The phenomenon of cavitation in grapevine...  
Unravelling implicated mechanisms

El fenómeno de la cavitación en vid...  
Descifrando los mecanismos implicados

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Originales: Recepción: 18/04/2017 - Aceptación: 07/11/2017

ABSTRACT

Cavitation is a physiological dysfunction that takes place in the xylem of water stressed plants. It leads to a loss of hydraulic conductance (kL) as the vessels are filled with air. This impacts water supply, water potential (ΨL) and canopy hydration. Stomatal closure is an effective response upon diminishing momentary or seasonal foliar hydraulic contents. Depending on each type of plant, stomata may close preventing catastrophic cavitations. This research intended to understand how stomatal control acts upon cavitation events in two contrasting grapevine varieties, Syrah and Grenache. A mechanistic was developed model based on the water and vapour fluxes, kL, stomata conductance (gs), and the vulnerability to cavitation of the xylematic tissue. The theoretical model explains how plants respond to drought and avoid catastrophic cavitation. Water stressed grapevines couple their gs with their kL in order to avoid embolism. It is not stomatal closure, by itsself, the controlling mechanism. Grapevines under mild water stress, do not need to completely close their stomata in order to avoid cavitation, therefore, photosynthesis is not completely impeded, and the cost in terms of carbon assimilation is less than expected for other species.

Keywords

Cavitation • stomatal conductance • hydraulic conductance • mechanistic model • Syrah • Grenache

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Resumen

La cavitation es una disfunción fisiológica que ocurre en el xilema de las plantas bajo déficit hídrico, y que entraña una pérdida de su conductancia hidráulica ($k_L$), cuando algunos vasos se llenan de aire. Esto incide negativamente sobre la oferta de agua y afecta el potencial hídrico foliar ($\Psi_f$) y la hidratación de la canopia. El cierre estomático es una respuesta efectiva ante la disminución del contenido hídrico. Dependiendo de la especie vegetal, los estomas suelen cerrarse para evitar la cavitation catastrófica. Mediante un modelo mecanístico, que se construyó teniendo en cuenta los flujos de agua y vapor, las $k_L$ y conductancia estomática ($g_s$), y la vulnerabilidad del xilema a cavitar; se probó que $g_s$ no es la única variable responsable de frenar la embolia. Se determinó que $g_s$ y $k_L$ están íntimamente asociadas y que este acople entre ambas conductancias es lo que frena la embolia. Se concluyó que, en la vid y bajo niveles de estrés hídrico moderado, no es necesario un cierre estomático para controlar la cavitation. Por esto, el mecanismo de control de la cavitation en la vid no conlleva un costo en términos de intercambio gaseoso.

Palabras clave

cavitation • conductancia estomática • conductancia hidráulica • modelo mecanístico • Syrah • Grenache

Introduction

Drought resistant crops have adaptive physiological and morphological traits that allow them to survive and grow under severe water deficit, resisting dehydration (14, 24). The origin of plant dehydration is embolism formation and catastrophic cavitation. In a dry soil, with low water potential, increasing xylem tension, triggers cavitation. This phenomenon consists on the formation of air bubbles inside the xylem vessels, and subsequently, the breakage of the water column (35). As consequence, the plant suffers a loss of hydraulic conductance ($k_L$) and desiccation (6, 11, 34, 35). Vulnerability curves relate the percentage loss of plant $k_L$ (PLC) or embolism to the increasing applied pressures that cause that drop of $k_L$. This pressure may be paralleled to the xylem tension, given that this positive pressure can be considered as equal, but opposite, to the negative pressure inside the xylem (1, 12, 31, 34).

It is well known that stomatal control prevents excessive water loss in an attempt to maintain $k_L$ and prevent desiccation under high evaporative conditions. Several authors have also concluded that stomatal adjustment limits cavitation (4, 8, 15, 19, 21, 28) and that the mechanism is subjected to hydraulic and hydromechanic laws (3, 4, 11). In general, grapevines have been considered as drought avoiding specie due to their efficient stomatal control (8). Most of the water that enters the plant (constituting $k_L$), leaves through opened stomata (depending on stomatal conductance, $g_s$) as transpiration ($E$) (9).

In this sense, many authors have already studied the relationship between $g_s$ and $k_L$, concluding that in most species, including grapevine, both conductances are tightly correlated (17, 18, 21, 29, 39). While aquaporins, in roots, act as the entrance valves for water (20, 28), stomata
in leaves act as water vapour exit valves that limit transpiration (E).

However, recent insights, cast doubt on the main role of stomatal closure on the embolism-avoidance strategy (38, 39), besides the fact that the actual involved mechanism is still not elucidated (2, 16).

In addition, grapevines have shown to own a highly resistant xylem (10) that cavitates at higher tensions than previously thought, keeping \( k_L \) between certain values before stomata respond. In this context, this research intended to study the cavitation phenomenon in grapevines and the mechanisms involved in its control. It tried to comprehend on a mechanistic manner, the stomatal functioning, its relation to the cavitation phenomenon, and the physical laws that rule them. This was achieved by complementing the construction of a functional and dynamic model with the comparison of two contrasting varieties, Syrah and Grenache, under two different water treatments, grown in pots, inside a greenhouse. These varieties were chosen because they have been reported as opposite in regards to stomatal behaviour, isohydric and anisohydric, respectively (8, 9). However, this classification is currently under strong debate (8, 14, 19, 21, 29). Given this controversy, tried to try the model as well as the varieties’ behaviour under these conditions.

**Model developing**

Several models have been developed describing and explaining the stomatal functioning (8). The model includes several sub-models and relates them in an attempt to explain embolism control by hydraulic traits in grapevine, adding the "vulnerability to cavitation element", and clarifying the coupling mechanism that achieves embolism control.

This model is based on the Ohm’s law analogue concept (37) that states that the flow \( (J_w) \) escaping through stomata, called transpiration (E), constitutes the impulsive force that drives water along the xylem vessels. This suction that occurs inside the xylem vessels is expressed in terms of water potential \( (\Psi; \text{MPa}) \). Finally, this \( \Psi \) difference between soil and leaves \( (\Delta\Psi) \) is what allows water to move from one place to the other (13, 37). 

\[
J_w = \frac{\Delta\Psi}{R} = k_L \Delta\Psi = E; \quad (1)
\]

\[
\Delta\Psi = \Psi_{\text{soil}} - \Psi_L \quad (2)
\]

where:

- \( J_w = E \), is transpiration \( \text{(mmol H}_2\text{O m}^{-2}\text{s}^{-1})} \)
- \( R = \text{hydraulic resistance} = 1/k_L \) \( (1/\text{mmol H}_2\text{O m}^{-2}\text{s}^{-1} \text{MPa}^{-1})) \)
- \( k_L = \text{hydraulic conductance} (\text{mmol H}_2\text{O m}^{-2}\text{s}^{-1} \text{MPa}^{-1}) \)
- \( \Delta\Psi = \text{water potential difference (MPa)} \)
- \( \Psi_{\text{soil}} = \text{soil water potential (MPa)} \)
- \( \Psi_L = \text{leaf water potential (MPa)} \)

Assuming that species like grapevines have null capacitance (25); (meaning that there is no water storage due to the water potential difference), \( J_w \) equals \( E \) (1), and may be expressed by Fick’s law as follows:

\[
E = \frac{(e_{sT(L)} - e_a)}{P_a (gs^{-1} + gb^{-1})} \quad (3)
\]

where:

- \( E = \text{transpiration (mmol H}_2\text{O m}^{-2}\text{s}^{-1})} \)
- \( (e_{sT(L)} - e_a)/Pa = q’ \) which is the difference of water vapour concentration between leaf and atmosphere, the vapour pressure gradient from leaf to air (dimensionless variable, it is a ratio of pressures).
- \( (gs^{-1} + gb^{-1}) = \text{sum of stomatal and boundary layer resistances (1/mmol H}_2\text{O m}^{-2}\text{s}^{-1})} \)
Then, by replacing (2) and (3) in (1), is obtained (4)

\[ \Psi_L = \Psi_{soil} - \frac{q'}{g_s^{-1} + g_s^{-1}} k_L \]  

where:

- \( q' \) = vapour pressure gradient from leaf to air.

Eq. (4) formalizes the relationship between \( \Psi_L \), \( q' \), \( g_s \) and \( k_L \).

The next step in this model development is to relate plant embolism (Emb) to the hydraulics described. Emb inversely depends on water potential (\( \Psi \)). For more negative values of \( \Psi \), higher percentages of Emb can be measured. In the model, Emb was interpreted by means of the mathematical adjustment of the sigmoid vulnerability curves of grapevines to a piece-wise defined function, shown as follows in figure 1.

In the first piece of the function, up to certain \( \Psi_L = \Psi_{L1} \), Emb equals zero (5). This part of the function is called "lag" and corresponds to a range of \( \Psi_L \) values where no embolism takes place. When \( \Psi_L \) diminishes because of increasing water deficit, exceeding \( \Psi_{L1} \), Emb linearly depends on \( \Psi_L \), and grows until maximum Emb - i.e. 100% - is achieved for \( \Psi_L = \Psi_{L2} \) (6). On or after \( \Psi_{L2} \), Emb equals 1 (or 100%; (7); figure 1).

\[ \begin{align*}
\text{Emb} &= 0 \quad \text{if } \Psi_L \leq \Psi_{L1} \\
\text{Emb} &= a + b \cdot \Psi_L \quad \text{if } \Psi_{L1} < \Psi_L \leq \Psi_{L2} \\
\text{Emb} &= 1 \quad \text{if } \Psi_L \geq \Psi_{L2}
\end{align*} \]

(5) \hspace{1cm} (6) \hspace{1cm} (7)

By trigonometry, equation (6) can mathematically be expressed as (8):

\[ \text{Emb} = \frac{\Psi_L - \Psi_{L1}}{\Psi_{L2} - \Psi_{L1}} \cdot \frac{\Psi_{L1}}{\Psi_{L2}} \quad \text{if } \Psi_{L1} \leq \Psi_L \leq \Psi_{L2} \]

Then, by replacing (4) in (6) is obtained the suffered Emb as output of the model, (Equation 9), for the part of the function in which Emb linearly depends on \( \Psi_L \).

\[ \text{Emb} = \frac{\Psi_{soil} - \Psi_{L1}}{\Psi_{L2} - \Psi_{L1}} \cdot \frac{\Psi_{L1}}{\Psi_{L2}} \quad \text{if } \Psi_{L1} \leq \Psi_L \leq \Psi_{L2} \]

(9)

The "lag" indicates the pressures under \( \Psi_{L1} \) where no embolism takes place. Note that \( \Psi \) are negative values.

El "lag" indica la presión bajo la cual no existe embolia. Los valores de \( \Psi \) son negativos.

**Figure 1:** Theoretical vulnerability curve for grapevines. \( \Psi_{L1} \), the pressure at which embolism starts, and \( \Psi_{L2} \), the pressure at which embolism reaches its maximum value.

**Figura 1:** Curva teórica de vulnerabilidad para vid. \( \Psi_{L1} \), la presión a la cual comienza la embolia, y \( \Psi_{L2} \), la presión a la cual existe embolía máxima.
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Values are individual measuremets. / Los puntos son valores individuales de medición.

Figure 2. Correlation between stomatal conductance ($g_s$, mmol H$_2$O m$^{-2}$ s$^{-1}$) and hydraulic conductance ($k_L$, mmol H$_2$O m$^{-2}$ s$^{-1}$ MPa$^{-1}$) for Syrah.

Figura 2. Correlación entre conductancia estomática ($g_s$, mmol H$_2$O m$^{-2}$ s$^{-1}$) y conductancia hidráulica ($k_L$, mmol H$_2$O m$^{-2}$ s$^{-1}$ MPa$^{-1}$) para Syrah.

The model also shows the feedback relationship between Emb and $k_L$. This feedback process states that Emb depends on $k_L$, while, at the same time, $k_L$ depends on Emb, as the former diminishes when the last rises.

Figure 3 (page 38), shows the dynamic mechanistic model that explains how the relationship between $g_s$ and $k_L$ controls embolism, and how embolism, in turn, modifies $k_L$ in a feedback loop. The $k_L$ (2) is the result of a $k_L$ before embolism ($k_Lbe$ or maximum $k_L$, 3), then affected by embolism (1). The $k_Lbe$ is a function of the time of day and $\Psi_{soil}$.

The model adopted several already existing models to fit into the general model.

The $g_s$ (5) is interpreted by the Buckley et al. model (2003); $g_b$ and leaf temperature ($T_L$) are expressed by the Campbell and Norman (2012) equations (4); $\Psi_{ng}$ (osmotic water potential of the guard cell), included in $g_s$ (5), is empirically expressed by Taiz and Zeiger (1998) and Tardieu and Simonneau (1998). The entry variables are $\Psi_{L1}$, $\Psi_{L2}$, time of day (hour), $\Psi_{sw}$, $T_a$, $e_a/P_a$, wind speed and solar radiation. The model output is Embolism (1).

Model parameterizing

To parameterize the model, it was designed an experiment with Grenache and Syrah plants, under field capacity and water stress. It was measured gas exchange and $\Psi_L$ during a complete day, from predawn to 18 h. The $k_L$ was calculated for each moment along the day, and a time dependent equation was then fitted to the data. Vulnerability curves were constructed and embolism achieved along the day was estimated.
The dynamic model was formulated to explain how embolism control takes place by stomatal and hydraulic coupling.

Las ecuaciones de la derecha se representan por números en el diagrama. Los mecanismos ambientales se formalizan mediante las ecuaciones de Campbell y Norman (1998) para la conductancia de la capa límite ($g_b$) y el déficit atmosférico ($q'$) que luego se incluyen en la ecuación de Buckley et al. (2005) para conductancia estomática ($g_s$) (5). La ecuación de Van den Honert (1948) (4) define el potencial hídrico foliar ($\Psi_L$) que luego definirá la conductancia hidráulica ($k_L$). La ecuación hidráulica antes de embolism ($k_{Lbe}$) formaliza la relación entre el $k_{Lmáx}$ y el $k_L$ después de la embolisa, siendo el formal sujeto a potencial hídrico del suelo ($\Psi_{soil}$) y el momento del día, según Vandeleur et al. (2009), (2) y (3). Finalmente, las ecuaciones expresando Embolism course son mostradas en (1). La ecuación (1) define Xylem Embolism basándose en los parámetros que caracterizan la vulnerabilidad de una especie definida. Nótese que este modelo integra modelos mecanísticos previos en uno solo, mediante la ecuación de embolism de la vid.

**Figure 3.** El modelo dinámico se construyó para explicar cómo la embolía es controlada por el acople entre la conductancia estomática e hidráulica.
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MATERIALS AND METHODS

**Vines and site**

The experiment was undertaken during the season 2012/2013 at the INTA’s Experimental Station, in Mendoza, Argentina.

A factorial experiment combining 2-year-old Syrah and Grenache grapevines and two water regimes was established on the summer of 2012. In quadruplicate, dormant own-rooted vines were removed from their 4-L pots and replanted on a sandy loam substrate on 15-L pots to allow good growth during the season.

Water regimes, named field capacity (FC) and water deficit (WD), were irrigated with 100% and 50% of the fraction of transpirable soil water (FTSW), respectively, as follow. Immediately after replanting all vines were irrigated to saturation and water treatments were applied; the FC treatment was watered every two days to maintain 100% FTSW whereas the WD treatment was left without irrigation for a week until it reached the targeted soil moisture of 50% FTSW (0.16 g/g). After the WD pots achieved the desired soil moisture, pots were watered every two days replenishing the transpired water. Water regimes were maintained for three months. Moisture was measured every two days using moisture probes (ECH2O EC-5 sensors, Decagon devices, USA). Vines were trained to one shoot and grown in a greenhouse with daily average temperature of 25°C and photosynthetic active radiation of 800 μmoles m\(^{-2}\)•s\(^{-1}\).

**Water potential and gas exchange measurements**

A portable photosynthesis system (CIRAS-2, PP Systems, Hertfordshire U.K) was used to measure instantaneous leaf gas exchange. Measurements were carried out every two hours from 6am and 6pm. The CO\(_2\) concentration of the incoming air was maintained at 375 μmol•mol\(^{-1}\). The same leaves used for gas exchange assessment were used to measure leaf water potential (Ψ\(_{L}\)), with the Scholander pressure chamber (Biocontrol, Córdoba, Argentina), using the procedure of Hsiao (37). Predawn water potential (Ψ\(_{pd}\)) was considered as a proxy to soil water potential (Ψ\(_{soil}\)).

Water vapour concentration at leaf temperature \(e_{sT(L)}/Pa;\ hPa\), was calculated via the equation of Teten.

\[
e_{sT(L)} = 6.11 \times \text{Exp} \left( \frac{17.502 \times T_L}{T_L + 240.97} \right) \quad (10)
\]

where:

- \(T_L\) = leaf temperature
- \(Hydraulic\ conductance\) was calculated through the Van den Honert law, with \(Ψ_{soil}\), \(Ψ_{L}\), and \(E\) for every assessed moment of the day.

Leaf embolism along the day (Emb) was estimated from the daily course curves of \(Ψ_{L}\) and the vulnerability curves, for each plant. Both curves were related and the positive pressures achieved for the vulnerability curves were directly linked to the \(Ψ_{L}\) measured along the day, assigning an embolism value to each moment and plant.

**Vulnerability curves**

The loss of \(k_H\) to increasing \(Ψ'_{L}\), *i.e.* cavitation, was studied by constructing vulnerability curves for each plant: after all water potential and gas exchange measurements finished, shoots were harvested and transported to the laboratory for vulnerability curves construction. Previously, every leaf was removed from the stem, and the cut surfaces were sealed with contact glue. Vulnerability
curves were constructed following the "Air Injection Long" method already described (10), using a double ended pressure sleeve connected to a Scholander pressure chamber (Biocontrol, Córdoba, Argentina). First, the shoots were flushed for 30 minutes using distilled, degassed 5% potassium hypochlorite (KClO) solution, removing all embolisms and obtaining maximum \(k_{\text{H}}\) (\(k_{\text{Hmax}}\)). Then, successive pressure cycles were imposed. The air pressure in the chamber was increased to a specific value and held for 10 minutes before it was reduced back to cero. Air pressure was successively increased to higher levels and hydraulic measures were taken, obtaining the \(k_{\text{H}}\) for each cycle. Percentage loss of conductivity (PLC) was calculated for each cycle relative to \(k_{\text{Hmax}}\).

\[
\text{PLC} = 100 \times \left(1 - \frac{k_i}{k_{\text{max}}} \right) = \frac{k_{\text{max}} - k_i}{k_{\text{max}}}
\]

**Statistics and data analysis**

Differences between treatments were assessed by multifactor and one-way ANOVA, followed by LSD test (\(p < 0.05\)) using StatGraphics Plus (Statistical Graphics Corp.; StatSoft, Inc., 2003). When homogeneity of variance was not reached, non-parametrical analyses were carried out.

**Results**

**Water relations: water potential and stomatal conductance**

The imposed water deficit had an evident effect on \(\Psi_{\text{L}}\). The water deficit (WD) treatments provoked a significantly lower (more negative) \(\Psi_{\text{L}}\) than those of the field capacity (FC) plants. For midday water potential (\(\Psi_{\text{md}}\)), interaction between treatments was significant (varieties vs. water treatments; \(p = 0.0333\)). As for stomatal conductance (\(g_s\)), the WD treatments had significantly lower values than the FC ones, for the whole day course (\(p = 0.01\); figure 4, page 41).

Once all vulnerability curves were obtained, each one was adjusted to a piece-wise defined function as described in the "Model developing" section: The first piece of the function, were no cavitation or embolism (Emb) has already happened, is called "lag" and equals cero for Emb; while the second part of the function, showing increasing cavitation, is fitted to a straight line equation (figure 1, page 36 and figure 5, page 41).

The ANOVA analysis for the straight line fitting parameters \(a\) (intercept) and \(b\) (slope) showed that neither statistically significant interaction, nor significant differences existed among varieties or water treatments (table 1, page 42). This means that under the experimental conditions, no xylem adaptation upon water stress occurred.

**Xylem embolism vs. water relations, throughout day**

Estimated embolism throughout the day was only achieved in four cases, because most of the plants had vulnerability curves with long lags that started from -1.5 MPa, while this \(\Psi_{\text{L}}\) value was generally not achieved in the greenhouse. As leaf embolism directly depends on \(\Psi_{\text{L}}\), both variables followed similar, though opposite daily courses. Lower \(\Psi_{\text{L}}\) corresponded to higher cavitation values (figure 6, page 42).

When it was observed the daily courses of embolism and \(g_s\) for the four plants that did cavitate, it was observed that there was no relation between both variables that could explain embolism control. Stomatal closure events (reduction on \(g_s\)) vs. embolism detention did not correlate throughout the day, meaning that stomatal conductance, per se, is independent of embolism (figure 7 page 42). This conclusion was already achieved theoretically by means of the model that clearly showed that the coupling between \(g_s\) and \(k_{\text{L}}\) (and not \(g_s\) alone), is, in fact, the controlling switch.
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Each point corresponds to the mean ± one SE for water deficit (WD; dotted line) and field capacity (FC; solid line) treatments.

Cada punto corresponde a la media ± un error para déficit hídrico (WD; línea de puntos) y capacidad de campo (FC; línea entera).

**Figure 4.** Daily course of stomatal conductance ($g_s$, mmol H$_2$O m$^{-2}$ s$^{-1}$) for Grenache (A) and Syrah (B).

**Figura 4.** Dinámica de la conductancia estomática a lo largo de un día ($g_s$, mmol H$_2$O m$^{-2}$ s$^{-1}$) para Grenache (A) y Syrah (B).

**Figure 5.** Vulnerability curve measured in Grenache under water stress [percentage loss of hydraulic conductance (PLC) vs. pressure ($\Psi_L$)]. The pointed $\Psi_{L1}$ and $\Psi_{L2}$ are the pressures at which xylem embolism starts and equals 100%, respectively. Under $\Psi_{L1}$ no embolism takes place.

**Figura 5.** Curva de vulnerabilidad medida en Grenache bajo estrés hídrico [porcentaje de pérdida de conductancia hidráulica (PLC) vs. presión ($\Psi_L$)]. Los valores de $\Psi_{L1}$ y $\Psi_{L2}$ señalados son las presiones a las que la embolia comenzó y alcanzó el 100%, respectivamente. Debajo de $\Psi_{L1}$ no existe embolia.
Table 1. Straight line fitting parameters b (slope) and a (intercept) for vulnerability curves in Grenache and Syrah cultivars, under field capacity (FC) and water deficit (WD). Data are means and p-values are from the ANOVA.

<table>
<thead>
<tr>
<th>Variable</th>
<th>b</th>
<th>a</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cultivar (C)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Grenache</td>
<td>-0.8855</td>
<td>-1.9221</td>
</tr>
<tr>
<td>Syrah</td>
<td>-0.8678</td>
<td>-1.4316</td>
</tr>
<tr>
<td>p value</td>
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<td>0.6209</td>
</tr>
<tr>
<td>Water treatment (W)</td>
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<td></td>
</tr>
<tr>
<td>FC</td>
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<td>-1.0930</td>
</tr>
<tr>
<td>WD</td>
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<tr>
<td>p value</td>
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<tr>
<td>p value (C × W))</td>
<td>0.8775</td>
<td>0.9983</td>
</tr>
</tbody>
</table>

Figure 6. Daily course of water potential ($\Psi_L$; solid line) and embolism (dotted line). Values are means from the three Syrah vines and the one Grenache plant suffering embolisms throughout the day.

Figura 6. Dinámica del potencial hídrico ($\Psi_L$; línea entera) y la embolia (línea punteada) a lo largo del día. Los valores son la media para tres plantas de Syrah y una planta de Grenache en condiciones de embolia.

Figure 7. Daily course of stomatal conductance ($g_s$; black squares) and embolism (black diamonds) for two vines that suffered embolism under no stress (A) and under water deficit (B). Clear independence between both curves is shown.

Figura 7. Cinética de la conductancia estomática ($g_s$; cuadros negros) y la embolia (diamantes blancos) a lo largo de un día para dos vides que sufren embolia bajo condiciones de riego suficiente (A) y bajo estrés hídrico (B). Notar la clara independencia entre ambas curvas.
Simulations

Figure 8, shows how embolism, $\Psi_L$, $g_s$ and $k_L$ behaved under water stress ($\Psi_{soil} = -0.2$), constant ambient circumstances and a variable guard cell osmotic adjustment ($\pi_g$). The $\pi_g$ modifies, in a significant manner, the stomatal adjustment (Buckley et al., 2003). In this case, it was modified $\pi_g$ by increasing its value by 20% for case A; and diminishing $\pi_g$ by 20% in case B. Notice that as $\Psi_L$ gets more negative and achieves -2 MPa, embolism starts and $g_s$ couples with $k_L$ ($\Delta g_s = 28$), allowing a maximum embolism of 45% for simulation B. For simulation A, $\Psi_L$ does not grow over the threshold value and no embolism takes place, the coupling is much less evident ($\Delta g_s = 16$).

![Figure 8. Simulations of embolism, $\Psi_L$, $g_s$ and $k_L$ under water stress ($\Psi_{soil} = -0.2$), constant ambient circumstances and variable guard cell osmotic adjustment ($\pi_g$). A: $\pi_g$ augmented by 20%. B: $\pi_g$ diminished by 20%.](image)
Discussion

These measurements and the ideated mechanistic model, demonstrated that embolism and $g_s$ are independent. Embolism depends on $g_s$ in addition to other physiological and ambient variables; like xylem vulnerability, $k_i$, difference on water vapour between leaf and atmosphere, and boundary layer conductance. In this study $g_s$ and $k_i$ were tightly associated ($R = 0.70$).

Through the model, it could be shown that the daily embolism restraint was linked to the variation that $g_s$ suffered in intimate relation with $k_i$, and not to $g_s$ itself. This tight relation between both conductances has already been widely observed in grapevine and trees (17, 21, 29, 39). This puts in evidence that $g_s$ responds to variations in $k_i$, and that both variables, in mutual interaction, control embolism in grapevines.

Apparently, $k_i$ and $g_s$ are related because, under drought, stomata operate allowing photosynthesis and preventing desiccation at the same time (7, 8). Consequently, $g_s$ must respond to $k_i$, since changes in $k_i$ influence plant and leaf water status (17). Therefore, the effect of $g_s$ as prime embolism restraint attributed in grapevines and other species, could be related to the fact that both conductances are strongly coordinated (21, 29, 39).

In relation to the generated model, it should be settled that the input variable $k_i$ is affected by the intrinsic embolism level, including a feedback process.

The model measures the phenomenon while $k_i$ grows, (and it grows despite of the portion of hydraulic conductivity that embolism captures, probably by the action of the root aquaporins) (36).

In fact, the model functions calculating embolism in a time $t$, from embolism in time $t-1$ (integrated in the input variable, $k_i$). This is incorrect in negative feedback mechanisms, as embolism control shows to be (30). It might also be probable that this embolism that affects $k_i$ is, partly, responsible for the stricter coupling of $g_s$ and $k_i$. Nardini and Salleo (2000), explained that in many species embolism cannot be completely avoided and that it could constitute the signal that stomata need to start closing up. This can be reinterpreted as follows: stomata actually respond to a lower $k_i$, caused by certain embolism formation, coupling itself to this changing $k_i$.

The obtained vulnerability curves were not different among treatments. Therefore, for the achieved water deficit, no xylem adaptation took place. It might be possible that differences among varieties were not evidenced because the stress levels achieved were not severe enough to generate these adaptation responses. It could also be possible that the three months period during which the plants lived was not long enough to let the xylem system anatomically adapt to the stressful situation. Besides, the possibility of discriminating vulnerability differences turns to be quite hard, given that the phenomenon shows great intrinsic variability in grapevines as in other species, (21, 22, 23, 35). For this study, variation coefficient for the "lag" value was 0.55.

Of great importance is to highlight that in grapevines, embolism control does not require complete stomatal closure, meaning that photosynthesis is not completely deprived, and the assimilation cost is not as high as expected for other species.
In this experiment, under severe water stress ($\Psi_{soil}=-0.2$ MPa, and 50% embolism) $g_s$ was significantly reduced but never achieved complete stomatal closure. In less stressful conditions, embolism is well controlled while stomata are maintained opened.

In 2011, Zufferey et al. (2011) found that stomata closed up only after 90% of embolism was achieved. This means that the plant can keep on photosynthesizing and, at the same time, avoid catastrophic cavitation (cavitation levels at which the plant cannot recover and dries). In this sense, it is one remarkable species that may require low amounts of water, and still produce quantity and quality of fruit, without the risk of suffering severe embolism events.

Acknowledgements
This work was supported by INTA EEA Mendoza, Argentina.
We thank Dr. Mark Matthews and Dr. Andrew McElrone (UC Davis, USA) for their critical reviews of the manuscript and valuable discussions.