similar climate change scenarios have been described in other parts of the world, for instance in the east-

ern Mediterranean basin (Ungar et al. 2013) and in the Southern United States (Seager et al. 2007). This situation underlines the need for studies on the water balance in plants from endangered ecosystems, in particular, the regulation of the hydric sta-

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tus and transpiration of the dominant woody vegetation. In other Mediterranean and semi-arid ecosystems, hydric and gas exchange parameters at the leaf level have successfully been used as indicators of the ecophysiological competence of forest species in a changing environment (Manzanera & Martínez-Chacón 2007).

Plant resistance to drought has been hypothesized to reflect two contrasting physiological strategies: drought avoidance and drought tolerance. The mechanisms of either approach, respectively called isohydric and anisohydric, involve control of stomata and water status (Tardieu & Simonneau 1998, McDowell et al. 2008). Isohydric plants reduce stomatal conductance in dry conditions, maintaining relatively constant leaf water potential regardless of drought intensity, for instance in *Prunus* sp. (Tardieu & Simonneau 1998). In contrast, anisohydric species allow leaf water potential to decrease with water availability; there are good examples among the Cupressaceae (Martínez-Vilalta et al. 2004). Both strategies enable plant survival under drought conditions. However, each has inherent drawbacks that can be fatal in certain conditions. Prolonged reductions of stomatal gas exchange expose isohydric species to carbon depletion, while extreme water potential gradients in anisohydric species result in xylem cavitation and embolism and subsequent hydraulic failure (McDowell et al. 2008).

Scots pine (*Pinus sylvestris* L.) is one of the most widely distributed forest species in the world, extending from the Siberian taiga to mountainous ecosystems in South-western Europe, where it reaches the boundary with semi-arid Mediterranean climate and may be more at risk of aridity in a climate change scenario (Martínez-Vilalta & Piñol 2002). Stone pine (*Pinus pinea* L.) is a Mediterranean pine species, present in many south European countries, adapted to semi-arid continental and coastal locations, from the sea-level to about 1000 m altitude. Stone pine prefers sandy acidic soils with good water drainage, and a local annual precipitation between 400 and 800 mm. Aleppo pine (*Pinus halepensis* Mill.) is another characteristic species of the Mediterranean basin, with preference for basic soils, which shows more resistance to drought, being capable of living with as little as 250 mm annual precipitation. In spite of the environmental importance of these pine species, there have been few comparative studies aimed at relating the different ecophysiological responses in whole-plant transpiration and in their adaptation to drought (Melzack et al. 1985). Moreover, transpiration of whole-tree water use may be estimated by means of sap flow measurements. One of the most common methods for measuring sap flow is the Thermal Dissipation Probe (TDP) technique, which measures sap flux density in the stem xylem and can then be converted to volumetric flow rate (Granier 1985). Sev-

eral studies have related the whole plant transpiration with gas exchange variables at the plant level and with micro-meteorological factors (Oliveras et al. 2003). Furthermore, the spectral features of radiation reflected or transmitted by plants have been used to retrieve their biophysical properties (Ustin et al. 2009), and the spectral behavior of the forest canopy has been successfully linked to physiological variables, for instance detecting forest decline before damage is visible (Hernández-Clemente et al. 2011).

The spectrum in the visible region (VIS: 400-700 nm) has been associated to high absorption of the photosynthetically active radiation (PAR) by plant pigments, mainly the chlorophylls, while reflectance in the near infrared wavelengths (NIR: 700-1350 nm) is determined by the structure of the crown. Water content is the main factor affecting the reflectance in the shortwave infrared wavelengths (SWIR: 1350-2500 nm – Jacquemoud et al. 2009). Spectral methods are based on the association of ecophysiological traits with spectral signatures from which the plant status is inferred, for instance through the inversion of radiative transfer models (Jacquemoud et al. 2009), or using spectral indices (Zarco-Tejada et al. 2005). In this study, we followed the second approach. Several remotely sensed vegetation indices have been used as ecophysiological indicators (Manzanera et al. 2013). These indices are associated with plant water stress (Suárez et al. 2008) or radiation use efficiency (Garbulsky et al. 2011). However, few studies have tried to relate sap flow rates and leaf-level gas exchange variables with spectral indices. Those studies have been restricted to the agricultural sector (Marino et al. 2014) and only very few are focused on the forestry and environmental sector.

A better understanding of the ecophysiology of young plants is important for successful nursery management and for the establishment of new plantations. Therefore, our objectives were: (1) to measure daily sap flow in plants from three pine species present in different ecosystems of the Mediterranean area, namely Scots pine (*Pinus sylvestris*), stone pine (*P. pinea*) and Aleppo pine (*P. halepensis*), and to examine their relationship with environmental conditions such as water availability and evaporative demand; (2) to describe and compare gas exchange responses to different water availabilities, defining and comparing the ecophysiological strategies to face drought adopted by the three pine species; (3) to measure hyperspectral reflectance and compare the spectral behavior of all three species in relation to water availability and to ecophysiological variables such as sap flow rate, water potential and leaf-level gas exchange variables; (4) to predict ecophysiological variables by means of the plant spectral behavior; and (5) to identify possible differences between ecophysiological attributes to explore management

implications in the context of a climate change scenario of increasing aridity.

Materials and methods

Plant materials

In all experiments, three pine species were used, i.e., *Pinus halepensis*, *P. pinea* and *P. sylvestris*. All the plants were ca. 10 years old, collected from a nursery. Eight plants per species were used, four of which were subjected to a high irrigation regime and four to a low irrigation regime. Thus, a total of 12 plants received a high irrigation dose and 12 a low irrigation dose. Besides, plants also received natural rainfall during the measurement period, although the amount was negligible (25.4 mm from May 10 to September 11). All plants were transferred to 15 L plastic pots (7 dm³ area) with a mixture (1:1) of peat and soil and placed in a nursery. Plants were allowed to establish good root contact with the substrate for 15 days before the experiment started. The experiment started on May 10, and ended on September 11. For each plant, the diameter was measured at the base of the stem and used to calculate the mean cross sectional area at the base of the stem. The total leaf area (Al, m²) of each plant was estimated using the diameter as independent variable in the allometric equations developed by Montero et al. (2005) for *P. sylvestris* and *P. pinea*, and by Mitsopoulos & Dimitrakopoulos (2007) for *P. halepensis*. Plant size at the beginning of the experiment is given in Tab. S1 (Supplementary material).

Experimental design and measurements

Two irrigation doses were applied throughout the experiment duration. The high dose consisted of 7 mm day⁻¹ through an automatic dripping system. One irrigation event was given per day, at 6 a.m. solar time. The low dose was 3.5 mm day⁻¹, in the same conditions.

Sap flow was determined using the thermal dissipation probe (TDP) technique (Granier 1985). Briefly, this system consists of two thermocouple probes which are inserted into the plant sapwood, one directly above the other, at a distance of 4 cm each other. The upper probe contains an electrical heater. The thermocouples measure the temperature difference (dT) between the heated probe and the sapwood temperature below. Then, dT will decrease as the flowing sap cools the heated needle. The dT variable and the daily maximum (dTm) at zero flow are used to estimate sap flux density and therefore whole-tree transpiration. In our experiment, a TDP30 Dynamax® (Houston, TX, USA) thermal dissipation probe was inserted in the stem of each plant, at about 10 cm height from the base. We checked that the sensing zone of the probe was in full contact with the xylem of the plant. Then the area between the upper and

lower probe was isolated with styrofoam eggs and shielded with aluminium reflective bubble foil cover to minimize natural temperature gradients and provide thermal insulation and mechanical protection, according to the vendor instructions. The TDP probes were connected to a Dynamax Probe 12 biomonitor. The upper probe of each sensor was supplied permanently with a constant power of 0.15 to 0.2 W, while the lower probe was unheated for reference. Sap flow data were recorded and estimated according to Granier (1985), from May 10th to September 11th 2012. From the hourly recorded dataset, two sap flow-based variables were calculated: mean sap flow of the growing season (F , g h⁻¹ – eqn. 1) and mean sap flow velocity (v , cm s⁻¹ – eqn. 2).

$$F = As \cdot 3600 \text{ s h}^{-1} \cdot v$$

where F is the sap flow (g h⁻¹) and As is the cross-sectional area of sap conducting wood (cm²). Given the age of the plants, we considered that the sapwood area equaled the whole cross section. The sap flow velocity v (cm s⁻¹) was estimated according to the empirical formula by Granier (1985 – eqn. 2):

$$v = 0.0119 \left(\frac{dT_m - dT}{dT} \right)^{1.231}$$

where dT is the measured difference in temperature between that of the heated probe, referenced to the lower non-heated probe, and dT_m is the daily maximum dT at zero flow.

Water potential was measured in all plants with a PMS pressure bomb at dawn and at midday. Three needles were measured per plant and the average water potential was calculated per plant. Measurements were taken on June 26th, August 17th and September 24th 2012, trying to represent the hydric status in the first month, mid-season and the end of the dry period, respectively.

Leaf-level gas exchange variables, *i.e.*, net assimilation (A_n), transpiration (E), stomatal conductance (g_s), internal CO₂ concentration (C_i), water use efficiency (WUE), intrinsic water use efficiency (IWUE), leaf-to-air vapor pressure gradient (VPG) and photosynthetic photon flux density (PPFD) were recorded with a Delta T Infrared Gas Analyzer (IRGA) and a conifer chamber, in the middle (August 20th, 2012) and the end (October 3rd) of the dry season, in outdoors environmental conditions. During measurements (12:00 to 16:00 h), the average PPFD ranged from 150 to 201 $\mu\text{mol m}^{-2} \text{ s}^{-1}$. Average leaf chamber CO₂ concentration and temperature ranged from 393 to 400 ppm and from 30 to 35.6 °C, respectively. After gas exchange measurements, the projected needle area was estimated with an image analyzer, and the gas exchange variables were referred to the measured needle area.

Spectral signatures were recorded from all 24 plants with an ASD FieldSpec 3[®] (Analytical Spectral Devices Inc., Boulder, CO, USA) field spectroradiometer. Measurements were always made in clear sky conditions, on June 25th and July 2nd, 2012, at midday, operating in the spectral range between 350 and 2500 nm with an average spectral resolution of 3 nm FWHM (Full-Width-Half-Maximum) and a sampling interval of 1.4 nm across the 350-1050 nm spectral range. The FWHM and the sampling interval for the 1051-2500 nm spectral range are 30 and 2 nm, respectively. A fibre optic cable, providing a field of view edge of 25°, was connected to the spectroradiometer to collect reflectance spectra in nadir direction. Three signatures were recorded per plant from their tops, and the average was used to estimate the spectral response. Before and between every series of three plant reflectance measurements, three white references were taken for calibration, by measuring the reflectance of a 1×1 m polystyrene panel, covered with a mixture of barium sulphate powder and white paint, as white standard. The ViewSpec Pro[®] (ASD) software was used for pre-processing reflectance spectra and then reflectance indices were derived. The reflectance responses of all 24 pines used in this experiment were tested for the species and the irrigation treatment. Reflectance in the range from 350 to 2500 nm wavelength was measured, and spectral indices were calculated to track changes in pigment concentration as a function of the water stress condition: green (531 nm), yellow (570 nm), red (670 nm), different Near Infra-Red (NIR) reflectances: NIR1 (760 nm), NIR2 (895 nm), NIR3 (900 nm), NIR4 (970 nm), NIR5 (1090 nm), NIR6 (1190 nm), two Short Wave Infra-Red: SWIR1 (1650 nm), SWIR2 (2217 nm); Normalized Difference Vegetation Index: (NDVI – Rouse et al. 1974), Physiological (or Photochemical) Reflectance Index (PRI – Garbulsky et al. 2011), Normalized Difference Infrared Index (NDII – Hardisky et al. 1983); different indices based on reflectance ratios: SWIR1/SWIR2, simple ratio (SR) = NIR5/Red (Rondeaux et al. 1996), SWIR1/Red, SWIR2/Red (see Tab. S2 in Supplementary material), and the Global Environmental Monitoring Index (GEMI – Pinty & Verstraete 1992), the latter being calculated according to the formula (eqn. 3):

$$GEMI = \eta \left(1 - 0.25 \eta \right) - \frac{\text{Red} - 0.125}{1 - \text{Red}}$$

where η is (eqn. 4):

$$\eta = \frac{2 \text{NIR5}^2 - \text{Red}^2 + 1.5 \text{NIR5} + 0.5 \text{Red}}{\text{NIR5} + \text{Red} + 0.5}$$

Micrometeorological variables were recorded with a Davis Vantage Pro2[™] (Davis Instruments, CA, USA) meteorological station every five minutes, during all the experiment duration. Temperature (°C), rel-

ative air humidity (rH, %), vapour pressure deficit (VPD), solar radiation (W m⁻²), wind speed (m s⁻¹), barometric pressure (hPa), rain rate (mm h⁻¹), soil moisture (kPa) at 5 and 15 cm depth and soil temperature (°C) at 5 and 15 cm depth were averaged per hour. Rainfall (mm) and reference evapotranspiration (ET_o, mm) were accumulated per hour. Reference evapotranspiration was estimated according to the modified Penman's equation as implemented by the California Irrigation Management Information System, CIMIS, including Net Radiation calculation (Snyder & Pruitt 1985).

Statistics

Tree species, leaf-level variables, irrigation treatments and dates of measurement were compared through multifactor Analysis of Variance (ANOVA). Water potential, sap flow, leaf-level gas exchange variables and spectral indices were also tested through correlation analysis. Significant variables were estimated with regression models. The statistical packages Statgraphics Centurion XVI[®] v. 16.1.17 (StatPoint Technologies, Warrenton, VA, USA) and STATISTICA v. 7.1 (StatSoft Inc., Tulsa, OK, USA) were used. Several reflectances and spectral indices were compared through ANOVA: green (531 nm), yellow (570 nm), red (670 nm), different Near Infra-Red (NIR) wavelengths: NIR1 (760 nm), NIR5 (1090 nm), NIR6 (1190 nm), two Short Wave Infra-Red: SWIR1 (1650 nm), SWIR2 (2217 nm); Normalized Difference Vegetation Index NDVI5, Photochemical Reflectance Index (PRI), Normalized Difference Infrared Index (NDII), Global Environmental Monitoring Index (GEMI); and different ratios: SWIR1/SWIR2, NIR5/Red, SWIR1/Red, SWIR2/Red.

Results

The leaf area (Al) per plant was compared before the beginning of the experiment between the two sets of plants which were going to be subjected to the different irrigation treatments by ANOVA, to test possible differences in size which could bias the results. However, no statistically significant differences were found among plants subjected to the irrigation treatments ($p = 0.999$), among the three species ($p = 0.460$) or the interaction between species and treatment ($p = 0.467$). Therefore, we assumed that differences in the results of the experiments carried out cannot be attributed to differences in leaf area among plants (see Tab. S1 in Supplementary material). On the other hand, stone pine plants had a significantly larger ($p \leq 0.0001$) cross sectional area (As) than the other two species, due to the intrinsic anatomical traits of stone pine. As a consequence, significant differences in the leaf area to cross-sectional sapwood area ratio ($Al:As$) were found ($p = 0.026$). The $Al:As$ ratio of stone pine was significantly lower than in Scots pine but it was not significantly different from that of Aleppo pine

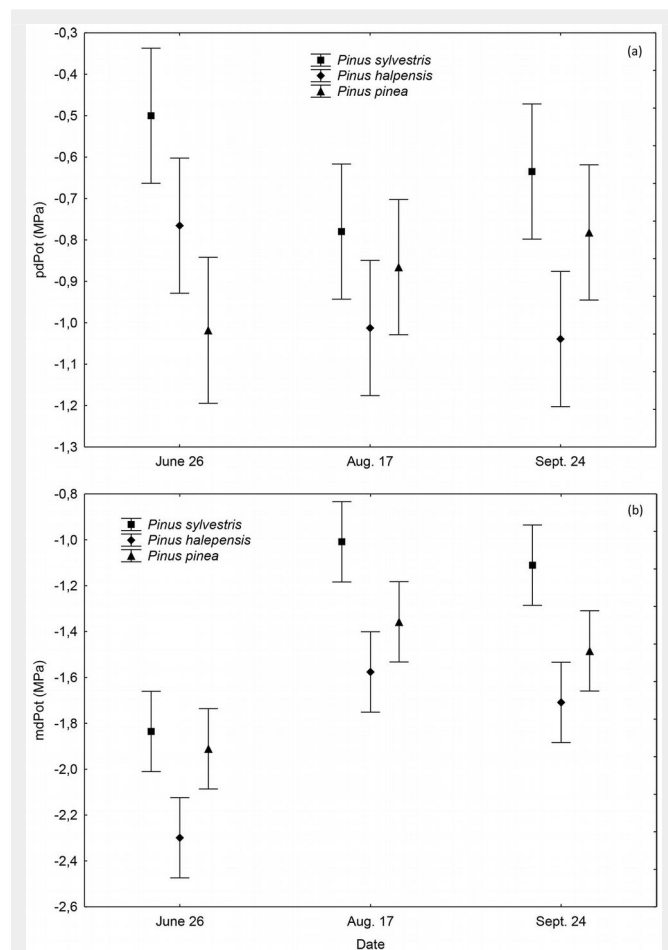


Fig. 1 - (a) Mean pre-dawn leaf water potential (pdPot, MPa) and (b) Midday water potential (mdPot, MPa) in *Pinus sylvestris*, *P. halepensis* and *P. pinea*. Plants were subjected to two irrigation regimes, high irrigation (7 mm day⁻¹) and low irrigation (3.5 mm day⁻¹) from June to September. Vertical bars denote 95% confidence intervals.

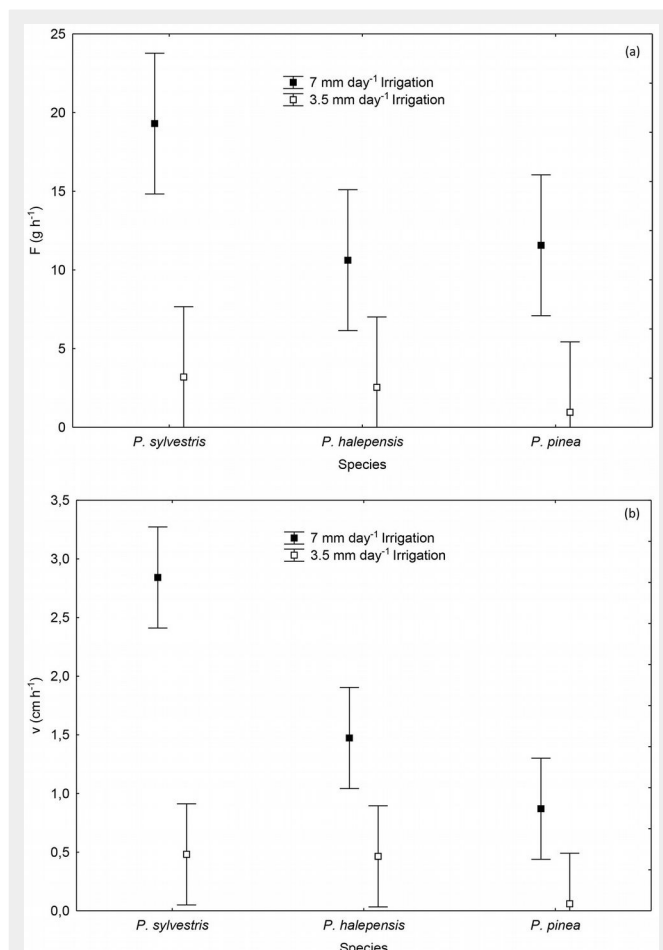


Fig. 2 - (a) Mean sap flow rate (F , g h⁻¹) and (b) mean sap-flow velocity (v , cm h⁻¹) in *Pinus sylvestris*, *P. halepensis* and *P. pinea* plants subjected to two irrigation regimes: high irrigation (7 mm day⁻¹) and low irrigation (3.5 mm day⁻¹). Vertical bars represent 95% confidence intervals.

(Tab. S1 in Supplementary material).

Observed climatic conditions over the study period (temperatures, precipitation, and air relative humidity) were compared with average conditions from the 1982-2010 period (Tab. S3 in Supplementary material), indicating that May, June and August were warmer and drier in the study period than the average, while July and September were close to the average.

Water potential

Significant differences in pre-dawn water potential were found for the interaction between species and date of measurement ($p = 0.020$). *Pinus sylvestris* showed the highest pre-dawn water potential as compared to *P. halepensis*, while *Pinus pinea* was lowest in June, and intermediate in August and September. In this period, the pre-dawn water potential was relatively stable (Fig. 1a).

On the other hand, midday water potential had statistically significant differences between species ($p \leq 0.001$) and between dates of measurement ($p \leq 0.001$ - Fig. 1b), but not between the irrigation treatments

nor for the interactions. *Pinus sylvestris* and *P. pinea* plants had the highest midday water potential and *P. halepensis* the lowest, without differences between treatments. Midday water potential was lowest on June 26th, especially for *Pinus halepensis* (-2.3 MPa) and recovered as the season advanced, in parallel with the decreasing temperature and ETo and the increasing RH (Fig. S1 in Supplementary material). *Pinus pinea* had an initial water potential (-1.91 MPa) closer to *Pinus sylvestris* (-1.84 MPa), but did not recover to the same high levels in August and September (Fig. 1b).

Sap flow measurements

Mean sap flow rate of the whole season was only significantly different among irrigation treatments ($p \leq 0.001$ - Fig. 2a) but not among species or for the interaction. On the other hand, mean sap-flow velocity showed significant differences for the irrigation treatment ($p \leq 0.001$), species ($p \leq 0.001$) and the interaction ($p = 0.003$ - Fig. 2b). As expected, sap flow rate and mean sap-flow velocity were higher in the more watered plants. *Pinus sylvestris* had higher

mean sap flow rate than the other two species with more irrigation, but the rate was reduced to approximately the same level as the other two pine species when less water was available (Fig. 2a). On the other hand, correlation analysis did not show significant relationships between water potential and sap flow either.

Leaf-level gas exchange variables

ANOVA of leaf-level gas exchange variables revealed significant differences in net assimilation rate (A_n) among species ($p = 0.044$) and dates of measurement ($p \leq 0.001$) but not for the interactions. During the adverse period of August, net assimilation was lower for all species, but as temperatures descended, *P. sylvestris* showed a better capacity of recovery by a higher assimilation rate than the other pines.

Transpiration rate (E , mmol m⁻² s⁻¹) was significantly lower in August than in October ($p \leq 0.001$) and for the interaction between species and irrigation treatment ($p = 0.035$ - Fig. S2a in Supplementary material). As for net assimilation, transpiration showed a depression in August, while in

Tab. 1 - Correlation coefficients between leaf-level gas exchange variables (An: net assimilation; E: transpiration rate; g_s : stomatal conductance; C_i : internal CO₂ concentration; VPG: vapor pressure gradient; IWUE: intrinsic water use efficiency), midday water potential (mdPot) and daily sap flow rate (Sap Flow) during August and September. Values of the second row for each variable are the significance values (prob, in italic). (*): $p \leq 0.05$; (**): $p \leq 0.01$; (***) : $p \leq 0.001$.

Parameter	E	g_s	C_i	mdPot	VPG	IWUE	Sap Flow
An	0.798 ***	0.848 ***	0.072	0.122	-0.633 ***	0.268	0.266
<i>prob</i>	<i>0.000</i>	<i>0.000</i>	<i>0.640</i>	<i>0.432</i>	<i>0.000</i>	<i>0.079</i>	<i>0.081</i>
E	-	0.897 ***	0.138	0.113	-0.574 ***	-0.089	0.209
<i>prob</i>	-	<i>0.000</i>	<i>0.373</i>	<i>0.463</i>	<i>0.000</i>	<i>0.566</i>	<i>0.174</i>
g_s	-	-	0.328 *	0.039	-0.765 ***	-0.158	0.306 *
<i>prob</i>	-	-	<i>0.030</i>	<i>0.802</i>	<i>0.000</i>	<i>0.305</i>	<i>0.043</i>
C_i	-	-	-	-0.100	-0.410 **	-0.515 ***	0.211
<i>prob</i>	-	-	-	<i>0.517</i>	<i>0.006</i>	<i>0.000</i>	<i>0.169</i>
mdPot	-	-	-	-	0.062	-0.037	0.447 **
<i>prob</i>	-	-	-	-	<i>0.691</i>	<i>0.814</i>	<i>0.002</i>

Tab. 2 - Linear correlation coefficients between meteorological variables and mean sap flow per species (*Pinus sylvestris*, *P. halepensis*, *P. pinea*) and irrigation treatment (high irrigation: 7 mm day⁻¹; low irrigation: 3.5 mm day⁻¹). (T): air temperature; (rH): relative humidity; (VPD): vapour pressure deficit; (Rad): solar radiation; (ETo): reference evapo-transpiration; (sylv-HighIrr): sap flow of *Pinus sylvestris* with high irrigation; (halep-HighIrr): sap flow of *P. halepensis* with high irrigation; (pinea-HighIrr): sap flow of *P. pinea* with high irrigation; (Sylv-LowIrr): sap flow of *Pinus sylvestris* with low irrigation; (halep-LowIrr): sap flow of *P. halepensis* with low irrigation; (pinea-LowIrr): sap flow of *P. pinea* with low irrigation. Values of the second row for each variable are the significance values (prob, in italic). (*): $p \leq 0.05$; (**): $p \leq 0.01$; (***) : $p \leq 0.001$.

Parameter	rH	VPD	Rad	ETo	sylv-HighIrr	halep-HighIrr	pinea-HighIrr	sylv-LowIrr	halep-LowIrr	pinea-LowIrr
T	-0.8197 *	0.9527 *	0.5529 *	0.6842 *	0.4874	0.3889	0.4085	0.1712	0.2635	-0.0739
<i>prob</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0001</i>
VPD	-	-	0.5446 *	0.7119 *	0.4493	0.3315	0.3553	0.0868	0.1999	-0.1176
<i>prob</i>	-	-	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>
Rad	-	-	-	0.9015 *	0.8286 *	0.7250 *	0.8068 *	0.5383 *	0.6831 *	0.1305
<i>prob</i>	-	-	-	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>
ETo	-	-	-	-	0.7248 *	0.5937 *	0.6136 *	0.3627	0.5232 *	-0.0311
<i>prob</i>	-	-	-	-	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0000</i>	<i>0.0965</i>

October *Pinus sylvestris* recovered with a higher E rate than the other species at the low irrigation treatment (Fig. S2a in Supplementary material). Stomatal conductance also showed significant differences among the dates of measurement ($p \leq 0.001$) and for the interaction between species and irrigation treatment ($p = 0.009$ – Fig. S2b in Supplementary material). Stomatal conductance followed the same pattern of An and E, as it was higher in October than in August. On the other hand, neither water use efficiency (WUE) nor intrinsic water use efficiency (IWUE) were statistically significant for any of the factors tested.

Correlations among water potential, sap flow and gas exchange variables

Correlation analysis among leaf-level gas exchange variables, water potentials and sap flow rates revealed a general good correlations between pairs of gas exchange variables, between water potentials, or between sap flow rates, but not between variables of different sets. Only An, E and g_s showed good correlations among them and with VPG, and a weak correlation was found between midday water potential and sap flow (Tab. 1). Remarkably, E did not show correlation with sap flow measurements. The best regression model for

predicting g_s using VPG as predictor variable was the square root-Y reciprocal-X model ($p \leq 0.000$, $r = 0.84$, $R^2 = 71.03\%$ – eqn. 5):

$$g_s = (-3.94985 + 30.1483 \cdot VPG^{-1})^2$$

Sap flow correlations with meteorological variables

Mean sap flow of each species and irrigation treatment were tested for correlations with meteorological data recorded during the experiment duration. Solar radiation had the strongest correlation with mean

sap flow, with differences among species (Tab. 2). The high irrigation treatment had better correlation than the low irrigation treatment. Also, ETo was moderately correlated with mean sap flow of the well-watered pines, but the correlation coefficients decreased for the less irrigated plants. Correlations of VPD with sap flow were weak (Tab. 2). As expected, some meteorological variables were strongly correlated: mean temperature (T) and relative humidity (rH), T and ETo, VPD and ETo. However, no good correlations were observed between them or between precipitation and sap flow, except for the men-

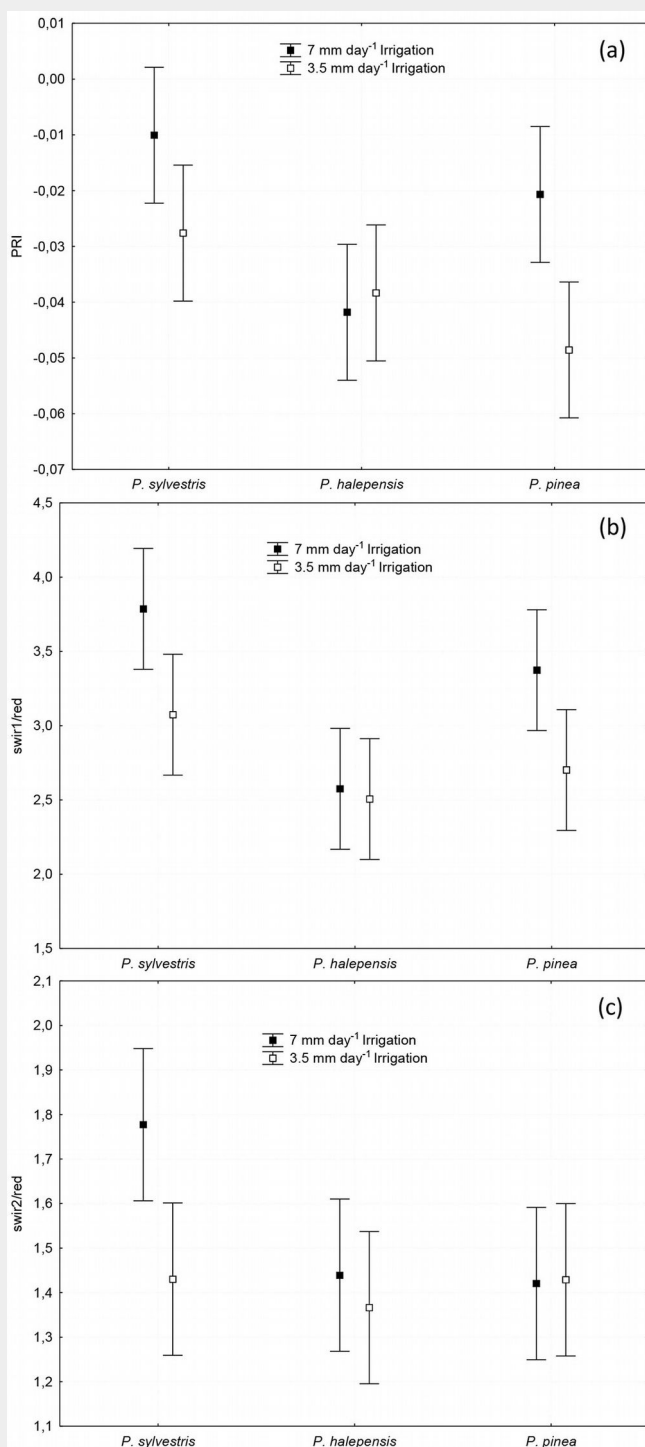
Tab. 3 - Mean sap flow rate (F, g h⁻¹) regression models, with solar radiation (W m⁻²) as the independent variable, in *Pinus sylvestris*, *Pinus halepensis* and *Pinus pinea* subjected to high irrigation (7 mm day⁻¹) or low irrigation (3.5 mm day⁻¹). All of them fitted to a double square root model, except for *P. halepensis* with low irrigation, which followed a square root-Y model. Model parameters: b, slope; a, y-intercept.

Species	Irrigation	Regression Model	r	adjusted R ²	p-value
<i>P. sylvestris</i>	7 mm day ⁻¹	eqn. 6; a = 0.55; b = 0.239	0.89	78.5%	0.000
<i>P. halepensis</i>	7 mm day ⁻¹	eqn. 6; a = 0.43; b = 0.165	0.79	62.32%	0.000
<i>P. pinea</i>	7 mm day ⁻¹	eqn. 6; a = 0.43; b = 0.182	0.86	74.09%	0.000
<i>P. sylvestris</i>	3.5 mm day ⁻¹	eqn. 6; a = 0.44; b = 0.07	0.63	39.46%	0.000
<i>P. halepensis</i>	3.5 mm day ⁻¹	eqn. 7; a = 0.38; b = 0.003	0.70	49.14%	0.000
<i>P. pinea</i>	3.5 mm day ⁻¹	eqn. 6; a = 0.54; b = 0.01	0.15	2.31%	0.000

Tab. 4 - Correlation coefficients between leaf-level gas exchange variables (An: net assimilation; E: transpiration rate; g_s : stomatal conductance; C_i : internal CO_2 concentration; VPG: vapour pressure gradient), and daily sap flow rate (Sap Flow), during September (Sep) and October (Oct). Values of the second row for each variable are the significance values (prob, in italic). (*): $p \leq 0.05$; (**): $p \leq 0.01$; (***) : $p \leq 0.001$.

Parameters	E (Oct)	g_s (Oct)	C_i (Oct)	VPG (Oct)	Sap Flow (Sep)
An (Oct)	0.891 ***	0.898 ***	-0.769 ***	-0.509 *	0.486 *
prob	0.000	0.000	0.000	0.015	0.022
E (Oct)	-	0.995 ***	-0.561 **	-0.563 **	0.303
prob	-	0.000	0.007	0.006	0.170
g_s (Oct)	-	-	-0.616 **	-0.611 **	0.360
prob	-	-	0.002	0.003	0.100
C_i (Oct)	-	-	-	0.614 **	-0.633 **
prob	-	-	-	0.002	0.002

Fig. 3 - (a): Photochemical Reflectance Index (PRI); (b): Short Wave Infra-Red (reflectance at wavelength 1650 nm; SWIR1) to Red (670 nm) ratio (SWIR1/Red); and (c): Short Wave Infra-Red (reflectance at wavelength 2217 nm; SWIR2) to Red (670 nm) ratio (SWIR2/Red), of *Pinus sylvestris*, *P. halepensis* and *P. pinea* subjected to high (7 mm day⁻¹) and low (3.5 mm day⁻¹) irrigation treatments. Vertical bars represent 95% Least Significant Difference (LSD) intervals.



tioned correlation of ETo with sap flow (Tab. 2). Significant regression models predicted sap flow responses with solar radiation as the independent variable for all species and treatments (Tab. 3). In all cases, the best fit was a double square root model (eqn. 6), except for *Pinus halepensis* subjected to the low irrigation treatment, which followed a quadratic model (eqn. 7):

$$F = (a + b \cdot Rad^{0.5})^2$$

$$F = (a + b \cdot Rad)^2$$

where a and b are the model parameters (Tab. 3).

Seasonal effect on leaf-level gas exchange and sap flow variables

Again, An, E and g_s were highly correlated ($r = 0.89$ to 0.99) while the correlation with the internal CO_2 concentration C_i was negative at the end of the growing season (Tab. 4). Sap flow rate was weakly associated with An. A positive correlation of C_i in October with VPG of the same day ($r = 0.61$) and a negative one with the sap flow rate of September ($r = -0.63$) was recorded (Tab. 4). A similar pattern has been observed for the other gas exchange variables: An, E and especially g_s show no correlation with VPG in August, but are negatively correlated with VPG in October (Tab. 4).

Analysis of spectral indices

The spectral reflectance responses of all 24 pines used in this experiment were tested for the species and the irrigation treatment (Fig. S3 in Supplementary material). The spectrum of the well-watered pines (high irrigation, 7 mm day⁻¹) was characterized by a low reflectance (< 0.12) in the visible region (VIS: 400-700 nm), while in the less irrigated pines (3.5 mm day⁻¹) this reflectance was slightly higher. Reflectance in the near infrared wavelengths (NIR: 700-1350 nm) was relatively higher (≥ 0.3). Gaps between the NIR and SWIR bands are spectral regions affected by the atmospheric moisture and these were omitted.

For green reflectance (531 nm - not shown) and yellow reflectance (570 nm - Fig. S4a in Supplementary material), significant differences were found in both wavelengths among species ($p \leq 0.000$) and irrigation doses ($p = 0.033$ for 531 nm and $p = 0.022$ for 570 nm), but not for the interaction ($p \geq 0.50$). Aleppo pine had a significantly higher green reflectance than the other pines, with slightly greater values for the lower watering. Similar results were obtained for red reflectance (670 nm), where significant differences were also found among species ($p \leq 0.000$) but not for the irrigation doses ($p = 0.06$) nor for the interaction ($p = 0.56$). Again, Aleppo pine had a significantly higher green reflectance than the other pines, without significant differences between irrigation treatments (Fig. S4b in Supplementary material).

Tab. 5 - Correlation indices and their significance level (prob, in italic) among reflectances, reflectance indices, leaf-level gas exchange variables, water potentials and sap flow-derived variables: mdPot, midday water potential; pdPot, pre-dawn water potential; Sap Flow rate; An, net assimilation rate. Variables with only non-significant correlations have been omitted for clarity. (*): $p \leq 0.05$; (**): $p \leq 0.01$; (***) : $p \leq 0.001$.

Parameter	mdPot (Jun)	Sap Flow rate	Monthly Flow (Jun)	mdPot (Aug)	Monthly Flow (Aug)	pdPot (Sep)	mdPot (Sep)	Monthly Flow (Sep)
green 531 nm	-0.573 **	-0.280	-0.283	-0.538 **	-0.353	-0.659 **	-0.623 **	-0.237
prob	<i>0.003</i>	<i>0.186</i>	<i>0.180</i>	<i>0.007</i>	<i>0.090</i>	<i>0.001</i>	<i>0.001</i>	<i>0.264</i>
yellow 570 nm	-0.587 **	-0.324	-0.329	-0.570 **	-0.400	-0.694 ***	-0.666 ***	-0.282
prob	<i>0.003</i>	<i>0.122</i>	<i>0.116</i>	<i>0.004</i>	<i>0.053</i>	<i>0.000</i>	<i>0.000</i>	<i>0.182</i>
red 670 nm	-0.523 **	-0.327	-0.343	-0.550 **	-0.396	-0.721 ***	-0.667 ***	-0.270
prob	<i>0.009</i>	<i>0.119</i>	<i>0.101</i>	<i>0.005</i>	<i>0.055</i>	<i>0.000</i>	<i>0.000</i>	<i>0.203</i>
SWIR2 2217 nm	-0.536 **	-0.210	-0.218	-0.562 **	-0.294	-0.714 ***	-0.646 **	-0.158
prob	<i>0.007</i>	<i>0.326</i>	<i>0.306</i>	<i>0.004</i>	<i>0.163</i>	<i>0.000</i>	<i>0.001</i>	<i>0.460</i>
NDVI5	0.493 *	0.294	0.319	0.513 *	0.339	0.735 ***	0.645 **	0.235
prob	<i>0.015</i>	<i>0.163</i>	<i>0.129</i>	<i>0.011</i>	<i>0.105</i>	<i>0.000</i>	<i>0.001</i>	<i>0.270</i>
PRI	0.394	0.623 **	0.642 **	0.542 **	0.672 ***	0.673 ***	0.723 ***	0.592 **
prob	<i>0.057</i>	<i>0.001</i>	<i>0.001</i>	<i>0.006</i>	<i>0.000</i>	<i>0.000</i>	<i>0.000</i>	<i>0.002</i>
NDII	0.445 *	0.157	0.175	0.458 *	0.214	0.666 ***	0.571 **	0.104
prob	<i>0.029</i>	<i>0.465</i>	<i>0.415</i>	<i>0.025</i>	<i>0.315</i>	<i>0.000</i>	<i>0.004</i>	<i>0.630</i>
swir1/red	0.380	0.583 **	0.603 **	0.434 *	0.604 **	0.645 **	0.643 **	0.541 **
prob	<i>0.067</i>	<i>0.003</i>	<i>0.002</i>	<i>0.034</i>	<i>0.002</i>	<i>0.001</i>	<i>0.001</i>	<i>0.006</i>
An (Oct)	0.345	0.444 *	0.447 *	0.475 *	0.442 *	0.517 *	0.672 ***	0.428 *
prob	<i>0.098</i>	<i>0.030</i>	<i>0.029</i>	<i>0.019</i>	<i>0.031</i>	<i>0.010</i>	<i>0.000</i>	<i>0.037</i>

For NIR reflectances, no statistical differences were found among species and water dose, except at 1190 nm wavelength, for which both the factor species ($p = 0.004$) and irrigation ($p = 0.03$) were significant. Again, *P. halepensis* showed greater reflectance with low irrigation than the other species (Fig. S4c in Supplementary material).

For SWIR1 (1650 nm) and SWIR2 (2217 nm), Aleppo pine had significantly higher reflectance ($p = 0.0012$ and $p = 0.005$, respectively) with no differences between Stone and Scots pine. On the other hand, Aleppo pine had a significantly lower NDVI5 than the other two species ($p = 0.003$).

The ANOVA of PRI showed significant differences between species ($p = 0.005$), water dose ($p = 0.009$) and the interaction of both factors ($p = 0.042$). In this case, Scots pine had a higher PRI than the other two species and the higher the water availability, the higher the PRI (Fig. 3a).

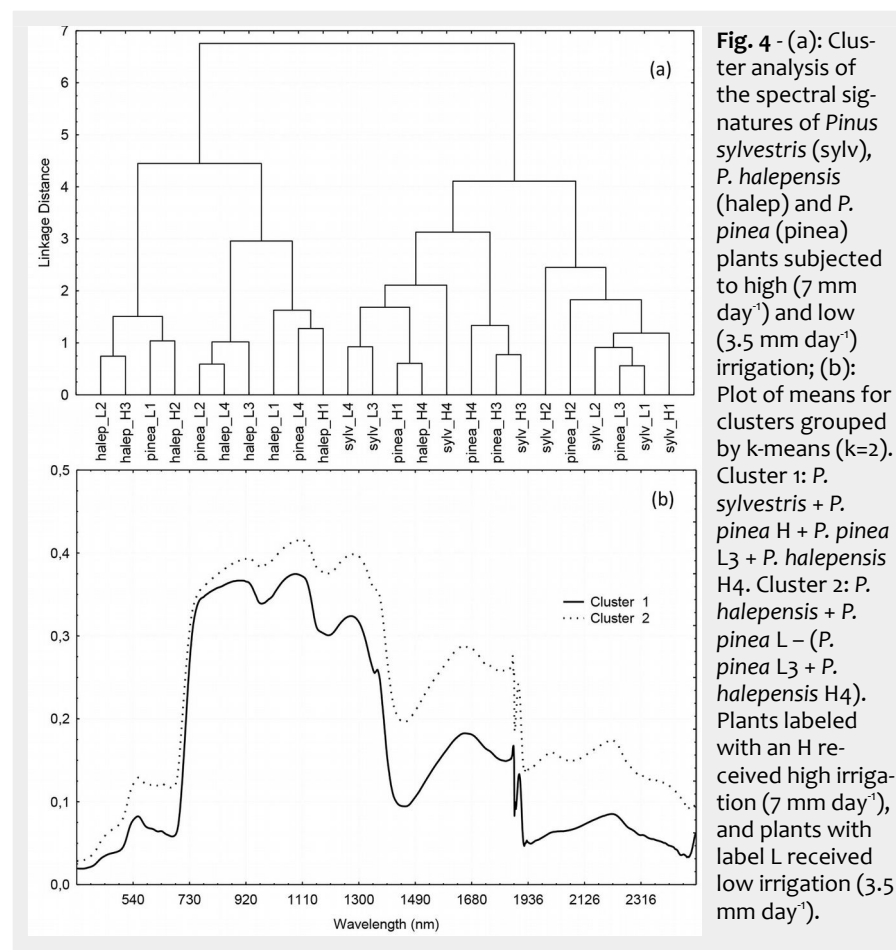
No statistical differences were found for the SWIR1/SWIR2 ratio between species, irrigation treatment nor the interaction ($p = 0.23$). On the other hand, The NDII of Aleppo pine (0.475) was significantly lower ($p = 0.0365$) than that of Scots pine (NDII = 0.599) and Stone pine (NDII = 0.633). Also, the NIR5/Red ratio index was significant for the species ($p = 0.025$), with higher values for both Stone pine (NIR5/Red = 7.219) and Scots pine (NIR5/Red = 6.849) than for Aleppo pine (NIR5/Red = 4.089).

The SWIR1/Red ratio was significant for the species ($p \leq 0.001$) and for the water dose ($p = 0.007$), but not for the interaction ($p = 0.20$ - Fig. 3b). The SWIR2/Red ratio was significant for the species ($p = 0.046$) but not for the water dose ($p = 0.054$) or for the interaction ($p = 0.10$).

Only this index has discriminated Scots pine from Stone pine with the high watering dose (Fig. 3c). On the other hand, GEMI was not significant for any factor or for their interaction ($p = 0.253$).

The multiple correlation analysis among reflectances, reflectance indices, leaf-level

gas exchange variables, water potentials and sap flow-derived variables, showed significant correlation indices (Tab. 5). Green reflectance (531 nm) showed moderate associations with midday water potential ($r = -0.573$) of June 26th, being slightly higher with the pre-dawn ($r = -0.659$) and midday



water potential ($r = -0.62$) of September. This effect could be explained by the cumulative action of drought, which was more severe at the end of the summer. Yellow reflectance (570 nm) also showed a moderate association with the midday water potential of June ($r = -0.59$ – Tab. 5), with the September water potential at dawn ($r = -0.69$) and at midday ($r = -0.66$). Slightly better correlations with pre-dawn water potential of September were found for red (reflectance at 670 nm, $r = -0.72$) SWIR2 ($r = -0.71$) and for NDVI5 ($r = 0.73$). For midday water potential in September, correlations were -0.67 for red (670 nm), -0.65 for SWIR2 and 0.645 for NDVI5 (Tab. 5).

PRI showed moderately good correlations with the total mean sap flow rate ($r = 0.62$), the monthly sap flow in June ($r = 0.64$), the sap flow in August ($r = 0.67$), and also with the pre-dawn ($r = 0.67$) and midday ($r = 0.72$) water potentials of September (Tab. 5). The SWIR1/red ratio did not provide higher correlations than PRI for the same variables. On the other hand, no good correlations were found for SWIR2/red, nor in general for other variables, such as E , g_s , WUE, IWUE and PPF (data were omitted from Tab. 5 for the sake of clarity and simplification).

Multivariate analysis of spectral signatures

Spectral signatures of all 24 plants were grouped by cluster analysis. Different linkage rules were compared: single linkage, complete linkage, unweighted and weighted pair-group average, unweighted pair-group centroid, weighted pair-group centroid, median and Ward's method. The best results were provided by the complete linkage rule, which discriminated two groups: one integrated by Aleppo pine plants (except for Aleppo pine plant H4) and Stone pine plants of the low irrigation treatment (except for Stone pine plant L3), and the other one formed by Scots pine plants, Stone pine plants of the high irrigation treatment and the two exceptions mentioned (Fig. 4a). Interestingly, there was no clear discrimination among Aleppo pine plants of either watering regime in the first group or among Scots pine plants of either irrigation treatments in the second group.

Similar results were obtained when spectral signatures were grouped by k-means (Fig. 4b). The average spectral signature of the cluster composed of Scots pine and highly watered Stone pine had a lower reflectance than the Aleppo pine and less watered Stone pine, for the spectral range of the study (from 350 to 2500 nm). This pattern difference was highest in the SWIR range, though it is also apparent in the visible green and red, and in the NIR wavelengths (Fig. 4b).

Discussion

Ecophysiological responses of young plants of three Mediterranean pine species (*Pinus sylvestris*, *P. pinea* and *P. halepensis*)

to low water availability have been investigated. We observed that the midday water potential was lower on June 26th than on August 17th and on September 24th (Fig. 1b). This rise of the midday water potential that occurred between June and August suggests that pines closed stomata when aridity reaches a threshold. Similar results were obtained by Oliveras et al. (2003) in adult individuals of *P. halepensis*, which also showed a more pronounced decrease in midday water potential as compared with *P. pinea*, while pre-dawn water potentials were not significantly different. These results were also in agreement with those reported for minimum needle water potential measured in severe drought field conditions, -2.6 MPa for *P. halepensis*, ca. -2.35 for *P. pinea*, and ca. -2.25 MPa for *P. sylvestris* (Martínez-Vilalta et al. 2004). Also, the water potentials for a 50% loss of xylem conductivity were, respectively, -3.1 MPa, -3.65 MPa and -3.57 MPa (Martínez-Vilalta et al. 2004). The low overall variation of minimum water potential as a general rule in pine species implies that structural and physiological adjustments take place to maintain water potentials within relatively narrow limits, in agreement with an isohydric behavior. These results suggest a homeostatic behavior in pines to avoid extremely negative xylem water potentials and to prevent embolism by closing stomata and/or reducing leaf area (Martínez-Vilalta et al. 2004). Our results corroborate those considerations, with similar water potential values, though not so excessively low, indicating a moderate water stress in the low irrigated plants (Fig. 1b).

Pinus sylvestris had a higher mean sap flow rate than the other two species with more irrigation but the rate was reduced to approximately the same level as the other two pine species when less water was available. This behavior, again, may be explained either by a superior stomatal conductance or by a more efficient water transport capacity. Indeed, it has been reported that *Pinus sylvestris* shows a higher wood-specific hydraulic conductivity than *P. halepensis* and *P. pinea* (Martínez-Vilalta et al. 2004). Our observations partly confirmed that *P. sylvestris* plants had higher Al:As ratio, which is related to hydraulic conductivity, than *P. pinea*, but not significantly different from that of *P. halepensis*. On the other hand, Martínez-Vilalta & Piñol (2002) found no differences among three pine species (*Pinus sylvestris*, *P. nigra* Arn. and *P. pinaster* Ait.) and their populations of NE Spain in maximum water transport capacity or vulnerability to embolism, corroborating that this homogeneity in hydraulic capacity is quite general in pines (Martínez-Vilalta et al. 2004). In *Pinus canariensis* Sweet, transpiration estimated by sap flow responded to canopy conductance, which was found to decline significantly with increasing evaporative demand and thus significantly reduced tree water loss, and this decrease was more pro-

nounced during the soil drought (Brito et al. 2014). These considerations suggest that other mechanisms, i.e., stomatal regulation, may play a major role in adaptation and acclimatization to drought stress in pine species (Irvine et al. 1998).

During the adverse period of August, net assimilation was lower for all species, but as temperatures decreased, *Pinus sylvestris* showed a better capacity of recovery by a higher assimilation rate than the other pines. In contrast, in an experiment with *Pinus halepensis*, Klein et al. (2011) found slow rates of CO_2 emission in trees under the highest drought intensity, i.e., $\approx 0.2 \mu\text{mol m}^{-2} \text{s}^{-1}$ at midday and $0.4 \mu\text{mol m}^{-2} \text{s}^{-1}$ in the late evening in the leaves of the less irrigated plants. Those plants later died, indicating that the CO_2 uptake threshold had been surpassed and carbohydrate reserves were depleted (Klein et al. 2011). Similarly in *P. sylvestris*, CO_2 depletion due to leaf mass reduction together with the tight stomatal closure in extremely dry conditions may lead to plant death, in spite of hydraulic adjustments that may allow short-term survival, for long-term resilience in response to drought is jeopardized (Poyatos et al. 2013). In our case, even in the driest period (August), average A_n was $0.33 \mu\text{mol m}^{-2} \text{s}^{-1}$ for *P. pinea*, 1.24 for *P. sylvestris*, and 1.49 for *P. halepensis*, indicating a non-extreme water limitation. In agreement with our results, Flexas et al. (2014) reported positive A_n at a lower leaf water potential in *P. halepensis* than in *P. pinea*. Stomatal conductance followed the same pattern of A_n and E , as it was higher in October than in August. The interaction results indicate that *P. halepensis* regulates stomata better than the other pines tested, limiting conductance with water shortage in greater measure than *Pinus pinea* and *P. sylvestris*. Leaf stomatal conductance also decreased in response to drought, in *P. halepensis* near the Negev desert (Klein et al. 2011). These authors also observed that g_s fell as midday leaf water potential declined and at -2.8 MPa conductance approached zero as a result of 99% stomatal closure. This tight regulation in *P. halepensis* is a characteristic of isohydric, water saving plant species, while *P. pinea* follows an intermediate pattern and *P. sylvestris* shows a tendency closer to the isohydric, water spending pattern, as indicated by the higher transpiration rates maintained by *P. sylvestris* with low water availability. On the other hand, this lower stomatal sensitivity of *Pinus sylvestris* may be in benefit of an increase in growth capacity. A_n , E and g_s showed good correlations among them, and of g_s with VPG ($r = -0.76$ – Tab. 1). The model response of g_s to VPG described by Eamus & Shanahan (2002) resembles our model in eqn. 5. Similar patterns of decreasing stomatal conductance with increasing VPG, while the correlation of net assimilation and transpiration with VPG are not so strong, have been described in *Pseudotsuga menziesii*

(Mirb.) Franco (Meinzer 1982), supporting the hypothesis that stomata respond to the water vapor demand favoring CO₂ uptake with respect to water loss.

Similar results of the relationship between solar radiation and sap flow were obtained in *Pinus sylvestris* (Gielen et al. 2010) and also in *Pinus cembra* L. of the Austrian Alps timberline, although VPG was the variable that best correlated with sap flow density (Wieser et al. 2014). A strong association was also found between sap flow-estimated transpiration and potential or reference evapo-transpiration in a *Pinus halepensis* stand with high soil water content, but this relationship was weaker with low water availability (Ungar et al. 2013). These results resemble our finding that sap flow correlated with ETo in the well irrigated plants, but the correlation coefficients were lower for all three species in the low irrigation treatment. Indeed, transpiration (estimated by sap flow) depends on both water availability and evaporative demand (ETo), and when water is a limiting factor, transpiration cannot keep pace with the ETo increase, as a probable consequence of stomatal closure.

Several authors have described pines, especially *P. halepensis*, as very sensitive to water deficit, showing a strong stomatal control under drought conditions (Baquedano & Castillo 2006), in agreement with our observations. This adaptation mechanism of *P. halepensis* by means of extremely conservative water use (water-saver), contribute in this species to high drought resistance. On the other hand, *Pinus halepensis* is highly vulnerable to embolism, with a narrow safety margin between stomatal closure and xylem dysfunction as small as 0.3 MPa (Klein et al. 2011). It has been hypothesized that *P. halepensis* has narrow tracheids and low hydraulic conductance to adjust to low water availability (Klein et al. 2011). This is probably why we observed lower sap flow and stomatal conductance rates than in the other species. Also, it has been reported that *P. halepensis* leaves are adapted to optimize stomatal conductance for avoiding carbon starvation as much as possible, resulting in leaf water potential approaching the danger zone for xylem collapse. That failure was usually avoided due to rapid stomatal responses. Therefore, *P. halepensis* is more efficient than most tree species in terms of using the carrying capacity of the xylem (Klein et al. 2011).

We also tested the relationships of leaf spectral reflectance with whole-plant transpiration, water relations and leaf-level gas exchange variables, finding that PRI was a better predictor of sap flow rate, i.e., whole plant transpiration, and of water potential than other spectral indices (Tab. 5). Although the PRI index was first proposed as a good indicator of photosynthetic activity and dissipation of excess energy not absorbed in the photochemical process (Garbulsky et al. 2011), we did not

observe a statistical relationship of PRI with net assimilation. Inter-specific differences in xanthophyll content may affect the role of PRI as indicator in drought stress conditions (Filella et al. 2004). Nevertheless, we found a significant relationship between PRI and the water status associated to the irrigation treatment, in agreement with the findings of Ripullone et al. (2011). Similar results were obtained in *Olea europaea*, where trees subjected to irrigation treatment had a higher PRI as compared to control rain-fed olive trees (Marino et al. 2014). Also significant differences in PRI value have been observed when comparing several species, indicating that PRI is species-specific (Ripullone et al. 2011). In *Quercus cocCIFera* it was concluded that, contrary to early reports, PRI is a sensible, indirect, non-destructive water stress indicator, even in plants experiencing intense drought (Peguero-Pina et al. 2008), confirming our finding. Our results also are in agreement with those observed in *Pinus sylvestris* and *P. nigra* afforestations, where PRI showed a moderately good association with water potential and stomatal conductance (Hernández-Clemente et al. 2011), and in *Olea europaea* trees, which also showed a linear relationship between PRI and sap flux density (Marino et al. 2014), though the association was stronger in our case (Tab. 5). A good correlation between PRI and pre-dawn water potential has been described in the short term, in a pool of forest species, i.e., *Arbutus unedo*, several *Quercus* species, *Castanea sativa*, *Fagus sylvatica* and *Populus x euroamericana* (Ripullone et al. 2011).

Characterization of ecosystem functioning and plant ecophysiology by means of associated spectral indices has been a major objective of remote sensing (Garbulsky et al. 2011). We identified simple spectral indices, such as PRI, NDVI, NDII, SR, SWIR/red and specific reflectances, for the discrimination of three dominant *Pinus* species present in many temperate and semi-arid Mediterranean ecosystems. We also found relationships between those spectral indices and ecophysiological features of the mentioned species, as tools opening possibilities for large-scale environmental studies. Previous studies reported the use of NIR and SWIR-based spectral indices associated with leaf water potential in *Pinus contorta* and *P. coulteri* (Cohen 1991). Close results were obtained in *Olea europaea* trees, also showing a good relationship between NDVI and water potential, for which R² = 0.668 was reported (Marino et al. 2014), similar to our results (Tab. 5). In contrast, Dziki et al. (2011) did not find significant correlations between water potential and various combinations of spectral reflectances, but the use of the first order spectral derivatives improved their results in orange trees.

Conclusions

In conclusion, ecophysiological responses

of young plants of three Mediterranean pine species (*Pinus sylvestris*, *P. pinea* and *P. halepensis*) to water availability have been investigated. Two irrigation treatments were compared: a high water dose (7 mm day⁻¹) and a low dose (3.5 mm day⁻¹). Significant differences have been recorded in water potential, sap flow and leaf-level gas exchange variables and in their interactions. Our results revealed different responses to low water availability among the three pine species studied. All species showed the isohydric, drought-avoiding strategy which is common in the genus *Pinus*, maintaining relatively high water potentials. Nevertheless, *P. halepensis* showed a strong water-saving pattern, having a tight stomatal closure with drought, even at the expense of affecting CO₂ uptake. On the other hand, stomatal regulation is less strict in *Pinus sylvestris*, closer to a water-spending behavior. *Pinus pinea* shows an intermediate behavior between the other two species. We may conclude that *Pinus sylvestris* may be endangered in a climate change context of more aridity in Mediterranean mountains. The search for inter- and intra-population variability, to explore more drought-tolerant individuals, more adapted ecotypes or other species should be considered in reforestation strategies. Alternatively, thinning may improve drought recovery and should be considered in forest management to improve drought tolerance in endangered forest species.

The spectral reflectance indices tested in this study may be valuable for fast, non-intrusive detection of water stress. We propose remotely sensed reflectance indices, in particular PRI, as promising predictive tools of the impact of drought on photosynthetic activity, water status and whole-plant transpiration.

Future applications of these techniques could be forest cover classifications in areas where *Pinus sylvestris*, *P. pinea* and *P. halepensis* are sympatric and need to be discriminated among them. We could discriminate the spectral signatures of *P. sylvestris* and *P. halepensis* by cluster analysis, and conclude that both species may be distinguished by hyperspectral sensors. On the other hand, *P. pinea* is intermediate between the other two species, and the hydric status of this pine can mimic the other two spectral patterns. With a high hydric availability, *P. pinea* spectral signature cannot be differentiated from that of *P. sylvestris*, and with less water, can be confused with *P. halepensis*. Presumably, in forested areas where *P. halepensis* and *P. pinea* are sympatric, the best moment for an adequate discrimination of the areas occupied by both species by hyperspectral imagery would be during the moist season. The findings of this experiment may help to quantify the impacts of early and mid-summer water deficit on Mediterranean pines and evaluate their potential responses to future climate regimes.

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This work is the result of collaboration in the Moncloa Campus of International Excellence. EM carried out gas exchange measurements, MR and JM carried out the spectro-radiometry measurements, JZ was in charge of the nursery and laboratory measurements, EMF and JM performed the statistical analysis, AGG, BP and AGA conceived the study and JM helped to draft the manuscript. This work was supported by the Spanish Ministry of Industry, Project FIT330221200610 and Ministry of Education and Science, Project AGL-2010-17505. We thank Prof. Alicia Palacios and Victor Cicuendez for their scientific advice. We also thank Prof. J.C. Robredo for providing the meteorological data.

Conflict of interests

The authors declare no conflict of interest.

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Supplementary Material

Tab. S1 - Experimental design and plant size at the beginning of the experiment of the pines used in the experiment.

Tab. S2 - Vegetation indices tested in *Pinus halepensis*, *P. pinea* and *P. sylvestris* plants subjected to irrigation treatments.

Tab. S3 - Meteorological data summary of the test period as compared to the 1981-2010 period.

Fig. S1 - Daily mean meteorological data registered from June to September.

Fig. S2 - Transpiration rate and stomatal conductance of *Pinus sylvestris*, *P. halepensis* and *P. pinea* subjected to two irrigation regimes, measured in August 21 and October 3.

Fig. S3 - Mean spectral signatures of *P. sylvestris*, *P. halepensis* and *P. pinea* plants subjected to high and low irrigation.

Fig. S4 - Visible and near-infrared spectral reflectances of *P. sylvestris*, *P. halepensis* and *P. pinea*.

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