Systems Analysis of Guard Cell Membrane Transport for Enhanced Stomatal Dynamics and Water Use Efficiency

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Stomatal transpiration is at the center of a crisis in water availability and crop production that is expected to unfold over the next 20 to 30 years. Global water usage has increased 6-fold in the past 100 years, twice as fast as the human population, and is expected to double again before 2030, driven mainly by irrigation and agriculture. Guard cell membrane transport is integral to controlling stomatal aperture and offers important targets for genetic manipulation to improve crop performance. However, its complexity presents a formidable barrier to exploring such possibilities. With few exceptions, mutations that increase water use efficiency commonly have been found to do so with substantial costs to the rate of carbon assimilation, reflecting the trade-off in CO₂ availability with suppressed stomatal transpiration. One approach yet to be explored in detail relies on quantitative systems analysis of the guard cell. Our deep knowledge of transport and homeostasis in these cells gives real substance to the prospect for reverse engineering of stomatal responses, using in silico design in directing genetic manipulation for improved water use and crop yields. Here we address this problem with a focus on stomatal kinetics, taking advantage of the OnGuard software and models of the stomatal guard cell recently developed for exploring stomatal physiology. Our analysis suggests that manipulations of single transporter populations are likely to have unforeseen consequences. Channel gating, especially of the dominant K⁺ channels, appears the most favorable target for experimental manipulation.

Stomata are pores that provide the major route for gaseous exchange across the impermeable cuticle of leaves and stems (Hetherington and Woodward, 2003). They open and close in response to exogenous and endogenous signals and thereby control the exchange of gases, most importantly water vapor and CO₂, between the interior of the leaf and the atmosphere. Stomata exert major controls on the water and carbon cycles of the world (Schimel et al., 2001) and can limit photosynthetic rates by 50% or more when demand exceeds water supply (Ni and Pallardy, 1992). Stomatal transpiration is at the center of a crisis in water availability and crop production that is expected to unfold over the next 20 to 30 years; indeed, global water usage has increased 6-fold in the past 100 years, twice as fast as the human population, and is expected to double again before 2030, driven mainly by irrigation and agriculture (United Nations Educational, Scientific and Cultural Organization, 2009).

Guard cell transport is integral to controlling stomatal aperture. Guard cells surround the stomatal pore and respond in a well-defined manner to an array of extracellular signals, including light, to regulate its aperture. Guard cells coordinate membrane transport within a complex network of intracellular signals (Willmer and Fricker, 1996; Blatt, 2000a, 2000b; Hetherington and Woodward, 2003; Shimazaki et al., 2007) to regulate fluxes, mainly of K⁺, Cl⁻, and malate, driving cell turgor and stomatal aperture. Our deep knowledge of these processes has made the guard cell the best known of plant cell models for membrane transport, signaling, and homeostasis (Willmer and Fricker, 1996; Blatt, 2000b; Roelfsema and Hedrich, 2010; Hills et al., 2012). This knowledge gives real substance to the prospect for reverse engineering of stomatal responses, using in silico design in directing genetic manipulation for improved crop yields, especially under water-limited conditions.

Water use efficiency (WUE; defined as the amount of dry matter produced per unit of water transpired) is directly related to stomatal function. Thus, at the practical level, stomata represent an important target for breeders interested in manipulating crop performance. A large body of data relates stomata, transpiration, and carbon assimilation (Willmer and Fricker, 1996; Farquhar et al., 2001; Hetherington and Woodward, 2003; Lawson et al., 2011). Several examples illustrate how manipulating of stomatal characteristics can affect WUE (Fischer et al., 1998; Rebetzke et al., 2002; Masle et al., 2005; Eisenach et al., 2012). With few exceptions,
RESULTS AND DISCUSSION

We previously developed the OnGuard software for quantitative dynamic systems modeling of the guard cell (Chen et al., 2012; Hills et al., 2012). OnGuard models incorporate explicitly all of the wealth of molecular, biophysical, and kinetic knowledge for guard cell transport, osmolyte metabolism, and H+ and Ca2+ buffering, and they couple this knowledge to guard cell volume, turgor, and stomatal aperture. OnGuard applications have yielded an abundance of counterintuitive, emergent characteristics in the function of wild-type guard cells of the crop Vicia and the model plant Arabidopsis (Arabidopsis thaliana; Chen et al., 2012; Wang et al., 2012). Applied to the Arabidopsis slac1 anion channel and ost2 ATPase mutants, the latter has also demonstrated new and unexpected connections between membrane transport, metabolism, and stomatal dynamics, uncovering a previously unrecognized homeostatic network that was subsequently verified experimentally (Wang et al., 2012; Blatt et al., 2013). Thus, OnGuard models incorporate the predictive power needed to guide reverse engineering of stomatal function starting with molecular manipulations in silico.

We carried out an analysis of stomatal opening and closing, using OnGuard to predict the consequences of targeted genetic manipulations. At present, OnGuard reflects the characteristics of stomata in isolation (Hills et al., 2012) and does not incorporate feedback from the transpiration stream nor from internal pCO2 in the leaf. As a consequence, kinetic relaxations of OnGuard models are prolonged, especially during opening, but are nonetheless scaled appropriately. The simulations presented here were designed to test the sensitivity to changes over a narrow range of parameters potentially tractable through genetic manipulation for each of the major transporters. We compared these outputs with similar analyses carried out for the wild-type (control) parameter set. For simplicity, each of the transporters was tested with population densities adjusted by factors of 0.5- and 2-fold. We followed a similar approach in varying the macroscopic gating characteristics of several voltage-sensitive ion channels. Outputs were quantified on the basis of the maximum rates of stomatal opening and closing, and on the dynamic range of apertures. Simulations used a standard diurnal cycle with a saw-tooth rise and fall in light over 12 h followed by 12 h of dark; as before, light was used as the driver for Suc and malate synthesis, and for the primary ATPase activities at the two membranes (Chen et al., 2012) employing, as a starting point, the Arabidopsis and Vicia models previously described (Hills et al., 2012; Wang et al., 2012). Analysis of the two models yielded similar results when scaled to the guard cell dimensions. To avoid duplication, only the results for the Arabidopsis model are reported here.

Manipulating Transporter Populations

These manipulations are comparable with variations in the steady-state levels of transporter expression such as might be achieved through moderated overexpression and RNA interference-mediated suppression. Figure 1A shows the simulation outputs for the wild type (control) and for 0.5x Kout and SLAC, equivalent to suppressing the population of outward-rectifying K+ channels (Hosy et al., 2003) and SLAC1 anion channels (Negi et al., 2008; Wang et al., 2012) in the guard cell to 50% of the wild type. The daylight period is marked by stomatal opening that progresses over the first 6 to 8 h and, as the light decayed to near its midpoint (K1/2) for activation of malate synthesis and the ATPases at its end, by stomatal closing (Chen et al., 2012; Wang et al., 2012). The Figure 1A inset also provides the outputs used for comparisons. Reducing the K+ and anion channel populations had visible effects on the dynamic range of stomatal movements. It also had substantial, although less obvious, effects on stomatal kinetics. These characteristics are summarized in Figure 1, B and C, and Figures 2 and 3 together with those for each of the other component transporters at the plasma membrane and tonoplast. A full description of each simulation and its outputs is unaddressed here.
The properties and functions of each of these transporters were summarized in Hills et al. (2012).
short, manipulations affecting $[\text{Ca}^{2+}]_i$ are predicted to have the most profound effects on aperture, again in a seemingly counterintuitive manner. We note that Conn et al. (2011) reported an opposing effect associated with the Arabidopsis cax1/cax3 double mutant lacking two of the dominant Ca$^{2+}$/H$^+$ exchangers that normally serve to sequester Ca$^{2+}$ in endomembrane compartments. This mutant reduced Ca$^{2+}$ accumulation in the leaf epidermis, the bulk of this in epidermal cells, and impaired stomatal dynamics and gas exchange. Their interpretation centered on elevated apoplastic [Ca$^{2+}$], which is known to reduce stomatal apertures (de Silva et al., 1996; Webb et al., 2001). A direct comparison is complicated by a lack of data on the Ca$^{2+}$ contents of the guard cells, and
by the fixed apoplastic [Ca\(^{2+}\)] in our simulations. Nonetheless, the much reduced dynamic range in simulated apertures is consistent with the experimental findings.

In simulations with the H\(^+-\)ATPase, a 2-fold increase in population at the plasma membrane displaced the dynamic range of apertures to higher values, whereas decreasing the number of pumps to 50% of the control was marginal in effect. Here the asymmetry is the direct consequence of the nonlinearity in pump capacity (Blatt, 1987, 1988) and its parallel to that anticipated for charge balance through H\(^+-\)coupled transport (Sanders et al., 1985; Blatt and Slayman, 1987; Blatt and Clint, 1989; Maathuis and Sanders, 1994; Hills et al., 2012) and the inward-rectifying K\(^+\) channels. Among others, the rise in aperture with H\(^+-\)ATPase activity is consistent with the known effects of locking the H\(^+-\)ATPase in an active state, either with the fungal toxin fusicoccin or in the ost2 mutant (MacRobbie, 1988; Fuglsang et al., 1999; Merlot et al., 2007).

A complementary pattern was evident in the predicted rates for stomatal opening and closing. Figures 2 and 3 show the predicted rates for 2- and 0.5-fold changes in the numbers of each transporter, and insets within each figure summarize the halftimes for opening and closing. Again, the consequences of any one manipulation are generally evident in both opening and closing rates, often in counterintuitive ways that connect through changes in cytosolic pH and [Ca\(^{2+}\)]. For example, elevating the H\(^+-\)ATPase population led to decreases in both initial opening and closing rates and substantial increases in the halftimes for both processes (Fig. 2). These effects arose, in part, from a small elevation in cytosolic pH and the enhanced H\(^+\) flux through coupled transport and malate metabolism, from plasma membrane hyperpolarization facilitating Ca\(^{2+}\) entry and [Ca\(^{2+}\)] elevation, and from the reduced dynamic range of apertures. Even more surprising, modest reductions in opening and closing rates were also observed when the H\(^+-\)ATPase density was reduced (Fig. 3). In both cases, the effects can be ascribed to 0.1- to 0.2-pH unit shifts in cytosolic pH and, most important, in changes to [Ca\(^{2+}\)], arising from the effects on the driving force for Ca\(^{2+}\) entry across the plasma membrane; in turn, these effects on cytosolic pH and [Ca\(^{2+}\)], alternately affected K\(^+\) flux through Kin and Kout and the balancing activities of the SLAC and R-type anion currents, much as documented by Wang et al. (2012).

It is noteworthy that several of the comparative predictions in Figure 2 are now confirmed. Coincident with the review of this article, independent experimental work was published by Wang et al. (2014), who demonstrated that a 3-fold overexpression of the AHA2 H\(^+-\)ATPase driven by a guard cell-specific promoter significantly enhanced stomatal conductance and opening in the light, and led to a small increase in carbon assimilation. Similar overexpression of two inward-rectifying K\(^+\) channels, KAT1, which is normally expressed in guard cells, and AKT1, which is primarily

**Figure 4.** Altering the voltage ranges activating the major ion channels at the plasma membrane in simulations identifies the K\(^+\) channels as targets for selectively enhancing stomatal opening and closing. Label signs (+ and −) refer to the midpoint voltage for gating (\(V_{1/2}\)) displacement from the control, either +18 mV or −18 mV. A, Dynamic range of apertures. Horizontal lines indicate the dynamic range of the control simulation for comparison. B, Initial rates of stomatal opening. The inset shows halftimes for opening. Horizontal lines in each case indicate the opening rate and halftime for the control simulation. C, Initial rates of stomatal closing. The inset shows halftimes for closing. Horizontal lines in each case indicate the closing rate and halftime for the control simulation.
found in roots, were without substantial effect on stomatal opening. They also found that overexpression of the H+-ATPase reduced WUE, suggesting that this enhanced activity had other consequences for stomatal behavior, possibly in altering stomatal closure (Lawson and Blatt, 2014).

Manipulating Channel Gating

The midpoint voltage ($V_{1/2}$) for the gating of voltage-sensitive channels is subject to experimental modification (Tao et al., 2010; Pless et al., 2011), and mutations affecting $V_{1/2}$ are known for KAT1 (C. Lefoulon, C. Grefen, and M.R. Blatt, unpublished data). We found that stomatal aperture was equally sensitive in simulation to altering this parameter in the dominant ion channels at the plasma membrane. Uniquely for the K+ channels, the effects were restricted primarily either to the opening or to the closing process. Most notable, manipulations of Kout to give an $\pm 18$ mV shift in $V_{1/2}$ reduced the aperture minimum without substantial change to its maximum and accelerated closing (Fig. 4A). A corresponding $+18$ mV shift in $V_{1/2}$ for Kin affected primarily the aperture maximum, again without a proportional change in the minimum; similarly, it accelerated the initial opening rate by roughly 35% compared with the control without affecting closing. For manipulations of both Kin and Kout, the increases in halftimes related directly to the corresponding expansion in the dynamic ranges of apertures. These findings contrast with altering the gating characteristics of the SLAC current, which had compound effects on both opening and closing kinetics when assessed against changes in the dynamic ranges of apertures. Much the same conclusion was drawn from analysis of Ca$^{2+}$ channel gating.

What defines the singular nature of Kin, and especially of Kout, to delimit the effects of manipulating their gating? Analysis of the underlying flux homeostasis for Kout, illustrated in Supplemental Figures S1 to S6, shows principally an absence of effect on either cytosolic pH or [Ca$^{2+}$]. Details are provided in the legends to Supplemental Figures S1 to S6, but can be summarized here. Kout normally activates together with anion currents by depolarization of the plasma membrane. Within the range of membrane voltages that activate Kout, changes to its activity have little impact on H+ or Ca$^{2+}$ flux across the plasma membrane, either through primary or secondary transport. Consequently, any changes in K+ current are balanced primarily by alterations in anion efflux. So virtually all of the anticipated benefit of increasing the Kout population is offset by a change in free-running voltage as K+ efflux adjusts with that of the anions. The voltage dependence of Kout and the anion channels are complementary, but for Kout this dependence is offset to voltages substantially positive of the anion channels, especially at millimolar K+ concentrations and above. In these circumstances, displacing $V_{1/2}$ for Kout by $\pm 18$ mV allows for K+ and anion fluxes to balance at marginally more negative voltages, thereby yielding an enhanced capacity for the combined ion efflux. Otherwise, changes to Kout gating had no substantial consequences for the Ca$^{2+}$ and voltage oscillations that are thought to drive solute flux and stomatal closure.

In conclusion, we suggest that the gating characteristics of the dominant K+ channels offer potential targets for genetic manipulation as a proof-of-principle in improving the WUE of plants. For reasons outlined above, it has been hypothesized that temporal mismatches between stomatal aperture and available light for photosynthesis significantly erode assimilation and WUE in fluctuating environments (Pearcy, 1990; Lawson et al., 2012; Lawson and Blatt, 2014). Thus, the challenge reduces to that of identifying, as candidates for manipulation, transporters that accelerate opening and closing, ideally both, without substantial constriction in the dynamic range of apertures. Achieving this goal has proven difficult, at least in simulation, because changes in transporter populations that were most effective in accelerating stomatal movements also led to counterintuitive and opposing effects on the complementary process, whether closing or opening. These findings lead us to discount simpler approaches that rely on altered expression of the wild-type transporters. It is important to recognize that these simulations are dissociated from feedback mechanisms affecting stomatal response through relative humidity, water supply, and pCO$_2$ within the leaf. In addition, the simulations do not account for the integrated sink strength of a crop (Baldocchi and Bowling, 2003), which may amplify the hysteresis in stomatal kinetics at leaf level by interactions between the multiple layers of a crop canopy. The consequences of manipulating channel gating as described here may well underestimate the impact on stomatal movements.

MATERIALS AND METHODS

OnGuard Modeling

The OnGuard software and model was driven through a diurnal 12-h-light/12-h-dark cycle as previously described (Chen et al., 2012) and all model outputs were derived from this cycle. Model parameters for Vicia (Filis et al., 2012) and Arabidopsis (Arabidopsis thaliana; Wang et al., 2012) were as described, the latter adjusted to reflect the physical dimensions of the Arabidopsis stomatal complex and transporter numbers scaled accordingly. Light sensitivity was assigned with a $K_{v,1/2}$ of 50 $\mu$mol m$^{-2}$ s$^{-1}$ solely to the plasma membrane H+-ATPase and Ca$^{2+}$-ATPase, the vacuolar V-type H+-ATPase, H+-PPase and Ca$^{2+}$-ATPase, and to Suc (and, hence, malic acid) synthesis in accordance with experimental observation (Chen et al., 2012). All other model parameters were fixed, the properties of the individual transporters and buffering reactions thus responding only to changes in model variables arising from the kinetic features encoded in the model. The OnGuard software and the Arabidopsis model used in these simulations are available at http://www.psrg.org.uk.

Supplemental Data

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

Supplemental Figure S1. Macroscopic outputs from the OnGuard model.

Supplemental Figure S2. K+ contents and analysis of K+ fluxes at the plasma membrane and tonoplast.

Supplemental Figure S3. Cytosolic and vacuolar pH, and H+ fluxes across the plasma membrane and tonoplast.
Supplemental Figure S4. Chloride contents and analysis of Cl\(^-\) fluxes at the plasma membrane and tonoplast.

Supplemental Figure S5. Malic acid synthesis, total malate (Mal) contents, and analysis of Mal\(^2+\) fluxes at the plasma membrane and tonoplast.

Supplemental Figure S6. Cytosolic and vacuolar [Ca\(^2+\)], cytosolic-free [Ca\(^2+\)]\(_{\text{free}}\), and analysis of Ca\(^2+\) fluxes across the plasma membrane and tonoplast.

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LITERATURE CITED

Balducci DD, Bowling DR (2003) Modelling the discrimination of \(^{13}\)CO\(_2\) above and within a temperate broad-leaved forest canopy on hourly to seasonal time scales. Plant Cell Environ 26: 231–244


Supplemental Figure S6. Cytosolic and vacuolar [Ca\(^2+\)], cytosolic-free [Ca\(^2+\)]\(_{\text{free}}\), and analysis of Ca\(^2+\) fluxes across the plasma membrane and tonoplast.