A simplified framework for fast and reliable measurement of leaf turgor loss point

Francesco Petruzzellis\textsuperscript{a,}*, Tadeja Savi\textsuperscript{b}, Giovanni Bacaro\textsuperscript{a}, Andrea Nardini\textsuperscript{a}

\textsuperscript{a} Dipartimento di Scienze della Vita, Università degli Studi di Trieste, via L. Giorgieri 10, 34127, Trieste, Italy
\textsuperscript{b} University of Natural Resources and Life Sciences, Vienna, Department of Crop Sciences, Division of Viticulture and Pomology, Konrad Lorenz Straße 24, A-3430, Tulln, Austria

\textbf{A R T I C L E I N F O}

\textbf{Keywords:}
Dewpoint hygrometer
Mechanistic traits
Osmotic potential
Water availability
Water potential

\textbf{A B S T R A C T}

Drought tolerance shapes the distribution of plant species, and it is mainly determined by the osmotic potential at full turgor (\(\pi_0\)) and the water potential at turgor loss point (\(\Psi_{tlp}\)). We provide a simplified framework for \(\pi_0\) and \(\Psi_{tlp}\) measurements based on osmometer determination of \(\pi_0\) (\(\pi_{0\_osm}\)). Specifically, we ran regression models to i) improve the predictive power of the estimation of \(\pi_0\) from \(\pi_{0\_osm}\) and morpho-anatomical traits; ii) obtain the most accurate model to predict \(\Psi_{tlp}\) on the basis of the global relationship between \(\pi_0\) and \(\Psi_{tlp}\). The inclusion of the leaf dry matter content (LDMC), an easy-to-measure trait, in the regression model improved the predictive power of the estimation of \(\pi_0\) from \(\pi_{0\_osm}\). When \(\pi_{0\_osm}\) was used as a simple predictor of \(\Psi_{tlp}\), discrepancies arose in comparison with global relationship between \(\pi_0\) and \(\Psi_{tlp}\). \(\Psi_{tlp}\) values calculated as a function of the \(\pi_0\) derived from \(\pi_{0\_osm}\) and LDMC (\(\pi_{0\_osm}\) and LDMC) were consistent with the global relationship between \(\pi_0\) and \(\Psi_{tlp}\). The simplified framework provided here could encourage the inclusion of mechanistically sound drought tolerance traits in ecological studies.

\textbf{1. Introduction}

Plant functional traits are defined as morphological, physiological, or phenological features measurable at the individual level, from the cell to the whole-organism (Violle et al., 2007). Recently, Brodribb (2017) suggested to distinguish “mechanistic” traits, which comprehends plant’s features clearly associated to a physiological process, from general functional traits (such as leaf mass per unit area), which rather represent “syndromes” that could be driven by several different physiological functions and associated trade-offs. Mechanistic traits have been increasingly included in trait-based studies and provided novel insights into several ecological processes, ranging from species assembly rules (Blackman et al., 2012; Brodribb et al., 2014), invasion of alien plant species (Petruzzellis et al., 2018), and vegetation dynamics under ongoing climate changes (Anderegg, 2015). As an example, hydraulic traits (e.g. \(\Psi_{50}\), the water potential inducing 50% loss of hydraulic conductivity, or \(K_s\), the stem specific hydraulic conductivity) have been used to model plant species distribution (Costa-Saura et al., 2016; Larter et al., 2017), and they were shown to correlate with growth rate and risk of mortality under drought (Anderegg et al., 2015; Choat et al., 2012; Fan et al., 2012).

Leaf water relation parameters have been recently proposed as predictors of the position of a species along the “fast-slow” whole plant economic spectrum (Blackman, 2018; Zhu et al., 2018), as they correlate to both leaf hydraulic and economic traits (Nardini and Luglio, 2014; Trifilò et al., 2016). Specifically, leaf osmotic potential at full turgor (\(\pi_0\)) and leaf water potential at turgor loss point (\(\Psi_{tlp}\)) are strongly linked to species-specific ability to tolerate leaf dehydration (Bartlett et al., 2012b) and consequently to sustain stomatal conductance, photosynthesis and growth even under water shortage conditions (Bartlett et al., 2016; Tognetti et al., 2000). In particular, \(\Psi_{tlp}\) indicates the water potential inducing loss of cell turgor pressure (Bartlett et al., 2012b), which is critical to maintain gas exchange and growth (Brodribb et al., 2003). In their recent analysis, Zhu et al. (2018) have reported that \(\Psi_{tlp}\) is correlated with leaf carbon investment, as species with lower \(\Psi_{tlp}\) tend to have higher leaf density (\(d_{leaf}\)) and leaf mass per unit area (LMA). Turgor loss point also correlates with habitat moisture, as species living in arid environments usually have lower values of \(\Psi_{tlp}\) than species living under higher water availability (Bartlett et al., 2012b; Lenz et al., 2006; Zhu et al., 2018). Given the correlation between \(\Psi_{tlp}\), hydraulic and economic traits and environmental features, the inclusion of the turgor loss point in ecological studies holds promises to provide important insights on ecological and evolutionary patterns in plants.

\* Corresponding author.
\textit{E-mail address:} fpetruzzellis@units.it (F. Petruzzellis).

https://doi.org/10.1016/j.plaphy.2019.03.043
Received 9 January 2019; Received in revised form 25 March 2019; Accepted 28 March 2019
Available online 03 April 2019
0981-9428/ © 2019 Elsevier Masson SAS. All rights reserved.
Ψ_{tlp} has been traditionally estimated from water potential isotherms (or pv-curves, Tyree and Hammel, 1972), i.e. by measuring the progressive decrease of the water potential and of the water content during leaf dehydration. This procedure is time-consuming, and this probably limited the inclusion of Ψ_{tlp} in studies involving large numbers of species/individuals and/or study sites. Recently, Bartlett et al. (2012b) have reported that the variation of Ψ_{tlp} both between and within species is mainly driven by changes in π₀, which reflects solute concentration in cells at full turgor. These two traits resulted highly correlated to each other, as species with lower Ψ_{tlp} also have lower values of π₀ (Bartlett et al., 2012b). Consequently, both traits could be considered as useful parameters to predict species drought tolerance, and π₀ could be used as a proxy of turgor loss point. Alternative methods have been proposed to obtain π₀, e.g. by directly measuring the osmotic potential of sap extracted from leaf tissues using a thermocouple psychrometer. In particular, rapid freeze and thaw of leaf samples, that induces cell disruption and the release of symplastic contents, is considered the most accurate procedure to measure π₀ with an osmometer (π₀_osm). Recently, Bartlett et al. (2012a) proposed a framework to predict both π₀_pv (osmotic potential at full turgor derived from pv-curves) and Ψ_{tlp} from π₀_osm measurements. In their analysis, the authors tested various models including different morpho-anatomical leaf traits, and they reported that models including bulk modulus of elasticity (ε) and d_leaf significantly improved the ability to predict π₀_pv from π₀_osm measurements. However, ε is generally derived from pv-curves, so that including this parameter in the derivation of π₀_pv does not represent a major advantage. Also, d_leaf measurements can be laborious and prone to errors as far as volume estimates are concerned. Hence, a simplified framework for estimation of π₀ would be useful for ecological studies.

In this study, we measured several functional traits as well as water relation parameters derived from pv-curves in 27 species, with the aim to provide a simple framework to estimate π₀_pv and Ψ_{tlp} from measurements of π₀_osm obtained with a dewpoint psychrometer. The specific aims were to i) obtain a model to predict π₀_pv on the basis of π₀_osm and easy-to-measure functional traits (like LMA or leaf dry matter content); ii) obtain the most accurate model to predict Ψ_{tlp} on the basis of the global relationship between π₀ and Ψ_{tlp}.

2. Materials and methods

2.1. Leaf traits measurements

To model the estimation of the osmotic potential at full turgor (π₀) and of the water potential at turgor loss point (Ψ_{tlp}) from π₀ values obtained with a dewpoint psychrometer (π₀_osm; see below), we selected 27 temperate and Mediterranean woody species (Table S1) with different levels of drought resistance. Species were sampled in natural habitats in the Karst region (NE Italy), or in the Botanical Garden of University of Trieste. Additional data were obtained from previous studies performed in our laboratory (see Table S1, Nardini et al., 2012; Savi et al., 2016a; Savi et al., 2016b; Savi et al., 2016c; Savi et al., 2017).

Three leaves were sampled from different individuals of each species to measure π₀_osm. Twigs were detached from branches and were rehydrated overnight. One leaf per twig was roughly crumpled up and sealed in cling film. Then, it was immersed in liquid nitrogen (LN₂) for 2 min. The leaf (still sealed in cling film) was then carefully ground and stored in sealed plastic bottles at −20 °C. Before measurements, samples were thawed at room temperature for 5 min while still sealed in cling film and in plastic bottles, to avoid evaporation effect on measurements (Bartlett et al., 2012a). Finally, π₀_osm was measured with a dewpoint psychrometer (Model WP4, Decagon Devices Inc., Pullman, Washington, USA).

Water potential isotherms (pv-curves) were measured to obtain reference values for Ψ_{tlp} and π₀_pv. Fresh leaves were rehydrated for 12 h with their petioles immersed in pure water and pv-curves were measured using the bench dehydration technique, by repeatedly measuring water loss and water potential with a balance and a pressure chamber (model 1505D, PMS Instruments, Albany, OR, USA), respectively, during progressive sample dehydration. Water potential (Ψ_{water}) and cumulative weight loss (WI) of leaves were measured until the relationship between 1/Ψ and WI became strictly linear, indicating the loss of cell turgor. Pv-curves were then elaborated according to Salerno (1983) to calculate the osmotic potential at full turgor (π₀_pv), the water potential at turgor loss point (Ψ_{tlp_pv}) and the modulus of elasticity (ε).

For each species, leaf morpho-anatomical parameters were measured on 6 leaves from the same individuals sampled for the measurements of π₀_osm and pv-curves. Specifically, we measured leaf thickness (Th, μm), leaf dry matter content (LDMC, mg g⁻¹), leaf mass per unit area (LMA, mg cm⁻²) and leaf density (d_leaf, g cm⁻³).

Th was measured after rehydrating leaves for 3 h using a digital calliper on three portions of the leaf (top, middle, bottom). Values were then averaged for each leaf.

LDMC and LMA were calculated as:

\[ \text{LDMC} = \frac{\text{Leaf dry weight}}{\text{Leaf turgid weight}} \]

\[ \text{LMA} = \frac{\text{Leaf dry weight}}{\text{Leaf area}} \]

Fresh leaves were first rehydrated for 3 h and leaf turgid weight was measured with an analytical balance. Leaves were scanned and leaf area was measured using the software Image J (Schneider et al., 2012). Leaves were finally oven-dried for 48 h at 72 °C and leaf dry weight was measured.

\[ d_{\text{leaf}} = \frac{\text{Leaf dry weight}}{\text{Leaf fresh volume}} \]

Leaf fresh volume was measured using a water displacement method (Hughes, 2005), while leaf dry weight was measured as described above.

2.2. Statistical analysis

The first aim of this study was to find an easy and fast method to measure the leaf osmotic potential at full turgor, based on the work of Bartlett et al. (2012a). To improve the predictive power of the estimation of π₀_pv from π₀_osm measurements, we ran a multiple linear regression model to predict π₀_pv (response variable) as a function of π₀_osm and the leaf traits described above. A Minimum Adequate Model (MAM) was obtained using package “glmmulti” (Calcagno, 2013) via minimization of the corrected Akaike information criterion (AICc) plus a backward procedure to avoid multicollinearity among selected explanatory variables. To compare results obtained by Bartlett et al. (2012a), we evaluated two other linear regression models, setting π₀_osm as the response variable. In a first model, only π₀_osm was set as the predictive variable, while in the second we considered both π₀_osm and ε as predictive variables. Models were then compared on the basis of their predictive power (adjusted R², R²_adj) and, in order to take in account the number of predictors included in the model, the AICc and their mean absolute error (MAE).

The second aim of this study was to test the ability to predict Ψ_{tlp_pv} both from π₀_osm measurements and the fitted values of the MAM (π₀,π₁) and compare models’ output with the global relationship between Ψ_{tlp} and π₀ described in Bartlett et al. (2012b). We first fitted two separated simple linear regression models on the data measured in the present study, considering Ψ_{tlp_pv} (response variable) as a function of π₀_osm or π₀_fresh, respectively. For each model, we calculated coefficient estimates and associated 95% confidence intervals (95% C.I.), R²_adj, AICc and MAE.

Because the number of species analyzed in Bartlett et al. (2012b) differed from the present study (248 and 27 respectively), we set up a bootstrap procedure (999 replicates) to obtain comparable values of
Table 1
Summary of the models predicting the osmotic potential at full turgor measured through pv-curves (\(\pi_{0_{pv}}\)) from osmotic potential at full turgor obtained with a dewpoint hygrometer (\(\pi_{0_{osm}}\)) alone, including leaf dry matter content (LDMC) or the modulus of elasticity of cell walls (\(e\)). \(R_{adj}^2\) = adjusted \(R^2\), AICc = Akaike informative criterion corrected for low number of observations. MAE = mean absolute error of the model.

<table>
<thead>
<tr>
<th>(\pi_{0_{pv}}) estimation</th>
<th>Estimate</th>
<th>Std. error</th>
<th>p-value</th>
<th>(R_{adj}^2)</th>
<th>AICc</th>
<th>MAE</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\pi_{0_{osm}} + \text{intercept})</td>
<td>(\beta)</td>
<td>0.681</td>
<td>0.143</td>
<td>&lt; 0.001</td>
<td>0.46</td>
<td>17.8</td>
</tr>
<tr>
<td></td>
<td>intercept</td>
<td>-0.434</td>
<td>0.240</td>
<td>0.08</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(\pi_{0_{osm}} + \beta_t \times \text{LDMC} + \text{intercept})</td>
<td>(\beta)</td>
<td>0.506</td>
<td>0.138</td>
<td>0.001</td>
<td>0.58</td>
<td>11.4</td>
</tr>
<tr>
<td></td>
<td>(\beta_t)</td>
<td>-0.002</td>
<td>0.001</td>
<td>0.007</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>intercept</td>
<td>0.013</td>
<td>0.258</td>
<td>0.96</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(\pi_{0_{osm}} + \beta_t \times e + \text{intercept})</td>
<td>(\beta)</td>
<td>0.654</td>
<td>0.226</td>
<td>0.17</td>
<td>0.53</td>
<td>16.1</td>
</tr>
<tr>
<td></td>
<td>(\beta_t)</td>
<td>-0.013</td>
<td>0.132</td>
<td>&lt; 0.001</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>intercept</td>
<td>-0.319</td>
<td>0.006</td>
<td>0.03</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Estimated coefficients. \(\pi_0\) and \(\Psi_{tlp}\) values of 27 randomly selected species from the dataset in Bartlett et al. (2012b) were chosen. From this selection, we fitted a simple linear regression model calculating coefficient estimates, 95% C.I., \(R_{adj}^2\), AICc and MAE. At the end of this bootstrap procedure, averaged values were calculated. Differences between \(\Psi_{tlp_{pv}}\) prediction from \(\pi_{0_{osm}}\), \(\pi_{0_{fit}}\), and the one derived from Bartlett et al. (2012b) were determined comparing 95% C.I. of the coefficient estimates and AICc. Specifically, predictions were assumed to differ if 95% C.I. of coefficient estimates did not overlap and if the difference between AICc values were > 2 (Burnham and Anderson, 2004). All statistical analyses were performed using R 3.4.1 (R Foundation for Statistical Computing, Vienna, AT).

3. Results

Species scientific name, abbreviation, and associated mean values of leaf traits are summarized in Table S1. Species that sustained higher leaf construction costs (higher LMA, LDMC and Th) also had higher drought resistance (lower \(\Psi_{tlp_{pv}}\)), as shown by correlation analysis reported in Table S2.

Although \(\pi_{0_{osm}}\) resulted a significant predictor \(\pi_{0_{pv}}\) (Table 1), it underestimated \(\pi_{0_{pv}}\) at less negative values and overestimated it at more negative values (Fig. 1). The best model to predict \(\pi_{0_{pv}}\) included \(\pi_{0_{osm}}\) and LDMC as predictive variables (Table 1). The inclusion of LDMC significantly improved the predictive power of the model, as the \(R_{adj}^2\) was higher and AICc were lower than those calculated on the model including only \(\pi_{0_{osm}}\) or the one including both \(\pi_{0_{osm}}\) and \(e\) (Table 1).

\(\pi_{0_{osm}}\) and \(\pi_{0_{fit}}\) were tested as predictors of \(\Psi_{tlp_{pv}}\). The parameters of the two derived models were compared with those derived from the relationship between \(\Psi_{tlp_{pv}}\) and \(\pi_{0_{pv}}\) reported in Bartlett et al. (2012b) and from the iterative procedure described above. The average \(\beta\) and intercept estimates calculated on a reduced number of species were not statistically different from those calculated including the whole dataset provided by Bartlett et al. (2012b) and on the models including \(\pi_{0_{osm}}\) and \(\pi_{0_{fit}}\) as predictive variables, as 95% C.I. overlapped each other (Table 2). However, the model including \(\pi_{0_{fit}}\) had higher predictive ability than the one including \(\pi_{0_{osm}}\) (Table 1), as the \(R_{adj}^2\) was higher and AICc and MAE were lower (Table 2). In addition, the model including \(\pi_{0_{osm}}\) led to overestimation of \(\Psi_{tlp_{pv}}\) for values < -2 MPa. As shown in Fig. 2 and Fig. 3, mean values of slope (\(\beta\)) and intercept calculated on the model including \(\pi_{0_{fit}}\) as predictive variable were much closer to those calculated on 27 randomly selected species from Bartlett et al. (2012b).
4. Discussion

As reported by Bartlett et al. (2012a), $\pi_{0,\text{osm}}$ significantly correlated with $\pi_{0,\text{pv}}$ (Fig. 1), but the regression line was different from the desired 1:1 relationship. A reason for this discrepancy is that osmometer-based measurements of $\pi_0$ could be biased by errors due to sample preparation. In fact, the disruption of cell walls could cause the dissolution of cell walls solutes that could lead to more negative $\pi_0$ values. On the other hand, symplastic fluids could be diluted by apoplastic water, leading to higher $\pi_0$ values. In this light, testing whether this prediction could be improved is fundamental to provide a solid framework for fast and reliable $\pi_0$ estimation. In the present study, we measured several leaf morpho-anatomical traits in order to enhance the predictive power of $\pi_{0,\text{pv}}$ from measurements done with an hygrometer ($\pi_{0,\text{osm}}$) on the basis of the framework proposed by Bartlett et al. (2012a).

The best model to predict $\pi_{0,\text{pv}}$ included $\pi_{0,\text{osm}}$ and LDMC, enhancing the predictive power of the model including only $\pi_{0,\text{osm}}$ as $R^2_{\text{adj}}$ was higher and AICc and MAE were lower (Table 1). As previously suggested by Bartlett et al. (2012a), the inclusion of LDMC in the predictive model could account for both errors associated to osmometer measurements. Higher values of LDMC are associated to greater cell wall investment, which in turn could improve the maintenance of relatively high water content, thus accounting for apoplastic dilution. On the other hand, higher LDMC could also reflect thicker cell walls or leaf with smaller but more numerous cells, thus accounting for solutes concentration enrichment derived from cell walls disruption.

In their analysis, Bartlett et al. (2012a) found that the best models to predict $\pi_{0,\text{pv}}$ included $\pi_{0,\text{osm}}$ and their interaction, or just $\epsilon$ and $\pi_{0,\text{osm}}$. In our analysis, $\delta_{\text{leaf}}$ was discarded during model selection and the model including $\epsilon$ had less predictive power than the model including LDMC. These discrepancies could be due to multiple factors. Both studies included a limited number of species (30 in Bartlett et al. (2012a) and 27 in the present study), adapted to different environments. Most of the species in the present study are typical of Mediterranean biomes, while most of the species in Bartlett et al. (2012a) originate from temperate and tropical biomes. Consequently, drought tolerance and turgor loss point could be driven by different morpho-anatomical features in the two sets of species. Moreover, whereas minimum and maximum $\pi_{0,\text{pv}}$ values were nearly the same between the two datasets, the distribution of density probability of $\pi_0$ was more skewed in Bartlett et al. (2012a) (Fig. S1), indicating a higher density of observations in a narrower range of $\pi_0$ values. However, a sort of consensus approach could be derived from these analyses. In both studies, the best model to predict $\pi_{0,\text{pv}}$ included traits which reflect leaf carbon investment ($\delta_{\text{leaf}}$, LDMC and $\epsilon$), suggesting that species that sustain higher leaf construction costs (denser and/or thicker leaves), and thus occupy the “slow-growing” space of leaf economic spectrum (Wright et al., 2004), also have higher drought resistance. The framework provided by Bartlett et al. (2012a) could allow to estimate $\pi_0$ and $\Psi_{\text{tp}}$ on a large number of species and samples strongly reducing the time needed for its measurement using pv-curves. The framework provided in this study further simplifies the model proposed by Bartlett et al. (2012a), as the measurement of LDMC is faster and simpler than the procedure for $\delta_{\text{leaf}}$ measurement.

$\pi_0$ and $\Psi_{\text{tp}}$ are strongly correlated with each other (Bartlett et al., 2012b) and thus, it is possible to estimate $\Psi_{\text{tp}}$ from measurements of $\pi_0$. A significant linear relationship between $\pi_0$ and $\Psi_{\text{tp}}$ was found in the regression models run on the data provided by the authors (Table 2). We used parameters estimates of this model as a reference to compare regression models with $\pi_{0,\text{osm}}$ or $\pi_{0,\text{fit}}$ as predictive variable. The model including $\pi_{0,\text{osm}}$ resulted very similar to the one obtained by Bartlett et al. (2012a):

$$\Psi_{\text{tp,v}} = 0.899 \pi_{0,\text{osm}} - 0.581$$ (5)
However, the regression model in eqn (5) had a lower predictive power and parameters’ estimates were slightly different than those calculated from the model run on data from Bartlett et al. (2012b) (Table 2). As shown in Fig. 2, we detected a discrepancy between the regression model considering π₀,osm as the predictive variable and the one calculated on data from Bartlett et al. (2012b). In particular, Ψ₀lp values < −2 MPa tended to be overestimated by eqn (2). On the contrary, the model considering π₀,fit as explanatory variable produced parameters’ estimates much closer to those obtained from data provided in Bartlett et al. (2012b) (Fig. 2) and no discrepancy was detected.

The number of studies including π₀ and Ψ₀lp estimation from osmometer/hygrometer measurement of π₀ rapidly increased in the last years (Maréchaux et al., 2015; Petruzelli et al., 2018, 2017), and it is likely that the number of species with associated Ψ₀lp values will increase as well. In this light, the standardization and the simplification of the framework for Ψ₀lp estimation is crucial to build a solid global dataset. To improve the predictive power of the estimation of Ψ₀lp, we suggest measuring LDMC as well as π₀,osm from leaves attached to the same twig or at least belonging to the same individual. To estimate π₀ and Ψ₀lp we then suggest applying the following equations:

\[
\pi_0_{fit} = 0.506\pi_0_{osm} - 0.002LDMC \text{ (expressed in mg g}^{-1}\text{)}
\]  

(6)

\[
\Psi_{0_{lp_{fit}}} = 1.313\pi_0_{fit} - 0.032
\]

(7)

Clearly, the inclusion of more species and π₀ values in this type of analysis is needed to furtherly optimize the framework for Ψ₀lp estimation.

Contributions

FP, TS and AN conceived and designed the study; FP and TS collected the data; FP, GB and AN analysed the data; FP, GB and AN wrote the manuscript, with the contribution of all authors.

Funding

This work is part of the project ‘Functional traits as a tool to predict invasive potential by alien species in different native communities’, funded by University of Trieste.

Conflicts of interest

The authors claim no conflict of interest.

Acknowledgments

We thank Roberto Alberti for help in collecting samples and Enrico Tordoni for helpful comments during data analysis.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.plaphy.2019.03.043.

References


Zhu, S.-D., Chen, Y.-J., Ye, Q., He, P.-C., Liu, Li.-H., Pu, P.-L., Jiang, G.-F., Cao, K.-F., 2018. Leaf turgor loss point is correlated with drought tolerance and leaf carbon economics traits. Tree Physiol. 38, 658–663.