## Thioredoxin-insensitive plastid ATP synthase that performs moonlighting functions

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The chloroplast ATP synthase catalyzes the light-driven synthesis of ATP and acts as a key feedback regulatory component of photosynthesis. Arabidopsis possesses two homologues of the regulatory  $\gamma$  subunit of the ATP synthase, encoded by the ATPC1 and ATPC2 genes. Using a series of mutants, we show that both these subunits can support photosynthetic ATP synthesis in vivo with similar specific activities, but that in wild-type plants, only  $\gamma_1$  is involved in ATP synthesis in photosynthesis. The  $\gamma_1$ -containing ATP synthase shows classical light-induced redox regulation, whereas the mutant expressing only  $\gamma_2$ -ATP synthase (gamera) shows equally high ATP synthase activity in the light and dark. In situ redox titrations demonstrate that the regulatory thiol groups on  $\gamma_2$ -ATP synthase remain reduced under physiological conditions but can be oxidized by the strong oxidant diamide, implying that the redox potential for the thiol/disulphide transition in  $\gamma_2$  is substantially higher than that for  $\gamma_1$ . This regulatory difference may be attributed to alterations in the residues near the redox-active thiols. We propose that  $\gamma_2$ -ATP synthase functions to catalyze ATP hydrolysis-driven proton translocation in nonphotosynthetic plastids, maintaining a sufficient transthylakoid proton gradient to drive protein translocation or other processes. Consistent with this interpretation, ATPC2 is predominantly expressed in the root, whereas modifying its expression results in alteration of root hair development. Phylogenetic analysis suggests that  $\gamma_2$  originated from ancient gene duplication, resulting in divergent evolution of functionally distinct ATP synthase complexes in dicots and mosses.

plant development

Light-driven linear electron flow (LEF) in photosynthesis stores energy in reduced NADPH. It is also coupled to the translocation of protons into the thylakoid lumen, generating an electrochemical gradient of protons, the proton motive force (pmf), across the thylakoid. The pmf in turn drives the synthesis of ATP from ATP and  $P_i$  via the  $CF_0$ - $CF_1$  ATP synthase (ATP synthase), storing energy in the form of the photophosphorylation potential. The plastid ATP synthase complex consists of nine different subunits. Four of these subunits (I, II, III<sub>14</sub>, and IV, also called b, b', c, and a) form the integral membrane  $CF_0$  subcomplex and the remaining five subunits make up the extrinsic  $CF_1$  subcomplex ( $\alpha_3$ ,  $\beta_3$ ,  $\gamma$ ,  $\delta$ , and  $\varepsilon$ ) that contains the catalytic sites of ATP synthase (1, 2). The  $CF_0$  subcomplex drives the synthesis of ATP at the  $CF_1$  subcomplex via the rotational-catalysis/binding-change mechanism (3–5).

The ATP synthase is regulated at several levels (6). It has been known for some time that the ATP synthase requires a substantial pmf to become activated (7–9). This pmf-activation is thought to involve interaction between the  $F_{\rm O}$  and  $F_{\rm 1}$  subcomplexes because detachment of  $F_{\rm O}$  activates ATP hydrolysis (10). A second level of regulation occurs in plants and green algae via thioredoxin-mediated thiol modulation of the  $\gamma$  subunit, altering the extent of pmf required to activate the complex (4). The structural basis

of the regulation is assigned to a sequence of nine amino acid residues containing two Cys residues (Cys199–Cys205, in *Arabidopsis thaliana* numbering) able to form an intrapeptide disulfide bond upon oxidation (11). In the light, electron flow from photosystem I (PSI) reduces thioredoxin-f via ferredoxin:thioredoxin oxidoreductase (12), which in turn reduces the  $\gamma$  subunit, forming an ATP synthase that is activated with a relatively low pmf of about 50 mV (13). After dark adaptation for tens of minutes, the  $\gamma$  subunit cysteine residues become oxidized, imposing a higher pmf requirement for activation, of about 100 mV, thus resulting in ATP synthase inactivation, preventing rapid hydrolysis of ATP. Redox-regulation via thioredoxin is highly sensitive to light, and acts as a "switch," rapidly transitioning between inhibition in the dark and full activity at even very low light intensities (14).

The ATP synthase is also fine-tuned in the light, during steadystate photosynthesis in response to changes in metabolism and/or environmental conditions, e.g., endogenous CO<sub>2</sub> levels or drought stress (15–17). The resulting changes in proton efflux adjust the steady-state pmf and subsequently affect the regulation of light capture and electron transfer. The mechanism of this fine tuning is not yet clear, though a number of mechanisms have been proposed, including drawing down of inorganic phosphate and phosphorylation (18).

Arabidopsis thaliana possesses two genes that appear to code ATP synthase  $\gamma$  subunits, ATPC1 and ATPC2 (At4G04640 and At1G15700), located on chromosomes 4 and 1, respectively (19, 20), and showing 73% sequence similarity (19). Both Arabidopsis  $\gamma$  homologues contain the domain responsible for the redox regulation of the enzyme (19), (Fig. S6). The current study aims to answer the intriguing questions: In which tissues are  $\gamma_1$  and  $\gamma_2$  localized? Can both ATPC homologues function in ATP synthesis? Are their regulatory and functional properties different? What functions might they serve in the plant?

## **Results and Discussion**

As described in detail below, this study used a series of *Arabidopsis* plant lines, wild type (Columbia), two mutant lines lacking *ATPC2*, *atpc2* (T-DNA inserted in chl1\_5402949), *atpc2*\_2 (T-DNA inserted in chl 1\_5404314), a mutant lacking *ATPC1* (*dpa1*) (21) complemented with 35S::*ATPC1*, and a mutant line lacking *ATPC1*but expressing *ATPC2* behind a 35S promoter (gamera).

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The  $\gamma_2$  Subunit Is Chloroplast Localized and Expressed at Very Low Levels. The primary sequences of  $\gamma_1$  (coded by ATPC1) and  $\gamma_2$  (coded by ATPC2) contain apparent N-terminal transit peptides of 50 and 60 amino acids, respectively, suggesting that  $\gamma_2$ , like  $\gamma_1$ , is targeted to the chloroplast. The subcellular localization of  $\gamma_2$  in chloroplast was confirmed by fluorescence microscopy on transiently transformed Arabidopsis protoplasts expressing  $\gamma_2$ -GFP-fusion protein (Fig. S1).

The *ATPC1* gene was found to be highly expressed in photosynthetic tissues, and disrupting this gene abolished photosynthetic growth implying that it is essential for photosynthesis (21, 22). In contrast, *ATPC2* was expressed at low levels in photosynthetic tissues, presumably insufficient to complement the *ATPC1* knockout (21) (See also Fig. S3). Previous mass spectroscopy-based proteome analysis (23) was able to detect the  $\gamma_2$  subunit in wild-type chloroplasts, but we were unable to detect it using immunoblot analysis (Fig. S2B and C), likely indicating very low expression (Fig. S2A) or photosynthesis (Fig. 1), implying that it plays little or no role in photosynthesis.

 $\gamma_2$  Can Form an Active ATP Synthase, Suppressing the Loss of  $\gamma_1$ . To determine whether  $\gamma_2$  could, if expressed, support ATP synthase activity, we generated plants containing exclusively \( \gamma\_2 \)—rather than  $\gamma_1$ -ATP synthase by expressing ATPC2 under control of the constitutive 35S promoter in the ATPC1 null mutant background (see Supporting Information) (21). The resulting mutant was termed gamera (gamma exchange-revised ATP synthase). Using Western blot analysis with antibodies specific to  $\gamma_1$  and  $\gamma_2$ (Fig. S2 B and C), we confirmed that gamera lacked detectable levels of the  $\gamma_1$  subunit, but gained  $\gamma_2$  subunit. On the other hand, leaves from wild type and the atpc2 mutant contained  $\gamma_1$ , but no detectable  $\gamma_2$ . Both  $\gamma_1$  in wild type and  $\gamma_2$  in gamera were found exclusively in the insoluble (membrane) fractions, indicating that these were likely bound into the ATP synthase complexes. All independent gamera transformants showed a lower than wild type accumulation of ATP synthase protein subunits, despite high levels of ATPC2 mRNA expression, suggesting that the  $\gamma_2$ -ATP synthase may be less stable than  $\gamma_1$ -ATP synthase (Fig. S2B and C).

The gamera mutant was able to grow photoautrophically (Fig. S24), indicating that higher (than wild type) expression levels of  $\gamma_2$  were able to functionally replace  $\gamma_1$  in the ATP synthase. However, the growth rates were somewhat slower in gamera than in wild type, resulting in rosette diameter 30% smaller in 4 wk old plants (Fig. S24).

**Photosynthetic Properties of Gamera.** Fig. 1*A* shows the dependence of LEF, measured by saturation-pulse chlorophyll fluorescence yield analysis (24) on photosynthetically active radiation (PAR) for wild type, atpc2 and gamera. The initial slope of LEF against PAR was only slightly (approximately 10%) reduced in gamera compared to wild type or atpc2, indicating that the maximal quantum efficiency of photosynthesis was similar for all three genotypes. This result is also consistent with our observation that atpc2 and gamera had minimal effects on maximal photosystem II (PSII) quantum efficiency estimated by saturating-pulse fluorescence yield changes in dark-adapted leaves (0.83  $\pm$  0.0019,  $0.83 \pm 0.0012$ , and  $0.80 \pm 0.0013$  for wild type, *atpc2*, and gamera, respectively). However, LEF saturated at about 100 µmol photons  $m^{-2}$  s<sup>-1</sup> in gamera compared to about 200  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> for wild type and atpc2, whereas gamera reached a maximum LEF of approximately 33% (approximately 20 µmol electrons  $m^{-2} s^{-1}$ ) compared to wild type and *atpc2*, indicating that gamera had a rate-limitation at steps beyond PSII, most likely at the ATP synthase.

We confirmed a lower in vivo ATP synthase activity for gamera by probing proton flux and light-induced pmf by measuring the decay of the electrochromic shift (ECS) (Fig. 1) as described

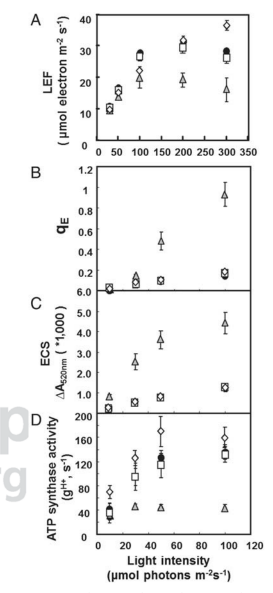


Fig. 1. Wild-type -Col (black circles), atpc2 (open squares), gamera (gray triangles), and dpa1 is complemented with 35S::ATPC1 (open diamond) were compared for differences in actinic light intensity-dependence of LEF (A), energy-dependent exciton quenching  $(q_E)$  (B), light-induced pmf (estimated by analysis of the electrochromic shift (ECS $_t$ ) (C), and proton conductivity across the thylakoid membrane  $(g_{H+})$  (D), based on ECS decay kinetics. Data is for attached leaves (n=3-4).

previously (15, 16, 25). The extent of the rapid decay of ECS upon switching off actinic light, termed ECS<sub>t</sub>, which reflects the light-driven thylakoid pmf was three- to fourfold larger in gamera than in wild type or atpc2 (Fig. 1C) suggesting that gamera generated a larger light-induced pmf than wild type or atpc2, consistent with a slow rate of proton efflux through the ATP synthase. Supporting this conclusion, gamera produced a substantially higher extent of energy-dependent exciton quenching ( $q_E$ ) than wild type or atpc2 (Fig. 1B), implying a more acidic lumen pH as a result of increased pmf.

The increased pmf in gamera could be attributed to a decrease in efflux of protons from the lumen. Fig. 1D plots the relative conductivity of the thylakoid membrane to protons  $(g_H^+)$ , which predominantly reflects ATP synthase activity and can be estimated by the decay kinetics of the ECS (15, 24). The  $g_H^+$  in atpc2 was similar to that in wild type, but remained consistently lower in gamera, indicating slower proton efflux. Western blot analysis

showed that ATP synthase  $\beta$ -subunit (Fig. S3B) decreased substantially in gamera compared to wild type and atpc2, implying that the total ATP synthase content, rather than decreased specific activity accounted for the lower  $g_H^+$  in gamera.

Analysis of *atpc1*-complemented with *ATPC1* (*dpa1* 35S::*ATPC1*) showed no differences from wild type in growth, photosynthetic performance, or in vivo ATP synthase activity (Fig. 1) indicating that the lowering of ATP synthase content or activity in gamera was due to substitution of *ATPC1* by *ATPC2*, rather than secondary effects of altering *ATPC* promoter regions.

 $\gamma_2$ -ATP Synthase Is Equally Active in Light- and Dark-Adapted Leaves. We analyzed ATP synthase activity in attached leaves by exciting PSI and PSII with short (100 µs), nonsaturating LED pulses to rapidly generate pmf, and probing the resulting proton flux through the ATP complex via the decay of the ECS absorbance change at 520 nm (13, 14) (Fig. 2 A–C). A few seconds after preillumination, ATP synthase activity of wild type remained active even at low pmf, as indicated by rapid nearly monophasic flashinduced ECS decay-with a fast phase of approximately 25 ms constituting approximately 80% of the decay, and a slower phase with lifetime longer than 250 ms, showing that the  $\gamma$  subunit was predominantly reduced. Upon extensive (>20 min). dark adaptation, the extent of the slow ECS decay phase markedly increased, reflecting oxidation of the γ subunit, which imposes an increased pmf requirement for activation of the ATP synthase. Owing to the slower decay of ECS in dark-adapted leaves, the contribution of the cytochrome  $b_6 f$  complex turnover to the electric field can be seen as a rise in ECS on the approximately 10 ms time scale. In light-adapted material, this phase is masked by the rapid decay of ECS. The extent of slow ECS decay phase showed a sigmoidal dependence on dark adaptation (Fig. 2D), with a pronounced lag phase from 0 to 10 min after preillumination, followed by a approximately 10 min half time decrease in activity. The sigmoidal oxidation kinetics were previously interpreted as reflecting the equilibration of  $\gamma$  subunit thiols redox state with that of a larger pool of redox carriers (14). Almost identical results were obtained for atpc2, indicating that  $\gamma_2$  is not required for regulation of ATP synthase. In contrast with wild type and atpc2, the ECS decay kinetics in gamera remained constant with a nearly monophasic decay showing halftime of about 21 ms, even after extensive (>60 min) dark-adaptation (Fig. 2B), indicating that  $\gamma_2$ -ATP synthase is not down-regulated by thiol oxidation in the dark.

**Equilibrium Redox Titrations of**  $\gamma_1$  **and**  $\gamma_2$ **-ATP Synthase.** In situ equilibrium redox titrations (22), were used to assess the redox properties of  $\gamma_1$ - and  $\gamma_2$ -ATP synthase in wild type and gamera, respectively (Fig. 3A). In wild type, the ECS decay was rapid at low redox potential and slow at high potential, where  $\gamma_1$  was expected to be reduced and oxidized, respectively. Assuming a stromal pH of 7.5, we estimate a redox midpoint potential for the  $\gamma_1$ subunit of approximately -356 mV (versus standard hydrogen electrode) with an n = 2 redox behavior for the thiol/disulfide interconversion, similar to that reported previously (22). By contrast, ECS decay in gamera remained constant with changing the redox potential (Fig. 3A) over the entire range accessible by varying reduced and oxidized DTT (approximately -300 to -420 mV versus standard hydrogen electrode). The maximal activity of the ATP synthase under reducing conditions, as judged by the ECS decay, was about twofold higher in the wild type than in gamera, presumably reflecting the overall differences in ATP synthase content (Figs. S2B and C).

To determine whether the lack of redox regulation of  $\gamma_2$ -ATP synthase was caused by a shift in the redox potential of the thiol/disulfide transition or by a change in the effects of such transition on the activity of the enzyme, we assayed the redox state of the  $\gamma_1$  and  $\gamma_2$  subunits directly, based on modification of free sulfhydryl groups with 4-acetamido-4'-maleimidylstilbene-2,2'-disulfonate

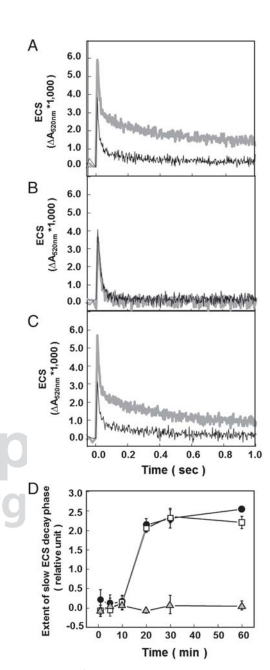
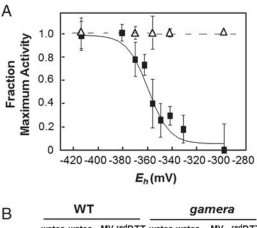


Fig. 2. Redox state of the ATP synthase probed by the decay of flash-induced ECS signals. Flash-induced relaxation kinetics of ECS were measured in wild type (A), gamera (B), and atpc2 (C) at 1 (thin black line) and 60 min (thick gray line) dark adaptation following after 2 min preillumination at  $100~\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> red actinic light. The extent of slow ECS decay phase of ECS kinetics after brief subsaturating actinic illumination in wild type (black circles), atpc2 (open squares), and gamera (gray triangles) (D). Data is for attached leaves with n=3.

(AMS) followed by differential separation by sodium dodecyl sulfate polyacrylamide gel electrophoresis (26) (Fig. 3B). Wild-type leaves infiltrated with water showed classical light-induced changes in redox state of  $\gamma_1$ , with the higher molecular weight reduced form, in light-adapted leaves. In contrast, we saw no light-induced changes in the apparent molecular weight of  $\gamma_2$  in gamera leaves infiltrated with water. To ensure maximal physiological differences in redox states, we repeated these experiments using two conditions designed to reflect the extremes of in vivo redox states: 1) oxidizing conditions in dark-adapted leaves in the presence of methyl viologen (MV) to prevent accumulation of electrons on thioredoxin (13); 2) reducing conditions in leaves illuminated with 300  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> actinic light and infil-

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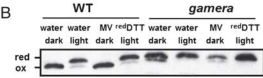


Fig. 3. Equilibrium redox titration (22) of thiol/disulfide regulatory groups in the  $\gamma$  subunit of the chloroplast ATP synthase in wild type (WT) (closed squares) and gamera (opened triangles) (A). In situ fractional ATP synthase activity was estimated by probing ECS decay kinetics ( $\Delta A_{520}$ ) as described in the text. (n=4–6). Detection of reduced (red) and oxidized (ox)  $\gamma$  subunit using gel shift assay, described in text (B). The redox states of  $\gamma_1$  and  $\gamma_2$  subunit thiols in wild type and gamera were visualized by AMS-labeling followed by SDS/PAGE and immnoblotting after treatment of leaf discs with water as control, MV, and reduced DTT under light (approximately 50  $\mu$ mol photons  $m^{-2}$  s $^{-1}$ ) or dark as described in Materials and Methods.

trated with DTT (Fig. 3B). As with water-infiltrated leaves,  $\gamma_1$  showed a higher molecular weight under reducing and not under oxidizing conditions, clearly indicating the expected redox transition that altered AMS binding. However, in gamera, the molecular weight of  $\gamma_2$  remained higher, even under these more extreme redox conditions, implying that its redox potential was shifted towards higher potentials with respect to  $\gamma_1$ . To determine whether  $\gamma_2$  is oxidizable under more oxidizing conditions, we infiltrated leaves with the strong thiol oxidant, diamide, and observed both oxidation of thiols (Fig. S3A) and a decrease in ATP synthase activity in vivo (Fig. S3B), indicating that  $\gamma_2$  is oxidizable but at a more oxidizing redox potential, probably out of the range set by the thioredoxin system in vivo.

Evidence That  $\gamma_2$ -ATP Synthases Have a moonlighting Function in Root **Morphogenesis.** The gene expression profiles of ATPC1 and ATPC2 in different tissues were investigated using the publicly available software Genevestigator (27) (Fig. S4). In leaves, ATPC1 expression was about 100-fold higher than that of ATPC2 consistent with the failure to observe accumulation of  $\gamma_2$  protein in wild-type leaves using  $\gamma_2$ -specific antibodies (Fig. S2B). However, in radicle or mature root tissue, ATPC2 expression was approximately equal to or above that of ATPC1, suggesting a function of  $\gamma_2$  in roots. To confirm the expression pattern of ATPC2, we constructed plants with  $\beta$ -glucouronidase (GUS) driven by ATPC1 (ATPC1-GUS) and ATPC2 promoters (Supporting Information). In contrast to ATPC1-GUS, which showed a strong staining over the whole seedling (Fig. S5), ATPC2-GUS expression was absent in the green tissues and appeared to be localized along the root and at the root-hypocotyl junction (Fig. 4 A-E), again suggesting a role of  $\gamma_2$  in root development.

We therefore analyzed the root morphology in wild type, gamera, and *atpc2* knock out lines. In 7 d old seedlings, two separate alleles of *atpc2* had developmentally immature root hairs. The *atpc2* mutant showed 60–80% fewer root hairs than wild type or gamera (Fig. 4F). The average length of root hair in *atpC2* was about 50% that of wild type. Strikingly, gamera, with over-

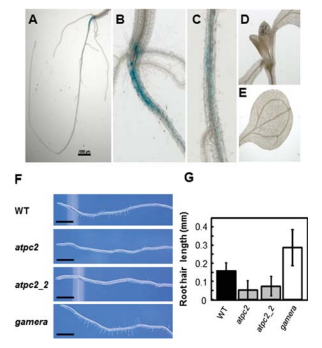


Fig. 4. Evidence for a function of ATPC2 in development. ATPC2::GUS staining in 10 d old seedlings in root system (A), hypocotyl-root junction (B), primary root (C), young leaves (D), and cotyledons (E). Representative photographs of root hairs of wild type (WT), two separate atpc2 lines, and gamera (F). Scale bars 1.00 mm. Root hair lengths measured in wild type, two atpc2 lines, and gamera (G) (n = 32-274).

expressed ATPC2, showed dramatically (approximately 2.5 times) longer root hairs than wild type (Fig. 4.F and G).

Although the precise function of  $\gamma_2$ -ATP synthase is currently unknown, our results indicate that it plays a role, either directly or indirectly, in development of nonphotosynthetic tissues, resulting in strong effects on root morphology.

Species Distribution, Phylogeny, and Structural Aspects of ATPC Genes. Comparison of sequenced genomes showed that dicots contain two or more homological duplicates (or multiplicates) of the  $\gamma$  subunit (Fig. S6). The monocots Zea mayes and Oryza sativa, the green algae Ostreococcus lucimarinus and Chlamydomonas reinhardtii, or cyanobacteria (see also below) showed only a single copy, suggesting a possible role of the "secondary-duplicate"  $\gamma_2$ -like subunit specifically in dicots. In all dicots, one homologue is clearly  $\gamma_1$ -like, possessing a regulatory loop region with two cysteine residues. In many cases, though, one or more of the " $\gamma_2$ -like" subunit duplicates have clear differences in the redox regulatory loop, probably rendering it redox-insensitive. For examples, grape (Vitis vinifera), poplar (Populus trichocarpa), and lotus (Lotus japonicus, Nelumbo nucifera) are missing one of the two regulatory cysteine residues, whereas tobacco (Nicotiana tabacum), barrel clover (Medicago truncatula), felon herb (Artemisia indica), apple (Malus pumila var. domestica), and tomato (Solanum lycopersicum) are missing both. Soybean (Glycine max), castor bean (Ricinus communis), and selaginella (Selaginella tamariscina)  $\gamma$ -homologues retain both cysteine residues, but also contains the same threonine to glutamate substitution in the regulatory loop region as in Arabidopsis  $\gamma_2$  (Fig. S6).

We investigated the possible evolutionary origins of the diversity of primary and secondary  $\gamma$  subunit homologues. The occurrence of duplicate (or multiplicate) genes coding for the  $\gamma$  subunit of the plastidic ATP sythase was screened in available genomes of uni- and multicellular representatives of viridiplantae and of the evolutionary ancestors of their plastids, i.e., the cyanobacteria. Multiple copies of *ATPC* were only detected in multicellular spe-

cies whereas both cyanobacteria and unicellular green algae contain only one ATPC gene strongly corroborating the hypothesis put forward above that the duplicate genes are expressed differentially to adapt different types of tissues to their specific requirements. Consequently, duplicate versions of the gene are found in many subgroups of multicellular virdiplantae down to the very early branching mosses. An evolutionary analysis shows a fargoing congruence between the phylogeny of these  $\gamma$  subunit genes and that of their parent species (Fig. S7).

The tree furthermore demonstrates that duplication of the ATPC gene had occurred frequently and independently during the evolution of multicellular plants. The two species of mosses, for example, as well as apple trees and glycines feature relatively recent duplication events within each of the mentioned lineages (Fig. S7). By contrast, an early duplication event apparently has given rise to two distinct ATPC genes in the ancestor of angiosperms with subsequent vertical inheritance of both copies towards most of the extant angiosperms. The resulting two clusters contain the above-analyzed ATPC1 and ATPC2 genes from Arabidopsis, respectively. The cluster encompassing the ATPC1 gene is substantially more compact (i.e., characterized by shorter branch lengths) than that containing the ATPC2 gene. This is in line with the ATPC1 gene operating under "standard" conditions in redox regulated ATP synthases, i.e., subjected to strong and unifunctional evolutionary constraints, whereas the secondary gene has evolved to fulfil different and possibly diverse functions in the various plants examined.

Conclusions: What Is the Function of  $\gamma_2$ -ATP Synthase in *Arabidopsis?* We found that the regulatory behavior of the chloroplast ATP synthase is markedly altered by substitution of  $\gamma_2$  by  $\gamma_1$ . Wild type and *atpc2* plants, which express only  $\gamma_1$ -ATP synthase in leaves, show classical light-dark regulation of ATP synthase activity via the thioredoxin system (Fig. 2). In contrast, gamera, which expresses exclusively  $\gamma_2$ -ATP synthase, shows light-dark-insensitive ATP synthase activity, due to a more positive redox potential for the regulatory thiol groups (Fig. 3*A*).

Because  $\gamma_2$ -ATP synthase is equally active in the light and dark, we propose that it may function to catalyze low levels of ATPdriven proton translocation in nonphotosynthetic tissues or in the dark, (i.e., proton pumping into the lumen driven by ATP hydrolysis) to maintain sufficient transthylakoid proton gradient when  $\gamma_1$ -ATP synthase is inactivated. The proton gradient across the thylakoid membrane is required for nonphotosynthesis processes, for example, ion transport or the TAT pathway for targeting plastid proteins into the thylakoid lumen or membrane (28–30) especially in nonphotosynthetic organelles such as etioplasts that develop without photosynthetic electron flow to reduce thioredoxin. It is also possible that  $\gamma_2$ -ATP synthase allows for ATP synthesis in nonphotosynthetic plastids, via a chlororespiratory pathway (31). In accord with these views, we showed that  $\gamma_2$ -ATP synthase is expressed in the root-hypocotyl junction (Fig. 4B), and altering its expression level had striking effects on root hair morphology (Fig. 4F and G). This is consistent with recent proteomic analysis that shows that, unlike most photosynthetic proteins and complexes, the ATP synthase is already present in dark-adapted etioplasts (32). Although a full investigation of the function of  $\gamma_2$ -ATP synthase in roots is beyond the scope of this paper, these observations support a role of  $\gamma_2$  not in photosynthesis but in development or maintenance/function of roots or other nonphotosynthetic tissues. Phylogenetic analyses (Fig. S7) likewise suggest the evolution of (possibly multiple) new "moonlight" functions for ATP synthase through divergent evolution of the  $\gamma$  subunits.

## **Materials and Methods**

Plant Materials and Growth Conditions. Wild-type Arabidopsis thaliana (ecotype Columbia) and ATPC2 T-DNA knockout lines, atpc2 (T-DNA is inserted in chl1\_5402949) and gamera in which ATPC2 overexpressed in the ATPC1 T-DNA knock out line (dpa1) (21) (see SI Materials and Methods) were grown

on soil under continuous light period at 20–30  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> at 22 °C for 4 wks, as described previously (21).

In Vivo Spectroscopic Assays. Maximal PSII quantum efficiency, LEF, and energy-dependent exciton quenching  $(q_E)$  were estimated from saturation-pulse chlorophyll a fluorescence yield measurements using the instrumentation and methods described previously (33). The value of steady-state fluorescence yield with all PSII centers closed,  $F_{M}$ , was obtained after at least 15 min illumination, at which point the photosynthetic rates and fluorescence parameters were stable. The value of  $F_{M}$  in fluorescence yield was estimated after 10 min of dark relaxation. Relative extents of steady-state, light-induced pmf (ECS<sub>t</sub>) and the conductivity of the ATP synthase to protons  $(g_{H+})$  were estimated from dark interval relaxation kinetics of absorbance changes associated with the ECS as described previously (15, 16). Flash-induced relaxation kinetics (FIRK) experiments were performed on infiltrated leaf discs described previously (17).

**Equilibrium Redox Titrations.** Redox titrations were performed in situ in fully expanded detached leaves vacuum-infiltrated with varying ratios of oxidized and reduced 20 mM DTT solutions for 30 min incubation in the dark, as described previously (22). The activity of the ATP synthase was then monitored using the FIRK of the ECS signal as described in (13) using the instrumentation described in (17). The equilibrium redox potential  $(E_h)$  for DTT at pH 7.5 was calculated according to the Nernst equation, using  $E_{m7.5} = -356$  mV for DTT.

**Determination of Redox States of γ Subunits.** The redox state of γ subunit of chloroplast ATP synthase was probed using the binding of AMS followed by nonreducing SDS/PAGE (26). To achieve oxidizing conditions, leaf 1 cm discs were infused with 0.1% Tween 20 and 20 mM Tricine containing 100 μM of MV followed by incubation for 30 min in darkness at room temperature. Reducing conditions were achieved by vacuum-infiltration of the leaf discs with 20 mM reduced DTT as a reducing agent (17) followed by illumination for 5 min by 100 μmol photons  $m^{-2}$  s<sup>-1</sup>.

Strongly oxidizing conditions were obtained by similar infiltration with 0, 0.5, or 5 mM diamide. After treatments, the discs were frozen in liquid nitrogen and ground with mortar and pestle, then extracted in 50 mM Tris·HCl (pH 8.0), 2 mM MgCl $_2$ , and 10 mM NaCl. The extractions were fractionated to soluble and insoluble protein by centrifugation. Protein precipitates were washed with acetone and dissolved in freshly prepared solution containing 1% SDS, 50 mM Tris·HCl (pH 8.0), and 15 mM AMS, as described in (26). Proteins were separated by SDS/PAGE using running buffer lacking reducing agent, 1% SDS, 50 mM Tris·HCl (pH 8.0), and 10% glycerol. Relative contents of  $\gamma$  subunits in wild type and gamera were estimated by Western blotting, as described in (33), using specific antibodies raised against  $\gamma_1$  and  $\gamma_2$  (see Fig. S2B). The antibody raised against the  $\gamma_1$  peptide reacts with both  $\gamma_1$  and  $\gamma_2$ .

**GUS Assay.** Seedlings were washed in water after soaking in 3.7% formaldehyde for 10 min. The seedlings were then incubated in 1 mM 5-bromo-4-chloro-3-indolyl  $\beta$ -D-glucuronide cyclohexylammonium salt, 100 mM potasium phosphate buffer (pH 7.0), 1 mM EDTA, 1 mM potassium hexacyanoferrate, 1 mM potassium hexacyanoferrate, and 0.3% Triton X-100 overnight at 37 °C. For visualization of blue staining the chlorophyll was removed by incubation in 70% ethanol (34).

**Observation of Root Phenotype.** Wild type, two homozygous *ATPC2* T-DNA knockout lines, atpc2 (T-DNA inserted in chl1\_5402949),  $atpc2_2$  (T-DNA inserted in chl 1\_5404314), and gamera were surface sterilized and germinated on Murashige and Skoog plates without sucrose in vertical transparent plates as in (21) . Plants were examined using light microscopy (Leica M165FC) after 5-d of growth. Tissue lengths were measured from recorded images using Image J (http://rsbweb.nih.gov/ij/). The lengths of root hairs were measured on the lower 5 mm of roots on four to five plants of each plant type (n=32-274).

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