Drought and Abscisic Acid Effects on Aquaporin Content Translate into Changes in Hydraulic Conductivity and Leaf Growth Rate: A Trans-Scale Approach

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The effects of abscisic acid (ABA) on aquaporin content, root hydraulic conductivity (Lp_r), whole plant hydraulic conductance, and leaf growth are controversial. We addressed these effects via a combination of experiments at different scales of plant organization and tested their consistency via a model. We analyzed under moderate water deficit a series of transformed maize (Zea mays) lines, one sense and three antisense, affected in NCED (for 9-cis-epoxy-carotenoid dioxygenase) gene expression and that differed in the concentration of ABA in the xylem sap. In roots, the mRNA expression of most aquaporin PIPs (for plasma membrane intrinsic protein) genes was increased in sense plants and decreased in antisense plants. The same pattern was observed for the protein contents of four PIPs. This resulted in more than 6-fold differences between lines in Lp_r under both hydrostatic and osmotic gradients of water potential. This effect was probably due to differences in aquaporin activity, because it was nearly abolished by a hydrogen peroxide treatment, which blocks the water channel activity of aquaporins. The hydraulic conductance of intact whole plants was affected in the same way when measured either in steady-state conditions or via the rate of recovery of leaf water potential after rewatering. The recoveries of leaf water potential and elongation upon rehydration differed between lines and were accounted for by the experimentally measured Lp_r in a model of water transfer. Overall, these results suggest that ABA has long-lasting effects on plant hydraulic properties via aquaporin activity, which contributes to the maintenance of a favorable plant water status.

During water deficit, abscisic acid (ABA) is involved in three strategies used by plants to avoid deleterious leaf dehydration. First, plants close stomata and decrease transpiration rate, with a consensus on the effect of ABA (Zhang and Davies, 1990a; Borel et al., 2001) but differences regarding its origin in the plant (Christmann et al., 2007; Endo et al., 2008). Second, plants decrease shoot growth in order to limit transpiration. The contribution of ABA to this reduction differs between studies, with either positive effects of ABA (Sharp, 2002; Sansberro et al., 2004; Thompson et al., 2007a) or negative effects (Zhang and Davies, 1990b; Ben Haj Salah and Tardieu, 1997; Bacon et al., 1998). Third, plants tend to control root water uptake and/or plant water status via root growth (Sharp, 2002) and root hydraulic conductivity (Lp_r, Kaldenhoff et al., 2008; Maurel et al., 2008).

The effects of soil water deficit and of ABA on Lp_r are controversial. Water deficit tends to decrease Lp_r (Lo Gullo et al., 1998; Zhang and Tyerman, 1999; North et al., 2004; Vandeleur et al., 2009), while ABA has the opposite effect in most studies (Morillon and Chrispeels, 2001; Thompson et al., 2007a). In a few studies, exogenous ABA had no effect or a negative effect on hydraulic conductivity (Wan and Zwiazek, 1999; Aroca et al., 2003), while in others a positive effect has been observed at both the cell level (Hose et al., 2000; Wan et al., 2004; Lee et al., 2005) and the whole root level (Quintero et al., 1999; Sauter et al., 2002; Schraut et al., 2005). However, these responses were transient (Hose et al., 2000) and were positive or negative depending on the ABA concentration (Beaudette et al., 2007).

Change in aquaporin mRNA and protein contents in response to water deficit and ABA is also a matter of debate (Kaldenhoff et al., 2008). ABA induces transcription factors that regulate the expression of PIPs (for plasma membrane intrinsic protein) aquaporins (Kaldenhoff et al., 1996; Shinozaki et al., 1998). However, exogenous ABA affects a larger number of PIP isoforms than water deficit (Jang et al., 2004), suggesting some degree of independence between ABA and drought signal transduction pathways (Mariaux et al., 1998). In addition, increase in PIP mRNA expression is...
often transient and dependent on ABA concentration (Zhu et al., 2005; Beaudette et al., 2007) and does not necessarily result in an increase in PIP protein content (Morrill and Chrispeels, 2001; Aroca et al., 2006).

Aquaporins play a key role in radial water transport in roots and leaves under both hydrostatic and osmotic gradients (Steedle, 2000; Tyerman et al., 2002; Maurel et al., 2008). The contribution of aquaporin-mediated water transport has been evaluated with inhibitors, namely mercuric salt (Martre et al., 2001), acid load (Tournaire-Roux et al., 2003), and hydrogen peroxide (H$_2$O$_2$; Ye and Steudle, 2006; Boursiac et al., 2008). These studies indicate that water transport by aquaporins accounts for 20% to 85% of the overall water transport depending on the species (Wan and Zwiazek, 1999; Barrowclough et al., 2000). Whereas the importance of aquaporins in Lp, has been demonstrated (Maurel et al., 2008), the resulting effect on overall plant conductance, leaf water potential, and leaf elongation rate is still poorly studied. Experiments using plants with modified aquaporin expression report differences in leaf water potential during the day (Lian et al., 2004) or during rewetting (Martre et al., 2002).

The purpose of this work was to test whether drought and ABA have consistent effects on plant hydraulic properties at different scales of plant organization, namely the abundance of aquaporin transcripts and proteins, the Lp, under both osmotic and hydrostatic gradients, the whole plant hydraulic conductance evaluated in steadily transpiring plants or upon rehydration, and the recovery of leaf elongation rate upon rehydration. Hence, we performed independent experiments at each scale of organization and then linked these scales with a model that allows the weighing of relative contributions of Lp and other possible causes on the whole plant hydraulic behavior.

To this aim, we have used maize (Zea mays) lines affected in the expression of the NCED/VP14 gene encoding the 9-cis-epoxycarotenoid dioxygenase enzyme, previously identified from the vp14 mutation (Tan et al., 1997). NCED/VP14 catalyzes the first specific step in ABA biosynthesis and affects ABA production when overexpressed (Thompson et al., 2000, 2007b) or underexpressed (Voisin et al., 2006). NCED/VP14 expression as well as NCED/VP14 protein content are indeed strongly correlated with ABA levels (Qin and Zeevaart, 1999). These lines were subjected to changes in soil water content, evaporative demand, or pressure on the root system, thereby affecting independently water deficit and the concentration of ABA in the xylem sap.

RESULTS

A Set of Maize Lines Transformed on the NCED/VP14 Gene Differed in [ABA]$_{xyl}$ Stomatal Conductance, and Transpiration Rate in Greenhouse Experiments

The genetic transformation was targeted to one NCED gene with sense and antisense constructs. It resulted in three antisense lines (AS1, AS2, and AS5) and one sense line (S). The concentrations of ABA in the xylem sap ([ABA]$_{xyl}$) differed significantly between AS, null transformants, and S plants in well-watered conditions as well as in moderate water deficit (Fig. 1A), consistent with the lower and higher transcript amounts of NCED/VP14 in AS and S plants, respectively (Supplemental Fig. S1). Null transformants resulting from each transformation event had insignificant differences in [ABA]$_{xyl}$. Therefore, they were pooled in all figures and are referred to as WT hereafter. Plants of the three AS lines had a low but still appreciable [ABA]$_{xyl}$ in both well-watered and dry

Figure 1. ABA concentration in the xylem sap (A), midday stomatal conductance (B; $g_s$), and leaf water potential (C) in three greenhouse experiments with high evaporative demand (experiments 1–3 in Table II; only days with nocturnal VPDs from 2 to 2.8 kPa were kept in the analysis). Two contrasting water regimes were compared, well watered and a mild water deficit with a soil water potential of −0.4 MPa, for three AS lines, one S line, and their null transformants (pooled and named WT). A, Xylem sap was harvested at the end of the night. B, Stomatal conductance was measured at noon. Because $g_s$ of WT plants differed between experiments, from 120 to 180 mmol m$^{-2}$ s$^{-1}$, it was normalized by the mean value in WT. C, Leaf water potential was measured at noon in nonexpanding leaves. Data were averaged from the three experiments (n > 10 in each condition and line). Error bars indicate confidence intervals at the 0.95 risk level. Bars associated with the same letter indicate nonsignificant differences (P > 0.05) in a Benjamini and Hochberg t test (Benjamini and Hochberg, 1995).
conditions (Fig. 1A; insignificant differences between AS lines), so their stomata closed under water deficit (Fig. 1B). The transpiration flow measured at midday in moderately droughted plants grown in the greenhouse was significantly higher in the three AS plants than in WT plants (Fig. 2A). Sense plants had the opposite behavior, with a high [ABA]$_{xyl}$ (>400 nM), a low stomatal conductance in both well-watered and droughted plants (Fig. 1, A and B), and a transpiration rate per unit leaf area 2-fold lower than that of WT plants (Fig. 2A). AS plants had comparable phenotypes to WT plants, except that they germinated more quickly and wilted slightly earlier upon water shortage. Because the three AS lines had similar behaviors, most studies were carried out in the AS5 line. S plants had a smaller leaf area than AS and WT plants, due to differences in leaf growth rate in well-watered conditions before the experiment.

**PIP Expression in Roots Was Highly Dependent on ABA Biosynthesis**

We compared by quantitative reverse transcription (RT)-PCR the expression levels of ZmPIP genes in roots of S, WT, and AS plants (Fig. 3). Plants were grown hydroponically and sampled in the early morning at the same phenological stage and root water potential (72-h polyethylene glycol [PEG] stress, −0.4 MPa) as in the water deficit treatments presented in Figures 1 and 2. Expression levels of the five PIP genes belonging to the PIP1 subgroup were significantly higher in S plants and lower in AS plants. The strongest effect was observed for ZmPIP1;2, ZmPIP1;3, and ZmPIP1;4, with a 9- to 10-fold difference in expression between AS and S plants ($P < 10^{-5}$). In the PIP2 subgroup, the expression levels of two PIP genes were increased in S plants (ZmPIP2;1 and ZmPIP2;2) and those of four PIP genes were decreased in AS plants (ZmPIP2;1, ZmPIP2;2, ZmPIP2;3, and ZmPIP2;6). Overall, for most PIP isoforms, the expression levels were affected by changes in ABA biosynthesis in a long-lasting way, with higher expression levels in S plants and lower levels in AS plants.

**Increased and Decreased ABA Biosynthesis Largely Affected the Protein Contents of Three PIPs in Roots and of Two PIPs in Leaves**

PIPs belonging to either PIP1 or PIP2 subgroups were chosen for further investigation using specific antibodies raised against each PIP (Fig. 4A). These were ZmPIP1;2, whose gene expression was the most increased in S plants and four proteins of the PIP2 subgroup; and ZmPIP2;1/2/2, ZmPIP2;5, and ZmPIP2;6, whose gene expression was the highest in maize roots in this study and in the study of Hachez et al. (2006). All ZmPIP immunoblot analyses revealed two major bands at about 28 and 55 kD, corresponding to monomeric and dimeric forms (Fig. 4A). Primary roots coming from AS lines showed strongly decreased amounts of the isoforms ZmPIP1;2 (−75%), ZmPIP2;1/2/2 (−43%), and ZmPIP2;5 (−49%; Fig. 4). Opposite effects on the amounts of the same proteins were observed in the S line, especially in ZmPIP1;2 (12-fold increase; Fig. 4). The intermediate band (around 40 kD) detected with ZmPIP2;6 antibodies corresponds to a cross-reacting unrelated protein (data not shown).

Leaves also showed differences in PIP protein amounts, although to a lesser extent than roots (Fig. 4B). S plants had higher amounts of ZmPIP1;2 and ZmPIP2;1/2/2 than WT plants (1.9- and 1.3-fold increases, respectively), with an opposite effect for AS plants (0.75 and 0.9). In contrast, no signals were detected from AS plants.
detected for ZmPIP2;5 and ZmPIP2;6, which were shown to be much less expressed in leaves compared with roots (Hachez et al., 2008).

**Increased and Decreased ABA Biosyntheses Affected the Lp, in Hydroponics, But This Effect Was Abolished by H₂O₂ Treatment**

Root systems were placed during the morning in a hydroponic solution, with a hydrostatic tension of −0.02 MPa applied to the hypocotyls of detopped plants. WT plants released a stable water flux for 40 min, which was multiplied by 3 and 0.5 in S and AS plants, respectively (Fig. 5A). H₂O₂ was then brought to the nutrient solution in order to decrease the hydraulic conductivity of the transcellular pathway (Ye and Steudle, 2006; Boursiac et al., 2008). This caused a steep decrease in water flux in both S and WT plants but had nearly no effect in AS plants. The water fluxes, therefore, were insignificantly different in the three treatments after the H₂O₂ treatment.

Differences in water flux were analyzed by measuring root hydraulic conductivities under hydrostatic (Lₚ) and osmotic (Lₚₒₛ) gradients of water potential. Lₚ was calculated as the slope of the relationship between the applied suction and the flux released by...
the root system at −0.02, −0.04, and −0.06 MPa. Lp,eg was calculated as the ratio between the free exudation flux and the gradient of osmotic potential between the nutrient solution and the sap released by the root system. Both Lp,h and Lp,eg were highly affected by the manipulation of ABA synthesis before the H2O2 treatment (Fig. 5, B and C). AS plants had a lower Lp,h and Lp,eg (−45% and −52%, respectively) than WT plants, while the S plants had higher Lp,h and Lp,eg (3- and 4-fold, respectively). Differences in Lp,h and Lp,eg between AS and WT plants were abolished after the H2O2 treatment, and those between S and WT plants were strongly reduced. The fact that flows were similar between WT and S plants after H2O2 treatment despite differences of Lp,h was due to a lower gradient of osmotic potential between the solution and the xylem sap in S plants.

Overall, these results show that the manipulation of ABA synthesis strongly affected the water flux through the root system via changes in the hydraulic conductivities under both hydraulic and osmotic gradients. This effect was strongly reduced or disappeared with the H2O2 treatment, with a drop in Lp,h that can be interpreted as the contribution of aquaporins to the water flux.

The Total Hydraulic Conductance between Soil and Leaves Increased with ABA Biosynthesis

The total hydraulic conductance between soil and leaves was estimated in a greenhouse experiment by dividing the water flux by the gradient of water potential between soil and leaves (Fig. 2C). Leaf water potential of AS plants was lower than that of WT plants in well-watered conditions as well as in water deficit (Fig. 2B; see also Fig. 1C), while transpiration rate was higher in AS than in WT plants (Fig. 2A). Sense plants had a significantly higher leaf water potential in all conditions, consistent with a lower transpiration rate. The three AS lines had lower hydraulic conductances, and the S line had a higher conductance than the WT plants (Fig. 2C). The differences in soil-plant conductance were not due to changes in root system architecture, because root length and area were very close in WT and AS plants and were lower in S plants than in WT plants (Fig. 2D). Hence, the higher hydraulic conductance observed in S plants in spite of a lower root area suggests a high Lp,h.

S and AS Lines Exhibited Marked Differences in Leaf Rehydration Half-Times and Recovery of Leaf Elongation Rate

We have evaluated the consequences of observed differences in Lp,h on the time courses of the recoveries of leaf water status and leaf elongation rate upon rehydration in a growth chamber experiment with soil-grown plants. Plants initially subjected to a soil water potential of −0.4 MPa and a vapor pressure deficit (VPD) of 2.5 kPa were rewatered and subjected to dark conditions at a VPD of 0.8 kPa that virtually stopped transpiration (time 0; Fig. 6). Before rewatering, leaf water potential differed between lines, consistent with experiments in
Leaf water potential recovered more rapidly in S plants and more slowly in AS plants compared with WT plants, with half-times of 0.5, 1.2, and 1.6 h, respectively (Fig. 6B, inset). Full recovery of leaf water potential occurred in 3, 5, and 7 h, respectively, and all lines reached a common water potential after 18 h. The recovery of leaf elongation rate also largely differed between the three lines, with the same trend as that of leaf water potential. It was more rapid than the recovery of leaf water potential, with half-times and times for full recovery of about 50% of those corresponding to leaf water potential. This difference in time course of recovery was consistently observed in three experiments in the growth chamber (data not shown).

The Differences in Time Course of Rehydration between Lines Are Accounted for by a Model Taking into Account the Measured Hydraulic Parameters

We have evaluated the relative contributions of $L_p$ and other possible causes of the differences in time courses presented in Figure 6, with the sensitivity analysis of a model. The model of stomatal control, biosynthesis of ABA, and water transfer is that of Tardieu and Davies (1993), widely tested since then and used by other groups (Dewar, 2002; Gutschick and Simonneau, 2002). It was combined with a model that calculates the water potential at leaf evaporating sites and with a module of capacitance, which allows the calculation of recovery rates (see “Materials and Methods’’). The parameters used in the model are presented in Table I, and the outputs are presented in Figure 7 for S, WT, and AS plants.

In transpiring plants (before time 0), leaf and xylem water potentials of simulated plants were lower in AS plants than in WT and S plants, because of a higher stomatal conductance that caused a higher water flux, the greenhouse (Figs. 1 and 2), with higher and lower values in S and AS plants, respectively, than in WT plants. Leaf elongation rates normalized by their maximum values under well-watered conditions for each line also differed before rewatering, with highest and lowest values for S and AS plants, respectively.

Leaf water potential recovered more rapidly in S plants and more slowly in AS plants compared with WT plants, with half-times of 0.5, 1.2, and 1.6 h, respectively (Fig. 6B, inset). Full recovery of leaf water potential occurred in 3, 5, and 7 h, respectively, and all lines reached a common water potential after 18 h. The recovery of leaf elongation rate also largely differed between the three lines, with the same trend as that of leaf water potential. It was more rapid than the recovery of leaf water potential, with half-times and times for full recovery of about 50% of those corresponding to leaf water potential. This difference in time course of recovery was consistently observed in three experiments in the growth chamber (data not shown).

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consistent with experimental data. Both this steady state and the recovery of leaf water potential after rehydration could be adequately simulated by the model without the necessity of additional parameters. In particular, a single value for the hydraulic conductance of leaf tissues \((g_{x-l})\) could be assumed for the WT, AS, and S lines. The model was then used to determine the contributions of several possible causes for the differences in time courses of leaf rehydration. (1) The elastic modulus had here a minor role because all studied lines had similar pressure-volume curves (Fig. 8). (2) The hydraulic conductance of the path between the xylem and leaf cells \((g_{x-l})\) could potentially have an important effect on the recovery of water potential according to the model. However, simulations with measured values of \(L_{pr}\) accounted for the whole differences between AS, WT, and S lines, leaving a marginal role or no role for differences in \(g_{x-l}\). This is consistent with the low differences in PIP amounts in leaves. Simulations were only slightly improved if a small difference in \(g_{x-l}\) was assumed between lines, but the effect was too small to justify different fitted values of \(g_{x-l}\) between lines (Table I). (3) The volume of water in the leaf tissues potentially has a large effect on the time courses of leaf water potential upon rehydration. It did not contribute to the difference between AS and WT plants, which had similar leaf areas and weights, but accounted for a large part of the difference between WT and S plants. When simulations were run with a common leaf water volume, the time course of rehydration still differed between WT and S plants, but with a half-time in S plants that increased to 1 h versus 0.5 h in experimental data (Fig. 9).

Overall, this sensitivity analysis suggests that the \(L_{pr}\) measured in detached root systems accounted for a large part of the differences between lines in whole plant conductance, both in steady-state transpiration and during rehydration. In AS plants, which presented no difference in leaf volume, the increase in half-time of rehydration could be entirely attributed to differences in \(L_{pr}\). Part of the difference between S and WT plants was due to a difference in leaf volume, but \(L_{pr}\) still accounted for 23% of the difference in half-time of rehydration.

**DISCUSSION**

Consistent effects were observed across different scales of plant organization, suggesting a simpler picture for the role of ABA on plant hydraulic properties than that presented in the introduction. This was probably because differences in ABA supply to the shoot were stable over a long period, thereby avoiding the complexity of transient effects of exogenous ABA application (Hose et al., 2000; Zhu et al., 2005), and because the effects of ABA and drought were not confused.

**Effects of Overproduction or Underproduction of ABA on Aquaporin Gene Expression and Protein Content**

ABA increased gene expression and protein content of most PIP isoforms and never decreased them. This is consistent with the results of Jang et al. (2004) obtained in roots of Arabidopsis \((Arabidopsis thaliana)\), in which exogenous ABA increased the expression of 12 PIPs, although one isoform was decreased by the same treatment. However, this is the first time, to our knowledge, that a long-lasting effect was observed, with a stable increase in expression levels for most PIPs, in opposition to the results of Zhu et al. (2005) in maize roots, in which an application of exogenous

![Figure 7](#)

**Figure 7.** Simulated time course of recovery of water potential in the soil, roots, xylem, and leaves in AS, WT, and S plants. Symbols represent experimental measurements of leaf water potential.

![Figure 8](#)

**Figure 8.** Relationship between leaf relative water content and leaf water potential in AS (AS5), WT, and S plants (experiment 4 in Table II). Each point shows one coupled value of relative water content and water potential corresponding to one leaf.
ABA caused an increase in expression of only one PIP (ZmPIP1;2) after 24 h. This difference is probably due to the fact that changes in ABA concentration were obtained through a transgenic approach in this study instead of the application of exogenous ABA.

The ABA effects on PIP expression resulted in differences between lines in PIP protein contents, in opposition to the data obtained by Morillon and Chrispeels (2001) in Arabidopsis leaves. However, we observed some discrepancies between mRNA and protein levels. ZmPIP2;6 did not show any significant variation at the protein level, although its mRNA expression profile was strongly correlated with endogenous ABA concentration. On the contrary, ZmPIP2;5 was found to have a fairly similar mRNA expression level in the different lines, whereas its protein level showed a clear correlation with ABA content. These observations suggest the existence of posttranscriptional mechanisms to regulate the amount of PIP proteins.

PIP1 gene expression was more affected than that of PIP2 in our study, especially that of ZmPIP1;2, resulting in protein content more affected in ZmPIP1;2 than in PIP2s. This is important in view of the regulating role of this PIP (Zelazny et al., 2007). Gene expression of the PIP2 subfamily was lower than that observed by Hachez et al. (2006) without osmotic or water stress.

This could be due to genotypic differences but also to an ABA-independent negative effect of drought on PIP2 expression, due to mild water stress (Jang et al., 2004).

### ABA Effect on Hydraulic Conductivity via Aquaporin Activity Modulation

The variation in the amount of ZmPIP isoforms in AS, WT, and S lines was correlated with the measured differences in Lp, indicating that PIP aquaporins play a crucial role in controlling Lp. ZmPIP2;1/2,2 and ZmPIP2;5 proteins were reported to be highly expressed in the exodermis and the endodermis, suggesting that they are involved in root radial water movement (Hachez et al., 2006). In addition, the detection of a polar localization of ZmPIP2;5 to the external periclinal side of epidermal cells indicates an important role in water transfer from the soil into the roots (Hachez et al., 2006). The increase in ZmPIP1;2 amount could also induce an increase in the water permeability of root cells. Although PIP1 proteins expressed alone in maize protoplasts were retained in the endoplasmic reticulum, their coexpression with PIP2s resulted in a relocalization to the plasma membrane due to physical interaction (Zelazny et al., 2007). When coexpressed in oocytes, this interaction enhanced the water membrane permeability (Chaumont et al., 2000; Fetter et al., 2004).

The fact that differences in Lp disappeared or were largely decreased after H2O2 treatment suggests that they were due to differences in aquaporin activity. Boursiac et al. (2008) showed that millimolar concentrations of H2O2, as applied in this work, enhance the accumulation of PIPs in intracellular structures, resulting in 60% inhibition of Lp. This difference in activity between lines was probably due, at least in part, to the difference in the amount of PIP isoforms in both AS and S plants. We do not rule out the possibility that ABA also had an effect on aquaporin gating, which is the most likely mechanism in short-term experiments with artificial ABA (Hose et al., 2000). However, the latter effect disappeared after 3 h in that study (Hose et al., 2000), suggesting that the effect of ABA on Lp, initially due to aquaporin gating, is linked to other mechanisms, including change in PIP content over

### Table II. Summary of experiments carried out in this study

| Experiment | Lines | Place | Medium | Tmin/Tmax | VPDmax | PFD | Measured Variable |
|------------|-------|-------|--------|-----------|--------|-----|-------------------|
| 1          | AS1, AS2, AS5, WT | Greenhouse | Soil | 19/26 | 2.5 | 500 | g_C [ABA], \(g_C\) |
| 2          | AS1, AS2, AS5, WT | Greenhouse | Soil | 19/30 | 3 | 700 | g_C [ABA], \(g_C\) |
| 3          | AS1, AS2, AS5, WT, S | Greenhouse | Soil | 21/29 | 2.8 | 660 | \(g_C\), \(g_C\), transpiration, root and leaf area |
| 4          | AS1, AS2, AS5, WT, S | Growth chamber | Soil | 28/28 | 2.8 | 400 | \(\psi\) and leaf elongation rate during rehydration |
| 5–8        | AS5, WT, S | Growth chamber | Hydroponics | 20/24 | 0.8 | 400 | Aquaporin expression and protein content, pressure-volume, Lp |
longer time scales. Therefore, the difference between the clear effect of ABA on Lp, observed here and the transient (Hose et al., 2000) or nonexisting (Wan and Zwiazek, 2001; Aroca et al., 2003) effects observed by other authors may be due to the difference in mechanisms between endogenous ABA and artificially fed ABA or to a rapid degradation of exogenously applied ABA.

Differences in Lp, Translate into Changes in Whole Plant Hydraulic Conductance

Two independent ways of evaluating whole plant hydraulic conductance, each of which has its drawbacks, gave consistent results. (1) The conductance calculated from the water flux and the gradient of water potential in the soil-leaves continuum differed between lines. It represents the overall water transport in plants and in the soil and can be affected by any difference in soil water potential (which has a large effect on soil hydraulic conductivity) or in root system architecture (which affects the distance that water has to cross from the soil to the nearest root). This was probably not the case here, because the soil water content did not differ between pots carrying plants of each line and because measured root areas were similar in AS and WT plants and lower in S plants. (2) The difference in the time course of leaf water potential upon rehydration also indicates a difference in overall plant conductance, although other differences between genotypes could also account for this effect. The model showed that the differences in Lp, measured in detached root systems were sufficient to account for the longer half-time of recovery of leaf water potential in AS plants. The shorter half-time of recovery observed in S plants was only partly due to measured differences in Lp, with a contribution of the leaf volume to the behavior of S plants.

Leaf Elongation Rate followed the Changes in Hydraulic Properties upon Rehydration

The effect of ABA on leaf growth via changes in aquaporin activity and Lp, is usually observed by the superposition of several effects of ABA at different time scales. In particular, leaf elongation rate during the night was faster and slower, respectively, in AS and S plants than in WT plants in both well-watered and water deficit treatments. This suggests an intrinsic negative effect of ABA on leaf elongation rate consistent with earlier results (Zhang and Davies, 1990b; Ben Haj Salah and Tardieu, 1997; Bacon et al., 1998) and caused a smaller plant size of S plants. During the day, a lower stomatal conductance and higher hydraulic conductivity in S plants compared with WT and AS plants translated into a higher leaf water potential and a higher leaf elongation rate normalized by its value in well-watered plants, as shown in Figure 6.

The complex situation described above led us to concentrate this study on the changes in leaf elongation rate during rehydration. The first surprising result was that growth recovery after rehydration had half-times of around 1 h. The recovery of leaf elongation rate was even faster than that of leaf water potential and responded to ABA production with half-times lower for the S line and higher for the AS line than for WT plants. The beneficial effect of ABA on Lp, therefore, had consequences on leaf water status and then on leaf elongation rate upon rewatering.

MATERIALS AND METHODS

Genetic Material

A series of transformed maize (Zea mays) lines were analyzed, one S line (S) and three AS lines (AS1, AS2, and AS5; already presented in the study of Voisin et al. (2006) with the names FCN-001a, FCN-002a, and FCN-005b, respectively). AS lines were engineered by transformation of the line A188 with Agrobacterium tumefaciens carrying the pRec318 superbinary vector (Ishida et al., 1996). The pRec318 vector was obtained after recombination between A. tumefaciens pSB1 superbinary vector and pBIOS518, which contains a 1.8-kb coding sequence of NCED/VP14 under the constitutive viral promoter CaMV35S and the upstream Nos terminator. pBIOS518 carries the BAR marker gene driven by the actin constitutive promoter and its first intron and is stopped by the Nos terminator. The sense transgenic line, PVC-002b (S), was engineered by biolistic and was obtained by cotransformation of the pCaMV35S/NCED/VP14 plasmid, which carries the NCED/VP14 sequence in the sense orientation under the CaMV35S constitutive promoter and upstream Nos terminator, and the pDM302 vector, which contains the following cassette: actin promoter and its first intron, marker gene BAR, and the Nos terminator (Gordon-Kamm et al., 1990). Primary transformants were grown in vitro, acclimated in the greenhouse (16-h day, 24°C, 80% relative humidity and 8-h night, 20°C, 100% relative humidity). They were then crossed with the line A188, and the resulting material (T1) was used in this study. It was checked by Southern blotting that transformed plants contained one or two copies of the transgene. To identify T1 plants carrying the insertion (50% of the plants), a PCR test (Tag Hot Start master mix; Qiagen) was performed in each studied plant on a 50-mg fresh weight leaf sample at the third leaf stage. Primers were designed for the NCED/VP14 gene (forward, 5′-AGTGTGTCCTCAGCACGATTCA-3′; reverse, 5′-CACGGACGATACCACA-3′). In all cases, nontransformed plants, sisters from transformed plants, were used as controls.

Plant Growth Conditions in Hydroponics

Maize seeds were placed in tubes with a wet sponge and germinated at 24°C in the dark and saturated air. After 3 to 5 d, the germinated seeds were placed in the growth chamber with their roots bathing in a continuously aerated solution with the following composition: 0.25 mM CaSO4, 0.8 mM KNO3, 0.6 mM KH2PO4, 0.2 mM MgSO4·7H2O, 0.4 mM NH4NO3, 2·10−3 mM MnSO4·4·10−3 mM ZnSO4, 0·4·10−3 mM CuSO4, 0·2·10−3 mM Na2MoO4·2H2O, 1·6·10−2 mM H2BO3, 0·04 mM Fe-EDDHA, and 2·5 mM MES, pH 5.5 to 5.8. The hydroponic solution was renewed every third to fourth day. Air temperature and relative humidity were measured at plant level every 30 s with two sensors (HMP35A; Vaisala Oy). The temperature of the meristematic zone was measured with fine copper-constantan thermocouples (0.2 mm diameter), inserted between the sheaths of leaves 2 and 3 of four to six plants per experiment. Photosynthetic photon flux density (PPFD) was measured every 30 s using two sensors (LI-190SB from Li-Cor and SOLEMS 01/012/012). All climatic data were averaged and stored every 15 min in a data logger (Campbell Scientific, LTD-CR10X Wiring Panel). Environmental conditions are summarized in Table II.

Plant Growth Conditions in the Greenhouse

Plants were grown in PVC columns (0.23 m diameter and 0.4 m height) containing a 4060 (v/v) mixture of filtered loamy soil (particle diameter ranging from 0.1 to 4 mm) and organic compost. Columns were filled with 10.5 kg of soil and sampled for measurement of water content at filling time. Seeds were sown at 2.5 cm depth and watered with water until the two-leaf stage.
and with a modified one-tenth-strength Hoagland solution after that. Environmental data were measured as above and are presented in Table II (experiments 1–3).

Soil water content was determined by weighing columns automatically every 15 min. Differences in weight were attributed to changes in soil water content, after correction for the increase in mean plant biomass as a function of phenological stage and for the effect of displacement transducers. A water-release curve of the soil was obtained by measuring the soil water potential of soil samples with different water contents, in the range 0.4 to 0.2 g cm⁻³ (VP4-T Dewpoint Meters; Decagon Devices), thereby allowing calculation of the mean soil water potential in each soil column every 15 min.

**Plant Growth Conditions in the Growth Chamber, Rewetting Experiment, and Measurement of Leaf Elongation Rate**

Plants with mild water deficit (soil water potential of −0.4 MPa) and growing in the greenhouse under high evaporative demand were transferred at noontime to a growth chamber with a moderate evaporative demand (Table II, experiment 3; VPD = 2.5 kPa, 28°C, PPFD = 400 μmol m⁻² s⁻¹). They were left to transpire under these conditions for 3 h, during which leaf water potential was measured with a pressure chamber (Soil Moisture Equipment) and leaf elongation rate of the sixth leaf was monitored every 15 min with rotational displacement transducers (601-1045 Full 360°). At time 0, plants were rewetted until retention capacity and placed in dark conditions at noontime to a growth chamber with a moderate evaporative demand (Table 2). Sap samples were then stored at −80°C until analysis. ABA content was calculated as the difference between weight and dry weight divided by the difference between fresh weight and dry weight.

The concentration of ABA was measured in sap samples obtained by pressurizing leaves in the pressure chamber in the early morning (experiments 1–3). Sap samples were then stored at −80°C until analysis. ABA concentration was then measured in crude samples of xylem sap by radioimmunoassay (Quarrie et al., 1988) as described previously (Barrieu and Simonneau, 2000).

**RNA Extraction and Quantitative RT-PCR (Experiment 8)**

RNA extraction was carried out as described by Hachez et al. (2006). Briefly, total RNA was extracted from thoroughly ground frozen root sections using an RNeasy Plant Extraction Minikit (Qiagen). DNase I digestion was performed on a column during RNA extraction according to the manufacturer’s recommendations. cDNA synthesis and real-time PCR were then performed as described by Hachez et al. (2006). Results were normalized using two maize internal control genes, α-tubulin (gi: 450292; Hachez et al., 2006) and actin (gi: 168493; forward primer, 5’-TGGCTCAGAAAGGCTTCTGG-3’; reverse primer, 5’-GCATCTCATGTGGAACATGC-3’). ZmPIP primers were those used by Hachez et al. (2006) targeting a 100-bp-long sequence from the 3’ untranslated region. NCED mRNA levels were assessed using 20-bp-long specific primers (forward primer, 5’-ATCAAGAGGCGGTTGACCTGTA-3’; reverse primer, 5’-GCATCTCCGTGCGCTTGAAC-3’). PCR efficiency of the NCED primers was checked and found to be appropriate.

### Protein Extraction and Analysis

Measurements were carried out on plants grown hydroponically until the six-leaf stage (experiment 5) and subjected to a 72-h water stress obtained with PEG (−0.4 MPa; PEG8000, 150 g L⁻¹). Root tips (4 cm) and the leaf elongation zone (6 cm) were collected during early morning, immediately placed in a tube immersed in liquid nitrogen, and stored at −80°C. To prepare the microsomal fraction, 1 g of tissue was ground in 1.5 mL of solution (250 mM sorbitol, 50 mM Tris-HCl [pH 8], and 2 mM EDTA) containing 0.6% polyvinylpyrrolidone, 0.5 mM dithiothreitol, and protease inhibitors (1 μg mL⁻¹ each of leupeptin, aprotinin, antipain, chymostatin, and pepstatin [Sigma]). All subsequent steps were performed at 4°C as described (Hachez et al., 2006). Fifteen micrograms of crude microsomal membranes was solubilized for 15 min at 60°C in a buffer (80 mM Tris-HCl, 2% SDS, 10% glycerol, 0.005% bromphenol blue, and 1% dithiothreitol), and the proteins were separated by SDS-PAGE on a 12% polyacrylamide gel. After electrophoresis, the gel was incubated for 5 min in semidyfer buffer (48 mM Tris, 39 mM gly, 20% methanol, and 0.375% SDS) before semidyfer transfer to a polyvinylidene difluoride (Millipore) membrane for 40 min at 23 V. Western-blot analysis was performed on the polyvinylidene difluoride membrane using antisera raised against the N-terminal peptides of ZmPIP1;2, ZmPIP2;1, ZmPIP2;5, and ZmPIP2;6 (Chauvinn et al., 2001). The antisera raised against ZmPIP2;1 also recognized ZmPIP2;2 (Hachez et al., 2006). The dilutions used were 1:1,000 for the ZmPIP1;2, ZmPIP2;5, and ZmPIP2;6 antisera and 1:1,000 for the ZmPIP2;1/2.2 antisera. Detection and protein quantification were carried out using a Kodak 4000R image station and the associated software. Three different exposure times were used per experiment for accurate protein quantification.

### Measurement of Lp, in Nutrient Solution

Measurements were carried out from 10 AM to 1 PM, while Lp was maximum (experiments 6 and 7). The free exudation rate of the excised seminal root system was measured by collecting exuded sap with a micro-pipette and weighing it in microtubes. The osmotic potentials of the sap and of the nutrient solution were measured with a vapor pressure osmometer (Vapro 5520; Wescor). A hydrostatically driven xylem sap flow was triggered in excised root systems by applying a vacuum-induced tension (Freund et al., 1998). The seminal root system was excised by sectioning the mesocotyl and then fixed tightly to a silicon tube using low-viscosity dental paste (President Light; Coltene Whaledent). The silicon tube was connected to a vacuum port equipped with a tension gauge. A two-valve system driven by the data logger allowed automatic control of the tension applied to the seminal roots. After 30 min, needed to stabilize the exudation rate, the water flux was...
measured every 10 min. The suction force applied to the root system was varied every 10 min in a standardized way (0, -0.02, -0.04, -0.06, -0.08, -0.04, -0.02, and 0 MPa). The water flow across the root system was measured with a water trap made of a 2-mL tube filled with dry cotton and inserted onto the tubing between the roots and the vacuum port. Water flow was measured by weighing the sap absorbed by the cotton that was renewed every 10 min. H₂O₂ treatment (2 mM) was applied to the root system after 40 min, and the depressurization protocol was then applied in the same way. At the end of each experiment, the root system was scanned and root area (A) was determined with an image analyzer.

The \( L_p \) under an osmotic gradient (\( L_{p,u} \)) was calculated as follows:

\[
L_{p,u} = J (\pi_{exp} - \pi_{cell})^{-1} \times A^{-1} \tag{1}
\]

where \( J \) is the water flux through the root system without depressurization, \( \pi_{exp} \) and \( \pi_{cell} \) are the osmotic potentials of the sampled sap and of the nutrient solution, respectively, and \( A \) is the area of the root system. \( L_p \) under a hydrostatic gradient (\( L_{p,h} \)) was calculated from the slope of the regression between water flow and the suction applied to the root system (\( df/\Delta \pi \)):

\[
L_{p,h} = (df/\Delta \pi) \times A^{-1} \tag{2}
\]

**Model of Water Transfer**

The model describes stomatal control, biosynthesis of ABA, and water transport is that of Tardieu and Davies (1993). This model calculates stomatal conductance, \( [ABA]_{xyl} \), water flux, root and the water potentials in transpiration sites from soil water potential, light intensity, and air VPD. The parameters used in calculations were those of the original paper, except for (1) the parameter that relates ABA biosynthesis to root water potential, which was calibrated for WT, AS, and S lines according to the measured \( [ABA]_{xyl} \) and (2) the root hydraulic conductance, which was calculated from the hydraulic conductivities presented in Figure 5 and the measured root area presented in Figure 2 (Table I).

Changes in the model were added to allow simulation of the water potential of leaf cells. The water potential at leaf evaporating sites (\( \Psi_{cell} \)) was calculated from the xylem water potential (\( \Psi_{xyl} \)), the water flux (\( J \)), and the conductance to the flux from xylem to leaves (\( g_{cel} \)):

\[
\Psi_{cell} = \Psi_{xyl} - J/g_{cel} \tag{3}
\]

This conductance was estimated by measuring leaf and xylem water potentials of maize plants at the same stage (Table I). Leaves presented a capacitance that was calculated from the pressure-volume curve presented in Figure 8 and the estimate of the leaf volume. They were a sink for water when their potential was lower than that of the evaporating sites and a source otherwise. The water flow corresponding to leaf growth during the considered 6 h of the simulation was negligible. Therefore, we solved the differential equation for calculating the cell water potential (\( \Psi_{cell} \)) and the water flux from the xylem to the leaf cells (\( J_{cel} \)):

\[
J_{cel} = dV_{cel}/dt = -g_{cel} (\Psi_{xyl} - \Psi_{cell}) \tag{4}
\]

where \( g_{cel} \) is the conductance of the pathway from xylem to leaf cells. At each time \( t \), the flux through roots and xylem was the sum of the transpiration flux (\( J \)) and of the water flux from the xylem to the leaf cells (\( J_{cel} \)), so

\[
\Psi_{cel} = \Psi_{xyl}(t) - R_p (t \times (J + J_{cel})) - R_t \times (J + J_{cel}) - (J_{cel} - J)/g_{cel} \tag{5}
\]

where \( R_p \) is the resistance to water flow in the soil, calculated as by Tardieu and Davies (1993), \( R_t \) the resistance to water flow in the root system, calculated from \( L_p \) measured experimentally and root area (Table I), and \( g_{cel} \) is as in Equation 4 (Table I).

A first calculation of \( J_{cel} \) was derived from Equations 4 and 5:

\[
J_{cel} = (g_{cel}^{-1} - \Psi_{xyl}/\Psi_{cell} - R_p \times J - R_t \times (J - J_{cel}))/\left(1 + L_p \times R_p + L_t \times R_t \times L_t/g_{cel}^{-1}\right) \tag{6}
\]

A second expression of \( J_{cel} \) was obtained from the relationship between \( V_{cell} \) and \( \Psi_{cell} \):

\[
V_{cel} = V_{sat} + \left(V_{res} - V_{sat}\right) \times (1/(1 + (\alpha - (\Psi_{cell} - \Psi_{sat})^{n})/(1 - n)) \tag{7}
\]

where \( V_{sat} \) is the leaf volume at saturation (early morning), \( V_{res} \) is the residual volume at the water potential at the end of the experiment, and \( \alpha \) and \( n \) are the parameters of a Van Genuchten equation fitted on the pressure volume curve. \( \Psi_{sat} \) was calculated as the difference in \( V_{cell} \) between two different times for the optimization process of resolution of the differential equation. \( \alpha \) was the only fitted parameter of the model. The elastic modulus of leaves was common to the three lines because the curves relating turgor to volume were indistinguishable in AS, S, and WT lines (Fig. 8).

**Supplemental Data**

The following materials are available in the online version of this article.

**Supplemental Figure S1.** Expression levels of the NCED/VP14 gene in roots in AS (ASS) and S plants relatively to their value in WT plants.

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