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Modelling reveals endogenous osmotic adaptation of storage tissue water potential as an important driver determining different stem diameter variation patterns in the mangrove species *Avicennia marina* and *Rhizophora stylosa*

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- **Background** Stem diameter variations are mainly determined by the radial water transport between xylem and storage tissues. This radial transport results from the water potential difference between these tissues, which is influenced by both hydraulic and carbon related processes. Measurements have shown that when subjected to the same environmental conditions, the co-occurring mangrove species *Avicennia marina* and *Rhizophora stylosa* unexpectedly show a totally different pattern in daily stem diameter variation.

- **Methods** Using in situ measurements of stem diameter variation, stem water potential and sap flow, a mechanistic flow and storage model based on the cohesion–tension theory was applied to assess the differences in osmotic storage water potential between *Avicennia marina* and *Rhizophora stylosa*.

- **Key results** Both species, subjected to the same environmental conditions, showed a resembling daily pattern in simulated osmotic storage water potential. However, the osmotic storage water potential of *R. stylosa* started to decrease slightly after that of *A. marina* in the morning and increased again slightly later in the evening. This small shift in osmotic storage water potential likely underlaid the marked differences in daily stem diameter variation pattern between the two species.

- **Conclusions** The results show that in addition to environmental dynamics, endogenous changes in the osmotic storage water potential must be taken into account in order to accurately predict stem diameter variations, and hence growth.

**Key words:** Functional–structural modelling, *Avicennia marina*, *Rhizophora stylosa*, stem diameter variation, dendrometer, osmotic water potential, endogenous control, mangrove, plant water relations, osmotic regulation, growth, mechanistic model, sap flow.

INTRODUCTION

Radial water transport between the xylem conductive elements and the surrounding storage tissues plays a key role in dynamic water transport in plants (e.g. *Steppe et al.*, 2006; Hölttä *et al.*, 2009; De Schepper and Steppe, 2010; De Swaef *et al.*, 2013b). Water in the storage tissues can buffer discrepancies between water demand and supply, avoiding hydraulic failure in the xylem, and allows turgor to build up. This turgor pressure ultimately enables irreversible growth, provided a specific threshold pressure is overcome (Lockhart, 1965). Moreover, radial water transport is indispensable for the functioning of the phloem (De Schepper *et al.*, 2013). To study this radial water transport, stem diameter variation (SDV) measurements have proven to be a powerful tool (e.g. Zweifel *et al.*, 2000, 2001; Peramaki *et al.*, 2001; Fereres and Goldhamer, 2003; Daudet *et al.*, 2005; De Swaef and Steppe, 2010; Betsch *et al.*, 2011; De Schepper *et al.*, 2012). Daily variations in stem diameter are the result of four main processes: (1) contraction and expansion of dead conducting elements because of the increase and relaxation of internal tensions; (2) reversible shrinking and swelling of living tissues in relation to different levels of tissue hydration; (3) irreversible radial stem growth; and (4) thermal expansion and contraction. Of these processes, reversible shrinking and swelling and irreversible radial stem growth have the most pronounced effects on SDV. Both processes are the result of the actual radial water transport caused by differences in water potential between the xylem and storage tissues (Daudet *et al.*, 2005).

Experimental studies on C3 and C4 plants have shown that stem diameters tend to decrease in the morning and increase in the afternoon, even though differences in the relative shrinking and swelling were noted depending on species, height within the tree and environmental conditions (e.g. Lovdahl and Odin, 1992; Sevanto *et al.*, 2002; Steppe *et al.*, 2008; De Swaef *et al.*, 2009). It has been commonly accepted that this daily pattern is caused by a water potential difference between xylem and storage tissues, originating from daily hydraulic and carbon-related processes within the tree. In the morning, the tension...
inside the xylem increases as transpiration starts, lowering the xylem water potential. When this xylem water potential drops below the storage water potential, water transport from the storage tissues to the xylem causes the stem diameter to decrease (Hinckley and Bruckerhoff, 1975). In the afternoon, when the atmospheric water demand decreases, water can again flow back into the storage tissues if the storage water potential is more negative than the rising xylem water potential, resulting in a stem diameter increase (Molz and Klepper, 1973). Even though the daily stem diameter pattern for CAM plants is reversed, with an increase in the morning and a decrease in the evening, it can still be explained by this same principle, because for CAM plants stomata are closed during the day and opened during the night (Matimati et al., 2012). Besides these hydraulic processes, SDV are influenced by plant carbon relations, including processes such as leaf and woody tissue photosynthesis, starch conversion to sugars, respiration, local production and accumulation of osmotic active compounds such as proline, glycinebetaine, mannitol and inorganic ions, and sink and source activity (Sevanto et al., 2003; Höttä et al., 2006; Naidoo, 2006; De Schepper and Steppe, 2010; Saveyn et al., 2010; Krauss and Ball, 2013; De Swaef et al., 2013a). De Schepper et al. (2010) showed that the timing of stem diameter shrinking was altered during sink activity manipulation while Sevanto et al. (2003) showed that the time lag between xylem and total stem diameter variation was related to photosynthetic carbohydrate assimilation, carbohydrate transport and sink activity. De Swaef et al. (2013b) reported measurements of the morning increase in tomato SDV, which they attributed to the phenomenon of root pressure, whereby active loading of solutes in the root xylem initiates osmotic water uptake (Kramer and Boyer, 1995).

Recently, M. W. Vandegehuchte et al. (unpubl. res.) discovered that, despite thriving in the same environment, two representatives of the two most dominant mangrove genera (Robert et al., 2009), Avicennia marina and Rhizophora stylosa, both C3 plants, showed opposite daily stem diameter patterns despite similar long-term trends. The aim of the current work was to assess and discuss the influence of the osmotic storage water potential on SDV and the implications for functional modelling. To this end, ecophysiological measurements conducted on A. marina and R. stylosa were applied in a mechanistic model based on the cohesion–tension theory.

**MATERIALS AND METHODS**

**Field site**

Measurements were conducted at the west coast of North Stradbroke Island, Queensland, Australia (27°27′06″S 135°25′08″E), near an old coastal sand dune island. The island is characterized by sandy soils and acidic water bodies intertwined by a complex mix of perched groundwater-fed freshwater lakes, swamps and creeks (Page et al., 2012). On this field site, three full-grown trees of both Avicennia marina and Rhizophora stylosa were chosen, located in proximity of each other to avoid tidal effects and spatial salinity gradients (Fig. 1, Table 1). The experimental data were gathered during winter, from 12 August to 11 September (day of year [DOY] 225–255) as during this period dry and mild climatic conditions were expected and the effect of rainfall could be excluded.

![Fig. 1. Schematic of the measured trees at the mangrove site on the west coast of North Stradbroke Island, showing the measured A. marina trees (grey circles), the measured R. stylosa trees (white circles) and the location of the weather station (MET).](https://academic.oup.com/aob/article-abstract/114/4/667/162358)

**Table 1. Diameter at breast height (m) of the measured mangrove trees**

<table>
<thead>
<tr>
<th>Tree</th>
<th>Av1</th>
<th>Av2</th>
<th>Av3</th>
<th>Rhi1</th>
<th>Rhi2</th>
<th>Rhi3</th>
</tr>
</thead>
<tbody>
<tr>
<td>0-194</td>
<td>0.306</td>
<td>0.181</td>
<td>0.188</td>
<td>0.117</td>
<td>0.172</td>
<td></td>
</tr>
</tbody>
</table>

**Environmental conditions**

Air temperature, relative humidity, shortwave solar radiation and wind speed were measured every minute and averages recorded every 10 min at 2 m above the soil surface. Rainfall was measured using a tipping-bucket rain gauge and recorded for every tip (HOBO Weather Station, Onset, Cape Cod, MA, USA). Vapour pressure deficit (VPD, kPa) was inferred from measured air temperature ($T_{air}$) and relative humidity (RH) according to Buck (1981). Soil salinity and water table depth were measured every minute and averages recorded every 10 min from DOY 245 onwards with an in situ pressure transducer (Aqua Troll 200, In-Situ, Fort Collins, CO, USA) installed in a piezometer, located close to the weather station at a depth of 0.25 m below the soil surface. Actual measured soil water electrical conductivity ($C_w$ in ms cm$^{-1}$) was converted to osmotic water potential $\Psi_w$ (MPa) based on McIntyre (1980):

\[
\Psi_w = -10^{1.091 \log(C_w) - 1.46}
\]  

**Plant variables**

Each tree was equipped with a dendroband at breast height (DBL60 Logging Band Dendrometer with a resolution of 1 µm, ICT international, Armidale, NSW, Australia), recording stem diameter every 10 min, from which stem diameter variations were derived. Temperature correction of the signals was not deemed necessary as temperature changes have no significant impact on the operation of the DBL60 (ICT international, Armidale, NSW, Australia). Sapflow+ sensors, registering heat velocity every 40 min, were installed at breast height at the south side of each tree (Vandegehuchte and Steppe, 2012). These heat velocities were then converted to sap flow based on wood core measurements of dry wood density and estimations of sapwood depth (Vandegehuchte and Steppe, 2013). Stem water potential was recorded with a stem psychrometer every 10 min (PSY-1 Stem Psychrometer, ICT International, Armidale, NSW, Australia).
Mechanistic stem diameter variation model

The mathematical flow and storage model of Steppe et al. (2006) was applied to enable the assessment of xylem and storage water potential dynamics based on stem sap flow and stem diameter variation measurements. Whereas in the work of Steppe et al. (2006) the aim of the model was to simulate SDV based on measured transpiration, here the objective was to derive the osmotic potential of the stem storage tissues based on the measured sap flow, SDV and soil water potential. In this model, the thickness of the stem storage compartment, encompassing the storage tissues throughout the stem, ΔS (m), was linked to the outer stem diameter $D_{out}$ (m) via an empirical relationship (Génard et al., 2001; Steppe et al., 2006):

$$\Delta S = a[1 - \exp(-b \times D_{out})]$$  \hspace{1cm} (2)

where $a$ and $b$ are empirical parameters. Based on measurements of bark thickness, as an approximation of ΔS, and stem diameter of the investigated A. marina and R. stylosa trees, estimates of $a$ and $b$ were obtained, with $a = 0.0035$ m and $b = 18.47$ m$^{-1}$ for A. marina and $a = 0.0153$ m and $b = 10.03$ m$^{-1}$ for R. stylosa, respectively. These parameters were assumed to be constant throughout the measurement period. By subtracting ΔS from the measured input variable $D_{in}$, the inner diameter of the stem $D_{in}$ could be calculated, as well as the water flow ($dW_{st}/dt$ in m$^3$ s$^{-1}$) in and out of the stem storage tissue, with $l$ the length of the stem, which was estimated for both species (2 and 4 m for R. stylosa and A. marina, respectively) (Fig. 2).

$$D_{in} = D_{out} - 2\Delta S$$  \hspace{1cm} (3)

$$V_{st} = \pi(l(D_{out} - \Delta S - \Delta S^2))$$  \hspace{1cm} (4)

The change in $V_{st}$ was directly linked to the storage tissue pressure potential $\Psi_{p,st}$, distinguishing between elastic expansion and contraction (if $\Psi_{p,st} > \Gamma$; eqn 5) or both elastic (el) and plastic (pl) growth (if $\Psi_{p,st} < \Gamma$; eqn 6) (Lockhart, 1965):

$$\left( \frac{dV_{st}}{dt} \right)_{el} = \frac{V_{st} \Psi_{p,st}}{\varepsilon} \text{ for } \Psi_{p,st} < \Gamma$$  \hspace{1cm} (5)

$$\left( \frac{dV_{st}}{dt} \right)_{el+pl} = \frac{V_{st} \Psi_{p,st}}{\varepsilon} + V_{st} \phi(\Psi_{p,st} - I) \text{ for } \Psi_{p,st} > \Gamma$$  \hspace{1cm} (6)

where $\phi$ is the extensibility of cell walls in relation to non-reversible dimensional changes (MPa$^{-1}$ s$^{-1}$), $\varepsilon$ is the bulk elastic modulus of living tissue in relation to reversible dimensional changes (MPa) and $\Gamma$ is the critical value (in MPa) of the pressure component ($\Psi_{p,st}$) that must be exceeded to produce irreversible growth in the storage compartment. For this threshold pressure $\Gamma$, different values have been observed, ranging from 0-1 to 0.9 MPa (Green et al., 1971; Green and Cummings, 1974; Hsiang and Xu, 2000). Based on the reasoning of Génard et al. (2001) that $\Gamma$ has to be higher for stem tissues than for the young tissues or individual cells on which most of the observed values are based, the upper value of 0.9 MPa was chosen for the simulations. As the bulk elastic modulus increases with turgor and cell size (Tyree and Jarvis, 1982; Dale and Sutcliffe, 1986), it was considered proportional to $D_{out}$ and $\Psi_{p,st}$:

$$\varepsilon = \varepsilon_0 D_{out} \Psi_{p,st}$$  \hspace{1cm} (7)

where $\varepsilon_0$ is a proportionality constant (Génard et al., 2001; Steppe et al., 2006).

The variables obtained from this stem diameter variation submodel could then be applied in the water transport submodel (Fig. 2). In this submodel, xylem water potential $\Psi_x$ (MPa) was derived from the root water potential $\Psi_r$ (considered proportional to the soil water potential $\Psi_{soil}$, applying a proportionality factor, $k_{soil}$), the measured sap flow $F_{s}$(mg s$^{-1}$) and the xylem resistance to flow $R_x$ (MPa s mg$^{-1}$). $R_x$ and $k_{soil}$ were calibrated based on stem water potential measurements, applying a 3-day moving window calibration to take temporal variability into account.

The mass flow of water to and from the storage compartment ($dW_{st}/dt$) was determined from the volumetric flow assuming a constant water density of 1000 kg m$^{-3}$ ($\rho_{w}$) and taking into account that only approximately 40 % of the total stem storage volume $V_{st}$ consisted of water (Steppe et al., 2006):

$$dW_{st}/dt = 0.4 \rho_{w} dV_{st}/dt$$  \hspace{1cm} (8)

The water potential of the storage tissue could be determined based on the flow to and from this tissue and the exchange resistance $R_{st}$ (MPa s mg$^{-1}$) between stem storage and stem xylem tissue. The latter was determined based on the assumption that the stem storage and xylem compartments were separated by a virtual membrane with radial hydraulic conductivity $L$ (m MPa$^{-1}$ s$^{-1}$; Steppe et al., 2006):

$$R_{st} = (\pi D_{in} L \rho_{w})^{-1}$$  \hspace{1cm} (9)

$$\Psi_{st} = \Psi_x - R_{st} \times dW_{st}/dt$$  \hspace{1cm} (10)

Based on the measured diameter changes, the changes in storage tissue volume could be derived (eqns 3 and 4). From these changes in storage tissue volume, the changes in water potential could be determined based on eqns (5) and (6) (Fig. 2). Knowing the pressure potential $\Psi_{p,st}$ of the storage tissue from the stem diameter variation submodel, the osmotic water potential of the storage tissue $\Psi_{\pi,st}$ could then be derived (Fig. 2).

$$\Psi_{\pi,st} = \Psi_{st} - \Psi_{p,st}$$  \hspace{1cm} (11)

From the derived $\Psi_{\pi,st}$ and $V_{st}$, a measure for the total amount of osmotically active compounds $N_{eq}$ (mol) in the storage tissue could be derived from the Van’t Hoff equation:

$$N_{eq} = (\Psi_{p,st}/RT)V_{st}$$  \hspace{1cm} (12)

As no distinction was made between the different compounds or their osmotic activity, $N_{eq}$ is referred to as a number of osmotic equivalents.

While $k_{soil}$ and $R_x$ could be assessed independently through model calibration using the measured xylem water potentials, the parameters $L$, $\phi$ and $\varepsilon_0$ remained unknown. For $L$, values obtained from cells of higher plant tissues between $1·10^{-10}$ and $1·67\times 10^{-4}$ m MPa$^{-1}$ s$^{-1}$ have been
mentioned (Dainty and Preston, 1963; Dale and Sutcliffe, 1986). For cell wall extensibility \(\phi\), Hsiao et al. (1998) mentioned a range from \(8.33 \times 10^{-6}\) to \(5.56 \times 10^{-5}\) for young plants, although for older tissues the extensibility is likely to be an order of magnitude lower (Génard et al., 2001). The elastic modulus \(\epsilon\) ranges from 0 to 30 MPa for higher plant tissues (Dainty and Preston, 1963; Dale and Sutcliffe, 1986), allowing a realistic range of 0 to 250 m\(^{-1}\) for \(\epsilon_0\) based on \(D_{\text{out}}\) and \(\Psi_{\text{p,st}}\) values. Table 2 shows the applied parameter values, based on Génard et al. (2001) and Steppe et al. (2006). To assess the influence of these parameter ranges on the total amount of osmotically active compounds \(N_{\text{eq}}\), a Monte Carlo uncertainty analysis was performed (for procedure see De Pauw et al., 2008b). To this end, a uniform continuous probability distribution function was assigned to each of these parameters using the above-mentioned parameter ranges (Table 2). From these distributions associated with the parameters, 1000 samples were generated using Latin hypercube sampling (e.g. De Pauw et al., 2008b; Helton and Davis, 2003). In a next step, the sampled parameter values were propagated through the model to generate the output uncertainty on the \(N_{\text{eq}}\) patterns for both species. The upper and lower limits of the uncertainty band were defined as the 95th and 5th percentiles of the resulting output probability distribution constructed from the 1000 different trajectories.
This uncertainty band was used to evaluate actual simulated $N_{eq}$. Additionally, a sensitivity analysis was conducted according to Brun et al. (2002), Steppe et al. (2006) and De Pauw et al. (2008a) to test the relative impact of the parameters as mentioned in Table 2 on the model output $N_{eq}$. The centralized relative sensitivity function $s_j$ of the model variable $y(N_{eq})$ towards the parameter $\theta_j$ ($L$, $e_0$, $\Gamma$, $\phi$) was calculated at every time instance $k$:

$$s_j = \frac{y(\theta_j + \Delta \theta_j) - y(\theta_j - \Delta \theta_j)}{2\Delta \theta_j} \times \frac{\theta_j}{y(\theta_j)} \tag{13}$$

where $\Delta \theta_j$ is taken as 1% of the parameter value $\theta_j$. As an indication of the relative importance of the parameters on the model output, sensitivity indices (SI) were calculated:

$$SI_j = \sqrt{\frac{1}{K} \sum_{k=1}^{K} s_{j,k}^2} \tag{14}$$

where $j$ is the $j$th parameter, $k$ is the time instance within a sensitivity function, $K$ is the number of time instances for a particular sensitivity function and $s_{j,k}^2$ is the squared value of the relative sensitivity (eqn 13) of the variable $y$ to the $j$th parameter at a certain time instance $k$.

### Table 2. Applied model parameters, chosen within the ranges mentioned in literature

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Uncertainty range</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Gamma$ (MPa)</td>
<td>0.9</td>
<td>0.1–0.9</td>
</tr>
<tr>
<td>$L$ (m MPa$^{-1}$ s$^{-1}$)</td>
<td>$2.85 \times 10^{-9}$</td>
<td>$1.1 \times 10^{-10}–1.67 \times 10^{-4}$</td>
</tr>
<tr>
<td>$\phi$ (MPa$^{-1}$ s$^{-1}$)</td>
<td>$2.34 \times 10^{-7}$</td>
<td>$1 \times 10^{-7}–5.56 \times 10^{-5}$</td>
</tr>
<tr>
<td>$e_0$ (m$^{-1}$)</td>
<td>150</td>
<td>0–250</td>
</tr>
</tbody>
</table>

The model, consisting of a set of algebraic and differential equations, was implemented, simulated and calibrated using the modelling and simulation software package PhytoSim (Phyto-IT BVBA, Mariakerke, Belgium). PhytoSim was also used for the uncertainty and sensitivity analyses.

### RESULTS

Given the similar patterns in the ecophysiological variables for the three trees of each species, the average of the three trees per species was considered for further analysis. This way, an investigation of the more general species’ water use was conducted rather than focusing on intra-species variability or sensor-specific effects.

In Fig. 3 a typical example of the microclimatic conditions is presented. Throughout the measurement period described in this study, no rain fell and the soil water potential hardly changed, with an average difference between daily minimal and maximal values of $0.10 \pm 0.03$ MPa. Overall, an average daily soil water potential of $-1.99 \pm 0.5$ MPa was obtained. This value was further applied throughout the model study. Figure 4 indicates that, despite thriving in the same environment and showing

![Fig. 3. Typical microclimatic conditions during the dry measurement period. The interval DOY 245–249 is shown as an example.](https://academic.oup.com/aob/article-abstract/114/4/667/162358/162358)

![Fig. 4. Measured stem diameter (A), heat velocity as indication of sap flow (B) and stem xylem water potential (C) for *A. marina* and *R. stylosa* for the same time period as shown in Fig. 3. The interval DOY 245–249 is shown as an example; similar patterns were obtained throughout the measurement period.](https://academic.oup.com/aob/article-abstract/114/4/667/162358/162358)
similar dynamics in heat velocity (directly related to sap flow) and xylem water potential pattern, the diurnal SDV pattern was totally different for *A. marina* and *R. stylosa*. While for *A. marina* the stem diameter started to decrease during the morning (often after a short morning increase, as depicted in Fig. 4) and increased again in the evening and during the night, the stem diameter of *R. stylosa* increased during the morning and started to decrease in the afternoon, while remaining more or less constant at night. This pattern was consistent throughout the measurement period. Applying these measured variables to the functional model (Fig. 2), an expected pattern of storage water potential was obtained (Fig. 5). While in *A. marina* the simulated xylem water potential was more negative than the storage water potential during the morning and vice versa during the evening, the opposite happened in *R. stylosa*. The storage pressure potentials obtained were highly linearly correlated to the stem diameter patterns for both species ($R^2 = 0.99$). When the storage osmotic potential and corresponding osmotic equivalents were derived from the storage water potential (eqns 11 and 12), a clear diurnal pattern was obtained for both species (Fig. 6). Note that, for *R. stylosa*, $N_{eq}$ started to increase slightly behind $N_{eq}$ of *A. marina* during the morning. The uncertainty analysis indicated that the $N_{eq}$ patterns obtained remained similar within the possible range of parameter values, as indicated in Table 2 (Fig. 7). Moreover, the effects of $e_0$ and $L$ on $N_{eq}$ were very limited, as indicated by the low sensitivity indices of 0.004 and 0.0006, respectively. The osmotic equivalent pattern obtained was insensitive to the parameters $I'$ and $\delta$ as long as the pressure potential of the osmotic storage tissue was lower than $I'$, as then the model only

![Fig. 5](https://academic.oup.com/aob/article-abstract/114/4/667/162358)

**Fig. 5** Model results showing the diameter input and xylem and storage water potential output for *A. marina* (A) and *R. stylosa* (B). The interval DOY 245–249 is shown as an example; similar patterns were obtained throughout the measurement period.

![Fig. 6](https://academic.oup.com/aob/article-abstract/114/4/667/162358)

**Fig. 6.** Modelled osmotic potential of the storage tissue (A) and derived osmotic equivalents of the entire storage pool (B) for *A. marina* and *R. stylosa*. The interval DOY 245–249 is shown as an example; similar patterns were obtained throughout the measurement period.
Introduction

The observed differences in SDV patterns for A. marina and R. stylosa point to differences in carbon- and/or ion-related processes, as the measured hydraulic variables for the two species were highly similar (Fig. 4; M. W. Vandegehuchte et al., unpubl. res.). Even though root pressure has been documented to lead to stem diameter increase in the morning (De Swaef et al., 2013b), this phenomenon seems inadequate to explain the R. stylosa SDV pattern as the diameter increased continuously when VPD increased and xylem water potential decreased (Figs 4 and 5). Nevertheless, root pressure might have occurred in both mangrove species when VPD was low in the early morning. From our modelling results, it is clear that osmotic adjustment in the storage tissues, whether because of sugar unload- ing from the phloem (Sevanto et al., 2003, 2011; De Schepper and Steppe, 2010; De Schepper et al., 2013) or local accumulation or synthesis of organic compounds such as proline, glycinebetaine and mannitol and inorganic ions such as sodium and potassium (Popp, 1984a; Popp et al., 1993; Zimmermann et al., 1994; Krauss et al., 2008), contributed to the observed SDV patterns (Figs 6, 7 and 9). The hypothesis that varying osmotic concentrations in the storage tissues influence the SDV dynamics has not only been confirmed in experimental research, but has also been successfully implemented in model studies (Sevanto et al., 2003; Hölttä et al., 2006; De Schepper and Steppe, 2010; De Swaef et al., 2013a). In this study it was remarkable that, for the two co-occurring species, only a slight shift in the pattern of osmotic equivalents (Figs 6 and 9) led to an entirely different SDV pattern (Figs 4A and 8). As for every model, the results obtained were dependent on the estimated parameters. The uncertainty analysis showed that during the night the uncertainty about simulated \( N_{\text{eq}} \) was larger than that during the day (Fig. 7). Because the uncertainty bands were narrow during the morning increase and evening decrease in \( N_{\text{eq}} \), it can be suggested that the morning increase in \( N_{\text{eq}} \) for R. stylosa started slightly after that of A. marina. Overall, for both species a clear diurnal pattern was obtained. The uncertainty analysis did not take possible temporal changes of the parameters into account. It has been shown that the radial hydraulic conductivity \( L \) may vary depending on the activity and/or abundance of putative aquaporins (Steppe et al., 2012). Nevertheless, in this study the sensitivity analysis indicated that \( L \) had little influence on the \( N_{\text{eq}} \) pattern. Moreover, even on those days when a steep decline in diameter occurred for each species (Fig. 8), the pattern of osmotic equivalents remained largely unaffected (Fig. 9). This raises the question of what the cause of these \( N_{\text{eq}} \) patterns might be.

In general, the effects of plant carbon status on SDV are somewhat slower than the effects of water status as plants seem to maintain a rather steady supply of carbon to the sinks thanks to temporary storage of carbohydrates under light conditions and remobilization under conditions of low or absent photosynthesis (Geiger et al., 2000; Komor, 2000; De Swaef et al., 2013a). Nevertheless, De Swaef et al. (2013a) indicated that the high correspondence between variations in measured photosynthetically active radiation and growth rate of tomato highlighted the effect of varying plant carbon status on SDV. They showed that, besides the instantaneous dynamics of the plant water relations, osmotic variations in the phloem caused by varying soluble carbohydrate content for tomato plants with limited starch reserves could
explain the measured SDV. These authors hypothesized that the lower the starch reserve in plants, the more pronounced the influence of carbon status on SDV, which might be an indication of carbon starvation. This fits the expectations of Komor (2000), who stated that the balance between carbon storage and phloem carbon loading is subject to adaptation to meet growth requirements under special circumstances, enabling the modulation of symplastic and apoplastic carbon transport pathways by environmental factors. Thus, carbohydrate transport to and from the phloem may partly explain the $N_{eq}$ patterns as modelled for $A. marina$ and $R. stylosa$. As it is clear from the long-term measurements (Fig. 8) that both mangrove species were subjected to stress, the loading of sufficient carbohydrates during the daytime might have been essential to sustain turgor in the phloem tissues and avoid cell lesion. For $R. stylosa$, it even seemed essential to enable growth, as only during the morning increase in $N_{eq}$ was stem diameter able to exceed the maximum diameter observed the previous day. Moreover, it is likely that during the dry measurement period, carbohydrate reserves were limited; following the hypothesis of De Swaef et al. (2013a), this could imply that, during more favourable conditions, the $N_{eq}$ patterns might be less pronounced, which could affect the osmotic water potential of the storage tissues and hence the SDV pattern. Long-term measurements during the wet season could confirm this.

Next to carbohydrate loading and unloading, compartmentalization of ions and the formation of low molecular weight carbohydrates as osmotic adjustors might play a role in $N_{eq}$ and SDV dynamics. While these processes have been mainly studied at leaf level (Popp, 1984a, b; Rada et al., 1989; Sobrado, 2005; Naidoo, 2006), it is not unlikely that they may also affect the storage tissues at stem level. However, to further investigate the cause of the $N_{eq}$ and SDV pattern it would be necessary to sample the storage tissues to analyse their non-structural carbohydrate and ionic composition over time. Additionally, the simultaneous measurement of xylem and over-bark stem diameter variations might provide evidence of the $N_{eq}$ pattern as the lag between these two diameter measurements indicates the transport of carbohydrates and sink activity (Sevanto et al., 2003).

As the model applied in this study did not include detailed structural information, many structural differences between $A. marina$ and $R. stylosa$ were not directly taken into account. However, these differences likely contributed to the difference in SDV pattern. The leaves of $R. stylosa$ are much more succulent and more vertically orientated, which may cause differences in the timing and rate of carbohydrate assimilation and transpiration. Additionally, $A. marina$ is characterized by a patchy network of phloem strings throughout the xylem, which has not been shown for $R. stylosa$ (Schmitz et al., 2008). Moreover, $A. marina$ is considered to be a salt excluder, while $R. stylosa$ is known to rely predominantly on salt filtration at root level (Ball, 1988). These differences influence xylem sap salinity, which may affect xylem hydraulic conductivity (Lopez-Portillo et al., 2005). Inclusion of these features in the model might provide new insights into the underlying cause of the differing SDV patterns, but would greatly increase model complexity and require additional measurements (e.g. photosynthesis, respiration, wood anatomical features, leaf angle and xylem sap osmotic concentrations).
Nevertheless, despite the exclusion of this detailed structural information, the mechanistic approach applied here remains valid. The results indicate that both mangrove species are characterized by an endogenous osmotic adaptation that might be a function of environmental stress. The gradual overall decline in stem diameter indicates, however, that this osmotic adaptation was insufficient to avoid long-term overall shrinking, especially during days with a steep decline in diameter. During these days, the decline in xylem water potential was less compensated by osmotic adaptive processes, resulting in higher water loss from the storage tissues in both species. Still, SDV and hence growth for these species are clearly determined by both environmental dynamics and endogenous control mechanisms. This points up the importance of correctly implementing carbon-driven and ionic osmotic adaptations in functional-structural plant models to allow accurate predictions of plant behaviour. Our results indeed show that very small differences in osmotically active compound regulation may have drastic influences on important plant physiological measurement variables such as stem diameter variations. More thorough knowledge of how these features influence stem diameter variations may result in new insights into why species differ in growth patterns and, hence, which strategies are more beneficial, given specific environmental conditions. Moreover, it may allow assessment of the relative importance of endogenous osmotic adaptations, such as carbohydrate loading and unloading, local production and accumulation of osmotically active compounds, and environmental dynamics to long-term growth.

Conclusions

The present research highlights the importance of endogenous osmotic adaptations for SDV and, by extension, radial stem growth, illustrated for the two common mangrove species A. marina and R. stylosa. Based on ecophysiological measurements and mechanistic modelling, the daily pattern of osmotically active compounds in the storage tissues was unveiled, indicating that only a minor difference in this pattern can cause large differences in daily SDV. Our results stress the importance of correct implementation of endogenous osmotic processes in functional plant models.

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