RESEARCH PAPER

Gating of aquaporins by light and reactive oxygen species in leaf parenchyma cells of the midrib of Zea mays

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Abstract

Changes of the water permeability aquaporin (AQP) activity of leaf cells were investigated in response to different light regimes (low versus high). Using a cell pressure probe, hydraulic properties (half-time of water exchange, $T_{1/2} \propto 1$/water permeability) of parenchyma cells in the midrib tissue of maize (Zea mays L.) leaves have been measured. A new perfusion technique was applied to excised leaves to keep turgor constant and to modify the environment around cells by perfusing solutions using a pressure chamber. In response to low light (LL) of 200 $\mu$mol m$^{-2}$ s$^{-1}$, $T_{1/2}$ decreased during the perfusion of a control solution of 0.5 mM CaCl$_2$ by a factor of two. This was in line with earlier results from leaf cells of intact maize plants at a constant turgor. In contrast, high light (HL) at intensities of 800 $\mu$mol m$^{-2}$ s$^{-1}$ and 1800 $\mu$mol m$^{-2}$ s$^{-1}$ increased the $T_{1/2}$ in two-thirds of cells by factors of 14 and 35, respectively. The effects of HL on $T_{1/2}$ were similar to those caused by H$_2$O$_2$ treatment in the presence of Fe$^{2+}$, which produced $\cdot$OH (Fenton reaction; reversible oxidative gating of aquaporins). Treatments with 20 mM H$_2$O$_2$ following Fe$^{2+}$ pre-treatments increased the $T_{1/2}$ by a factor of 30. Those increased $T_{1/2}$ values could be partly recovered, either when the perfusion solution was changed back to the control solution or when LL was applied. 3mM of the antioxidant glutathione also reversed the effects of HL. The data suggest that HL could induce reactive oxygen species (ROS) such as $\cdot$OH, and they affected water relations. The results provide evidence that the varying light climate adjusts water flow at the cell level; that is, water flow is maximized at a certain light intensity and then reduced again by HL. Light effects are discussed in terms of an oxidative gating of aquaporins by ROS.

Key words: Aquaporin, cell pressure probe, glutathione, hydraulic conductivity, hydrogen peroxide, light, oxidative gating, reactive oxygen species, Zea mays.

Introduction

Plants experience various regimes of light intensity, which is an important issue during photosynthesis. In response to light, stomata open to fix carbon diffusing in as CO$_2$, and plants lose substantial amounts of water using the same pathway. Therefore, plants developed strategies to manage the use of water efficiently in response to changes in the light regime. It has been shown that the overall hydraulic conductance of leaves is affected by light (Sack et al., 2002; Lo Gullo et al., 2005; Nardini et al., 2005; Tyree et al., 2005; Sack and Holbrook, 2006; Cochard et al., 2007). In part, this could have happened either by a change of the conductance of the vascular system (Zwieniecki et al., 2001) or by changes of water permeability of the living cells of xylem parenchyma or bundle sheath (cell $L_p$; Nardini et al., 2005; Tyree et al., 2005). The change in cell $L_p$ in response to light was speculated to occur via a gating of water channel activity sitting in the cell membranes (aquaporins, AQPs). However, there has been, to date, no direct evidence of a gating of AQPs by light. According to Cochard et al. (2007), the level of AQP transcripts increased in response to light. Kim and Steudle (2007) recently started to fill the gap by measuring changes in cell $L_p$ in response to light. In leaf cells of maize, they found that turgor as well as light affected cell $L_p$. To separate the effects of light from those of turgor, they kept the turgor constant and found that, when light was varied at low absolute values of light intensity, AQP activity increased...
as the light intensity increased, which was in agreement with earlier results obtained at the whole leaf level (see above).

There is increasing evidence that many environmental factors affect AQP s (Steudle, 2000; Maurel and Chrispeels, 2001; Tournarie-Roux et al., 2003; Lee et al., 2005). Light could be one of them. It is plausible that a gating of AQPs by light may involve an oxidative gating of AQP activity (Henzler et al., 2004; Ye and Steudle, 2006). As photosynthesis produces O2 and reduction equivalents, damage by light may involve an oxidative gating of AQP activity (Henzler et al., 2004; Ye and Steudle, 2006). It is plausible that a gating of AQPs by reactive oxygen species (ROS) may be anticipated by reactions involving the partial or complete reduction of O2 and the production of ROS (such as superoxide, H2O2, and the hydroxyl radical -OH; Foyer and Noctor, 2000; Dietz, 2008). In the Chara internode and maize roots, an oxidative gating of AQPs has been demonstrated by Henzler et al. (2004) and Ye and Steudle (2006). These authors showed that AQPs could be closed by H2O2/OH, and that closure was reversible. According to Aroca et al. (2005), treatment with 100 µM H2O2 decreased the root hydraulic conductance in a chilling-sensitive, but not in a chilling-tolerant maize genotype. Oxidative gating may be a common response to different kinds of stresses (Pastori and Foyer, 2002; Xiong et al., 2002), and it may provide appropriate adjustments in water relations (Ye and Steudle, 2006).

In the present study, the experiments of Kim and Steudle (2007) have been extended from low to high levels of light intensity to provide further insights into the mechanism(s) of a gating of cell Lp via changes of AQP activity. Measured cells were parenchyma cells in the midrib of maize leaves (Westgate and Steudle, 1985; Wei et al., 1999). These cells are (i) located in the vicinity of photosynthetically active cells, stomata, and xylem vessels; (ii) easy to puncture due to their large size; and (iii) also suitable for long-term measurements in single cells (up to 6 h), since the midrib tissue could be well fixed, and has, therefore, been used as a model system by Kim and Steudle (2007). A cell pressure probe was applied to access cell Lp. The perfusion of excised leaves allowed infiltration of leaf tissue via cut xylem vessels to adjust apoplastic concentrations and to apply H2O2 and the antioxidant glutathione (GSH). Most importantly, turgor could be kept constant by infiltration, i.e., this variable affecting AQP activity was excluded during light treatments. Different effects of light at two light regimes were investigated [low light (LL), up to 650 µmol m−2 s−1; and high light (HL), 800 µmol m−2 s−1 and 1800 µmol m−2 s−1]. In response to LL, cell Lp increased. HL regimes, however, caused a decrease, as did the infiltration of solutions of H2O2/OH. The data are discussed in terms of gating of AQPs by ROS.

Materials and methods

Plant material

Maize (Zea mays L. cv. symphony) plants were grown from caryopses in soil in a greenhouse of Bayreuth University as described by Kim and Steudle (2007). In the lab, where the experiments were performed, the ambient light intensity was ~5 µmol m−2 s−1 (20–25 °C; relative humidity = 30–60%). Experiments were conducted on 4- to 8-week-old plants, which were 0.8–1.2 m tall and had about eight leaves. The parenchyma cells used in the pressure probe experiments were located in the midrib region 100–200 mm behind the tip of leaves. Cells were measured at a distance of 100–200 µm from the abaxial surface of the midrib, i.e. in the same range as those used by Kim and Steudle (2007). They usually contained no chlorophyll, but they were close to photosynthetically active cells (~50 µm away; see cross-section in Fig. 1 of Kim and Steudle, 2007). Third or fourth leaves from the plants were used for experiments. Leaf blades were cut to a length of ~0.3–0.4 m. About 40 mm of the leaf tip was cut and removed to enhance transport through xylem vessels by perfusion.

Experimental set-up using a CPP

A Leitz manipulator (Wetzler, Germany) was used to carry a cell pressure probe (CPP; Kim and Steudle, 2007). It was fixed on a thick iron plate placed on a heavy stone table to minimize vibrations during the CPP measurements. Using magnetic bars, a cut leaf was fixed upside down on a metal sledge to expose the midrib securely for measurements of cell hydraulics. The basal cut end of the leaf was encased in a pressure chamber (Fig. 1), where it was immersed in...
a solution of defined composition (see below). The micro-
capillary attached to the CPP was filled with silicone oil (oil
type AS4 from Wacker, Munich, Germany). It had a fine
tip of up to ~8 µm in diameter. When midrib cells were
punctured, a meniscus formed between the cell sap and oil
within the tip and tended to move away from the cell. Using
the metal rod of the CPP, the meniscus was re-adjusted at
a position close to the surface of the midrib. This restored
cell sap volume close to its original value. The pressure
transducer of the CPP measured turgor pressure ($P$, which
was recorded by a computer. To investigate the hydraulic
conductivity of cell membranes ($L_p$), hydrostatic relaxations
of turgor were induced by instantaneously moving the
meniscus to another position and keeping it there. To avoid
effects of large pressure pulses (‘energy injection’; Wan
et al., 2004), peak sizes of hydrostatic relaxations were
< 0.1 MPa. The half-times of hydrostatic relaxation, i.e. the
time taken for pressure pulses to reach half of the original
peak ($T_{1/2}$), were inversely proportional to $L_p$. Further
details of the function of the CPP can be found in previous
studies (e.g. Steudle, 1993; Henzler and Steudle, 1995). In
this study, $T_{1/2}$ was used as a direct measure of changes of
$L_p$ to reduce the effect of error propagation when calculating
$L_p$did not change significantly during measure-
ments (Kim and Steudle, 2007). To avoid variations
between cells, relative changes of $T_{1/2}$ rather than absolute
changes were often compared. Treatments were performed
on individual cells. This required measurement of hydraulic
parameters of a given cell for up to 4 h.

Perfusion of leaves with solutions of defined
composition (defined apoplastic environment)
The apoplastic environment of cells was varied during
treatments, for example by infiltration of AQP inhibitors.
In order to perfuse a leaf tissue at constant turgor pressure
during illumination and to provide a certain ionic apop-
plastic environment, the pressure chamber (see above) was
provided with different solutions, which were infiltrated at
a pressure of 0.1–0.2 MPa above atmospheric. When pressurizing the basal cut end of the xylem, guttation
droplets appeared at the leaf margin and at the cut surface
of the leaf outside of the pressure chamber. To test how
long it took for the perfusion solution to move across the
tissue, 70 mM H$_2$O$_2$ solution was injected in the chamber.
The appearance of H$_2$O$_2$ in the other cut end and leaf
margins was qualitatively tested with KI and starch
solution. Droplets in the midrib of the cut end were
collected and added to KI–starch solution. The presence of
H$_2$O$_2$ in the droplets changed the solution to a blue colour.
According to the tests, the time required for the solution to
move across the tissue and to modify the apoplastic solution
was 5–20 min, depending on the pressure applied to the
pressure chamber and resistance in the water pathway of the
leaf. During 1 h perfusion at 0.2 MPa, the amount of

solution that passed through the leaf tissue was of an order
of magnitude which was similar to the volume of the excised
leaf. Assuming the apoplast was 10% of the leaf volume
(Kosala Ranathunge, University of Bonn, personal commu-
nication), this was equivalent to 10 times the volume of the
apoplast. The perfusion solution could be exchanged via
a tube leading into the chamber, which could be closed by
a valve, when the chamber was pressurized. This allowed
exchange of the apoplastic solution quickly and also for the
solution to be changed back to the original solution during
measurements with an individual cell. To minimize variability
between cells, it was intended to measure effects and
reversibility on individual cells.

Illumination experiments
In order to run the CPP experiments, both the micro-
capillary and tissue near the cell punctured had to be
illuminated using an Osram halogen lamp (150 W, Xen-
ophot HLX, Munich, Germany) through glass fibre optics
(Scott, Mainz, Germany). The light intensity at the tissue
level was ~50 µmol m$^{-2}$ s$^{-1}$. The $T_{1/2}$s were continuously
measured after a cell was punctured. Using the glass fibre
optics, the light intensity could be increased to values as
high as 200 µmol m$^{-2}$ s$^{-1}$ (LL). Usually, the effects of light
treatment were measured following changes of $T_{1/2}$. To
apply light intensities which are similar to that in the field
during a bright day (~2000 µmol m$^{-2}$ s$^{-1}$; Nobel, 1999),
a screen projector as used for powerpoint presentations
(AstroBeamX211 with 200 W UHP lamp, A+K, UK) was
positioned at a distance of 200–300 mm from the specimen.
This was usually turned on for 15 min to produce light
intensities of between 800 µmol m$^{-2}$ s$^{-1}$ and 1800 µmol m$^{-2}$
s$^{-1}$ at the tissue level [as regulated by the distance of the light
source from the leaf surface (HL)]. The UV content of the
light source (UV-A and UVB of the spectrum from 290 nm to
390 nm) was measured by a UV light meter (PeakTech,
Ahrensburg, Germany). It was verified that the light source
contained less UV light (1–2 W m$^{-2}$) than the sunlight
measured outside on a bright day (PAR=1900 µmol m$^{-2}$

s$^{-1}$, UV=47 W m$^{-2}$). Relaxations ($T_{1/2}$s) were continuously
measured during light and other treatments.

Perfusion solutions
As a control, leaves were perfused with a solution of
0.5 mM CaCl$_2$ solution, which resulted in a constant turgor
pressure for at least 4 h in a given cell, even under
conditions of high rates of transpiration such as occur at
HL. Effects of H$_2$O$_2$ were measured by changing from
0.5 mM CaCl$_2$ to solutions that contained either 20 mM or
70 mM H$_2$O$_2$ in addition to the CaCl$_2$. In the presence of
natural levels of Fe$^{2+}$, this should produce -OH (Fenton
reaction: H$_2$O$_2$+Fe$^{2+}$→Fe$^{3+}$+OH–+OH). In order to en-
hance the level of -OH, leaves were perfused with 0.5 mM
CaCl$_2$+3 mM FeSO$_4$ for 1–2 h, and then the perfusion
solution was changed to 0.5 mM CaCl$_2$+20 mM H$_2$O$_2$ to
increase the oxidative stress. To see the recovery from stress
treatment, solutions were changed back to 0.5 mM CaCl₂. To demonstrate the effects of the antioxidant GSH, leaves were perfused by 3 mM GSH and 0.5 mM CaCl₂, and the perfusion solution was then changed back to the control solution. The solution containing GSH had a pH of 3.5. Other solutions had a pH of 4.8, which was similar to the known apoplastic pH of leaves in different plant species (4.7–5.1; Felle and Hanstein, 2007).

Results

Half-times of hydrostatic relaxations, \( T_{1/2} \propto 1/L_p \)

From the pressure relaxations, \( L_p \) values could be worked out, when the elastic modulus (\( e \)) was also measured. However, in most cases, \( T_{1/2} \)s (inversely proportional to cell \( L_p \)) were taken as a measure of cell \( L_p \) because \( e \) did not change during treatments for a given cell (Kim and Steudle, 2007). In most cases, there was no transient effect of puncturing on \( T_{1/2} \) (inversely proportional to \( L_p \)) as observed with young maize roots (Wan et al., 2004). When there was such an effect, it took 10 min, which was sufficient to achieve a constant \( T_{1/2} \). At the ambient light intensity of 50 \( \mu \text{mol m}^{-2} \text{s}^{-1} \), the \( T_{1/2} \)s of cells from leaves infiltrated with 0.5 mM CaCl₂ ranged from 0.3 s to 35 s (mean±SD, 4.8±7.7 s, \( n=31 \) cells; Fig. 2). The variability of \( T_{1/2} \) was also known in the same tissue of intact maize plants (Kim and Steudle, 2007). However, >75% of those cells (24 out of 31 cells with 0.5 mM CaCl₂) had \( T_{1/2} \)s of <4 s, and one-third of cells had \( T_{1/2} \)s between 1.0 s and 2.0 s (12 out of 31 cells; Figs 2, 3A). Cells infiltrated with 0.5 mM CaCl₂/3 mM FeSO₄ had \( T_{1/2} \)s similar to those infiltrated with 0.5 mM CaCl₂ (range¼0.8–17 s; mean±SD, 3.6±3.8 s, \( n=24 \) cells; 21 out of 24 cells had a \( T_{1/2} \) <4 s; Figs 2, 3B). Usually, cells probed from one leaf had similar \( T_{1/2} \)s. For example, six cells probed from the same leaf had \( T_{1/2} \)s ranging from 0.9 s to 2.3 s. The large \( T_{1/2} \)s occasionally measured were observed in a few leaves and they were probably caused by a closure of AQPṣ, even in the absence of inhibitors or HL. Those \( T_{1/2} \)s could be reduced by light treatment of up to 650 \( \mu \text{mol m}^{-2} \text{s}^{-1} \), according to the earlier results of Kim and Steudle (2007).

Responses to \( \cdot \text{OH} \) (Fe²⁺ and H₂O₂)

When adding 20 mM H₂O₂ to the reference solution (0.5 mM CaCl₂) at a light intensity of 50 \( \mu \text{mol m}^{-2} \text{s}^{-1} \), there was no effect on \( T_{1/2} \) (four cells tested). Most probably, this means that Fe²⁺ in apoplasts was not adequate to produce a sufficiently high level of \( \cdot \text{OH} \) (Fenton reaction; \( \text{H}_2\text{O}_2+\text{Fe}^{2+}=\text{Fe}^{3+}+\text{OH}^-+\cdot \text{OH} \)). However, when the concentration of H₂O₂ was raised to 70 mM, \( T_{1/2} \) increased by a factor of 4. When perfusing with 3 mM FeSO₄+0.5 mM CaCl₂, \( T_{1/2} \)s were similar to those when using only 0.5 mM CaCl₂ (see above). There was a marked increase in \( T_{1/2} \) when 20 mM H₂O₂ solutions were perfused following 2 h treatments with Fe²⁺ (Fig. 3C). It can be seen

Fig. 2. Frequency histogram of half-time of water exchange, \( T_{1/2} \) (inversely proportional to cell \( L_p \)). More than 80% of cells from excised leaves which were perfused with either 0.5 mM CaCl₂ or 0.5 mM CaCl₂+3 mM FeSO₄ had \( T_{1/2} \)s of <4 s. There were a few leaves showing only large \( T_{1/2} \)s.

Fig. 3. Representative relaxation curves used to measure half-times of water exchange, \( T_{1/2} \). (A) The half-time of cells from excised leaves which were perfused with 0.5 mM CaCl₂ typically ranged between 1.0 s and 2.0 s. A reversible oxidative gating of AQPṣs was demonstrated by following an individual cell in B to D. (B) Half-times of cells in the presence of Fe²⁺ were similar to those in CaCl₂ and this was used as the control. (C) On the same cell as in B, addition of 20 mM H₂O₂ produced OH⁻ by the Fenton reaction and caused a substantial increase in \( T_{1/2} \) by a factor of 27, i.e. \( L_p \) was reduced by the same factor. (D) Subsequent exchange to 0.5 mM CaCl₂ to remove radicals resulted in a partial recovery of \( T_{1/2} \) (\( L_p \)), i.e. the effect was reversible, at least to some extent.
from the figure that $T_{1/2}$ increased by a factor of as much as 14 most likely due to the action of ·OH (Fig. 3C). Changing back to 0.5 mM CaCl$_2$ again reduced $T_{1/2}$ by 50% within 30 min, to a value of 700% of the original (Fig. 3D). Recovery was observed for only up to 30 min, because long-term measurements following oxidative responses and recovery in individual cells were demanding; however, there could have been further recovery during long-term measurements. It was stressed that, during measurements, turgor pressure was kept constant. Similar experiments showing inhibition by H$_2$O$_2$/Fe$^{2+}$ treatment and partial recovery were repeated in three different cells. Overall, $T_{1/2}$ increased due to the treatment by a factor of 30, and recovered to 600% of the original value within 30 min after changing back to the control perfusion solution (Fig. 4). More data showing substantial increases in $T_{1/2}$ due to H$_2$O$_2$/FeSO$_4$ treatment are shown for other cells in Fig. 5A (see below).

Responses to low light of 200 μmol m$^{-2}$ s$^{-1}$

The response to LL (200 μmol m$^{-2}$ s$^{-1}$) was measured in cells which originally had a low $T_{1/2}$ of ~1 s in the FeSO$_4$/CaCl$_2$ solution, but were then inhibited to have a large $T_{1/2}$ by the addition of 20 mM H$_2$O$_2$. When cells exhibited long $T_{1/2}$s in the presence of H$_2$O$_2$/FeSO$_4$, LL of 200 μmol m$^{-2}$ s$^{-1}$ caused a significant reduction of $T_{1/2}$ by a factor of 7 within 30 min (Fig. 5A, n=3 cells). In one experiment, $T_{1/2}$ increased by a factor of 4 caused by treatment with 70 mM H$_2$O$_2$ could be recovered to its original value by LL of 200 μmol m$^{-2}$ s$^{-1}$ (data not shown). For cells which already had a large $T_{1/2}$ of >2 s in the control solution (AQPs already closed), LL treatment reduced the $T_{1/2}$ (Kim and Steudle, 2007). It can be seen from Fig. 5B that $T_{1/2}$s were reduced by a factor of 2 within 30 min ($n$=4 cells). In one of those four cells, $T_{1/2}$ levelled off to a small value within 30 min of light treatment, but not in the others. Longer light treatments may further reduce $T_{1/2}$. It can be seen from these results that although light effects were substantial, there was a considerable variability in the LL responses.
Responses to high light of 800 μmol m⁻² s⁻¹ and 1800 μmol m⁻² s⁻¹

Although there was an overall trend of a reduction of $T_{1/2}$ following HL treatment, this was statistically not significant. As during the LL treatment, there was a substantial variability between cells. For example, HL of 800 μmol m⁻² s⁻¹ and 1800 μmol m⁻² s⁻¹ reduced the $T_{1/2}$ during the first 5–10 min of the 15 min period of illumination, but then $T_{1/2}$ increased substantially by factors of 14 and 35, respectively (Fig. 6A, n=3–4 cells; type-1 response). In total, 10 cells showed an increase in $T_{1/2}$ due to HL. The 15 min treatment was chosen as the maximum treatment which could be applied at constant turgor and with stable cells. Longer HL treatments caused a continuous decrease in turgor pressure, which indicated damage to the cells. In this respect, punctured cells could have been more prone to damage in the presence of light stress than other cells (see Discussion). In eight out of 10 cells, the $T_{1/2}$s remained large when light was turned off for 30 min, which was the maximum period of time measured. There were, however, cells, which were hardly affected by HL, as seen in Fig. 6B (n=5 cells; type-2 response). In those cells, there was only a 37–86% reduction within 15 min. Those five cells were from two leaves, and no cells in those leaves showed a type-1 response. Maximum temperature changes on the leaf surface due to illumination with 800 μmol m⁻² s⁻¹ and 1800 μmol m⁻² s⁻¹ were 3 °C and 7 °C, respectively.

Responses to GSH

Cells pre-treated with the antioxidant GSH were exposed to a 15 min period of illumination at the intensity of 1800 μmol m⁻² s⁻¹. As seen in Fig. 7, a cell which had been perfused with 3 mM GSH+0.5 mM CaCl₂ for 0.5–1.0 h increased its $T_{1/2}$ by a factor of 46 by HL, as in the absence of GSH. However, in the presence of GSH, $T_{1/2}$ recovered back to its original level within 15 min when the light was turned off (n=5 cells). This was different from treatment in the absence of GSH. Pre-treatment with 3 mM GSH of 24 h duration had pronounced ameliorative effects. In contrast to short treatments, $T_{1/2}$s showed no significant increase with HL (n=4 cells). To confirm that the lack of response was due to the presence of GSH rather than a coincidence originating from the variable responses between cells (one-third of cells did not react to HL; see above), by following one cell tests were conducted to determine whether the $T_{1/2}$ in control solution increased in response to HL, and then that the addition of GSH caused recovery of the $T_{1/2}$ to its original value. On the same cell, light treatment in the presence of GSH caused only a temporary increase in $T_{1/2}$, which eventually recovered to the original value at the ambient light intensity (data not shown). In the reverse type of experiment using 3 mM GSH, there was no response of the $T_{1/2}$ by 15 min HL treatment following treatment with 3 mM GSH solution for 3 h. On the same cell, the exchange of GSH solution by the reference solution and waiting for its complete removal (~1 h) caused a substantial increase of $T_{1/2}$ by 15 min HL treatment (data not shown). Overall, the results indicate that there was a clear ameliorative effect of GSH on cell $Lp$ ($T_{1/2}$). It is unlikely that the effect of the GSH solution is due to changes in pH (from 4.8 to 3.5), since cytosolic acidification is known to reduce water permeability by a gating of AQPs (Tournaire-Roux C et al., 2003). The effects of HL may be related to oxidative stress in the presence of HL (see Discussion).

Fig. 6. Two types of response to high light (HL) treatment. (A) HL of 800 μmol m⁻² s⁻¹ (black bars, n=4 cells) and 1800 μmol m⁻² s⁻¹ (grey bars, n=3 cells) increased the $T_{1/2}$ at constant turgor in CaCl₂ solution. The increase in $T_{1/2}$ began ~10 min after HL was turned on. The largest $T_{1/2}$s caused by HL were significantly larger than those measured at ambient light (AL) intensity before HL treatments ($P<0.05$, t-test). During 15 min after the light was switched off, $T_{1/2}$s remained large, i.e. not reversible within 15 min. (B) There were cells in which $T_{1/2}$ did not increase but decreased due to the effect of HL of 1800 μmol m⁻² s⁻¹ (grey bars, n=5 cells). Values are means ± SD and are shown as relative changes. The absolute values of $T_{1/2}$ are shown in the inset.
Cells pre-treated with 3 mM GSH+0.5 mM CaCl\(_2\) for 0.5–1.0 h show an increase in T\(_{1/2}\) due to HL as in the absence of GSH (as in Fig. 5). In the presence of GSH, in contrast, there was a recovery within 15 min after the light was switched off. Cells pre-treated with GSH for 24 h (grey bars, n=4 cells) did not show an increase in T\(_{1/2}\) by HL (P<0.05, t-test). Values are means ±SD and are shown as fold changes. The absolute values of T\(_{1/2}\) are shown in the inset.

**Discussion**

The results of this study indicate that HL intensity inhibits the AQPs in perfused leaves of maize plants. This extends earlier findings of Kim and Steudle (2007), who showed that, at constant turgor, LL intensity had an ameliorative effect on cell Lp, most probably by acting on AQPs. The treatment of the tissue with oxidants (H\(_2\)O\(_2\) and ·OH as produced by the Fenton reaction; H\(_2\)O\(_2\)+Fe\(^{2+}\)=Fe\(^{3+}\)+OH·+OH·-) had an effect similar to HL. Perfusion with a solution of the antioxidant GSH increased cell Lp (reduced T\(_{1/2}\)) in cells having a long T\(_{1/2}\). The presence of the antioxidant tended to prevent the inhibition by HL. This may indicate that there was indeed an action of ROS on AQPs in the leaf caused by HL. Similar findings of an oxidative gating of AQPs have been shown with Chara internodes and maize roots (Henzler et al., 2004; Ye and Steudle, 2006). In Chara, H\(_2\)O\(_2\) in the presence of Fe\(^{2+}\) caused a reversible oxidative gating and reduced cell Lp by >90%. In the presence of rapidly permeating solutes, anomalous (negative) osmosis could be observed when AQPs were closed (Henzler et al., 2004). Ye and Steudle (2006) showed that in root cells, AQ activity was reduced by a factor of 9. At the whole root level, the reduction was smaller by a factor of 3, as expected from the composite transport model of the root. In the present study, effects of HL on cell Lp in leaves were shown for the first time, suggesting that this may also be related to an oxidative gating. It is known that during light stress, ROS develop in leaves by the partial reduction of oxygen or from hydrogen peroxide produced in metabolic reactions (Foyer and Noctor, 2000).

It may be argued that the huge effects caused by HL on cell Lp (AQP activity) could be an artefact caused by the fact that cells had to be punctured to measure water relations parameters, and that these cells were more susceptible to stress. If true, the effects on the ‘intact’ system could have been different. In principle, this may be true, but is unlikely, because cells punctured by the CPP had stable turgor for up to 6 h, indicating high membrane integrity and stability. Also, punctured cells showed reversible responses during treatments (light, inhibitors, and GSH), as expected. This was true, although there was a substantial variability between cells which could have been caused by other factors (see below). It appears that there is, at present, no alternative technique for measuring light responses at the level of individual, intact tissue cells, which is completely non-invasive.

Because a molecular analysis of AQPs was not provided, the interpretation of the present data in terms of a gating of AQPs may be questioned. In the present study, what was directly measured was the water permeability of parenchyma cells in the midrib of cut leaves from maize plants (cell Lp). However, it would be difficult to find an alternative plausible interpretation different from the one offered. In maize, AQPs were classified into four different groups of proteins, as plasma membrane intrinsic proteins (PIPs), tonoplast intrinsic proteins, Nod26-like intrinsic proteins, and small and basic intrinsic proteins (Chaumont et al., 2001). To date, there have been no data showing the effects of light on any of these AQPs. In walnut, the transcript abundance of two AQPs (JrPIP2,1 and JrPIP2,2) in response to light was correlated to the overall leaf hydraulic conductance (K\(_{\text{leaf}}\); Cochard et al., 2007). The activation of AQPs by light has been speculated from K\(_{\text{leaf}}\) measurements (Nardini et al., 2005; Tyree et al., 2005). Kim and Steudle (2007) started to fill the gap between the two levels by measuring changes in cell Lp in response to light. All the currently available data attribute changes in cell Lp to a gating of AQPs. The results of this study support this interpretation, namely by involving an oxidative gating of AQPs in addition to the light effect. HL decreased cell Lp most probably by closure of AQPs, whereas LL aided the recovery of cell Lp to a high value by opening of AQPs. The closure of AQPs by HL could be tightly connected to an oxidative gating by ROS produced during HL treatment (see below).

Earlier it was found that both turgor and light affect cell Lp, hence a perfusion technique in cut leaves was applied to keep the turgor constant. Kim and Steudle (2007) showed that, as transpiration increased during light treatment, turgor decreased, and this resulted in a decrease of cell Lp. Turgor had to be kept constant to separate the effects of turgor from those of light. The perfusion technique used in the present study allowed this. Furthermore, leaves could be perfused with solutions of defined composition. For example, when using the perfusion technique, either H\(_2\)O\(_2\) or the...
antioxidant GSH could be perfused. In the future, this technique could be used further to test other solutes such as heavy metals or solutions with different pH or pKₐ. It is unlikely that the perfusion of leaf tissue with solutions saturated with oxygen could have caused the cells to be deprived of oxygen and carbon dioxide, but this point needs to be clarified in further experiments.

The light intensities used in this study were up to 1800 µmol m⁻² s⁻¹, which is comparable with outdoors on a bright day (~2000 µmol m⁻² s⁻¹; Nobel, 1999). It is known that HL produces ROS. ROS are unstable partly reduced oxygen species, produced as by-products of photosynthesis (singlet oxygen, H₂O₂, hydroxyl radical, superoxide; Foyer and Noctor, 2000; Jiang and Zhang, 2001). In Arabidopsis, Fryer et al. (2002) could visualize the generation of ROS in response to a light intensity of 600 µmol m⁻² s⁻¹ by digital imaging. ROS are known to cause oxidative damage such as lipid peroxidation, denaturation of proteins, and DNA mutation (Jiang and Zhang, 2001). Plants have protective mechanisms to get rid of stresses caused by ROS by an adjustment of antioxidants such as ascorbate (ASC) or GSH. In the present study, HL treatment resulted in a decrease of cell Lp, which was similar to that obtained by exogenous H₂O₂ (Fe²⁺) applied by perfusion. This supports the idea of an action of ROS during HL. In addition, the effects of HL were greater, when the Fe²⁺ level in the tissue was elevated. Further support for an action of ROS was derived from the fact that perfusion of the tissue with the antioxidant GSH protected AQPs from inhibition, probably by reducing ROS. Although the amounts of GSH which reached the inside of the cells were probably small (Gukasyan et al., 2002), the intracellular ratio GSHred/GSHox, which determines the redox potential, could have been high and sufficient to reduce ROS. From the findings of Henzler et al. (2004) on Chara internodes, it is known that ROS (including ·OH) react on AQPs, even when ROS are present at a very low concentration. Overall, the different circumstantial evidence suggested that the response of cell Lp to HL was caused by ROS.

Available data on overall water transport in leaves in response to light (Kleaf; see Introduction) suggest a trend for light increasing Kleaf. Light intensities used in those studies ranged from 400 µmol m⁻² s⁻¹ to 1400 µmol m⁻² s⁻¹ for different herbaceous and woody species (Sack et al., 2002, 1200 µmol m⁻² s⁻¹; Nardini et al., 2005, 400 µmol m⁻² s⁻¹; Tyree et al., 2005, 1000–1200 µmol m⁻² s⁻¹; Cochard et al., 2007, 1400 µmol m⁻² s⁻¹). At first sight this seems to contradict the present findings (i.e. a decrease in cell Lp at HL of 800 µmol m⁻² s⁻¹ and 1800 µmol m⁻² s⁻¹). However, this may not be true for the following reasons. (i) There is an increase of cell Lp at lower light intensity (Fig. 5A; Kim and Steudle, 2007). It may be that there is a maximum cell Lp in response to light intensity, which may be species dependent. This has to be determined in future studies. (ii) Kleaf may incorporate effects other than those related to membranes (Zwieniecki et al., 2001; Tyree et al., 2005). (iii) So far, increases of Kleaf have been demonstrated using the ‘high pressure flow meter’ (HPFM), where high pressures were applied during infiltration, which, in part, was different from the present measurements (Sack et al., 2002, 0.5–0.6 MPa; Nardini et al., 2005, 0.15 MPa; Tyree et al., 2005, 0.3–0.5 MPa; Cochard et al., 2007, 0.2 MPa). Using figleaf gourd, Lee et al. (2008) measured midrib cell Lp in response to a light intensity of 300 µmol m⁻² s⁻¹. Those authors observed a decrease in Lp. However, they did not control turgor, which may have decreased as transpiration increased with light.

There was a wide variability in response to light between cells. Light effects may be different not only for different species but also for tissues. For example, in Arabidopsis, singlet oxygen and superoxide production were primarily located in mesophyll tissue whereas hydroxyl peroxide accumulation was localized in the vascular tissues (Fryer et al., 2002). Maize leaf cells are known to have differential intercellular partitioning of GSH metabolism (Noctor et al., 2002). Glutathione reductase was localized only in leaf mesophyll cells, but other antioxidant enzymes could be restricted to bundle sheath cells. In the present study, there was also some variability in response to HL in that one-third of cells did not show stress response to HL. Although the former possibility cannot be neglected, there is an indication that this variability in response to HL could have originated from differences of individual leaves. All the five cells measured from two leaves showed no stress response. Those leaves could have been more resistant to HL. Problems related to variability of cells were in part solved by measuring the effects of treatments in individual cells, which required measurements to be taken for at least 1–4 h. Further investigations on the response of different types of cells to light (oxidative) stress are required. Although there have been attempts to work out the contribution of vessels and of non-vascular tissue in the overall measurements, it is still necessary to work out, in greater detail, the main hydraulic resistances in leaf water transport and how they depend on light intensity (Cochard et al., 2004; Nardini et al., 2005; Sack et al., 2005).

Although the exact mechanism(s) of how ROS react with AQPs are not yet known, it has been shown that AQPs reversibly close in the presence of ROS (Henzler et al., 2004; Ye and Steudle, 2006). ROS could be present in different amounts at different light levels. At LL intensity, the level of ROS could be low and ROS could be reduced by the antioxidants present such as ASC or GSH. However, regardless of the precise mechanisms of the action of ROS, the data indicate that there are interactions between light climate and water relations, which should be of key importance during photosynthesis. To maximize photosynthesis, enough water, carbon dioxide, and light should be gained. The open/closed state of stomata is regulated not only by light but largely by the water status of leaves. Stomata can only be kept open when there is enough water uptake. Plants need to manage the resources to maximize productivity. As the present results show, the management of water resources may be a complex process involving
many factors required to maximize productivity, which requires an interaction between light intensity and water flow. The present data indicate that LL may promote water flow, but that, at HL, water flows are down-regulated using ROS as messengers, which may be a common ‘alarm’ signalling system to provide defences against harmful environmental challenges (Pastori and Foyer, 2002).

In conclusion, the study presents the first evidence of an inhibition of cell \( Lp \) by HL, which was most probably caused by an action of ROS on AQPs. HL responses of parenchyma cells in the midrib of the maize leaf were similar to those caused by \( \text{H}_2\text{O}_2/\text{H}_2\text{O}_2 \) treatment. On the other hand, the antioxidant GSH had an ameliorative effect. Unlike HL, AQP activity increased at LL, which was in agreement with earlier results from leaves of intact plants. There should be an optimal light intensity to maximize water flow across leaf cells, but enhanced water flow could be inhibited at a certain light intensity. One may speculate that, by acting on the redox status of leaves, the light climate directly interacts with the water status of plants in a way which is different from that via stomata. It involves an action on the water supply of leaves by triggering cell \( Lp \) (AQP activity).

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References

Aroca R, Amodeo G, Fernandez-Illescas S, Herman EM, Chaumont F, Chrispeels MJ. 2005. The role of aquaporins and membrane damage in chilling and hydrogen peroxide induced changes in the hydraulic conductance of maize roots. *Plant Physiology* 137, 341–353.

Chaumont F, Barrieu F, Wojcik E, Chrispeels MJ, Jung R. 2001. Aquaporins constitute a large and highly divergent protein family in maize. *Plant Physiology* 125, 1206–1215.

Cochard H, Nardini A, Coll L. 2004. Hydraulic architecture of leaf blades: where is the main resistance? *Plant, Cell and Environment* 27, 1257–1267.

Cochard H, Venisse J-S, Barigha TS, Brunel N, Herbette S, Guilliot A, Tyree MT, Sakr S. 2007. Putative role of aquaporins in variable hydraulic conductance of leaves in response to light. *Plant Physiology* 143, 122–133.

Dietz KJ. 2008. Redox signal integration: from stimulus to networks and genes. *Physiologia Plantarum* 133, 459–468.

Felle HH, Hanstein S. 2007. Probing apoplastic ion relations in *Vicia faba* as influenced by nutrition and gas exchange. In: Sattelmacher B, Horst WJ, eds. The apoplastic of higher plants: compartment of storage, transport and reactions. Dordrecht, The Netherlands: Springer, 295–306.

Foyer CH, Noctor G. 2000. Tansley Review No. 112. Oxygen processing in photosynthesis: regulation and signalling. *New Phytologist* 146, 359–388.

Fryer MJ, Oxborough K, Mullineaux PM, Baker NR. 2002. Imaging of photo-oxidative stress responses in leaves. *Journal of Experimental Botany* 53, 1249–1254.

Gukasyan HJ, Lee VHL, Kim KJ, Kannan R. 2002. Net glutathione secretion across primary cultured rabbit conjunctival epithelial cell layers. *Investigative Ophthalmology & Visual Science* 43, 1154–1161.

Henzler T, Steudle E. 1995. Reversible closing of water channels in Chara intermedes provides evidence for a composite transport model of the plasma membrane. *Journal of Experimental Botany* 46, 199–209.

Henzler T, Ye Q, Steudle E. 2004. Oxidative gating of water channels (aquaporins) in Chara by hydroxyl radicals. *Plant, Cell and Environment* 27, 1184–1195.

Jiang MY, Zhang JH. 2001. Effect of abscisic acid on active oxygen species, antioxidative defence system and oxidative damage in leaves of maize seedlings. *Plant and Cell Physiology* 42, 1265–1273.

Kim YX, Steudle E. 2007. Light and turgor affect the water permeability (aquaporins) of parenchyma cells in the midrib of leaves of *Zea mays*. *Journal of Experimental Botany* 58, 4119–4129.

Lee SH, Chung GC, Steudle E. 2005. Low temperature and mechanical stresses differently gate aquaporins of root cortical cells of chilling-sensitive cucumber and -resistant figleaf gourd. *Plant, Cell and Environment* 28, 1191–1202.

Lee SH, Zwiazek JJ, Chung GC. 2008. Light-induced transpiration alters cell water relations in figleaf gourd (*Cucurbita ficifolia*) seedlings exposed to low root temperatures. *Physiologia Plantarum* 133, 354–362.

Lo Gullo MA, Nardini A, Trifilo P, Salleo S. 2005. Diurnal and seasonal variations in leaf hydraulic conductance in evergreen and deciduous trees. *Tree Physiology* 25, 505–512.

Maurel C, Chrispeels MJ. 2001. Aquaporins. A molecular entry into plant water relations. *Plant Physiology* 125, 135–138.

Nardini A, Salleo S, Andri S. 2005. Circadian regulation of leaf hydraulic conductance in sunflower (*Helianthus annuus* L. cv Margot). *Plant, Cell and Environment* 28, 750–759.

Nobel PS. 1999. Physicochemical and environmental plant physiology. San Diego: Academic Press.

Noctor G, Gomez L, Vanacker H, Foyer CH. 2002. Interactions between biosynthesis, compartmentation and transport in the control of glutathione homeostasis and signalling. *Journal of Experimental Botany* 53, 1283–1304.

Pastori G, Foyer CH. 2002. Common components, networks, and pathways of cross-tolerance to stress: the central role of ‘redox’ and abscisic acid mediated controls. *Plant Physiology* 129, 460–468.

Sack L, Holbrook NM. 2006. Leaf hydraulics. *Annual Review of Plant Biology* 57, 361–381.

Sack L, Melcher PJ, Zwieniecki MA, Holbrook NM. 2002. The hydraulic conductance of the angiosperm leaf lamina: a comparison of...
three measurement methods. Journal of Experimental Botany 53, 2177–2184.

Sack L, Tyree MT, Holbrook NM. 2005. Leaf hydraulic architecture correlates with regeneration irradiance in tropical rainforest trees. New Phytologist 167, 403–413.

Steudle E. 1993. Pressure probe techniques: basic principles and application to studies of water and solute relations at the cell, tissue and organ level. In: Smith JAC, Griffiths H, eds. Water deficits: plant responses from cell to community. Oxford: BIOS Scientific Publishers, 5–36.

Steudle E. 2000. Water uptake by roots: effects of water deficit. Journal of Experimental Botany 51, 1531–1542.

Tournaire-Roux C, Sutka M, Javot H, Gout E, Gerbeau P, Luu DT, Bligny R, Maurel C. 2003. Cytosolic pH regulates root water transport during anoxic stress through gating of aquaporins. Nature 425, 393–397.

Tyree MT, Nardini A, Salleo S, Sack L, Omari BE. 2005. The dependence of leaf hydraulic conductance on irradiance during HPFM measurements: any role for stomatal response? Journal of Experimental Botany 56, 737–744.

Wan XC, Steudle E, Hartung W. 2004. Gating of water channels (aquaporins) in cortical cells of young corn roots by mechanical stimuli (pressure pulses): effects of ABA and of HgCl2. Journal of Experimental Botany 55, 411–422.

Wei C, Tyree MT, Steudle E. 1999. Direct measurement of xylem pressure in leaves of intact maize plants. A test of the cohesion–tension theory taking hydraulic architecture into consideration. Plant Physiology 121, 1191–1205.

Westgate ME, Steudle E. 1985. Water transport in the midrib tissue of maize leaves. Plant Physiology 78, 183–191.

Xiong LM, Schumaker KS, Zhu JK. 2002. Cell signaling during cold, drought and salt stress. Plant Cell 14(suppl.), S165–S183.

Ye Q, Steudle E. 2006. Oxidative gating of water channels (aquaporins) in corn roots. Plant, Cell and Environment 29, 459–470.

Zwieniecki MA, Melcher PJ, Holbrook NM. 2001. Hydrogel control of xylem hydraulic resistance in plants. Science 291, 1059–1062.