

The importance of cuticular permeance in assessing plant water use strategies

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Abstract

Accurate understanding of plant responses to water stress is increasingly important for quantification of ecosystem carbon and water cycling under future climates. Plant water use strategies can be characterized across a spectrum of water stress responses, from tight stomatal control (isohydric) to distinctly less stomatal control (anisohydric). A recent and popular classification method of plant water use strategies utilizes the regression slope of predawn and midday leaf water potentials, σ , to reflect the coupling of soil water availability (predawn leaf water potential) and stomatal dynamics (daily decline in leaf water potential). This type of classification is important in predicting ecosystem drought response and resiliency. However, it fails to explain the relative stomatal responses to drought of *Acer sacharrum* and *Quercus alba*, improperly ranking them on the spectrum of isohydricity. We argue this inconsistency may be in part due to the cuticular conductance of different species. We used empirical and modeling evidence to show that plants with more permeable cuticles are more often classified as anisohydric; the σ values of those species were very well correlated with measured cuticular permeance. Furthermore, we found that midday leaf water potential in species with more permeable cuticles would continue to decrease as soils become drier, but not in those with less permeable cuticles. We devised a diagnostic parameter, Γ , to identify circumstances where the impact of cuticular conductance could cause species misclassification. The results suggest that cuticular conductance needs to be considered to better understand plant water use strategies and to accurately predict forest responses to water stress under future climate scenarios.

Keywords

cuticle, ecohydrology, drought, isohydricity, stomata, water stress

This is the author's manuscript of the article published in final edited form as:

Lanning, M., Wang, L., & Novick, K. A. (2020). The importance of cuticular permeance in assessing plant water–use strategies. *Tree Physiology*, 40(4), 425–432. <https://doi.org/10.1093/treephys/tpaa020>

Introduction

Vegetation is a dominant factor controlling the hydrological cycle (Jasechko et al., 2013; Lanning et al., 2019; Schlesinger & Jasechko, 2014; Wang, Good, & Caylor, 2014). Diverse vegetation responses to water stress have important consequences in ecosystem carbon and water cycles. Plants respond to water stress in a multitude of ways. At the leaf level, some plants tightly control their stomata in response to drought (isohydric), others don't (anisohydric), and a spectrum of incrementally different responses exist between these two end members (Klein, 2014; Oren et al., 1999; Tardieu, Lafarge, & Simonneau, 1996). A quantitative definition of plant water use strategies which compliments empirical observations has been highly sought after within the plant physiology, ecosystem ecology, and ecohydrology communities. Of these attempts, one of the more popular ways to rank water use strategies is that proposed by Martinez-Vilalta et al. 2014 which leveraged existing measurements of predawn leaf water potential (Ψ_{PD}) and midday leaf water potential (Ψ_{MD}) to define a quantitative representation of isohydricity (Martinez-Vilalta et al., 2014). The slope (σ) of a regression between Ψ_{PD} and Ψ_{MD} (e.g., $\sigma = \partial\Psi_{MD}/\partial\Psi_{PD}$) defines the position of a particular species along the isohydric continuum. By definition, σ has mathematical meaning in describing the response of stomata to increased soil water deficit (Martinez-Vilalta et al. 2014, see proof and references within), *id est*, the larger the absolute value of σ , the more anisohydric the species is.

This definition of isohydricity, while useful and easy to apply, has recently come under criticism for its failure to fully capture links between leaf water potential and stomatal functioning across spatial and temporal gradients (Hochberg, Rockwell, Holbrook, & Cochard, 2018; Martínez-Vilalta & Garcia-Forner, 2017). For example, this definition cannot satisfactorily explain the relative stomatal dynamics of two eastern United States tree species groups known to possess different drought response strategies: *Acer* species (specifically, *Acer sacharrum* and *Acer rubrum*) and *Quercus* species (specifically, *Quercus alba* and *Quercus rubra*). Loewenstein and Pallardy (1998) report that both maples and oaks are anisohydric species, which is consistent with σ based classification. However, subsequent empirical work suggests that oaks are substantially more anisohydric than maples (Kannenbergh, Novick, & Phillips, 2019; Roman et al., 2015; Thomsen et al., 2013; Yi, Dragoni, Phillips, Roman, & Novick, 2017). The *Acer* species mentioned earlier have been observed to maintain a more consistent leaf water potential (Ψ_{leaf}) by reducing stomatal conductance under drought conditions, whereas the *Quercus* species do not do this (Roman et al., 2015). Instead, *Quercus* maintains sap flux and stomatal conductance values similar to non-drought conditions (Roman et al., 2015; Yi et al., 2017), likely supplemented by deeper water sources as has been observed in other studies (Matheny et al., 2016; Thomsen et al., 2013). However, rooting depth alone cannot explain the decoupling between σ and stomatal dynamics, as deeper roots should promote more stationarity in both Ψ_{leaf} and conductance, whereas the evidence suggests that high conductance is sustained despite substantial variability in mid-day oak Ψ_{leaf} . In other words, leaf water potential based classification methods contradict observed stomatal behavior in oaks and maples.

A reason for this paradox may be failing to consider cuticular permeance, sometimes expressed as cuticular conductance, which has been described as a parameter that needs consideration by Martínez-Vilalta & Garcia-Forner, 2017, a paper critical of Martinez-Vilalta et al. 2014. We propose that cuticular permeance influences plant water use strategies and that cuticular permeance variation (and minimum conductance generally) provides a mechanistic explanation of how σ can be a successful indicator of isohydricity in one location but fail in others.

The consequence of cuticular water loss – a modeling perspective

Water loss via the leaf is primarily but not exclusively through stomatal pore, as the cuticle of leaf can also conduct water. The composition and arrangement of molecules which make up the cuticle transmit and absorb water, presenting an interface between the atmosphere and leaf water stores within the plant (Kerstiens, 1996, see references within). This interface is dynamic and has different transmissivities as the leaf dries when water deficit arises internally (e.g., low xylem water potential) or externally (e.g., high vapor pressure deficit, VPD) (Kerstiens, 1996). A mathematical representation of leaf water potential incorporating the potential for substantial cuticular water loss may be expressed as:

$$\psi_{leaf} = \psi_{soil} - a \frac{g_t * VPD}{K} - \rho gh \quad , \quad (1)$$

where Ψ_{soil} is soil water potential, K is whole plant hydraulic conductance, ρgh is gravitational head loss (assumed negligible), g_t is total conductance (stomatal conductance (g_s) + cuticular conductance ($g_{cuticle}$)), and a is a constant. For the sake of simplicity, cuticular conductance and cuticular permeance will be used interchangeably. However, it is important to note the difference in units ($m\ s^{-1}$ vs. $mmol\ m^{-2}\ s^{-2}$, respectively) and that water movement within the cuticle is driven by a concentration gradient whereas water moving from the cuticle to the atmosphere has meteorological dependencies too (e.g., temperature, relative humidity, and as a product VPD; discussed in Kerstiens, 1996).

Cuticular conductance is part of total leaf conductance, but is not typically considered important as stomatal conductance (g_s) is dominant most of the time (Duursma et al., 2018; Kerstiens, 1996). As drought intensifies, g_s is reduced while $g_{cuticle}$ may be unchanged, which would cause the ratio of $g_{cuticle}:g_t$ to increase. The magnitude of this error is important when evaluating the isohydricity of a given species based on Ψ_{leaf} measurements. To illustrate this, we modeled declines in Ψ_{leaf} under a simulated drought according to Equation 1 allowing g_t to be simulated for each instance (i) as follows:

$$g_t = g_{ref}(1 - 0.6 \ln(VPD)) * \left(1 - c_g \frac{\psi_s}{\min(\psi_s)}\right) + g_{cuticle} \quad , \quad (2)$$

where g_{ref} is a reference conductance rate for well-watered conditions and $VPD = 1$ kPa. The term $(1 - 0.6 \ln(VPD))$ reduces conductance as a function of increasing VPD, using the formulation proposed by Oren et al., (1999). The $\left(1 - c_g \frac{\psi_s}{\min(\psi_s)}\right)$ term reduces conductance as a simple linear function of relative soil water potential, where c_g is a sensitivity parameter related to the degree of isohydricity (see Martinez-Vilalta et al. 2014, Novick et al. 2019). It should be equal to 0 for a perfectly anisohydric species, and larger values represent more isohydric species. To illustrate the importance of $g_{cuticle}$, drought conditions were modeled and Ψ_{leaf} was estimated for an anisohydric ($c_g = 0.4$; I_{40}) and distinctively more isohydric ($c_g = 0.6$; I_{60}) species. The subscript here represents the degree of isohydricity. $g_{cuticle}$ was set to literature reported values converted from cuticular permeance (as per Kerstiens 1996) of *Quercus alba* (representing I_{40}) and *Acer sacharrum* (representing I_{60} ; Figure 1). As $g_{cuticle}$ was set as a constant, it is important to note that our simulation assumes that changes to the ratio of $g_{cuticle}:g_t$ is only determined by changes in g_s (Equation 1). In reality $g_{cuticle}$ (and generally minimum conductance, g_{min}) will change as drought intensifies and temperature increases (see discussions in: Duursma et al., 2018; Kerstiens, 1996, 2006). To evaluate the importance of $g_{cuticle}$, we simulated g_t for I_{60} using the reported high cuticular conductance (I_{60H} ; $6.97\ mmol\ m^{-2}\ s^{-1}$) for *Acer sacharrum*, and also simulate g_t using g_s only (I_{60g_s}). We used the averaged reported value of $g_{cuticle}$ for *Quercus alba* (I_{40A} ; $2.56\ mmol\ m^{-2}\ s^{-1}$; Figure 1). We calculated a simulated σ for each modeled scenario (Figure 1).

Relative Ψ_{leaf} reductions in isohydric and anisohydric species can vary substantially simply due to cuticular conductance (Figure 1). Our modeling results indicate that when cuticular conductance is considered, an isohydric species with a permeable cuticle (e.g., I_{60H}) could demonstrate Ψ_{leaf} and g_t reflective of a distinctly more anisohydric species (e.g., I_{40A}) under representative non-drought conditions (Figure 1). Simulated σ values were remarkably similar between species with distinct drought responses if cuticular conductance is considered (Figure 1). For example, the simulated σ for I_{40A} and I_{60H} were identical (Figure 1). However, when g_{cuticle} was not included there was clear difference (Figure 1). Including cuticular conductance in Ψ_{leaf} models illustrates how the conventional calculation and interpretation of σ can be influenced by traits that do not factor into the definition of σ (Figure 1). Specifically, assuming cuticular conductance is negligible while comparing species with different cuticle properties may significantly bias the estimation of σ .

The consequence of cuticular water loss - empirical evidence

To assess the influence of cuticular permeance on σ using empirical observations, we combined and averaged measurements of cuticular permeance from two sources (Kerstiens, 1996; Schuster, Burghardt, & Riederer, 2017) and extracted the paired measurements of Ψ_{PD} and Ψ_{MD} of any overlapping species in Martinez-Vilalta et al. (2014) (Table 1). It should be noted that some measurements of cuticular conductance are more representative of the sum of g_{cuticle} and a non-negligible flux of water from incompletely closed stomata (i.e., g_{min} , see Kernsteins 1996 and Duursma et al., 2018). A relationship between cuticular conductance and changes in minimum conductance are discussed later in this section.

We focused on angiosperms only due to the differences in stomatal function and xylem structure between angiosperms and gymnosperms. For example, guard cell sensitivity to abscisic acid (ABA) over drought conditions is different between the two clades (discussed in Brodribb & McAdam, 2017). In response to rises in VPD, angiosperms upregulate ABA synthesis within minutes causing fast stomatal closure, likely due to high speed gene expression at a critical step in ABA synthesis (McAdam, Susmilch, & Brodribb, 2016). Within the gymnosperm clade there is not a uniform control of stomatal conductance. Instead, there are reports of more passive regulation of g_s with no significant increase in ABA (McAdam & Brodribb, 2015), other instances where sufficient ABA concentrations to induce stomatal closure took more than six hours to reach (McAdam & Brodribb, 2014). Such differences would complicate our analysis which uses observations from multiple time scales (days to months) and make interpreting results subject to error.

Measurements of leaf water potential conducted on seedlings, branches, exclusively riparian species with constant groundwater access (e.g., some *Populus* species) and under potentially biasing treatments (e.g., greenhouse, increased salinity) were excluded to assess the relationship between permeance and σ . Additionally, if Ψ_{PD} and Ψ_{MD} were present, but could not be obviously extracted as a pair, these were also excluded. Ranges of Ψ_{leaf} for the species presented here may not reflect exactly what was used in the analysis by Martinez-Vilalta et al. (2014). Therefore, we made a modified estimate of the degree of isohydricity with the same method in Martinez-Vilalta et al. (2014) using our extracted Ψ_{leaf} data from articles which fit our criteria (hereafter σ_{calc}). For discussion purposes σ_{calc} was used interchangeably with the reported σ values in Martinez-Vilalta et al. (2014) as their relationships with cuticular permeance were not different (Figure 2).

In addition to the calculations of σ , we introduced a new parameter, Γ , defined as the correlation (r) between $\Psi_{\text{PD}} - \Psi_{\text{MD}}$ and Ψ_{PD} , when $\Psi_{\text{PD}} > \Psi_{\text{MD}}$, similar in principle to an analysis by Meinzer et al. (2016) (Figure 2). The Γ value enhances understanding that can be gained using σ in a few important ways. First, the calculated Γ values for each species are confined to a range (-1 to 1) whereas σ is unbounded. Second, Γ provides information not only about the direction of the relationship between pre-dawn and mid-day leaf

water potential, but also provides information about the covariability between the two. This is important, because σ , as traditionally defined (e.g., Martinez-Vilalta et al. 2014), does not directly accommodate the influence of vapor pressure deficit, among other factors, on the dynamics of Ψ_{MD} (Novick, Konings, & Gentine, 2019). Assuming all water flux from the leaf is through the stomata, and that σ is an appropriate indicator of isohydricity (e.g., influence of VPD and other variables is small), then more negative Γ values should represent a more ‘wasteful’ physiotype within the traditional framework, meaning that as Ψ_{PD} declines Ψ_{MD} will also decline, reflecting an anisohydric water use strategy and/or higher cuticular permeance. Positive Γ values represent a ‘conservative’ physiotype where Ψ_{MD} will be more stationary as soil drying progresses, likely related to species with an isohydric water use strategy and/or lower permeance. In both of these examples, σ and Γ should be relatively well coupled across species and sites. However, if VPD or other factors substantially affect the relationship between Ψ_{PD} and Ψ_{MD} , then Γ and σ will be less coupled from one species to the next. Thus, considering both metrics, if they lead to similar conclusions, can increase confidence that results are not driven by spurious excursions in Ψ_{MD} , for example due to passing fronts that may affect VPD more than Ψ_{PD} . It should be noted that such representation of isohydricity may only be relevant for species at the extremes of the isohydric continuum, thus Γ should only be used as a diagnostic parameter. Likewise, both Γ and σ could also be sensitive to non-cuticular impacts (e.g., cavitation).

Calculated Γ and the reported σ values from Martinez-Vilalta et al. (2014) were compared against the extracted values of cuticular permeance from Kersteins (1996) and Schuster et al. (2017) using simple linear regression. Cuticular permeance and Γ had a predictable relationship (Figure 3). Species with the lowest permeance of the dataset also had the highest Γ , meaning that at low Ψ_{PD} , Ψ_{MD} is not likely to change (Figure 2, Table 1). Conversely, the species with the highest cuticular permeance had the lowest Γ , indicating continued leaf dehydration as Ψ_{PD} decreases (Figure 3, $R^2 = 0.86$, $p = 0.02$). σ values for all the study species were ranked the same way and were well described by the extracted cuticular permeance values, though *Fagus sylvatica* deviated slightly from the overall line trend causing the relationship with σ_{calc} to be marginally significant (Table 1, Figure 3, $R^2=0.86$ and 0.71 , $p = 0.02$, 0.07 for σ and σ_{calc} , respectively). The relationship between cuticular permeance and σ observed in this analysis suggests there may be an inherent classification bias for species at the extremes of the permeance spectrum (Figure 3). The values of σ and Γ seem to be well characterized, if not constrained, by the cuticular permeance of the plant and thus may not independently provide an ecologically relevant mechanism as to how the plant survives drought or how such a response will affect carbon and water cycling (Figure 3).

Concurrent measurements of cuticular conductance and leaf water potential under natural or controlled conditions were not available in literature. The natural variability of cuticular permeance of an average cuticle is not known for any of the species included in this analysis, most of which are important hardwood trees common to the northeastern United States. All of these limit the robustness of our conclusions about the importance of cuticular permeance on plant water use. However, besides our own analyses, multiple observations implicate cuticular permeance as an important and dynamic component of leaf hydrology. For example, a study by Boyer, Wong, & Farquhar, 1997 shows that leaf age can cause a substantial increase in permeance, increasing from 10% of g_s to nearly 30% of g_s in *Vitis vinifera* L. (grapes), thus imposing leaf level drought stress in older fraction of the canopy earlier than the younger fraction. There is also evidence that a drying cuticle can have enough ‘mechanical advantage’ over nearby stomata to pull them open, increasing g_{min} (Buckley 2005). As discussed in detail by Kersteins (1996) and Duursma et al. (2018), some reported measurements of $g_{cuticle}$ may actually be more representative of g_{min} , which is important since the dehydration of the cuticle could lead to higher g_{min} . Growth conditions associated with dry leaves (e.g., high temperature) can also lead to reduced g_{min} as shown by Duursma et al., (2018) in *Eucalyptus parraattensis*, however, the impact of higher temperature on g_{min} or $g_{cuticle}$ across

species is largely unassessed, preventing any complete understanding of plant transpiration response under future climates.

Conclusions and future directions

Based on modeling exercises and empirical analysis, our results indicate that cuticular properties are important in regulating plant water use and should be considered for a more holistic evaluation of the links between gas exchange and leaf water potential during water stress going forward. To aid in such efforts, Γ could serve as a good quality control parameter prior to using σ as a classifier of isohydricity. For example, if the Γ value of two species is similar it may be appropriate to classify them using σ values. However, if the species with a larger σ value also has a strong negative Γ value (e.g., *Acer sacharrum*), Ψ_{leaf} measurements alone should be used with caution. In such circumstances, it would be best to determine the relative isohydricity by comparing stomatal conductance and Ψ_{leaf} in a variety of environmental conditions, not relying on the relationship between Ψ_{PD} and Ψ_{MD} alone.

Though this study was written to include discussions of the isohydric framework, it is important to emphasize that there appears to be a close relationship between Ψ_{leaf} declines, permeance, and plant water stress, regardless the actual utility of plant isohydricity classifications. It is expected that droughts will occur under warmer conditions and that VPD in many plant dominated ecosystems will likely increase (Novick et al., 2016). Additionally, land-atmosphere feedbacks can prolong arid atmospheric conditions (Zhou et al., 2019), causing variations in g_{cuticle} and g_{min} , meaning that our findings and modeled examples might underestimate the influence of cuticular conductance in future climate scenarios as our analysis keeps g_{cuticle} constant. Plant acclimatization to warmer conditions also will likely impact g_{cuticle} and g_{min} (Duursma et al., 2018)—another instance where water and carbon cycling will be impacted in the future but empirical data is lacking at present. Based on our current analyses and literature information, the seemingly small effect of cuticular conductance could play a large role in both how vegetation responds to drought and in forest management decisions, which will rely on accurate representations and predictions of forest response to drought. To meet that need, it will be important for researchers to begin incorporating measurements and estimations of cuticular conductance in their experiments.

Acknowledgments

We acknowledge support from Division of Earth Sciences of National Science Foundation (NSF EAR-1554894) and from the Agriculture and Food Research Initiative program (2017-67013-26191) of the USDA National Institute of Food and Agriculture. We thank the comments from two anonymous reviewers, which significantly improved the quality of the manuscript.

Authors' Contributions

ML and LW conceptualized the main research questions. ML collected data and performed the data analyses. ML and LW wrote the first draft. KAN contributed modeling analyses and revised manuscript.

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Table 1 Complied species level values of cuticular permeance, σ (slope of regression of Ψ_{MD} vs Ψ_{PD}), σ_{calc} (same as σ but from mature trees in a natural setting), and Γ (the correlation of the regression between Ψ_{PD} - Ψ_{MD} and Ψ_{PD}) as well as the source of the leaf water potential (Ψ_{leaf}) data. Values from non-significant regression marked with 'NS'.

Species	Letter Code	Cuticular Permaince ($ms^{-1} \times 10^{-5}$)	σ^{***}	σ_{calc}	Γ	Sampling Resolution	Ψ_{leaf} Data Sources
<i>Acer saccharum</i>	AS	8.13 ^{***}	1.127	1.46	-0.58	multiple days	Loewenstein and Pallardy 1998
<i>Fagus sylvatica</i>	FS	5.28 ^{***}	0.681	0.39	0.63	one day, multiple days	Ismael Aranda, Gil, and Pardos 2000; I. Aranda, Gil, and Pardos 2005
<i>Quercus alba</i>	QA	7.4 [*]	0.918	1.28	-0.32 ^{NS}	multiple days	Loewenstein and Pallardy 1998
<i>Quercus ilex</i>	QI	3.6 [*]	0.652	0.58	0.71	various	Martínez-Vilalta et al. 2002, 2003; Serrano and Peñuelas 2005; Cubera and Moreno 2007b, 2007a
<i>Quercus petra</i>	QP	5.55 ^{**}	0.829	1.09	-0.03 ^{NS}	various	Ismael Aranda, Gil, and Pardos 2000; I. Aranda, Gil, and Pardos 2005; Breda et al. 1993

* Kerstiens 1996

** Schuster, Burghardt, and Riederer 2017

***Martinez-Vilalta et al. 2014

Figure legends

Figure 1 Modeled leaf water potential (top) and total leaf conductance (g_t , bottom) for simulated I_{60} and I_{40} plants according to equations 1 and 2. The subscripts '60' and '40' refer to the degree of isohydricity the modeled plant represents as described in section 1. The simulations were made using the g_s only (I_{60g_s}) and reported high (I_{60H} ; $6.97 \text{ mmol m}^{-2} \text{ s}^{-1}$) estimations of cuticular conductance for *Acer saccharum* and the reported estimation of cuticular conductance for *Quercus alba* (I_{40A} ; $2.56 \text{ mmol m}^{-2} \text{ s}^{-1}$). Simulated σ values are the slopes of the line equation reported in the top figure.

Figure 2 The correlations between measured pre-dawn leaf water potential (Ψ_{PD}) and midday leaf water potential (Ψ_{MD}) (σ) as well as between $\Psi_{PD}-\Psi_{MD}$ and Ψ_{PD} (Γ) for different species. The filled circles correspond to the data used to calculate σ and the open circles correspond to the data used to calculate Γ . Regression lines are shown to indicate direction and please note the difference in x-axis scales.

Figure 3 Observed relationships between σ (solid line), σ_{calc} (dashed line), Γ (dotted line), and cuticular permeance. The two-letter codes above correspond to the following individual species and are listed in Table 1: QI = *Quercus ilex*; FS = *Fagus sylvatica*; QP = *Quercus petraea*; QA = *Quercus alba*; AS = *Acer saccharum*.

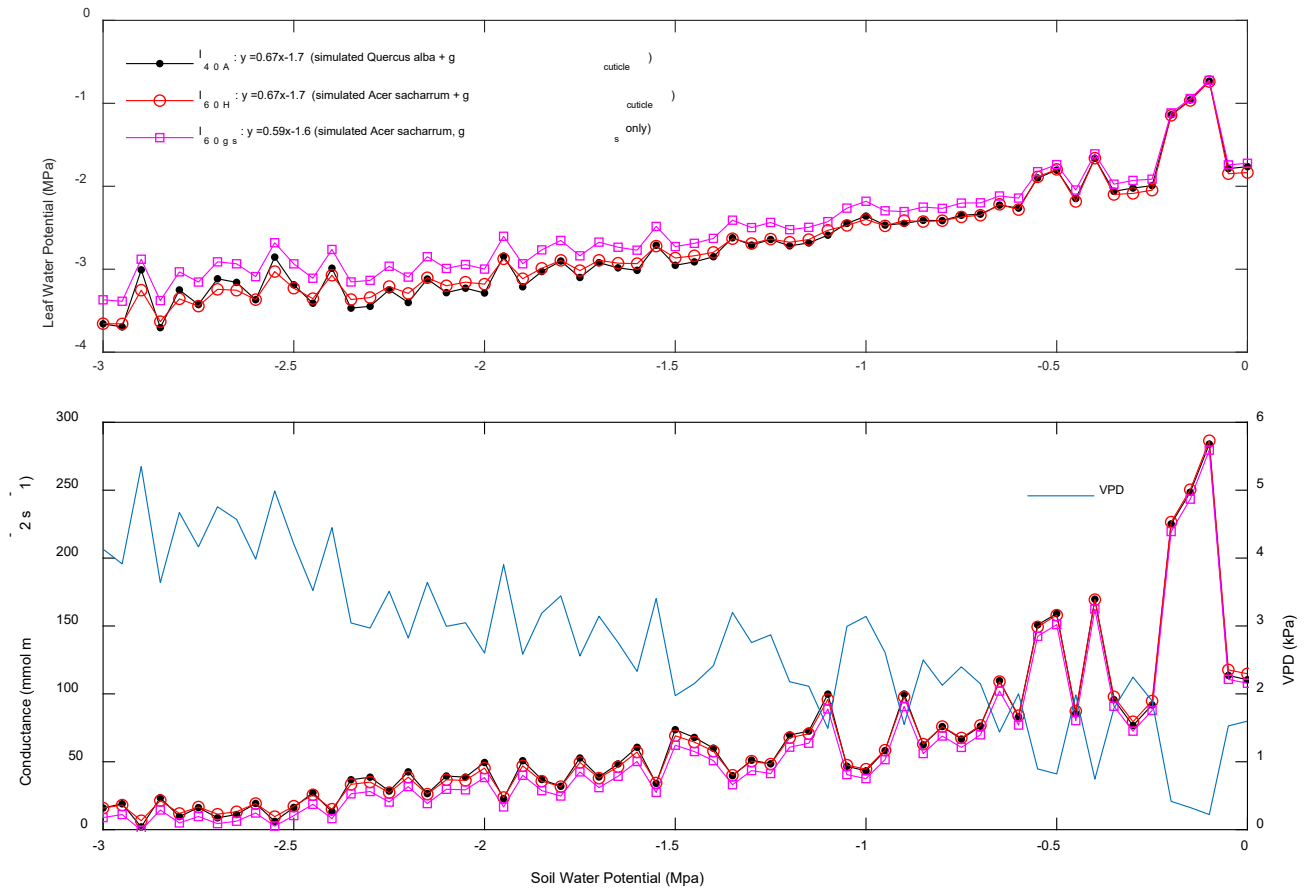


Figure 1

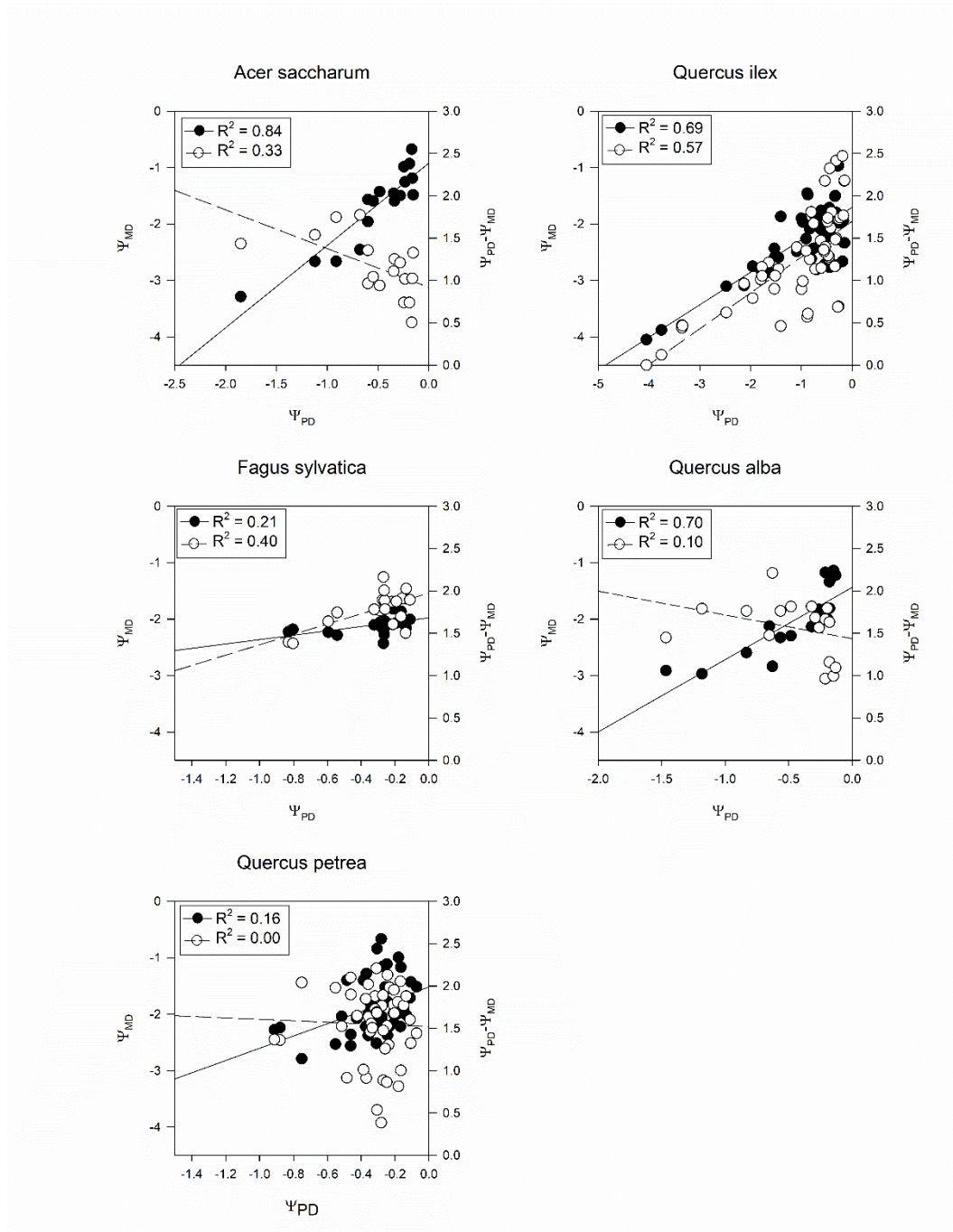


Figure 2

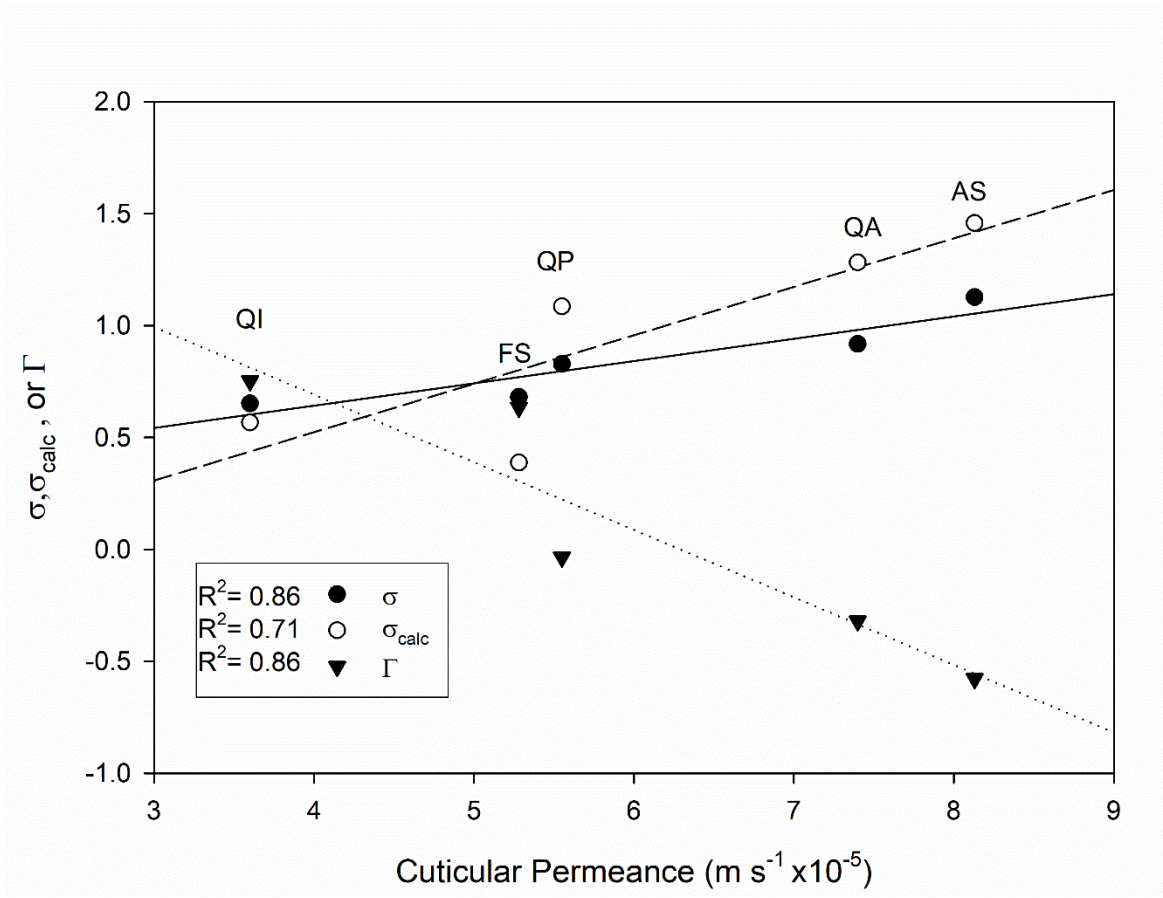


Figure 3