Flexibility in the Energy Balancing Network of Photosynthesis Enables Safe Operation under Changing Environmental Conditions

Berkley J. Walker 1,2,*, David M. Kramer 1,3, Nicholas Fisher 1 and Xinyu Fu 1

1 Plant Research Laboratory, Michigan State University, East Lansing, MI 48823, USA; kramerd8@msu.edu (D.M.K.); nefisher@msu.edu (N.F.); fuxinyu2@msu.edu (X.F.)
2 Department of Plant Biology, Michigan State University, East Lansing, MI 48823, USA
3 Department of Biochemistry and Molecular Biology, Michigan State University, East Lansing, MI 48823, USA

* Correspondence: berkley@msu.edu; Tel: +517-355-3928

Received: 17 January 2020; Accepted: 15 February 2020; Published: 1 March 2020

Abstract: Given their ability to harness chemical energy from the sun and generate the organic compounds necessary for life, photosynthetic organisms have the unique capacity to act simultaneously as their own power and manufacturing plant. This dual capacity presents many unique challenges, chiefly that energy supply must be perfectly balanced with energy demand to prevent photodamage and allow for optimal growth. From this perspective, we discuss the energy balancing network using recent studies and a quantitative framework for calculating metabolic ATP and NAD(P)H demand using measured leaf gas exchange and assumptions of metabolic demand. We focus on exploring how the energy balancing network itself is structured to allow safe and flexible energy supply. We discuss when the energy balancing network appears to operate optimally and when it favors high capacity instead. We also present the hypothesis that the energy balancing network itself can adapt over longer time scales to a given metabolic demand and how metabolism itself may participate in this energy balancing.

Keywords: energy balancing; cyclic electron flux; malate valve; photorespiration; photosynthesis; C3 cycle

1. Introduction

Photosynthetic organisms must match energy supply from the light reactions with metabolic demands to enable safe, flexible and efficient photosynthesis. Because of the interdependency between energy supply and metabolic demand, it is valuable to consider this linked energy network and not as a series of separate metabolic processes. This energy balancing network integrates ATP and reductant supply and metabolic demand to allow plants to efficiently and safely harvest energy from the sun under dynamic conditions (Figure 1). From this perspective, we will first discuss the basic mechanisms of energy balancing before presenting the demand for energy balancing under a variety of conditions. We will then discuss how the structure and efficiencies of the energy balancing network are poised to provide and turnover ATP and reducing power under a variety of conditions before exploring how the energy balancing network responds to long-term changes in metabolic demand.
The core reactions of oxygenic photosynthesis involve a process called "linear electron flow" (LEF), which produce reducing power (referred to generically as NAD(P)H, in red) and ATP (yellow). Metabolic demand comprises the primary ATP and NADPH consuming processes in C3 plants, the C3 cycle, photorespiration (PR), nitrate assimilation (N) and the remaining metabolic sinks or CEF (in yellow). Energy Balancing is Essential for Safe and Optimal Photosynthetic Systems.}

2. Energy Balancing is Essential for Safe and Optimal Photosynthetic Systems

The light reactions of photosynthesis provide the chemical energy needed for plant metabolism. The core reactions of oxyogenic photosynthesis involve a process called "linear electron flow" (LEF), in which light energy is used to extract electrons from water and transfer them to NADP⁺ while generating ATP from ADP and Pᵢ [1,2], as detailed in Figure 2. These core processes store energy in two forms; ATP and NADPH. Extracting electrons from water and transferring them to NADP⁺, energy is stored in the two redox half reactions $4H^+ + O_2/H_2O \rightarrow NADP^+ + H^+/NADPH$. In addition, the transfer of electrons results in the formation of the proton motive force (pmf), an electrochemical gradient of protons across the thylakoid membrane, which is dissipated by the ATP synthase to fuel the formation of ATP from ADP and Pᵢ. The pmf is the sum of two energetic components; an electric field component ($\Delta \psi$) and the free energy stored in a chemical gradient of protons ($\Delta \rho \mathrm{H}$). Vectorial electron transfer from the lumenal to the stromal face of the thylakoid membrane, within photosystem II (PSII) and the cytochrome $b_6f$ complex by the Q-cycle mechanism (reviewed in [3]) and photosystem I (PSI) results in the formation of $\Delta \psi$. Both $\Delta \psi$ and $\Delta \rho \mathrm{H}$ are energetically equivalent drivers of the ATP synthase [4,5], but have very different impacts on photophysiological processes, as discussed below. One important feature of LEF is that it produces ATP and NADPH in a fixed stoichiometry, likely 2.6 ATP to 2 NADPH, or 1.28 ATP/NADPH [6].
"photosynthetic control" (reviewed in [21,22]) and subsequent PSI photodamage. Lumen acidification is, in turn, modulated by several processes that respond to the physiological state of the cell [1]. When metabolic demand is low, the activity of the ATP synthase is also down-regulated to slow proton ejection to PSI by slowing the rate of PQH$_2$ oxidation by the cytochrome $b_6f$ complex. The acidification of the lumen (indicated by the H$^+$ in the red box) activates violaxanthin deepoxidase (VDE) which converts violaxanthin (V) to zeaxanthin (Z) and protonates the PsbS protein, which triggers the photoprotective dissipation of light energy by the qE (black arrow). Lumen pH also regulates electron flow (red box with -'') to PSI by slowing the rate of PQH$_2$ oxidation at the cytochrome $b_6f$ complex.

The chloroplast must also balance the output of energy into the ATP and NADPH pools to perfectly match metabolic demands. The pool sizes of ATP and NADPH are small relative to the high fluxes of energy from the light reactions. Thus, any imbalance in the production and consumption of ATP or NADPH can rapidly lead to "metabolic congestion," depletion or buildup of metabolic intermediates, leading to the accumulation of high energy intermediates of the light reactions within seconds [7–10]. On the other hand, if too little ATP and NADPH are produced metabolic demand is energy limited, meaning that central metabolism is sub-optimal. The "correct" output of ATP and NADPH is a moving target since metabolic demand for ATP and NADPH changes dynamically based on environmental and physiological contexts (See below and [11]). The supply of ATP and NADPH must be matched with demand both in total capacity and stoichiometrically, and therefore plants have evolved mechanisms for regulating total energy output and fine tuning ATP/NADPH production ratios.

To regulate total energy production, chloroplasts partition light energy between photochemical processes which generate ATP and NADPH (LEF) and the energy dissipating process of "non-photochemical quenching" (NPQ) [12–17]. When metabolic demand for energy is less than current supply, the major form of NPQ, termed qE (for 'energy dependent' quenching), is triggered by acidification of the lumen (i.e., by the ΔpH component of pmf), through activation of violaxanthin deepoxidase, which catalyzes the conversion of violaxanthin to antheraxanthin and zeaxanthin [18], and through protonation of the antenna protein PsbS [19,20]. The ΔpH component of pmf also down-regulates electron flow by slowing plastoquinol (PQH$_2$) oxidation by the cytochrome $b_6f$ complex, preventing accumulation of electrons on highly reducing components of PSI, a process called "photosynthetic control" (reviewed in [21,22]) and subsequent PSI photodamage. Lumen acidification is, in turn, modulated by several processes that respond to the physiological state of the cell [1]. When metabolic demand is low, the activity of the ATP synthase is also down-regulated to slow proton efflux, increasing pmf and down-regulation of the light reactions [1,23–27]. The fraction of pmf stored in the

Figure 2. Basic Z-scheme model for the electronic and protonic circuits of the light reactions of photosynthesis, and the pmf paradigm for regulation of the light reactions. Scheme of linear electron flow (LEF) in oxygenic photosynthesis, in which light energy is captured by light harvesting complexes associated with photosystem II (PSII) and photosystem I (PSI), which initiates electron flow (orange arrows) from PSII, through the cytochrome $b_6f$ complex, plastocyanin (PC), to PSI and ferredoxin (Fd) and finally to NADPH. Also shown is the formation of the LEF is coupled to proton flow (blue arrows) at PSII and the cytochrome $b_6f$ complex, storing energy in the thylakoid proton motive force (pmf). Transfer of electrons from the lumenal to the stromal side of the thylakoid forms a transmembrane electric field (Δψ, blue arrow), while proton uptake from the stroma and deposition in the lumen lead to the formation of a transthylakoid pH gradient (ΔpH, red arrow), which together drive the synthesis of ATP from ADP + P$_i$ at the thylakoid ATP synthase, storing energy in ΔAGTP. The acidification of the lumen (indicated by the H$^+$ in the red box) activates violaxanthin deepoxidase (VDE) which converts violaxanthin (V) to zeaxanthin (Z) and protonates the PsbS protein, which triggers the photoprotective dissipation of light energy by the qE (black arrow). Lumen pH also regulates electron flow (red box with -'') to PSI by slowing the rate of PQH$_2$ oxidation at the cytochrome $b_6f$ complex.
ΔpH or Δψ is modulated to adjust its regulatory impact of a particular pmf [8,26,28]. The responses of qE to lumen pH may also be modulated by altering the expression of qE-related components [29–31]. Quantitatively, the dynamic range of NPQ is large, able to effectively partition from <5% to >80% of absorbed light energy to or away from energy production within tens of minutes. Importantly, even though increased light induces NPQ and decreases photochemical efficiency, the increase in total absorbed photons often more than compensates for this reduction and total LEF increases to safely produce sufficient NADPH to meet metabolic demand. Note that NPQ can only modulate total NADPH production from LEF with no change to the production stoichiometry of 1.28 ATP/NADPH.

3. The Structure of the Energy Network Simplifies ATP and NADPH Balancing

Once the total demand for NADPH is satisfied via the interplay between LEF and NPQ, other processes fine tune ATP/NADPH production ratios to match metabolic demand precisely. Downstream metabolism of an illuminated leaf (discussed in more detail below) requires ATP/NADPH ratios above 1.5, meaning that extra ATP is needed to achieve energy balancing. By poising metabolic demand at a higher ATP/NADPH ratio than that produced by LEF, the system can first produce the necessary NADPH, with coupled baseline production of ATP, before then producing the supplemental ATP needed for the specific metabolic context. This greatly simplifies the requirements of energy balancing since ATP and NADPH production ratios do not need to be independently re-adjusted following changes in total demand and total production capacity can be adjusted first based on a single factor (NADPH demand) before supplemental processes overcome the ATP deficit. In higher plants, three mechanisms are proposed to supply the additional ATP: (1) cyclic electron flux around PSI (CEF), (2) the malate valve and (3) the Mehler reaction. All three of these mechanisms have received extensive coverage in past reviews [6,32–34], and so we will focus on their basic mechanisms and relevance to the particular focus of this perspective.

4. Introduction to Supply-Side Mechanisms for Energy Balancing

4.1. Cyclic Electron Flux around Photosystem I

CEF contributes to the transthylakoid pmf without net production of NADPH by cycling electrons from photoexcited PSI via ferredoxin (Fd) back into the thylakoid plastoquinone (PQ) pool via the activity of Fd:PQ reductases (PQR) and the cytochrome b6f complex [35]. Aside from ATP generation, the proton gradient generated by CEF may also serve a photoprotective function by triggering qE (‘energy dependent’) NPQ, although CEF in itself is not essential for this process [8,36]. Many of the details of the electron transport pathways of CEF remain obscure. At least three PQR pathways have been postulated to function in CEF, which may operate in an organism-specific manner; (i) the antimycin A-sensitive Fd:PQ reductase (FQR), which has been proposed to be associated with the PGR5 and/or PGRL1 proteins [37–39], (but see [8,40–43] for additional viewpoints); (ii) the respiratory Complex I-like NADPH/Fd:PQ dehydrogenase (NDH) [44–47] and iii) direct reduction of b6f-bound PQ through Qi-associated FNR/Fd via b6f hemes b6fi [41,48–50]. Of these CEF pathways, those utilizing the proton motive NDH complex is likely to be the most energetically efficient, with a net H+ /2e− ratio of 8 [46], with the PGR5/PGRL1 and b6f Qi pathways yielding an H+ /2e− ratio of 4 by virtue of the (b6f-associated) Q-cycle alone [35].

The NDH pathway is, for the most part, associated with plant (and cyanobacterial) CEF, as this enzyme is absent from the majority of algal genera, although it should be noted that it is also absent from certain orchids, cacti and gymnosperms [51]. In general, the electron flux through CEF during steady-state photosynthesis in healthy, non-stressed C3 plants is considered to be small compared to LEF (i.e., ≤15%) [8,9,50], although it is likely to be (significantly) up-regulated during environmental stress like drought or during the induction of photosynthesis in dark-adapted plants, conditions under which increased ATP demand may be expected [48,52,53]. Nevertheless, this small flux is of vital importance for balancing the ATP and NADPH demands of metabolic supply and demand.
Furthermore, CEF is likely to be of particular importance to C4 photosynthetic species and aquatic algae to generate ATP and proton/ion gradients necessary for the carbon-concentrating mechanisms of these organisms [46,54–56].

4.2. The Malate Valve

The malate valve operates to adjust cellular ATP/NADPH supply by shuttling reducing power from the chloroplast to other organelles like the mitochondria via malate/oxaloacetate shuttles [32,57,58]. In the chloroplast, NADPH reduces oxaloacetate to malate via chloroplastic malate dehydrogenase (MDH). This malate is then exported from the chloroplast where it can be oxidized to form NADH in the cytosol, peroxisome or mitochondria via organelle-specific MDH enzymes. Reducing power shuttled to the mitochondria can fuel mitochondrial electron transport following transfer through the full complement of the electron transport complex proteins, generating additional pmf and ATP, or through only a portion of the electron transport complex proteins by dissipation of electrons via the alternative oxidase (AOX) or alternative mitochondrial electron carrier proteins. In all cases, the net effect is to increase ATP/NADPH supply either by decreasing NADPH or by simultaneously decreasing NADPH and increasing ATP. Chloroplastic NADP-MDH operates under tight light regulation via the Fd-thioredoxin (Fd-Trx) system, suggesting a role in photosynthetic energy balancing [59,60]. Importantly, the malate valve offers a way to “trade” NADPH for ATP by diverting reducing equivalence directly into mitochondrial electron transport.

4.3. The Mehler Peroxidase Reaction (Water–Water Cycle)

The term water–water cycle was coined by Asada (1999) [33] to indicate a process wherein electrons from LEF are extracted from water at the oxygen evolving complex of PSII, through the intermediate electron transfer chain, to PSI and to O₂, reforming H₂O. In plants, most of the O₂ reduction occurs by the Mehler peroxidase reaction (sometimes referred to as the water–water cycle (WWC)), electrons are transferred from low-potential donors (most probably F(X/A/B) within PSI to molecular oxygen, bypassing terminal NADP⁺ reduction, and producing superoxide. The resulting reactive oxygen species are rapidly detoxified by the activities of superoxide dismutase and the plastid ascorbate peroxidases [33]. This results in H⁺ translocation through water oxidation and the Q-cycle, without parallel NADPH production, increasing ATP/NADPH supply. Note, however, that PSI-involvement is not a strict requirement of a WWC (i.e., the ‘traditional’ Mehler peroxidase reaction), and the activity of the plastid terminal (plastoquinol) oxidase, also serves to bypass NADPH production, albeit at low capacity [61]. Alternative forms of the WWC are also found in moss, algae and cyanobacteria, where flavodiiron proteins function as NADPH-dependent oxygen reductases [62]. While the WWC may be important under certain stress conditions, current evidence suggests that it does not operate at significant rates under a variety of conditions (i.e., <5% of LEF in tobacco when the C3 was inhibited [63]), and so it will not be further considered in the context of energy balancing [64,65].

5. Metabolic Demand for ATP and NADPH

While plant metabolism employs ATP and NADPH in a myriad of biochemical reactions, the vast majority of ATP and NADPH flux in an illuminated leaf enters metabolic networks at relatively few nodes of central metabolism, most notably CO₂ assimilation and related processes, making it possible to reasonably estimate total ATP/NADPH demand [66]. Some reactions require reductive energy from alternative redox carriers (i.e., Fd or NADH) but for convenience in calculation and discussion, we will refer to reductive demand in terms of NADPH (2 e⁻) equivalents. The fixation of each CO₂, and subsequent regeneration of the C3 cycle intermediates requires 3 ATP and 2 NADPH for a total demand of 1.5 ATP/NADPH [67]. In C3 plants growing under ambient conditions, the next largest demand for ATP and NADPH is photorespiration, which results from the molecular fixation of O₂ by the first enzyme of the C3 cycle (rubisco, [66,68]). Photorespiration requires 3.5 ATP and 2 NADPH for
complete operation, meaning that as photorespiration increases relative to CO$_2$ fixation, ATP/NADPH demand increases as well.

Altering the relative rates of photorespiration and carbon fixation will alter the relative demands for ATP and NADPH. Rates of rubisco carboxylation ($V_c$) and oxygenation ($V_o$) determine downstream rates and energy requirements for carbon fixation and photorespiration respectively. Since $V_c$ and $V_o$ in C3 plants are constrained by rubisco kinetics, rates of each can be estimated for a given rate of net CO$_2$ exchange (A) and CO$_2$ and O$_2$ concentration to calculate subsequent ATP and NADPH demand [68–72]. While these calculations have been presented in part across many publications, we compile them all herein to make their use more convenient for the non-specialist to use measured gas exchange data to calculate $V_c$, $V_o$, ATP and NADPH demand, and extra ATP needed above that provided from LEF (see Appendix A). This quantitative framework requires several simplifying assumptions, but these estimates are close enough to show the magnitude of fluxes and relative impact between conditions.

While carbon fixation and photorespiration comprise the largest portion of central metabolic demand, other metabolic processes such as nitrate assimilation requires a significant contribution. Nitrogen assimilation in leaves involves nitrate reduction into nitrite by nitrate reductase (NR) in the cytosol, translocation of nitrite to chloroplast where it is reduced to ammonium by nitrite reductase (NiR), followed by ammonium assimilation into amino nitrogen via the glutamine synthetase (GS)-glutamine-2-oxoglutarate aminotransferase (GOGAT) pathway in the chloroplasts [73]. Nitrate assimilation to glutamine requires 5 NAD(P)H and 1 ATP. Specifically, reduction of one molecule of nitrate (oxidation state +5) to ammonium (oxidation state −3) requires eight electrons (equivalent to four NADPH), whereas the production of a glutamate via the GS-GOGAT pathway requires an additional two electrons (equivalent to 1 NADPH) and 1 ATP [66]. The reducing power required by the plastidic NiR and GS-GOGAT is supplied from photosynthetic electron transport via the reduced Fd. Higher rates of nitrate assimilation in the light than in the dark [74] reflects the tight connection between photosynthetic metabolism and nitrate assimilation. The reducing power needed for nitrate reduction via the cytosolic NR could be provided by the plastidic NAD-driven malate valve [75]. The NADPH demand for nitrate assimilation is estimated to range from ~0.35 to 3 µmol m$^{-2}$ s$^{-1}$ on an area basis, based on the nitrate assimilation rate measured by prior studies [76,77]. These rates of nitrate reduction would require ~2.5%–23% of total LEF in the sample dataset examined in Table 1, making nitrate assimilation a significant electron sink in terms of total electron flux.

Lipid biosynthesis represents another sink for NADPH and ATP consumption in plants, but quantitative estimates of its magnitude have not been reported. Lipids, being an important structural component of membranes, constitute approximately 5% to 10% of the dry weight of vegetative cells of plants [78]. The major constituents of lipids are fatty acids, which can represent up to 10% of the chemical energy of leaves on a biomass basis [79]. The synthesis and breakdown of fatty acids occur constitutively during leaf development. As much as 4% of total fatty acid content in leaves is degraded per day [80]. The turnover of fatty acids is exceeded by the rate of de novo fatty acid synthesis in non-senescent leaves. The net fatty acid accumulation generally increases during leaf expansion, with a rate ranging from 0.16 to 8 µmol carbon atoms mg$^{-1}$ chlorophyll h$^{-1}$ [80–83]. Plant de novo fatty acid synthesis is an energy-demanding process occurred in plastids. ATP drives the first committed step of fatty acid synthesis, the formation of malonyl-CoA from acetyl-CoA catalyzed by acetyl-CoA carboxylase. Reducing power in the form of NADPH and NADH is required for the two reductases involved in each round of fatty acid synthesis [78]. The predominant carbon source of plastidic acetyl-CoA is pyruvate, which is generated from photosynthetically fixed 3-phospho-D-glycerate (3-PGA) via the intermediate phosphoenolpyruvate. For every molecule of palmitic acid (16:0) produced, eight molecules of acetyl-CoA (generation of each acetyl-CoA from 3-PGA regenerates one ATP and one NADH), seven molecules of ATP, and 14 molecules of NAD(P)H are needed, resulting in the consumption of six molecules of NAD(P)H and surplus of one ATP collectively. Based on the total fatty acid content measured in Arabidopsis and Brachypodium distachyon (40 µg cm$^{-2}$ leaf area, [83]), we estimate that the NADPH demand to maintain the 4% turnover rate of fatty acids is ~0.5
µmol m\(^{-2}\) s\(^{-1}\), which represents approximately 2% and 0.5% of the total NADPH demand under low light and high light, respectively (Table 1). Due to the small pool size of fatty acids in young leaves, the NADPH demand for fatty acid synthesis would be even smaller in the developing leaves. Although the NADPH demand for fatty acid synthesis is relatively small, this process is highly dependent on light and subject to redox regulation [84]. Nevertheless, while up to 2% of total NADPH demand has potential implications to some situations, this is insufficient to significantly affect calculations for total leaf energy balancing.

| Table 1. Requirements for energy production for the supply and demand of the energy balancing network under low and high light in Nicotiana tabacum. For metabolic demand, shown are rates of CO\(_2\) assimilation (A), intercellular and chloroplastic CO\(_2\) concentration, rates of rubisco carboxylation (v\(_c\)) and oxygenation (v\(_o\)), rates of nitrate reduction (V\(_n\)), rates of lipid production (V\(_l\)) and total ATP and NADPH demand. For energy supply shown are photosynthetically active radiation (PAR), measured rates of electron transport through PSII (LEF) and PSI (J\(_{PSI}\)), rates of linear electron flux needed to provide sufficient NADPH for metabolic demand (LEF\(_{pred}\)), ATP produced from LEF\(_{pred}\) (ATP\(_{LEF}\)) and the ATP deficit. For energy balancing, shown are the electron and photon demands for the ATP deficit to be provided by CEF via the NDH, FQR or b5f pathways or the malate valve. Details for these calculations found in the text. Values taken from Miyake et al. 2005 [1] indicated with a star (*), with remaining values calculated or assumed herein. For these calculations R\(_s\), Γ\(^*\) and g\(_{m}\) were assumed to be 1.5 µmol CO\(_2\) m\(^{-2}\) s\(^{-1}\), 4.7 Pa and 6 µmol CO\(_2\) Pa CO\(_2\)^\(-1\) m\(^{-2}\) s\(^{-1}\). |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| **High Light (1100 PAR)** | **Low Light (150 PAR)** |
| Metabolic demand | Energy supply |
| A (µmol CO\(_2\) m\(^{-2}\) s\(^{-1}\)) | 21.3 * | 5.7 * |
| Intercellular CO\(_2\) (Pa) | 23.0 * | 25.0 * |
| Chloroplastic CO\(_2\) (Pa) | 19.5 | 24.1 |
| v\(_c\) (µmol CO\(_2\) m\(^{-2}\) s\(^{-1}\)) | 30.1 | 8.9 |
| v\(_o\) (µmol O\(_2\) m\(^{-2}\) s\(^{-1}\)) | 14.5 | 3.5 |
| v\(_n\) (µmol N m\(^{-2}\) s\(^{-1}\)) | 1.5 | 0.5 |
| v\(_l\) (µmol N m\(^{-2}\) s\(^{-1}\)) | 0.3 | 0.1 |
| Total ATP demand (µmol ATP m\(^{-2}\) s\(^{-1}\)) | 143 | 40 |
| Total NADPH demand (µmol NADPH m\(^{-2}\) s\(^{-1}\)) | 97 | 27 |
| Total ATP/NADPH ratio | 1.47 | 1.45 |
| Energy balancing requirements via CEF |
| NDH e\(^-\) (µmol m\(^{-2}\) s\(^{-1}\)) | 43.3 | 11.1 |
| NDH photons (µmol m\(^{-2}\) s\(^{-1}\)) | 43.3 | 11.1 |
| FQR/b5f e\(^-\) (µmol m\(^{-2}\) s\(^{-1}\)) | 86.6 | 22.2 |
| FQR/b5f photons (µmol m\(^{-2}\) s\(^{-1}\)) | 86.6 | 22.2 |
| Energy balancing requirements via malate valve |
| e\(^-\) (µmol m\(^{-2}\) s\(^{-1}\)) | 9.3 | 2.4 |
| Photons (µmol m\(^{-2}\) s\(^{-1}\)) | 18.5 | 4.7 |

6. Determining the Efficiency of EnergyBalancing Mechanisms

As an autotrophic organism, the energy that fuels metabolism in plants is derived ultimately from absorbed photons, providing a metric by which to gauge the efficiency of an energy balancing mechanism. Photon use efficiency has thus provided a logical objective function for approaches that assume photoautotrophs use light energy optimally (i.e., [85]), but given the massive amount of absorbed energy that is dissipated as NPQ under high light, it is not clear that light energy is always limiting to growth. Additionally, given the dynamic fluctuations in energy demand and light supply.
many plants face under growing conditions, it is likely that the capacity for a given energy balancing mechanism may become more important than its efficiency when light energy supply is adequate. In this section we outline the photon costs of various energy balancing mechanism and incorporate them into a quantitative framework. We then use this framework to examine past work and hypothesize that the energy balancing network operates in a high or low-efficiency mode based on light availability.

To examine the energy requirements for energy balancing under various light conditions, the ATP and NADPH demand for the C3 cycle and photorespiration has been determined from past work which paired concurrent gas exchange with measurements of electron flux through PSII and PSI (Figure 1, Table 1 and [86]).

Different pathways of CEF have different costs for energy balancing, depending on how many H\(^+\) are pumped per electron excited by an absorbed photon. As outlined above, the highest efficiency CEF pathway proceeds through NDH, which pumps 4 H\(^+\)/2 e\(^-\) (Table 2). The FQR and b\(_{6f}\) pathways have identical yields of 2 H\(^+\)/2 e\(^-\). Since 14 H\(^+\) are required to generate 3 ATP in the chloroplast, CEF has an ATP/photon or e\(^-\) stoichiometry of 0.43 via NDH and 0.21 via FQR or b\(_{6f}\). Additionally, the e- and photon demand for energy balancing can be calculated for the low and high-light conditions presented in Table 1 and data from Miyake et al., 2005. Under low light, 11 or 22 \(\mu\)mol photons m\(^{-2}\) s\(^{-1}\) are needed to produce the ATP needed to balance energy supply via NDH or FQR/b\(_{6f}\) pathways, respectively, or between 7% and 15% of the total incident light (Table 1). Under high light, this requirement drops to 4%–8%. To gain a more complete picture of the relative energy cost of these mechanisms under their respective light conditions, the photon demand can be expressed in terms of actual absorbed photon energy that enters photochemistry by adding the rates of flux through PSII and PSI. Interestingly, this recalculation reveals that as light level increases, a greater percentage of photon energy absorbed and passed through the photosystems would need to be partitioned to CEF processes for energy balancing, specifically 11%–22% under low light and 13%–27% under high light. Under high light, however, energy from more photons is dissipated as NPQ compared to low light. If energy to NPQ is considered as excess, this means that there is more excess energy under high light that can be used to drive CEF. Specifically, based on the data from Miyake et al. 2005 [86], energy from only 48 \(\mu\)mol photons m\(^{-2}\) s\(^{-1}\) was dissipated via NPQ under low-light conditions, but energy from 776 \(\mu\)mol photons m\(^{-2}\) s\(^{-1}\) was dissipated via NPQ under high-light conditions. These numbers reveal that while photon energy could be limiting to drive CEF under low-light conditions, there appears to be enough surplus photon energy available under high light to drive CEF. This is expected because if NPQ limits excitation of PSII, it should also limit flux to PSI through LEF, but it will not necessarily limit PSI electron flow of electrons through CEF.

| Table 2. Energy requirements and efficiencies of CEF pathways and the malate valve to produce supplemental ATP. Shown are the number of absorbed photons used for the calculation of each pathway. Further details and assumptions for calculations are found in the text. |

| CEF Pathways | NDH | FQR | b\(_{6f}\) | Malate Valve |
|--------------|-----|-----|--------|-------------|
| **Chloroplast** | | | | |
| e-/photons | 1 | 1 | 1 | 0.5 |
| H\(^+\)/e\(^-\) | 2 | 1 | 1 | 3 |
| ATP/H\(^+\) | 0.21 | 0.21 | 0.21 | 0.21 |
| ATP/photon in chloroplast | 0.43 | 0.21 | 0.21 | 0.32 |
| **Mitochondria** | | | | |
| H\(^+\)/e\(^-\) | - | - | - | 5 |
| ATP/H\(^+\) | - | - | - | 0.27 |
| ATP/photon in mitochondria | - | - | - | 0.68 |
| **Total ATP/photon** | 0.43 | 0.21 | 0.21 | 1.00 |
The energetics of the malate valve are more difficult to assess given the added complexity of transport and flexibility of mitochondrial electron transport. The initial energetics and efficiency of the malate valve are tied to LEF; eight photons produce two NADPH and 12 H\(^+\), resulting in 2.57 ATP. The energetics following the transport of the reducing power of 2 NADPH into mitochondrial electron transport and ATP generation depend on the e\(^-\)/H\(^+\) and, more generally, the e\(^-\)/ATP efficiency of the mitochondria. For our theoretical evaluation of malate valve energetics, we will first assume mitochondrial electron transport operates optimally and each electron contributes maximally to the proton gradient, passing through Complex I, III and IV to pump 10H\(^+\)/2e\(^-\). To produce ATP, these protons pass through a ring of c-subunits of ATPase, with each full rotation producing three ATP and the number of H\(^+\) per rotation depending on the number of c-subunits present in the ring [87–91]. We assume the number of c-subunits is the same as found in animal cells since there is no available data on plant mitochondrial c-subunit number, requiring eight H\(^+\)/3 ATP, although in yeast there are 10 c-subunits [92]. Since each molecule of ATP synthesized requires the (electroneutral) transport of one P\(_i\) with the associated (electrogenic) ADP\(^3-\)/ATP\(^4-\) exchange activity of the mitochondrial adenine nucleotide translocase (equivalent to the uptake of an additional proton per molecule of ATP synthesized) [93,94], the final stoichiometry is 11 H\(^+\)/3 ATP, making a theoretically maximum ATP/oxygen ratio of 2.7 [95]. This stoichiometry is closely matched in experimental measurements of the ratio of 2.6 ADP/oxygen consumed in intact mitochondria in potato [96], suggesting that these stoichiometries reasonably approximate mitochondrial respiration in plants despite the highly-branched potential of mitochondrial electron transport. Therefore, for every two NADPH (4 e\(^-\)) that are processed via the malate valve, 5.45 ATP are produced in the mitochondria. The above discussion focuses specifically on the ATP produced via mitochondrial respiration fueled by electrons provided from the light reactions, we recognize that some ATP may be produced in the light from pyruvate produced during “dark-type” glycolysis. Exact rates of glycolysis-supplied mitochondrial respiration in the light are not available, but estimates from CO\(_2\) gas exchange indicate these rates are rather small compared to net assimilation and lower than rates measured in the dark [97–99], suggesting that the bulk of ATP generated in the mitochondria may come from other sources (such as the Mehler valve), but more information is needed to explore this in more detail.

To integrate the production stoichiometries into a complete malate valve cycle, the costs of transporting ATP from the mitochondria back into the chloroplast where it is primarily needed for the phosphorylation of C3 and photorespiratory cycle intermediates must also be considered. Transport of ATP from the mitochondria proceeds via the ADP/ATP translocase [100,101] and into the chloroplast via the plastidic ADP/ATP transporter [102,103].

The above energetics determine that the malate valve is a highly efficient ATP producer on a photon basis. For every eight photons of light energy, 2.57 ATP are produced in the chloroplast and 5.45 are produced in the mitochondria for a total ATP/photon ratio of 1, much higher than the 0.21–0.43 determined for CEF (Table 2). This high ATP/photon ratio means that much less absorbed light energy is needed for energy balancing assuming low and high-light conditions (Table 1). Specifically, only 4.7 and 18.5 \(\mu\)mol photons m\(^{-2}\) s\(^{-1}\) were needed under low and high light, respectively (Table 1). This comprises only 3.28 and 1.7% of incident irradiance for the low and high light intensities.

7. With an Efficient Malate Valve, Why is CEF Important?

Given the theoretically greater efficiency of the malate valve, how can we explain the commonly observed participation of CEF in energy balancing? We propose that the real energy cost of CEF will depend on the light intensity and other factors. At low light, when energy capture is strongly limited by the number of photons captured by the photosystems, activating CEF will require diverting energy from LEF, limiting the overall efficiency of energy capture. However, as light input nears saturation imposed by downstream reactions, PSII efficiency drops substantially, either by decreased efficiency of antenna (increased NPQ), or by increases in the fraction of closed PSII centers. In this case, activating CEF may have little effect on the efficiency of LEF (because it is already light saturated) but will increase...
total energy capture, albeit with a higher fraction stored in ATP/NADPH. Indeed, experimentally, CEF appears to play a larger role in energy balancing under high, but not low irradiances when there is limiting energy available from absorbed photons [72,86].

For example, when high light intensities drive light-saturated photosynthetic rates, CEF shifts proportionally in response to changing CO₂ to cover ATP deficient predicted from gas exchange [72]. These measurements were made across CO₂ concentrations and reveal the capacity for CEF to respond to changes in leaf energy demand (Figure 3). We refer to the measured change of CEF in response to changing energy balancing requirements as the “dynamic range of CEF”. By comparing the measured dynamic range of CEF to the change predicted from gas exchange modeling, the degree by which CEF participates in energy balancing can be determined. The dynamic range of CEF can run into a self-regulating upper limit when the high rates of CEF increase the ΔpH and initiate qE-dependent NPQ, as occurs in shade leaves exposed to increasing light [104]. This could serve as a protective mechanism, when CEF increases ΔpH to sufficient levels, light harvesting is down-regulated and further energy mismatch is avoided when the capacity for energy balancing by CEF is reached. Interestingly, the dynamic range of CEF was minimal in response to changing energy demanding conditions when measured under low light, indicating that processes other than CEF (e.g., the malate valve) accomplish energy balancing under these low flux conditions [72].

![Figure 3. Comparison of the measured relative rate of cyclic electron flux (CEF: circle symbols) to the predicted change in CEF required to match ATP/NADPH supply with demand across CO₂ concentration (line). Shown are n = 3–4 ± SE. This data is a replotted subset of measurements from Walker et al. (2014) [72].](image-url)

A role for “low-light, high-efficiency” and “high-light, low-efficiency” energy balancing networks is further supported by flux balance analysis of photosynthetic systems and in mutant lines deficient in CEF. Flux balance analysis of photosynthetic systems that are optimized for energy production per photon of absorbed light predict the malate valve to be the optimal mechanism of energy balancing unless the additional costs of enzymatic interconversions are introduced into the model [105]. This is also supported in work using a modified flux balance analysis approach, which weights flux solutions based on pathway complexity [106]. As light intensity increases, absorbed light energy is actively released as NPQ, indicating that under high light, the system is no longer light limited and the energy
balancing network could trade the more light-optimal malate valve for CEF. This position is supported by work showing that CEF is critical for plant growth under high, but not low-light conditions [107].

It is not clear what the advantages of CEF might be over the malate valve, but the speed and flexibility of CEF may provide an explanation. The operation of the complete malate valve requires tight coordination of enzymatic and transport activity between the chloroplast and mitochondria, limiting the dynamic range of its energy balancing capacity over short time scales. The malate valve also requires the careful coordination of two electron transport chains in separate organelles, further complicating the upregulation of this pathway under greater energy balancing demand. CEF occurs only in the chloroplast, simplifying the signaling network required to up- or down-regulate ATP production. By contrast, CEF is likely regulated by stromal ATP levels as well as stromal redox state [108] and thus may also be more rapidly responsive to alterations in energy demands, e.g., during induction or rapid changes in light, CO₂, etc. whereas the malate valve appears to require (potentially slower) redox activation (see below). Thus, having at two routes of ATP/NADPH balancing, provides photosynthetic systems with greater flexibility to balance diverse metabolic imbalances as well as providing optimal efficiency under low light (malate valve) or more rapid/responsive responses (CEF) pathways. We further hypothesize that the baseline requirements for energy balance are achieved by the more light-optimal malate valve. This baseline activity satisfies the needs for energy balancing until greater capacity is needed, such as occurs under higher light regimes. Under high light, CEF acts as a highly flexible stop-gap to allow energy balancing to occur under dynamic conditions.

8. Demand-Side Energy Balancing Processes

While there is much focus on how supply-side reactions mediate energy balancing, there is less focus on how metabolic demand itself changes. For example, under increased ATP/NADPH demand, metabolism could either increase the supply of ATP via supply-side mechanisms like CEF or the malate valve or reprogram metabolism itself to use the supply more optimally. A simple example of this is in the redox regulation of the C3 cycle, where multiple redox post-translational modifications may tune activity to available reducing power availability [109].

There is additional evidence for this demand-side reprogramming to achieve energy balancing in the unique link between nitrate assimilation and photorespiration. C3 plants have lower nitrate assimilation rates when photorespiration is reduced through altered atmospheres [76,77,110–112]. This link could be explained if under ambient conditions, the increased demand for ATP/NADPH imposed by photorespiration is offset by increased rates of nitrate assimilation, which has a much lower ATP/NADPH demand. This would achieve ATP/NADPH balance not exclusively by increased ATP supply, but by repartitioning demand-side processes themselves. Interestingly, expression of nitrate assimilation genes increase in NADP-MDH mutants, suggesting that nitrate assimilation could be a compensatory response to achieve energy balancing when the malate valve is disrupted [75]. NADP-MDH mutants also show improved growth on nitrate-rich media [75,113]. These experiments demonstrate that nitrate assimilation and the malate valve may cooperate to maintain a baseline level of energy balancing, increasing the complexity of the energy balancing network.

9. The Acclimation of Energy Balancing Networks to Long-Term Change in Energy Demand

As mentioned above, plants must cope in the long term to changing ATP and NADPH demand to achieve energy balance. It is unclear whether the same mechanisms balance energy mismatches under long time scales as occur under shorter time scales. Furthermore, it is unknown to what degree energy balancing networks poise to a given demand and how this poise acclimates to changing demands. The acclimation of the energy balancing network can be investigated experimentally either via transition experiments or by examining mutants with an altered network capacity that forces flux through alternative facets of the network. We will first discuss the potential for acclimation of supply-side processes to changing energy demand before outlining how metabolic demand itself may acclimate to changes in energy balancing requirements.
According to our model of the two-component supply-side energy balancing system, malate valve activity should scale with excess ATP demand over long-term transitions to optimally use absorbed light. The largest driver of excess ATP demand for any condition is increased light, and so this model predicts that malate valve activity should increase with light. Indeed there appears to be a light and dark malate valve cycle, with the dark cycle relying on plastidic NAD-MDH and the light shuttle using plastidic NADP-MDH [58]. The switch to the NADP-MDH cycle is mediated through the light-dependent Fd-Trx system [114,115]. This activation (at least for NADP-MDH in isolated Pea chloroplasts) occurs within 10–20 min, and so activation of this component is likely integrated in our short-term measurements [116]. Activation occurs even more rapidly during a high-light transition [117]. This is an effective regulatory strategy at short time scales, since it activates malate valve activity when there is a surplus of NADPH, as when the C3 cycle is not consuming it fast enough and reduces it when there is too much NADP\(^+\) [58]. It is also likely that as the malate valve is reaching full capacity, CEF plays a role in vivo during very short time scales, at least in C3 plants [118].

Other factors increase malate valve capacity over longer time scales. For example, after transfer to sustained high light, NADP-MDH expression and protein levels increase, suggesting increased capacity of the malate valve following hours of exposure to the new condition [119]. Interestingly, the same response is not observed when photoperiod increases, suggesting that a photosynthetic steady-state solution must be found and that the effect is not cumulative.

Interestingly, chloroplastic nadh-mdh mutants show no phenotype, even under stress conditions, potentially due to additional compensatory redox strategies [75,113]. It is important to note that the malate valve shuttles reducing power not only between the chloroplast and mitochondria, but also the peroxisome during photorespiration [120]. In contrast to chloroplastic nadh-mdh mutants, mitochondrial nadh-mdh mutants lacking both MDH isoforms (mmdh1mmdh2) show lowered photosynthetic rates and growth rates [121]. These decreases were likely due to impaired shuttling of reducing power from the mitochondrion to the peroxisome via the malate valve to provide the reducing power for hydroxypyruvate reduction in photorespiration, a viewpoint supported using \(^{13}\)C flux analysis of mmdh1 [122]. Indeed mmdh1mmdh2 show reduced photorespiratory capacity, but the reduced growth and photosynthesis is not explained strictly by decreased availability of reductant to photorespiration since mutants lacking the peroxisomal MDH isoforms show an even more subtle phenotype than mmdh1mmdh2 [123,124]. This work with mitochondrial MDH indicates that the malate valve is not strictly required for energy balancing, but it is important for optimal photosynthesis and long-term growth.

The capacity for CEF itself may also increase over longer time scales to allow for increased energy balancing demand. Notably, NDH and FQR content change under different growth conditions [51]. The ratio of PSI and PSII re-proportions when plants are grown under light regimes with outputs that favor PSI or PSII. After hours or days, this results in changes to the actual stoichiometry of PSI and PSII photosystems, in green algae [125–127] and plants [128]. This re-proportioning also occurs days following transition between different light qualities, which can increase the capacity for CEF [129]. At short time scales, repartitioning of light energy between PSI and PSII occurs when reduced PQ builds up and triggers the phosphorylation of the PSII light harvesting complex. These then migrate to PSI to balance out energy capture [130]. While these state transitions occur in response to long-term differences in energy supply (changes in light regimes), it is not clear if this happens in response to long-term changes in energy demand. Such a change would predict that as conditions decrease in the ratio of photorespiration, there should be a decrease in demand for CEF and, therefore, a decrease in the PSI/PSII ratio.

Measurements of PSI/PSII from plants grown under conditions of different ATP/NADPH demand did not indicate that the capacity for CEF change with energy demand via changes in photosystem stoichiometry. Specifically, there was no difference in PSI/PSII in aspen trees exposed to elevated CO\(_2\) (560 PPM) over a single season following 5 years of elevated high CO\(_2\) treatment [131]. However, this increase in CO\(_2\) is not expected to change the demand for CEF by all this much (~1% of LEF
change). Additionally, micro-array work in soybeans exposed to 550 PPM also show no difference in photosystem expression, but interestingly show an increased expression of a mitochondrial ATP/ADP antiporter [132]. Overall, these findings do not point clearly to the acclimation of the capacity of CEF in response to changing energy demand, but the treatments resulted in relatively modest changes in energy demand and CEF was not evaluated specifically. There is clearly room for more work examining this question specifically.

10. Conclusions

The energy-balancing network comprises a flexible set of possibilities that enable partitioning through pathways with different ATP and NADPH production stoichiometries that require different amounts of light energy. We hypothesize that the network partitions flux through high-efficiency pathways (e.g., the Malate valve) when light is limiting and high-efficiency pathways (e.g., CEF) when light is abundant. Furthermore, the energy balancing network may adapt to long-term energy demand through enzyme expression.

Author Contributions: This article was prepared with the following contributions: Conception, B.J.W. and D.M.K.; Formal Analysis, B.J.W.; Writing — original draft, B.J.W., D.M.K., N.F., and X.F.; writing—review and editing, B.J.W., D.M.K., N.F., and X.F.; Funding acquisition, B.J.W. and D.M.K. All authors have read and agreed to the published version of the manuscript.

Funding: This work was supported by the Division of Chemical Sciences, Geosciences and Biosciences, Office of Basic Energy Sciences of the United States Department of Energy [Grant DE-FG02-91ER20021].

Acknowledgments: We would like to thank Yair Shachar-Hill for discussions on flux balance analysis work examining the malate valve and cyclic electron flux.

Conflicts of Interest: The authors declare no conflict of interest.

Abbreviations

| Abbreviation | Full Name |
|--------------|-----------|
| $\Delta p$H | The gradient potential component of the proton motive force |
| $\Delta \psi$ | Electric force component of the proton motive force |
| A | Net CO$_2$ assimilation |
| AOX | Alternative oxidase |
| CEF | Cyclic electron flux around photosystem I |
| Fd | Ferredoxin |
| Fd-Trx | Ferredoxin-thioredoxin system |
| FQR | Ferredoxin:plastoquinone reductase |
| GS | Glutamine synthetase |
| GOGAT | Glutamine-2-oxoglutarate aminotransferase |
| LEF | Linear electron flow |
| NDH | NADPH/Ferredoxin:plastoquinone dehydrogenase (NDH) |
| pmf | Proton motive force |
| MDH | Malate dehydrogenase |
| NiR | Nitrite reductase |
| NPQ | Non-photochemical quenching |
| NR | Nitrate reductase |
| PQ | Plastoquinone |
| PQH$_2$ | Plastoquinol |
| PQR | Ferredoxin:plastoquinone reductases |
| PSI | Photosystem I |
| PSII | Photosystem II |
| $q_E$ | Energy-dependent quenching |
| $R_l$ | Respiration in the light |
| $V_c$ | Rate of rubisco carboxylation |
| $V_o$ | Rate of rubisco oxygenation |
Appendix A

Calculating ATP and NADPH Ratios from Gas Exchange Data

While the theory behind calculating ATP and NADPH demand ratios for rubisco carboxylation ($V_c$) and oxygenation ($V_o$) from gas exchange data has been published on previously [6,72], the underlying derivation and final equations have been admittedly lacking. Here, we present a complete derivation, complete with underlying assumptions, appropriate to allow the non-specialist to apply these calculations to their gas exchange data. We will not attempt a full derivation of the underlying biochemical model of leaf photosynthesis, but will instead refer to the relevant equations directly as presented completely previously [133,134].

The cornerstone equation for modelling net CO$_2$ assimilation (A) is the mass balance subtracting from rates of carbon fixation via $V_c$ and rates of CO$_2$ loss from photorespiration and respiration in the light ($R_l$). Rates of photorespiratory CO$_2$ loss are calculated by multiplying CO$_2$ loss per rubisco oxygenation (usually assumed to be 0.5) by $V_o$ and total A is represented by Equation (2.1) in von Caemmerer 2000 [133]:

$$A = V_c - 0.5V_o - R_l \quad (A1)$$

Since our goal is to use measured rates of A to estimate $V_c$ and $V_o$, and subsequent ATP and NADPH demand, it becomes convenient to express Equation (A1) in terms of one unknown variable ($V_c$) and then solve for $V_o$. Equation (A1) is expressed in terms of $V_c$ in principle by combining rubisco specificity for CO$_2$ relative to O$_2$ (S$_{c/o}$) with measured gas concentrations to determine what catalytic rates of $V_c$ and $V_o$ would produce the measured A. This is accomplished based on the following relationships (Equations (2.16) and (2.18) from von Caemmerer 2000 [133])

$$\phi = \frac{V_o}{V_c} \quad (A2)$$

$$\phi = \frac{2\Gamma^*}{C_c} \quad (A3)$$

where $C_c$ is the partial pressure of CO$_2$ at the site of rubisco catalysis and $\Gamma^*$ is the CO$_2$ compensation point in the absence of $R_l$ defined as

$$\Gamma^* = \frac{0.5O}{S_{c/o}} \quad (A4)$$

where O is the oxygenation partial pressure. Note that since O is part of the definition of $\Gamma^*$, it must be scaled according to the measurement concentration if an altered oxygen background is used during the experiment. In using Equation (A1), $R_l$ is assumed or independently measured under the experimental conditions using a variety of gas exchange approaches and treated as a constant [98,135,136]. With Equations (A1)–(A3) we are able to represent the relationship between A and $V_c$ with no other unknown variables

$$V_c = \frac{A + R_l}{1 - \Gamma^*} \quad (A5)$$

The solution for $V_c$ can then be used with Equation (A1) to solve for $V_o$.

$$V_o = \frac{V_c - A - R_l}{0.5} \quad (A6)$$

With $V_c$ and $V_o$ determined from the above, the rate of demand for ATP ($V_{ATP}$) and NAD(P)H ($V_{NADPH}$) can then be determined based on the requirements for the C3 cycle (3 ATP and 2 NADPH) and photorespiration (3.5 ATP and 2 NAD(P)H, [66,67]) according to

$$V_{ATP} = 3V_c + 3.5V_o \quad (A7)$$
And
\[ V_{\text{NADPH}} = 2V_c + 2V_o \]  
(A8)

Of course additional energy demanding processes can be added to Equations (A6) and (A7) to determine total leaf energy demand [72], but we have limited these calculations to those most directly measured using gas exchange.

It should be noted that several of the constants assumed above require additional interpretation depending on the species and conditions they are measured under. These calculations depend on \( C_c \) to account for the chloroplastic supply of \( \text{CO}_2 \), but standard gas exchange measurements can only practically resolve the concentration of \( \text{CO}_2 \) in the intercellular airspace (\( C_i \)). \( C_i \) can be converted to \( C_c \) assuming a simple linear conductance using Fick’s law as

\[ C_c = C_i - \frac{A}{g_m} \]  
(A9)

where \( g_m \) is the mesophyll conductance to \( \text{CO}_2 \) diffusion. Selecting an appropriate \( g_m \) to use experimentally is complicated since it varies by species, temperature and the underlying theory used for it estimation [137–145]. Fortunately, under most conditions, \( V_{\text{ATP}} \) and \( V_{\text{NADPH}} \) are not extremely sensitive to small errors in \( g_m \) assumptions, but a sensitivity analysis can be performed to confirm that the findings of a study are robust. Note that stomatal conductance has a similar impact on changing \( C_i \) for a given photosynthetic rate. Since stomata close during drought, this means that the ratio \( V_o/V_c \) increases under these water-limiting conditions, increasing metabolic demand for ATP/NADPH. Additionally, the temperature response of \( \Gamma^* \) should be accounted for in addition to its linear dependence on \( O \) [138].

References
1. Miyake, C.; Miyata, M.; Shinzaki, Y.; Tomizawa, K.-I. CO\(_2\) Response of Cyclic Electron Flow around PSI (CEF-PSI) in Tobacco Leaves-Relative Electron fluxes through PSI and PSII Determine the Magnitude of Non-photochemical Quenching (NPQ) of Chl Fluorescence. Plant Cell Physiol. 2005, 46, 629–637. [CrossRef]
2. Kramer, D.M.; Avenson, T.J.; Edwards, G.E. Dynamic flexibility in the light reactions of photosynthesis governed by both electron and proton transfer reactions. Trends Plant Sci. 2004, 9, 349–357. [CrossRef]
3. Ort, D.R.; Yocum, C.F. Light reactions of oxygenic photosynthesis. In Oxygenic Photosynthesis: The Light Reactions; Ort, D.R., Yocum, C.F., Eds.; Kluwer Academic Publishers: Dordrecht, The Netherlands, 1996; pp. 1–9.
4. Cape, J.L.; Bowman, M.K.; Kramer, D.M. Understanding the cytochrome \( bc \) complexes by what they don’t do. The Q-cycle at 30. Trends Plant Sci. 2006, 11, 46–55. [CrossRef]
5. Fischer, S.; Graber, P. Comparison of DpH- and Dy-driven ATP synthesis catalyzed by the H(+)‑ATPases from Escherichia coli or chloroplasts reconstituted into liposomes. FEBS Lett. 1999, 457, 327–332. [CrossRef]
6. Hangarter, R.P.; Good, N.D. Energy thresholds for ATP synthesis in chloroplasts. Biochim. Biophys. Acta 1982, 681, 396–404. [CrossRef]
7. Kramer, D.M.; Evans, J.R. The importance of energy balance in improving photosynthetic productivity. Plant Phys. 2011, 155, 70–78. [CrossRef] [PubMed]
8. Avenson, T.J.; Kanazawa, A.; Cruz, J.A.; Takizawa, K.; Ettinger, W.E.; Kramer, D.M. Integrating the proton circuit into photosynthesis: Progress and challenges. Plant Cell Environ. 2005, 28, 97–109. [CrossRef]
9. Avenson, T.J.; Cruz, J.A.; Kanazawa, A.; Kramer, D.M. Regulating the proton budget of higher plant photosynthesis. Proc. Natl. Acad. Sci. USA 2005, 102, 9709–9713. [CrossRef] [PubMed]
10. Cruz, J.A.; Avenson, T.J.; Kanazawa, A.; Takizawa, K.; Edwards, G.E.; Kramer, D.M. Plasticity in light reactions of photosynthesis for energy production and photoprotection. J. Exp. Bot. 2005, 56, 395–406. [CrossRef] [PubMed]
11. Amthor, J.S. From sunlight to phytomass: On the potential efficiency of converting solar radiation to phyto-energy. New Phytol. 2010, 188, 939–959. [CrossRef]
12. Blankenship, R.E.; Tiede, D.M.; Barber, J.; Brudvig, G.W.; Fleming, G.; Ghirardi, M.; Gunner, M.R.; Junge, W.; Kramer, D.M.; Melis, A.; et al. Comparing Photosynthetic and Photovoltaic Efficiencies and Recognizing the Potential for Improvement. Science 2011, 332, 805–809. [CrossRef] [PubMed]

13. Müller, P.; Li, X.-P.; Niyogi, K.K. Non-photochemical quenching. A response to excess light energy. Plant Physiol. 2001, 125, 1558–1566. [CrossRef] [PubMed]

14. Anderson, B.; Barber, J. Mechanisms of photodamage and protein degradation during photoinhibition of photosystem II. In Photosynthesis and the Environment; Baker, N.R., Ed.; Kluwer Academic Publishers: Dordrecht, The Netherlands, 1996; pp. 101–121.

15. Styring, S.; Virgin, I.; Ehrenberg, A.; Andersson, B. Strong light photoinhibition of electron transport in photosystem II. Impairment of the function of the first quinone acceptor, QA. Biochim. Biophys. Acta 1990, 1015, 269. [CrossRef]

16. Aro, E.; Virgin, I.; Andersson, B. Photoinhibition of photosystem II. Inactivation, protein damage and turnover. Biochim. Biophys. Acta 1993, 1143, 113–134. [CrossRef]

17. Baker, N.R.; Bowyer, J.R. Photoinhibition of photosynthesis from molecular mechanisms to the field. In Environmental Plant Biology Series; Davies, W.J., Ed.; Bios Scientific Publishers: Institute of Environmental and Biological Sciences; Bios Scientific Publishers: Institute of Environmental and Biological Sciences, Division of Biological Sciences, University of Lancaster: Oxford, UK, 1994; pp. 1–471.

18. Demmig-Adams, B.; Adams-III, W.W. The role of xanthophyll cycle carotenoids in the protection of photosynthesis. Trends Plant Sci. 1996, 1, 21–26. [CrossRef]

19. Eskling, M.; Emanuelsson, A.; Akerlund, H.-E. Enzymes and mechanisms for violaxanthin-zeaxanthin conversion. In Regulation of Photosynthesis; Aro, E.-M., Anderson, B., Eds.; Kluwer Academic Publishers: Dordrecht, The Netherlands, 2001; Volume 100, pp. 806–816.

20. Li, X.P.; Muller-Moule, P.; Gilmore, A.M.; Niyogi, K.K. PsbS-dependent enhancement of feedback de-excitation protects photosystem II from photoinhibition. Proc. Natl. Acad. Sci. USA 2002, 99, 15222–15227. [CrossRef]

21. Niyogi, K.K.; Li, X.-P.; Rosenberg, V.; Jung, H.-S. Is PsbS the site of non-photochemical quenching in photosynthesis? J. Exp. Bot. 2004, 56, 375–382. [CrossRef]

22. Takizawa, K.; Kanazawa, A.; Cruz, J.A.; Kramer, D.M. In vivo thylakoid proton motive force. Quantitative non-invasive probes show the relative lumen pH-induced regulatory responses of antenna and electron transfer. Biochim. Biophys. Acta 2007, 1767, 1233–1244. [CrossRef]

23. Tikhonov, A.N. The cytochrome b6f complex at the crossroad of photosynthetic electron transport pathways. Plant Physiol. Biochem. 2014, 81, 163–183. [CrossRef]

24. Cruz, J.A.; Avenson, T.J.; Takizawa, K.; Kramer, D.M. The contribution of cyclic electron flux (CEF1) to formation of proton motive force (pmf). In Proceedings of the 13th International Congress of Photosynthesis: Fundamental Aspects to Global Perspectives, Lawrence, KA, USA, 29 August–3 September 2005; pp. 1033–1035.

25. Kanazawa, A.; Kramer, D.M. In vivo modulation of nonphotochemical exciton quenching (NPQ) by regulation of the chloroplast ATP synthase. Proc. Natl. Acad. Sci. USA 2002, 99, 12789–12794. [CrossRef]

26. Avenson, T.; Cruz, J.; Kramer, D. Modulation of CF0-CF1 ATP synthase conductivity and proton motive force (pmf) partitioning regulate light capture. In Proceedings of the 13th International Congress of Photosynthesis: Fundamental Aspects to Global Perspectives, Lawrence, KA, USA, 29 August–3 September 2005; pp. 575–577.

27. Avenson, T.; Cruz, J.A.; Kramer, D. Modulation of energy dependent quenching of excitons (qE) in antenna of higher plants. Proc. Natl. Acad. Sci. USA 2004, 101, 5530–5535. [CrossRef] [PubMed]

28. Cruz, J.A.; Sacksteder, C.A.; Kanazawa, A.; Kramer, D.M. Contribution of Electric Field (∆ψ) to Steady-State Transthylakoid Proton Motive Force (pmf) in Vitro and in Vivo. Control of pmf Parsing into ∆ψ and ∆pH by Ionic Strength. Biochemistry 2001, 40, 1226–1237. [CrossRef] [PubMed]

29. Kramer, D.M.; Cruz, J.A.; Kanazawa, A. Balancing the central roles of the thylakoid proton gradient. Trends Plant Sci. 2003, 8, 27–32. [CrossRef]

30. Zhang, R.; Cruz, J.A.; Kramer, D.M.; Magallanes-Lundback, M.; DellaPenna, D.; Sharkey, T.D. Heat stress reduces the pH component of the transthylakoid proton motive force in light-adapted intact tobacco leaves. Plant Cell Environ. 2009, 32, 1538–1547. [CrossRef] [PubMed]

31. Murchie, E.H.; Niyogi, K.K. Manipulation of photoprotection to improve plant photosynthesis. Plant Physiol. 2011, 155, 86–92. [CrossRef]
32. Li, X.P.; Gilmore, A.M.; Caffarri, S.; Bassi, R.; Golan, T.; Kramer, D.; Niyogi, K.K. Regulation of photosynthetic light harvesting involves intrathylakoid lumen pH sensing by the PsbS protein. *J. Biol. Chem.* **2004**, *279*, 22866–22874. [CrossRef]

33. Scheibe, R. Malate valves to balance cellular energy supply. *Physiol. Plant* **2004**, *120*, 21–26. [CrossRef]

34. Asada, K. The water-water cycle in chloroplasts: Scavenging of active oxygens and dissipation of excess photons. *Annu. Rev. Plant Biol.* **1999**, *50*, 601–639. [CrossRef]

35. Alric, J.; Johnson, X. Alternative electron transport pathways in photosynthesis: A confluence of regulation. *Curr. Opin. Plant Biol.* **2017**, *37*, 78–86. [CrossRef]

36. Strand, D.D.; Kramer, D.M. Distinct Energetics and Regulatory Functions of the Two Major Cyclic Electron Flow Pathways in Chloroplasts; Caister Academic Press: Poole, UK, 2016.

37. Ishikawa, N.; Endo, T.; Sato, F. Electron transport activities of Arabidopsis thaliana mutants with impaired chloroplastic NAD(P)H dehydrogenase. *J. Plant Res.* **2008**, *121*, 521–526. [CrossRef]

38. Munekage, Y.; Hojo, M.; Meurer, J.; Endo, T.; Tasaka, M.; Shikanai, T. PGR5 Is Involved in Cyclic Electron Flow around Photosystem I and Is Essential for Photoprotection in *Arabidopsis*. *Cell* **2002**, *110*, 361–371. [CrossRef]

39. DalCorso, G.; Pesaresi, P.; Masiero, S.; Aseeva, E.; Schünemann, D.; Finazzi, G.; Joliot, P.; Barbato, R.; Leister, D. A Complex Containing PGR1 and PGR5 Is Involved in the Switch between Linear and Cyclic Electron Flow in *Arabidopsis*. *Cell* **2008**, *133*, 273–285. [CrossRef] [PubMed]

40. Hertle, A.P.; Blunder, T.; Wunder, T.; Pesaresi, P.; Pribil, M.; Armbruster, U.; Leister, D. PGR1 Is the Elusive Ferredoxin-Plastoquinone Reductase in Photosynthetic Cyclic Electron Flow. *Mol. Cell* **2013**, *49*, 511–523. [CrossRef] [PubMed]

41. Nandha, B.; Finazzi, G.; Joliot, P.; Hald, S.; Johnson, G.N. The role of PGR5 in the redox poising of photosynthetic electron transport. *Biochim. Biophys. Acta Bioenerg.* **2007**, *1767*, 1252–1259. [CrossRef] [PubMed]

42. Joliot, P.; Johnson, G.N. Regulation of cyclic and linear electron flow in higher plants. *Proc. Natl. Acad. Sci. USA* **2011**, *108*, 13317–13322. [CrossRef] [PubMed]

43. Fisher, N.; Kramer, D.M. Non-photochemical reduction of thylakoid photosynthetic redox carriers in vitro: Relevance to cyclic electron flow around photosystem I? *Biochim. Biophys. Acta Bioenerg.* **2014**, *1837*, 1944–1954. [CrossRef]

44. Mosebach, L.; Heilmann, C.; Mutoh, R.; Gäbelein, P.; Steinbeck, J.; Happe, T.; Ikegami, T.; Hanke, G.; Kurisu, G.; Hippler, M. Association of Ferredoxin:NADP+ oxidoreductase with the photosynthetic apparatus modulates electron transfer in *Chlamydomonas reinhardtii*. *Photosynth. Res.* **2017**, *134*, 291–306. [CrossRef]

45. Burrows, P.A.; Sazanov, L.A.; Svab, Z.; Maliga, P.; Nixon, P.J. Identification of a functional respiratory complex in chloroplasts through analysis of tobacco mutants containing disrupted plastid ndh genes. *EMBO J.* **1998**, *17*, 868–876. [CrossRef] [PubMed]

46. Shikanai, T. Chloroplast NDH: A different enzyme with a structure similar to that of respiratory NADH dehydrogenase. *Biochim. Biophys. Acta Bioenerg.* **2016**, *1857*, 1015–1022. [CrossRef]

47. Strand, D.D.; Fisher, N.; Kramer, D.M. The higher plant plastid NAD(P)H dehydrogenase-like complex (NDH) is a high efficiency proton pump that increases ATP production by cyclic electron flow. *J. Biol. Chem.* **2017**, *292*, 11850–11860. [CrossRef]

48. Laughlin, T.G.; Bayne, A.N.; Trempe, J.-F.; Savage, D.F.; Davies, K.M. Structure of the complex I-like molecule NDH of oxygenic photosynthesis. *Nature* **2019**, *566*, 411–414. [CrossRef] [PubMed]

49. Joliot, P.; Joliot, A. Cyclic electron transfer in plant leaf. *Proc. Natl. Acad. Sci. USA* **2002**, *99*, 10209–10214. [CrossRef] [PubMed]

50. Strobel, D.; Choquet, Y.; Popot, J.-L.; Picot, D. An atypical haem in the cytochrome b6f complex. *Nature* **2003**, *426*, 413–418. [CrossRef] [PubMed]

51. Alric, J.; Pierre, Y.; Picot, D.; Lavernge, J.; Rappaport, F. Spectral and redox characterization of the heme ϵ of the cytochrome b6f complex. *Proc. Natl. Acad. Sci. USA* **2005**, *102*, 15860–15865. [CrossRef]

52. Strand, D.D.; D’Andrea, L.; Bock, R. The plastid NAD(P)H dehydrogenase-like complex: Structure, function and evolutionary dynamics. *Biochim. J.* **2019**, *476*, 2743–2756. [CrossRef]

53. Joliot, P.; Béal, D.; Joliot, A. Cyclic electron flow under saturating excitation of dark-adapted *Arabidopsis* leaves. *Biochim. Biophys. Acta Bioenerg.* **2004**, *1656*, 166–176. [CrossRef]
54. Kohzuma, K.; Cruz, J.A.; Akashi, K.; Hoshiyasu, S.; Munekage, Y.N.; Yokota, A.; Kramer, D.M. The long-term responses of the photosynthetic proton circuit to drought. Plant Cell Environ. 2009, 32, 209–219. [CrossRef]

55. Kubicki, A.; Funk, E.; Westhoff, P.; Steinmüller, K. Differential expression of plastome-encoded ndh genes in mesophyll and bundle-sheath chloroplasts of the C4 plant Sorghum bicolor indicates that the complex I-homologous NAD(P)H-plastoquinone oxidoreductase is involved in cyclic electron transport. Planta 1996, 199, 276–281. [CrossRef]

56. Takabayashi, A.; Kishine, M.; Asada, K.; Endo, T.; Sato, F. Differential use of two cyclic electron flows around photosystem I for driving CO2-concentration mechanism in C4 photosynthesis. Proc. Natl. Acad. Sci. USA 2005, 102, 16898–16903. [CrossRef]

57. Luckner, B.; Kramer, D.M. Regulation of cyclic electron flow in Chlamydomonas reinhardtii under fluctuating carbon availability. Photosynth. Res. 2013, 117, 449–459. [CrossRef]

58. Scheibe, R. Maintaining homeostasis by controlled alternatives for energy distribution in plant cells under changing conditions of supply and demand. Photosynth. Res. 2019, 139, 81–91. [CrossRef] [PubMed]

59. Selinski, J.; Scheibe, R. Malate valves: Old shuttles with new perspectives. Plant Biol. 2019, 21, 21–30. [CrossRef]

60. Fickenscher, K.; Scheibe, R. Limited proteolysis of inactive tetrameric chloroplast NADP-Malate dehydrogenase produces active dimers. Arch. Biochem. Biophys. 1988, 260, 771–779. [CrossRef]

61. Ocheretina, O.; Harnecker, J.; Rother, T.; Schmid, R.; Scheibe, R. Effects of N-terminal truncations upon chloroplast NADP-malate dehydrogenases from pea and spinach. Biochim. Biophys. Acta BBA Protein Struct. Mol. Enzymol. 1993, 1163, 10–16. [CrossRef]

62. Krieger-Liszkay, A.; Feilke, K. The Dual Role of the Plastid Terminal Oxidase PTOX: Between a Protective and a Pro-oxidant Function. Front. Plant Sci. 2016, 6, 1147. [CrossRef] [PubMed]

63. Allahverdiyeva, Y.; Suorsa, M.; Tikkanen, M.; Aro, E.-M. Photoprotection of photosystems in fluctuating light intensities. J. Exp. Bot. 2014, 66, 2427–2436. [CrossRef] [PubMed]

64. Ruuska, S.A.; Badger, M.R.; Andrews, T.J.; von Caemmerer, S. Photosynthetic electron sinks in transgenic tobacco with reduced amounts of Rubisco: Little evidence for significant Mehler reaction. J. Exp. Bot. 2000, 51, 357–368. [CrossRef]

65. Heber, U. Irrungen, Wirrungen? The Mehler reaction in relation to cyclic electron transport in C3 plants. Photosynth. Res. 2002, 73, 223–231. [CrossRef]

66. Driever, S.M.; Baker, N.R. The water–water cycle in leaves is not a major alternative electron sink for dissipation of excess excitation energy when CO2 assimilation is restricted. Plant Cell Environ. 2011, 34, 837–846. [CrossRef]

67. Noctor, G.; Foyer, C. Review article. A re-evaluation of the ATP:NADPH budget during C3 photosynthesis: A contribution from nitrate assimilation and its associated respiratory activity? J. Exp. Bot. 1998, 49, 1895–1908. [CrossRef]

68. Edwards, G.E.; Walker, D.A. C3, C4: Mechanisms, and Cellular and Environmental Regulation, of Photosynthesis; Blackwell Scientific Publications: Oxford, UK, 1983.

69. Walker, B.J.; VanLoocke, A.; Bernacchi, C.J.; Ort, D.R. The costs of photorespiration to food production now and in the future. Annu. Rev. Plant Biol. 2016, 67, 107–129. [CrossRef] [PubMed]

70. Farquhar, G.D.; von Caemmerer, S.; Berry, J.A. A biochemical model of photosynthetic CO2 assimilation in leaves of C3 species. Planta 1980, 149, 78–90. [CrossRef] [PubMed]

71. von Caemmerer, S.; Farquhar, G.D. Some relationships between the biochemistry of photosynthesis and the gas exchange of leaves. Planta 1981, 153, 376–387. [CrossRef] [PubMed]

72. Sharkey, T.D. Estimating the rate of photorespiration in leaves. Physiol. Plant. 1988, 73, 147–152. [CrossRef]

73. Walker, B.J.; Strand, D.D.; Kramer, D.M.; Cousins, A.B. The response of cyclic electron flow around photosystem I to changes in photorespiration and nitrate assimilation. Plant Phys. 2014, 165, 453–462. [CrossRef]

74. Mascles-Daubresse, C.; Daniel-Vedele, F.; Dechorgnat, J.; Chardon, F.; Gaufichon, L.; Suzuki, A. Nitrogen uptake, assimilation and remobilization in plants: Challenges for sustainable and productive agriculture. Ann. Bot. 2010, 105, 1141–1157. [CrossRef]

75. Reed, A.J.; Canvin, D.T. Light and Dark Controls of Nitrate Reduction in Wheat (Triticum aestivum L.) Protoplasts. Plant Phys. 1982, 69, 508–513. [CrossRef]
76. Selinski, J.; Scheibe, R. Lack of malate valve capacities lead to improved N-assimilation and growth in transgenic A. thaliana plants. *Plant Signal. Behav.* 2014, 9, e29057. [CrossRef]

77. Rachmilevitch, S.; Cousins, A.B.; Bloom, A.J. Nitrate assimilation in plant shoots depends on photosynthesis. *Proc. Natl. Acad. Sci. USA* 2004, 101, 11506–11510. [CrossRef]

78. Bloom, A.J.; Burger, M.; Asensio, J.S.R.; Cousins, A.B. Carbon Dioxide Enrichment Inhibits Nitrate Assimilation in Wheat and Arabidopsis. *Science* 2010, 328, 899–903. [CrossRef]

79. Ohlrogge, J.; Browse, J. Lipid biosynthesis. *Plant Cell* 1995, 7, 957–970. [CrossRef] [PubMed]

80. Troncoso-Ponce, M.A.; Cao, X.; Yang, Z.; Ohlrogge, J.B. Lipid turnover during senescence. *Plant Sci.* 2013, 205–206, 13–19. [CrossRef] [PubMed]

81. Bao, X.; Focke, M.; Pollard, M.; Ohlrogge, J. Understanding in vivo carbon precursor supply for fatty acid synthesis in leaf tissue. *Plant J.* 2000, 22, 39–50. [CrossRef] [PubMed]

82. Pollard, M.; Ohlrogge, J. Testing Models of Fatty Acid Transfer and Lipid Synthesis in Spinach Leaf Using in Vivo Oxygen-18 Labeling. *Plant Physiol.* 1999, 121, 1217–1226. [CrossRef]

83. Bonaventure, G.; Bao, X.; Ohlrogge, J.; Pollard, M. Metabolic responses to the reduction in palmitate caused by disruption of the FATB gene in Arabidopsis. *Plant Physiol.* 2004, 135, 1269–1279. [CrossRef]

84. Yang, Z.; Ohlrogge, J.B. Turnover of Fatty Acids during Natural Senescence of Arabidopsis, *Brachypodium*, and Switchgrass and in Arabidopsis β-Oxidation Mutants. *Plant Physiol.* 2009, 150, 191–1989. [CrossRef]

85. Geigenberger, P.; Kolbe, A.; Tiessen, A. Redox regulation of carbon storage and partitioning in response to light and sugars. *J. Exp. Bot.* 2005, 56, 1469–1479. [CrossRef]

86. Shastri, A.A.; Morgan, J.A. Flux Balance Analysis of Photoautotrophic Metabolism. *Biotechnol. Prog.* 2005, 21, 1617–1626. [CrossRef]

87. Von Ballmoos, C.; Cook, G.M.; Dimroth, P. Unique Rotary ATP Synthase and Its Biological Diversity. *Annu. Rev. Biophys.* 2008, 37, 43–64. [CrossRef]

88. Von Ballmoos, C.; Wiedemann, A.; Dimroth, P. Essentials for ATP Synthesis by F1F0 ATP Synthases. *Annu. Rev. Biochem.* 2009, 78, 649–672. [CrossRef]

89. Junge, W.; Sielaff, H.; Engelbrecht, S. Torque generation and elastic power transmission in the rotary FOF1-ATPase. *Nature* 2009, 459, 364–370. [CrossRef] [PubMed]

90. Ferguson, S.J. ATP synthase: From sequence to ring size to the P/O ratio. *Proc. Natl. Acad. Sci. USA* 2010, 107, 16755–16756. [CrossRef] [PubMed]

91. Silverstein, T. The mitochondrial phosphate-to-oxygen ratio is not an integer. *Biochem. Mol. Biol. Educ.* 2005, 33, 416–417. [CrossRef] [PubMed]

92. Watt, I.N.; Montgomery, M.G.; Runswick, M.J.; Leslie, A.G.W.; Walker, J.E. Bioenergetic cost of making an adenosine triphosphate molecule in animal mitochondria. *Proc. Natl. Acad. Sci. USA* 2010, 107, 16823–16827. [CrossRef] [PubMed]

93. Ferguson, S.J.; Sorgato, M.C. Proton Electrochemical Gradients and Energy-Transduction Processes. *Annu. Rev. Biochem.* 1982, 51, 185–217. [CrossRef]

94. Nicholls, D.G. Mitochondrial ion circuits. * Essays Biochem.* 2010, 47, 25–35.

95. Bhatla, S.C.; Lal, M.A. ATP Synthesis. In *Plant Physiology, Development and Metabolism*; Springer: Singapore, 2019; pp. 115–172.

96. Haupt-Herting, S.; Klug, K.; Fock, H.P. A new approach to measure gross CO₂ fluxes in leaves. Gross CO₂ assimilation, photorespiration, and mitochondrial respiration in the light in tomato under drought stress. *Plant Phys.* 2001, 126, 388–396. [CrossRef]

97. Villar, R.; Held, A.A.; Merino, J. Comparison of methods to estimate dark respiration in the light in leaves of two woody species. *Plant Phys.* 1994, 105, 167–172. [CrossRef]

98. Kromer, S. Respiration During Photosynthesis. *Annu Rev. Plant Physiol Plant Mol. Biol* 1995, 46, 45–70. [CrossRef]

99. Villar, R.; Held, A.A.; Merino, J. Comparison of methods to estimate dark respiration in the light in leaves of two woody species. *Plant Phys.* 1994, 105, 167–172. [CrossRef]

100. Kunji, E.R.S.; Aleksandrova, A.; King, M.S.; Majd, H.; Ashton, V.L.; Cerson, E.; Springett, R.; Kibalchenko, M.; Tavoulari, S.; Crichton, P.G.; et al. The transport mechanism of the mitochondrial ADP/ATP carrier. *Biochim. Biophys. Acta BBAMol. Cell Res.* 2016, 1863, 2379–2393. [CrossRef]

101. Palmieri, F.; Monné, M. Discoveries, metabolic roles and diseases of mitochondrial carriers: A review. *Biochim. Biophys. Acta BBAMol. Cell Res.* 2016, 1863, 2362–2378. [CrossRef]
Plants 2020, 9, 301

102. Heldt, H.W. Adenine nucleotide translocation in spinach chloroplasts. FEBS Lett. 1969, 5, 11–14. [CrossRef]
103. Schunemann, D.; Borchert, S.; Flugge, U.I.; Heldt, H.W. ADP/ATP Translocator from Pea Root Plastids (Comparison with Translocators from Spinach Chloroplasts and Pea Leaf Mitochondria). Plant Phys. 1993, 103, 131–137. [CrossRef] [PubMed]
104. Huang, W.; Yang, Y.-J.; Hu, H.; Zhang, S.-B. Different roles of cyclic electron flow around photosystem I under sub-saturating and saturating light intensities in tobacco leaves. Front. Plant Sci. 2015, 6. [CrossRef] [PubMed]
105. Cheung, C.Y.M.; Ratcliffe, R.G.; Sweetlove, L.J. A Method of Accounting for Enzyme Costs in Flux Balance Analysis Reveals Alternative Pathways and Metabolite Stores in an Illuminated Arabidopsis Leaf. Plant Phys. 2015, 169, 1671–1682. [CrossRef] [PubMed]
106. Shameer, S.; Ratcliffe, R.G.; Sweetlove, L.J. Leaf Energy Balance Requires Mitochondrial Respiration and Export of Chloroplast NADPH in the Light. Plant Phys. 2019, 180, 1947–1961. [CrossRef] [PubMed]
107. Munekage, Y.; Hashimoto, M.; Miyake, C.; Tomizawa, K.-I.; Endo, T.; Tasaka, M.; Shikanai, T. Cyclic electron flow around photosystem I is essential for photosynthesis. Nature 2004, 429, 579–582. [CrossRef]
108. Strand, D.D.; Fisher, N.; Davis, G.A.; Kramer, D.M. Redox regulation of the antimycin A sensitive pathway of cyclic electron flow around photosystem I in higher plant thylakoids. Biochim. Biophys. Acta Bioenerg. 2016, 1857, 1–6. [CrossRef]
109. Michelet, L.; Zaffagnini, M.; Morisse, S.; Sparla, F.; Pérez-Pérez, M.E.; Francia, F.; Danon, A.; Marchand, C.; Fermani, S.; Trost, P.; et al. Redox regulation of the Calvin–Benson cycle: Something old, something new. Front. Plant Sci. 2013, 4. [CrossRef]
110. Bloom, A.J.; Caldwell, R.M.; Finazzo, J.; Warner, R.L.; Weissbart, J. Oxygen and Carbon Dioxide Fluxes from Barley Shoots Depend on Nitrate Assimilation. Plant Physiol. 1989, 91, 352–356. [CrossRef]
111. Bloom, A.J.; Smart, D.R.; Nguyen, D.T.; Searles, P.S. Nitrogen assimilation and growth of wheat under elevated carbon dioxide. Proc. Natl. Acad. Sci. USA 2002, 99, 1730–1735. [CrossRef]
112. Bloom, A.J.; Burger, M.; Kimball, B.A.; Pinter, P.J. Nitrate assimilation is inhibited by elevated CO₂ in field-grown wheat. Nat. Clim. Chang. 2014, 4, 477–480. [CrossRef]
113. Hebbelmann, I.; Selinski, J.; Wehmeyer, C.; Goss, T.; Voss, I.; Mulo, P.; Kangasjärvi, S.; Arro, E.-M.; Oelze, M.-L.; Dietz, K.-J.; et al. Multiple strategies to prevent oxidative stress in Arabidopsis plants lacking the malate valve enzyme NADP-malate dehydrogenase. J. Exp. Bot. 2011, 63, 1445–1459. [CrossRef] [PubMed]
114. Buchanan, B.B. The Ferredoxin-Thioredoxin System: A Key Element in the Regulatory Function of Light in Photosynthesis. Bioscience 1984, 34, 378–383. [CrossRef] [PubMed]
115. Buchanan, B.B.; Balmer, Y. REDOX REGULATION: A Broadening Horizon. Annu. Rev. Plant Biol. 2005, 56, 187–220. [CrossRef] [PubMed]
116. Scheibe, R. Thioredoxinin in pea chloroplasts: Concentration and redox state under light and dark conditions. FEBS Lett. 1981, 133, 301–304. [CrossRef]
117. Scheibe, R.; Stitt, M. Comparison of NADP-malate dehydrogenase activation, QA reduction and O₂ evolution in spinach leaves. Plant Physiol. Biochem. 1988, 26, 473–481. [CrossRef]
118. Joliot, P.; Joliot, A. Cyclic electron flow in C₃ plants. Biochim. Biophys. Acta Bioenerg. 2006, 1757, 362–368. [CrossRef]
119. Becker, B.; Holtgreve, S.; Jung, S.; Wunrau, C.; Kandlbinder, A.; Baier, M.; Dietz, K.-J.; Backhausen, J.E.; Scheibe, R. Influence of the photoperiod on redox regulation and stress responses in Arabidopsis thaliana L. (Heynh.) plants under long- and short-day conditions. Planta 2006, 224, 380–393. [CrossRef]
120. Bauwe, H.; Hagemann, M.; Fernie, A.R. Photorespiration: Players, partners and origin. Trends Plant Sci. 2010, 15, 330–336. [CrossRef]
121. Tomaz, T.; Bagard, M.; Pracharoenwattana, I.; Linden, P.; Lee, C.P.; Carroll, A.J.; Stroher, E.; Smith, S.M.; Gardestrom, P.; Millar, A.H. Mitochondrial malate dehydrogenase lowers leaf respiration and alters photorespiration and plant growth in Arabidopsis. Plant Phys. 2010, 154, 1143–1157. [CrossRef] [PubMed]
122. Lindén, P.; Keech, O.; Stenlund, H.; Gardeström, P.; Moritz, T. Reduced mitochondrial malate dehydrogenase activity has a strong effect on photorespiratory metabolism as revealed by 13C labelling. J. Exp. Bot. 2016, 67, 3123–3135. [CrossRef] [PubMed]
123. Cousins, A.B.; Pracharoenwattana, I.; Zhou, W.; Smith, S.M.; Badger, M.R. Peroxisomal malate dehydrogenase is not essential for photorespiration in Arabidopsis but its absence causes an increase in the stoichiometry of photorespiratory CO₂ release. *Plant Phys.** 2008, 148, 786–795. [CrossRef]

124. Cousins, A.B.; Walker, B.J.; Pracharoenwattana, I.; Smith, S.M.; Badger, M.R. Peroxisomal hydroxyypyruvate reductase is not essential for photorespiration in arabidopsis but its absence causes an increase in the stoichiometry of photorespiratory CO₂ release. *Photosynth. Res.** 2011, 108, 91–100. [CrossRef]

125. Anderson, J.M.; Chow, W.S.; Park, Y.-I. The grand design of photosynthesis: Acclimation of the photosynthetic apparatus to environmental cues. *Photosynth. Res.** 1995, 46, 129–139. [CrossRef] [PubMed]

126. Fujita, Y. A study on the dynamic features of photosystem stoichiometry: Accomplishments and problems for future studies. *Photosynth. Res.** 1997, 53, 83–93. [CrossRef]

127. Melis, A.; Murakami, A.; Nemson, J.A.; Aizawa, K.; Ohki, K.; Fujita, Y. Chromatic regulation inChlamydomonas reinhardtii alters photosystem stoichiometry and improves the quantum efficiency of photosynthesis. *Photosynth. Res.** 1996, 47, 253–265. [CrossRef] [PubMed]

128. Pfannschmidt, T.; Allen, J.F.; Oelmüller, R. Principles of redox control in photosynthesis gene expression. *Physiol. Plant.** 2001, 112, 1–9. [CrossRef]

129. Dietzel, L.; Bräutigam, K.; Pfannschmidt, T. Photosynthetic acclimation: State transitions and adjustment of photosystem stoichiometry—Functional relationships between short-term and long-term light quality acclimation in plants. *FEBS J.** 2008, 275, 1080–1088. [CrossRef]

130. Allen, J.F. Protein phosphorylation in regulation of photosynthesis. *Biochim. Biophys. Acta Bioenerg.** 1992, 1098, 275–335. [CrossRef]

131. Gupta, P.; Duplessis, S.; White, H.; Karnosky, D.F.; Martin, F.; Podila, G.K. Gene expression patterns of Melis. *Plant Cell Environ.* 2005, 275, 1–9. [CrossRef]

132. Ainsworth, E.A.; Rogers, A.; Vodkin, L.O.; Walter, A.; Schurr, U. The E. *Photosynth. Res.* 1996, 47, 253–265. [CrossRef] [PubMed]

133. Von Caemmerer, S. Steady-state models of photosynthesis. *Photosynth. Res.* 1995, 46, 129–139. [CrossRef] [PubMed]

134. Von Caemmerer, S. Biochemical Models of Leaf Photosynthesis; CSIRO: Collingwood, Australia, 2000; Volume 2.

135. Allen, J.F. Protein phosphorylation in regulation of photosynthesis. *Biochim. Biophys. Acta Bioenerg.* 1992, 1098, 275–335. [CrossRef]

136. Laisk, A. Kinetics of photosynthesis and photorespiration in C₂ plants. *Nauka Mosc.* 1977. (In Russian)

137. Laisk, A.; Loreto, F. Determining photosynthetic parameters from leaf CO₂ exchange and chlorophyll fluorescence. *Plant Physiol.* 1996, 110, 903–912. [CrossRef] [PubMed]

138. Bernacchi, C.J.; Portis, A.R.; Nakano, H.; von Caemmerer, S.; Long, S.P. Temperature response of mesophyll conductance. Implications for the determination of Rubisco enzyme kinetics and for limitations to photosynthesis in vivo. *Plant Phys.* 2002, 130, 1992–1998. [CrossRef]

139. Tazoe, Y.; von Caemmerer, S.; Estavillo, G.M.; Evans, J.R. Using tunable diode laser spectroscopy to measure carbon isotope discrimination and mesophyll conductance to CO₂ diffusion dynamically at different CO₂ concentrations. *Plant Cell Environ.* 2011, 34, 580–591. [CrossRef]

140. Gilbert, M.E.; Pou, A.; Zwieniecki, M.A.; Holbrook, N.M. On measuring the response of mesophyll conductance to carbon dioxide with the variable J method. *J. Exp. Bot.* 2012, 63, 413–425. [CrossRef]

141. Tholen, D.; Ethier, G.; Genty, B.; Pepin, S.; Zhu, X.-G. Variable mesophyll conductance revisited: Theoretical background and experimental implications. *Plant Cell Environ.* 2012, 35, 2087–2103. [CrossRef]

142. Evans, J.R.; von Caemmerer, S. Temperature response of carbon isotope discrimination and mesophyll conductance in tobacco. *Plant Cell Environ.* 2013, 36, 745–756. [CrossRef]

143. Jahan, E.; Amthor, J.S.; Farquhar, G.D.; Trethewan, R.; Barbour, M.M. Variation in mesophyll conductance among Australian wheat genotypes. *Funct. Plant Biol.* 2014, 41, 568–580. [CrossRef]
144. von Caemmerer, S.; Evans, J.R. Temperature responses of mesophyll conductance differ greatly between species. *Plant Cell Environ.* **2014**, *38*, 629–637. [CrossRef] [PubMed]

145. Loreto, F.; Harley, P.C.; Di Marco, G.; Sharkey, T.D. Estimation of mesophyll conductance to CO$_2$ flux by three different methods. *Plant Phys.* **1992**, *98*, 1437–1443. [CrossRef] [PubMed]

© 2020 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).