The Contribution of Photosynthesis to the Red Light Response of Stomatal Conductance1[OA]

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To determine the contribution of photosynthesis on stomatal conductance, we contrasted the stomatal red light response of wild-type tobacco (Nicotiana tabacum ‘W38’) with that of plants impaired in photosynthesis by antisense reductions in the content of either cytochrome b6/f complex (anti-b/f plants) or Rubisco (anti-SSU plants). Both transgenic genotypes showed a lowered content of the antisense target proteins in guard cells as well as in the mesophyll. In the anti-b/f plants, CO2 assimilation rates were proportional to leaf cytochrome b6/f content, but there was little effect on stomatal conductance and the rate of stomatal opening. To compare the relationship between photosynthesis and stomatal conductance, wild-type plants and anti-SSU plants were grown at 30 and 300 μmol photon m⁻² s⁻¹ irradiance (low light and medium light [ML], respectively). Growth in ML increased CO2 assimilation rates and stomatal conductance in both genotypes. Despite the significantly lower CO2 assimilation rate in the anti-SSU plants, the differences in stomatal conductance between the genotypes were nonsignificant at either growth irradiance. Irrespective of plant genotype, stomatal density in the two leaf surfaces was 2-fold higher in ML-grown plants than in low-light-grown plants and conductance normalized to stomatal density was unaffected by growth irradiance. We conclude that the red light response of stomatal conductance is independent of the concurrent photosynthetic rate of the guard cells or of that of the underlying mesophyll. Furthermore, we suggest that the correlation of photosynthetic capacity and stomatal conductance observed under different light environments is caused by signals largely independent of photosynthesis.

Stomata function as hydraulic valves on the surface of aerial parts of plants, with the guard cells that surround each pore rapidly adjusting their turgor to optimize photosynthetic CO2 uptake and minimize transpirational water loss from leaves. Stomata respond to a variety of signals, either received from the environment or produced within the plant, which lead to changes in the activities of ion or solute channels regulating guard cell turgor. Stomatal opening is induced by low CO2 concentrations, high light intensity, and high humidity, and closing is promoted by high CO2 concentrations, darkness, drought, and the plant hormone abscisic acid (Outlaw, 2003).

In C₃ species, stomatal opening in response to light is thought to be induced by distinct mechanisms depending on the wavelength of incident light. Blue light is perceived directly by phototropins (Kinoshita et al., 2001; Doi et al., 2004) and activates a signaling cascade that results in fast stomatal opening under background red light (Shimazaki et al., 2007). The opening response of stomata to red light requires higher irradiance than blue light and shares characteristics of photosynthesis in its action spectra in the red region (Sharkey and Raschke, 1981). Furthermore the red light response can be abolished by 3-(3,4-dichlorophenyl)-1,1-dimethyleurea, a PSII inhibitor in whole leaf, epidermal strips, and guard cell protoplasts (Sharkey and Raschke, 1981; Tominaga et al., 2001; Olsen et al., 2002; Messinger et al., 2006). Using isolated guard cell protoplasts, Tominaga et al. (2001) showed that 3-(3,4-dichlorophenyl)-1,1-dimethyleurea inhibited proton pumping in red light, suggesting that guard cell chloroplasts provide ATP required for H⁺ pumping in the guard cell plasma membrane.

It has also been suggested that the guard cell response to red light is in part an indirect response to red-light-driven intracellular CO2 uptake in the mesophyll (Roelfsema et al., 2002). For example, Roelfsema et al. (2006) have shown that chloroplast-containing guard cells in albino sections of variegated leaves do not respond to photosynthetically active radiation, but are sensitive to blue light and CO2, bringing into question a direct role of guard cell photosynthesis on red-light-mediated stomatal opening in intact leaves.

With the exception of the orchid Paphiopedilum, guard cells from all species studied to date contain chloroplasts. Chlorophyll fluorescence measurements (Cardon and Berry, 1992; Goh et al., 1999; Lawson et al., 2002, 2003) and biochemical and immunolocalization experiments (Ueno, 2001; Zeiger et al., 2002) suggest that guard cell chloroplasts have the capacity for electron transport, Rubisco-mediated CO2 assimilation, and photorespiration. Guard cell photophosphorylation has been postulated to be a significant energy source driving stomatal assimilation rates and stomatal conductance in both genotypes. Despite the significantly lower CO2 assimilation rate in the anti-SSU plants, the differences in stomatal conductance between the genotypes were nonsignificant at either growth irradiance. Irrespective of plant genotype, stomatal density in the two leaf surfaces was 2-fold higher in ML-grown plants than in low-light-grown plants and conductance normalized to stomatal density was unaffected by growth irradiance. We conclude that the red light response of stomatal conductance is independent of the concurrent photosynthetic rate of the guard cells or of that of the underlying mesophyll. Furthermore, we suggest that the correlation of photosynthetic capacity and stomatal conductance observed under different light environments is caused by signals largely independent of photosynthesis.

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opening in red light (Tominaga et al., 2001). These results have suggested a role of guard cell photosynthesis in the red light response of stomata; however, the mechanism underpinning this link has remained elusive.

Across species and under a variety of growth conditions, plants regulate their transpiration and photosynthetic rates in parallel, maintaining a balance between the stomata-mediated supply of CO2 to the mesophyll chloroplasts and their photosynthetic demand for CO2. This results in the conservation of the ratio of intercellular (Ci) to ambient (Ca) CO2 partial pressures (pCO2) within the leaf (Wong et al., 1979, 1985; Hetherington and Woodward, 2003). This empirical direct correlation between photosynthesis and stomatal conductance was central to initial models of stomatal control of photosynthesis (Farquhar and Wong, 1984; Ball et al., 1987) and has been carried over to more recent models (Jarvis and Davies, 1998; Dewar, 2002; Buckley et al., 2003). However, the underlying regulatory mechanism is still unclear. It has been proposed that guard cells sense the metabolic status of the mesophyll via a diffusible factor that is a product of photosynthetic activity in the mesophyll (Wong et al., 1979; Lee and Bowling, 1992) and that stomatal aperture would be inversely proportional to the pool size of such metabolites (Farquhar and Wong, 1984). Possible metabolites include ATP, NADPH, or ribulose 1,5-bisphosphate (RuBP), the concentration of which depends strongly on the balance between chloroplast electron transport and the carboxylation reaction catalyzed by Rubisco. However, exogenous ATP added to isolated epidermis did not stimulate opening (Lee and Bowling, 1992). Alternatively, guard cells could respond to photosynthetic demand by direct sensing of Ci (Mott, 1988; Roelfsema et al., 2002).

Contrary to the predictions of the above-mentioned models, transgenic plants with impairments in different steps of the photosynthetic process can maintain normal stomatal conductances, resulting in elevated Ci values casting some doubt on the extent of the control of stomatal movements by Ci (Hudson et al., 1992; Lauerer et al., 1993; Stitt and Schulze, 1994). In antisense plants that accumulate only 10% to 15% of the wild-type levels of Rubisco and show a proportional decrease in CO2 assimilation rate, chlorophyll fluorescence measurements have shown that the antisense reduction of the target gene was effective in guard cells as well as in the mesophyll. However, those plants also maintained wild-type values of conductance under ambient CO2 concentrations and a light source with a mix of red and blue light (von Caemmerer et al., 2004).

Most of the gas-exchange measurements made on transgenic plants with impaired photosynthesis have so far been made under white light and this raises the question of whether stomatal conductance would be affected when red light is the only source of illumination. This report examines the contribution of photosynthetic activity to the stomatal response to red light in intact plants. We contrasted the red light response of stomata of wild-type tobacco (Nicotiana tabacum ‘W38’) with that of antisense plants impaired in photosynthetic CO2 assimilation either by a decrease in chloroplast electron transport rate and ATP synthesis or by a decrease in Rubisco activity and ATP consumption, and we find that these impairments do not affect stomatal conductance. To further explore the relationship between photosynthesis and stomatal conductance, we also examined the stomatal response of wild-type and antisense small subunit of Rubisco (anti-SSU) plants to growth irradiance. Remarkably, despite the large difference in photosynthetic rates, the transpiration machinery of wild-type and anti-SSU plants responded in the same manner to the different light growth conditions.

RESULTS

Cytochrome f and Rubisco Content in the Epidermis of Wild-Type and Transgenic Tobacco

We used three different phenotypes of tobacco, the wild type and two lines with low photosynthetic CO2 assimilation rates, generated by antisense technology: anti-b/f plants, which carry an antisense construct directed against the Rieske iron-sulfur subunit of the chloroplast cytochrome b,f complex (Price et al., 1998), and homozygous anti-SSU plants with 10% to 15% of the wild-type content of Rubisco (Ruuaska et al., 1998). To minimize developmental effects of the transgenes, our plants were grown under conditions that provided similar rates of growth for the wild type and transgenic genotypes (see “Materials and Methods”). Figure 1 shows that the known leaf phenotype of the transgenic plants is also expressed in the epidermal tissue. As reported previously (Price et al., 1998), the segregating population of anti-b/f plants exhibited a variety of cytochrome b,f content, as estimated from leaf content of cytochrome f, ranging from 100% to less than 10% of wild-type levels. Three representative examples of plants with leaf cytochrome f content that was 51%, 28%, and 7% of wild type, respectively, are shown in Figure 1. Anti-SSU plants, which are homozygous for a single insertion of the transgene (Hudson et al., 1992), showed the previously observed drastic reduction in Rubisco levels, with only 17% of the wild-type Rubisco SSU content when quantified by immunoblotting (Fig. 1). The antisense transgenes were driven by the CaMV 35S promoter, thus it was expected that both transgenic genotypes would show reduced levels of the target genes in all green cell types, including guard cells. Epidermal tissue prepared from anti-b/f plants with low leaf cytochrome b,f content showed a corresponding decrease in the level of cytochrome f, indicating that the antisense effect of the transgene was also operational in guard cells (Fig. 1). As with the anti-b/f plants, the effect of the anti-SSU transgene was also present in the epidermis, where Rubisco content was 10% of that present in guard cells of the wild type (Fig. 1). Compared to the wild type, anti-b/f plants showed a slight decrease (15% on average) in the amount of Rubisco in...
whole leaves and the whole leaf cytochrome \( f \) content of anti-SSU plants was also decreased (Fig. 1). This is consistent with published data (Jiang and Rodermel, 1995).

Photosynthetic Rates and Stomatal Conductances under Red Measuring Light in Wild-Type and Anti-\( b/f \) Plants

We used red light in our gas-exchange experiments to induce photosynthesis independently of any stimulation of the blue light response of stomata, which is mediated by phototropins (Shimazaki et al., 2007). Attached leaves were equilibrated to ambient \( p \text{CO}_2 \) (362 \( \mu \text{bar} \)) in the gas-exchange chamber in the dark for at least 20 min before red light of 1,000 \( \mu \text{mol photons m}^{-2} \text{s}^{-1} \) was turned on. Wild-type plants gradually attained an average steady-state net rate of \( \text{CO}_2 \) assimilation of 7.5 \( \mu \text{mol photons m}^{-2} \text{s}^{-1} \), whereas anti-\( b/f \) plants exhibited net rates of \( \text{CO}_2 \) assimilation ranging from wild-type values to only 0.8% of the wild-type rate (Fig. 2A). There was a direct correlation between the steady-state photosynthetic rates under red light and cytochrome \( b/f \) content in all plants (Fig. 3A). Both the wild-type and anti-\( b/f \) plants showed variability in the rate of stomatal opening and the maximal stomatal conductance in red light (Fig. 2B). We found that stomatal conductance in both wild type and transgensics did not always reach a complete steady state under red light and therefore reported the maximal conductance. There was little effect of low photosynthetic rate on stomatal parameters, and even the anti-\( b/f \) plant with the lowest photosynthetic rate in the range (0.8% of wild type) showed a maximal conductance that was 50% of the average wild-type value (Fig. 2, A and B).

Except for plants with wild-type rates of \( \text{CO}_2 \) assimilation, the anti-\( b/f \) plants maintained a ratio of intercellular to ambient \( \text{CO}_2 \) concentration (\( C_i/C_a \)) higher than in the wild type (Fig. 2C). There was a marked proportionality of \( C_i/C_a \) and cytochrome \( b/f \) content, with the anti-\( b/f \) plant with the lowest photosynthetic
rate showing a $C_i/C_a$ ratio close to 1 (Fig. 3C). Figure 4 shows the maximal stomatal conductance and half-times of stomatal opening of wild type and a large number of individual anti-$b/f$ plants plotted against their steady-state CO$_2$ assimilation rates in red light. Although the presence of the antisense Rieske iron-sulfur protein transgene reduced photosynthetic rates in the anti-$b/f$ plants, their maximal stomatal conductance and rate of stomatal opening remained remarkably similar to that of the wild type when measured with light that stimulates photosynthesis but not the blue light photoreceptors that mediate stomatal opening.

Figure 3. Leaf gas-exchange parameters as a function of cytochrome $b_6f$ complex content in wild-type (black circles) and anti-$b/f$ (white circles) tobacco. A, CO$_2$ assimilation rate. B, Stomatal conductance. C, The ratio of intercellular to ambient CO$_2$ ($C_i/C_a$). Leaf cytochrome $b_6f$ complex content was determined from immunoblots such as the one shown in Figure 1 and is expressed as a fraction of the maximum wild-type value. Gas-exchange experiments were performed as described in Figure 2. Each point corresponds to a leaf from a different plant.

Figure 4. Relationship between maximal stomatal conductance and CO$_2$ assimilation rate in wild-type and anti-$b/f$ tobacco measured under red light. A, Stomatal conductance. B, Half-times of stomatal opening. Experimental conditions were the same as for Figure 2. Each point corresponds to a different plant. Black circles, wild-type plants; white circles, anti-$b/f$ plants. Half-times were calculated as the time taken to reach half the maximal conductance from the time the light was turned on.

Photosynthetic Rates and Stomatal Conductances under Red Measuring Light in Wild-Type and Anti-SSU Plants and the Effect of Growth Irradiance

To gain further insight on the relationship between stomatal conductance and photosynthesis, we contrasted the red light response of stomata in wild-type and anti-SSU plants that contained between 10% and 15% of wild-type Rubisco content. These transgenic plants do not show the light sensitivity and phenotypic instability of the anti-$b/f$ plants and can be grown under a broader range of irradiances. To compare with our results on anti-$b/f$ plants, we grew the anti-SSU plants at the same low light (LL) intensity (25–35 μmol photons m$^{-2} s^{-1}$) and also chose a growth medium light (ML) intensity (300 μmol photons m$^{-2} s^{-1}$). This allowed us to contrast the effect of growth irradiance on stomatal conductance in wild-type and photosynthetically impaired plants. Figure 5 shows the kinetics of gas exchange in leaves of wild-type and anti-SSU plants at ambient pCO$_2$.
(362 μbar) during a transition from darkness to 1,000 μmol photons m⁻² s⁻¹ of red light. Attached leaves were equilibrated to ambient pCO₂ (362 μbar) in the gas-exchange chamber in the dark for at least 20 min before red light of 1,000 μmol photons m⁻² s⁻¹ was turned on. Wild-type plants grown in LL conditions gradually attained a steady-state rate of net CO₂ assimilation of 8.9 ± 0.19 μmol m⁻² s⁻¹, whereas anti-SSU plants grown under the same conditions reached a steady-state CO₂ assimilation rate of only 1.64 ± 0.3 μmol m⁻² s⁻¹ (Fig. 5A). A 10-fold increase in irradiance during growth increased CO₂ assimilation rate in the wild type, to 19.1 ± 1.3 μmol m⁻² s⁻¹ in ML-grown plants. The anti-SSU plants reached a CO₂ assimilation rate of 6.8 ± 0.2 μmol m⁻² s⁻¹ when grown in ML. Stomatal conductance in both sets of plants reached steady-state values within 50 min of the onset of red light illumination (Fig. 5B). The maximal stomatal conductance in ML-grown plants was 0.34 ± 0.05 and 0.288 ± 0.03 mol m⁻² s⁻¹ for wild type and anti-SSU, respectively, and was higher than that of LL-grown plants, which had conductances of 0.16 ± 0.06 and 0.13 ± 0.02 mol m⁻² s⁻¹ for wild type and anti-SSU, respectively. However, the differences in stomatal conductance between wild-type and anti-SSU plants at either light intensity were nonsignificant (P = 0.05). The low CO₂ assimilation rates and relatively unchanged stomatal conductances in anti-SSU plants resulted in higher Cᵢ/Cₐ ratios for the transgenic plants than for the wild type (Fig. 5C). The initial transient lowering of Cᵢ/Cₐ results from the fact that CO₂ assimilation rate increases more rapidly with irradiance than stomatal conductance.

Effect of Growth Irradiance on Stomatal Density and Index in Wild-Type and Anti-SSU Plants

The drastic increase in stomatal conductance in plants grown at ML compared with LL was caused by increases of stomatal density by a factor of approximately 2 in both the abaxial and the adaxial surfaces of the leaf, irrespective of plant genotype (Fig. 6A). In the abaxial leaf surface, the stomatal index rose by 33% and 25% in wild-type and anti-SSU plants, respectively, when grown at ML, whereas the top side of the leaves showed an increase in stomatal index of 50% in both genotypes when grown at ML (Fig. 6A). Compared to LL growth conditions, both types of plants showed a slight decrease in the size of pavement cells when grown in ML. Stomatal dimensions did not vary with different growth light intensity or genotype (data not shown). There was no significant effect of genotype on the stomatal conductance, calculated relative to the combined number of stomata on both leaf surfaces (Fig. 6B).

Relationship between Photosynthetic Rate and Stomatal Conductance in Wild-Type and Transgenic Tobacco

Because it has been shown that there can be a strong correlation between CO₂ assimilation rates and stomatal conductance over a range of growth conditions and leaf ages (Wong et al., 1979; for review, see Hetherington and Woodward, 2003), we were interested to see how our data on transgenic tobacco would fit with the expected linear trend. Figure 7 (black circles) shows the covariation of stomatal conductance and CO₂ assimilation rates for young wild-type tobacco plants grown in environmental cabinet conditions, at a CO₂ concentration of 1,000 μmol mol⁻¹. Differences in growth light intensity and plant-to-plant variation produced a range of wild-type net rates of CO₂ assimilation between approximately 7 and 20 μmol m⁻² s⁻¹. In these plants stomatal conductance was directly proportional to net CO₂ assimilation rates. Transgenic plants with reduced photosynthetic rates, caused either by decreased Rubisco content (Fig. 7, white circle, square, and diamond) or by low cytochrome b₆f complex (Fig. 7, triangles), maintain stomatal conductances higher than expected from their low CO₂ assimilation rates and thus break the linear relationship of conductance and photosynthetic rate observed for the wild type.

Light Response of CO₂ Assimilation Rate and Stomatal Conductance

We also examined the fluency response of CO₂ assimilation rate and stomatal conductance to varying intensities of red light in wild-type and anti-SSU plants (Fig. 8). Leaves from ML-grown plants were acclimated in the dark for a minimum of 20 min before the red light was turned on for 30 min at each irradiance. Stomatal opening continued even after 50 min in the light and we decided to take measurements at a defined time interval of 30 min. The CO₂ assimilation rate was similar for wild-type and anti-SSU plants at LL but saturated for the anti-SSU plants at a low rate around 300 μmol photons m⁻² s⁻¹, whereas it continued to increase for wild-type leaves (Fig. 8A). Stomatal conductance on the other hand was similar for wild-type and anti-SSU plants (Fig. 8B). The largest increase in conductance occurred in the first step from dark to 50 μmol photons m⁻² s⁻¹; however, conductance continued to increase up to 1,500 μmol photons m⁻² s⁻¹ in both genotypes and the response was distinctly biphasic. The different response of CO₂ assimilation rate and stomatal conductance to irradiance in the anti-SSU plants resulted in greater ratios of Cᵢ/Cₐ compared with wild type (Fig. 8C). The humidity of the chamber was not controlled after the initial adjustment and led to a decrease in leaf-to-air vapor pressure difference that was, however, similar in wild-type and anti-SSU plants (Fig. 8D).

DISCUSSION

The Red Light Response of Stomata in Transgenic Tobacco with Impaired Photosynthesis

We have used transgenic tobacco with low capacity for either chloroplast electron transport (anti-b₆f plants)
or CO₂ fixation capacity (anti-SSU plants) to probe the contribution of photosynthetic capacity to stomatal opening in red light. In both types of transgenic plants, there was an effective decrease in the amount of the proteins targeted by antisense technology in the guard cells, as shown by immunoblotting (Fig. 1). Thus, it is expected that guard cells from these transgenic plants will share at least some of the deficiencies in photosynthetic performance that have been described before for whole leaves. In fact, von Caemmerer et al. (2004) have observed a strong correlation of photosynthetic performance in guard cells and mesophyll cells of anti-SSU plants when comparing measurements of guard cell chlorophyll fluorescence with that of the underlying mesophyll.

The two transgenic genotypes have contrasting phenotypes. The antisense RNA decrease in Rubisco content has been shown to cause an imbalance between the capacity of the photosynthetic carbon reduction cycle to fix CO₂ and the chloroplast’s capacity for electron transport, resulting in an increase in the pool size of RuBP and ATP (Quick et al., 1991b; Hudson et al., 1992), and zeaxanthin (Ruuska et al., 2000a). Conversely, in the anti-b/f plants, low cytochrome b/f content and hence low electron transport rates cause a decrease in RuBP content and an altered redox state (Price et al., 1998; Ruuska et al., 2000a) together with a lowered capacity for zeaxanthin formation via the xanthophyll cycle (Hurry et al., 1996). Despite the differences in photosynthetic properties and rates, we observed no difference in steady-state stomatal conductance and stomatal opening in red light between wild type and the transgenic plants. Our results clearly indicate that in intact attached leaves, the response of stomata to a dark-to-light transition utilizing red light as irradiance under ambient pCO₂ is independent of the concurrent photosynthetic rate of the guard cells or of the underlying mesophyll (Figs. 2–5).

The opening response of stomata to red light has frequently been linked to photosynthesis because the stomatal response saturates at similar irradiance to photosynthesis and can be abolished by PSII inhibitors. Our results confirm that stomatal conductance continues to increase with increasing red irradiance in both wild-type and anti-SSU plants (Fig. 8). The fact that stomatal conductance continued to respond to increasing irradiance in the anti-SSU plants although the CO₂ assimilation rate was saturated at low irradiance also suggests that the stomatal response to red light is not linked to the response of the CO₂ assimilation rate to red light.

A reduction in the b/f content in our transgenic line leads to a near linear decrease in CO₂ assimilation rate (Fig. 3) as has been previously observed (Price et al., 1998). It is thus surprising that this reduction of chloroplast electron transport mediated by the reduction of the cytochrome b/f content does not have a proportional effect on the stomatal red light response and strongly suggests that the red light response of stomata is not quantitatively linked with chloroplast electron transport of guard cells or the mesophyll. All our transgenic plants by necessity have some chloroplast electron transport of guard cells or the mesophyll. All our transgenic plants by necessity have some chloroplast electron transport as they can be grown autotrophically. Thus, we cannot exclude the possibility that a complete inhibition of guard cell chloroplast electron transport is required to decrease the extent of the stomatal red light response.

The lack of a stomatal phenotype in the anti-b/f plants suggests that routes other than photophosphorylation can provide the energy required for stomatal opening. The importance of guard cell respiration as

Figure 5. A to C, Kinetics of CO₂ assimilation rate (A), leaf conductance (B), and the ratio of intercellular to ambient CO₂ (C/Cₐ; C) in wild-type and anti-SSU plants with 10% to 15% of wild-type Rubisco during illumination of dark acclimated leaves at an irradiance of 1,000 μmol photons m⁻² s⁻¹ of red light. Plants were grown under an irradiance of 30 μmol photons m⁻² s⁻¹ (LL, squares) or 300 μmol photons m⁻² s⁻¹ (ML, circles). During gas-exchange measurements, light was turned on at time 0. Leaves were acclimated in the dark for a minimum of 20 min before the measurements. Experimental conditions were the same as for Figure 2. Data are the means of measurements on four different plants; error bars represent ±SE and are not shown if smaller than the symbols. Black symbols, wild-type plants; white symbols, anti-SSU plants.
an energy source to drive opening has been pointed out (Parvathi and Raghavendra, 1995). Recent experiments on plants with reduced tricarboxylic acid cycle activity but normal chloroplast electron transport rates support the suggestion that mitochondrial function is necessary to maintain optimal stomatal opening and transpiration rates (Nunes-Nesi et al., 2007).

**Is There a Link between Photosynthetic Processes and the Red Light Response of Stomata?**

The fact that the transgenic plants used in this study maintain normal conductances but low photosynthetic rates results in higher than wild-type $C_i$ values for a given ambient $CO_2$ partial pressure (Figs. 3 and 5). The lack of sensitivity of guard cells to $C_i$ has been observed in transgenic plants with low Rubisco or cytochrome $b_6/f$ content before (Quick et al., 1991b; Price et al., 1998; von Caemmerer et al., 2004). However, because those experiments were performed under white light or a red/blue light source, they did not rule out the possibility of an equal, direct blue light stimulation of opening in wild-type and transgenic plants, which could be independent of photosynthesis. To our knowledge, this is the first report to use red light to address this question in intact plants. The nature of the red light response mechanism remains unresolved. The experiments in this article would argue that the effect of red light absorption and utilization by photosynthesis on conductance is not as direct as previously thought. Perhaps another, so far unidentified, photoreceptor is involved. However, the opening response to red light intensity (Fig. 8) clearly shows that no matter how light perception is achieved, stomatal conductance does respond to high light flux levels in a manner not dis-

![Figure 6](image_url)  
**Figure 6.** Number of stomata on the leaf surface of wild-type (dark gray bars) and anti-SSU (light gray bars) tobacco plants as a function of growth irradiance and stomatal conductance normalized by stomatal numbers. Plants were grown under an irradiance of 30 $\mu$mol photons m$^{-2}$ s$^{-1}$ (LL) or 300 $\mu$mol photons m$^{-2}$ s$^{-1}$ (ML). A, Stomatal density. B, Stomatal index. C, Stomatal conductance normalized by stomatal numbers. Data represent mean values ± se from four different plants.

![Figure 7](image_url)  
**Figure 7.** Relationship between stomatal conductance and $CO_2$ assimilation rate in wild-type and transgenic tobacco plants impaired in photosynthesis either by a decrease in electron transport rates (anti-$b/f$ plants) or in Rubisco function (anti-SSU plants). Plants were grown under elevated $CO_2$ in environmentally controlled chambers and conductance and photosynthesis measurements were performed under ambient $CO_2$. Black circles, wild type; white triangles, anti-$b/f$ plants; white diamond, mean ± se ($n = 4$) from LL-grown anti-SSU plants; white square, mean ± se ($n = 4$) of ML-grown anti-SSU plants; white circle, mean ± se ($n = 5$) from ML-grown anti-SSU plants assayed in red-blue light (von Caemmerer et al., 2004). Arrows link data from anti-SSU plants with the mean ± se of four to five wild-type plants grown and assayed under identical conditions at the same time. The solid and dashed lines represent the linear regression fit of all wild-type data ($y = 0.0217 \pm 0.00069 \times x; R = 0.90$), and LL-grown wild-type and anti-$b/f$ data as shown in Figure 4A ($y = 0.1209 \pm 0.0159 + 0.00514 \pm 0.029 \times x; R = 0.34$), respectively. Each data point not showing error bars corresponds to an individual plant. Error bars represent ± se.
similar to photosynthesis, although there is a distinctly biphasic nature to it with both low and high light response regions. Clearly, more research needs to be done to find alternative mechanisms to explain the red light response of stomata.

Recent mathematical models that attempt to link guard cell photosynthesis with stomatal function hypothesize that the response of stomatal conductance is controlled by the balance between electron transport capacity and Rubisco capacity, and zeaxanthin and ATP have been proposed as possible metabolic links (Zhu et al., 1998; Buckley et al., 2003). In our transgenic lines the balance between electron transport and Rubisco capacity has been perturbed in opposite directions, and our results suggest that the pool size of either metabolite is not the main determinant of stomatal opening under red light. Due to their low electron transport rates relative Rubisco capacity, anti-\(b/f\) plants have a substantially decreased zeaxanthin pool (Hurry et al., 1996); however, their maximal stomatal conductance and stomatal opening rate are similar to those of the wild type (Fig. 4). Conversely, anti-SSU plants have a decreased Rubisco relative to electron transport capacity and increased levels of zeaxanthin (Ruuska et al., 2000b); nevertheless, stomatal conductance is similar to wild type. A logical interpretation of our results is that in general they do not support the hypothesis of a direct link between the stomatal response to \(\text{CO}_2\) and the photosynthetic process.

Suc, either synthesized inside the guard cell or imported from the apoplast, has been proposed to play a key role as an osmoregulatory solute in stomatal movements (Talbott and Zeiger, 1998; Outlaw, 2003). Low leaf sugar content has been reported in anti-SSU plants (Quick et al., 1991a), even under elevated \(\text{CO}_2\) growth conditions (Masle et al., 1993). We do not know whether these transgenic plants can maintain their apoplastic Suc concentrations despite the lowered bulk leaf concentrations; however, the lack of a stomatal phenotype in anti-SSU plants would suggest that metabolites other than Suc can act as osmoregulators during stomatal opening.

Correlation between Photosynthetic Capacity and Stomatal Conductance

We used growth light intensity as the environmental variable with which to investigate the effect of low photosynthetic rate in the commonly observed comodulation of stomatal conductance and photosynthesis by environmental conditions (Hetherington and Woodward, 2003). Wild-type and anti-SSU plants responded to an increase in growth irradiance from LL to ML by doubling their \(\text{CO}_2\) assimilation rates (Fig. 5A) and their stomatal conductance (Fig. 5B) and density (Fig. 6). For wild-type plants this resulted in a strong linear correlation between \(\text{CO}_2\) assimilation rates and conductance, as expected (Fig. 7). Although the response to growth irradiance was similar in wild-type and anti-SSU plants, the latter maintained a high stomatal conductance relative to their decreased \(\text{CO}_2\) assimilation rates under the two light conditions, showing that stomatal conductance and photosynthetic rate can be uncoupled by genetic manipulation of Rubisco content (Fig. 7). These results are in agreement with previous findings on transgenic plants with impairments in photosynthesis due to antisense decreases in the levels of Rubisco and other PCR cycle enzymes (Hudson et al., 1992; Lauerer et al., 1993; Haake et al., 1998; Muschak et al., 1999). In contrast to these observations in plants with low photosynthetic rates, the correlation between stomatal conductance and photosynthetic rate was apparently maintained in transgenic plants with decreased mitochondrial respiration (Nunes-Nesi et al., 2007).
CONCLUSION

Our study illustrates the power of the transgenic approach in unraveling correlative links to reveal mechanistic connections. The results show that the red light response of stomata may not be linked to photosynthesis and that further work is required to discover the nature of the red light receptor. Furthermore, we have shown that the environmentally induced correlation between stomatal conductance and photosynthetic capacity so frequently observed must be caused by signals not directly related to photosynthesis. The results have major implications for our understanding of stomatal function and demonstrate that photosynthetic metabolism can be manipulated with minimal coupling to stomatal function and aperture. This means that if plants can be genetically engineered for improved photosynthesis this should also lead to improved plant water use efficiency.

MATERIALS AND METHODS

Plant Material and Growth Conditions

Wild-type and transgenic tobacco plants (Nicotiana tabacum ‘W38’) were grown in 0.25-L pots in seed-raising soil containing approximately 2 g/L of a slow-release fertilizer (Osmocote; 15:4:8:10:8.1:2 N:P:K:Mg and trace elements, B, Cu, Fe, Mn, Mo, Zn; Scotts Australia) and bottom-watered daily. Two types of transgenic tobacco were used: (1) anti-SSU plants, which contain 10% to 15% of the wild-type Rubisco content and are used in the experiments. Transgenic and wild-type plants had similar leaf sizes and number of leaves at the time of measurement.

Preparation of Epidermal Fragments, Protein Extraction, and Immunoblotting

A fraction enriched in epidermal tissue was prepared by adapting the method of Kopka et al. (1997). A young expanding leaf was picked, the major veins were removed and discarded, and the rest was blended with 250 mL of chilled distilled water with a Sorvall Omni Mixer blender at maximum speed, with four pulses of 30 s each and waiting 30 s between pulses. The resulting epidermal fragments were rinsed with 300 mL of chilled distilled water on a 100- to 149-μm Nytal mesh to rid them of contaminating mesophyll cells. The epidermal fragments were drained of excess water, disrupted by grinding with mortar and pestle in liquid nitrogen for 3 min, and stored at −80°C until later use. The resulting fraction was highly enriched in epidermis compared with mesophyll cells (less than one mesophyll cell per 200 stomata was routinely observed under the compound microscope).

Measurement of Leaf Gas Exchange

Gas-exchange measurements were made with a LI-6400 portable gas-exchange system (LI-COR), equipped with a red light-emitting diode light
source with a maximum emission peak centered at 670 nm (LI-6400-02, LI-COR). This light source was fitted on the standard 6-cm² clamp-on leaf chamber. Sample pCO₂, flow rate, and temperature were kept constant at 362 pbar, 500 μmol s⁻¹, and 25°C, respectively. Leaves were equilibrated in the gas-exchange leaf chamber in darkness, at an initial humidity of 19 mbar for a minimum of 20 min before measurements. For light response curves, the conditions were the same, and after the initial 20 min of dark adaptation the intensity of red light was increased in steps of 30-min duration. The average atmospheric pressure was 950 mbar. Gas-exchange parameters were calculated using the equation derived by von Caemmerer and Farquhar (1981).

Determination of Stomatal Numbers

Stomatal numbers were determined from the same or similar leaves as used for gas-exchange measurements, from impressions taken from both sides of the leaves with dental silicone (Optosil-Xantopren; Heraeus Kulzer). Stomata and epidermal cells were counted from positives made from the impressions with nail polish, in 10 different fields of view per leaf, with a compound microscope using a magnification of 200-fold. Digital photographs of each field were taken and cells were counted and measured with Image J software (http://rsb.info.nih.gov/ij/).

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