Far-Red Chlorophyll Fluorescence Radiance Tracks Photosynthetic Carbon Assimilation Efficiency of Dark Reactions

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Abstract: Solar-induced chlorophyll fluorescence (SIF) observations from space have shown close relationships with terrestrial photosynthesis rates. SIF originates from the light reactions of photosynthesis, whereas carbon fixation takes place during the dark reactions of photosynthesis. Questions remain regarding whether SIF is able to track changes in the efficiency of the dark reactions in photosynthesis. Using concurrent measurements of leaf-scale gas exchange, pulse amplitude-modulated (PAM) fluorescence, and fluorescence spectral radiances, we found that both far-red fluorescence radiances and PAM fluorescence yields responded rapidly to changes in photosynthetic carbon assimilation due to changes in environmental factors or induced stomatal closure under constant light conditions. Uncertainties in outgoing and incoming irradiance mismatch for SIF measurements may very likely obscure the contributions of the dark reactions, thereby causing the inconsistent findings previously reported, which were no change in far-red SIF and PAM fluorescence yields after clear reductions in the photosynthetic carbon assimilation efficiency of dark reactions. Our results confirm that high-quality SIF measurements have the potential to provide insights into the dark reactions of photosynthesis. This study is particularly relevant for better interpreting satellite SIF observations that are obtained under roughly constant overpass times and relatively stable light intensities.

Keywords: solar-induced chlorophyll fluorescence; gross primary productivity; far-red chlorophyll fluorescence; dark reactions; CO₂ concentration; air temperature; stomatal closure

1. Introduction

Nearly all lives on Earth ultimately depend on energy from the sun. Photosynthesis is the only process that can convert light energy to chemical energy [1]. Photosynthesis consists of two sets of reactions: light reactions and dark reactions (the carbon reactions). In the light reactions, the photon energy absorbed by photosystem II (PSII) is consumed in three different pathways: photochemistry, fluorescence, and heat loss (nonphotochemical quenching, NPQ) [2,3]. In the photochemistry pathway, the absorbed light energy is used to remove an electron from H₂O, generating linear electron transport from PSII to photosystem I (PSI) [4]. During this electron transport process, absorbed light energy is converted into chemical energy, producing NADPH (the reduced form of nicotine adenine dinucleotide phosphoric acid) and ATP (adenosine triphosphate) [5]. Meanwhile, about 1–2% of absorbed light energy is re-emitted as chlorophyll a fluorescence (ChlF) [6,7].
The probability of partitioning an absorbed photon in a given pathway is dependent on the amount of absorbed light and the efficiency of plant photochemical processes (e.g., the photochemical quantum yield of PSII ($\Phi_{\text{PSII}}$) and the fluorescence quantum yield ($\Phi_{\text{ChlF}}$)) [7]. In the dark reactions, the energy stored in ATP and NADPH is then consumed to assimilate CO$_2$ and produce carbohydrates, primarily glucose [8]. The processes in the dark reactions do not directly require light but are strongly regulated by temperature, water, nutrients, and CO$_2$ from the environment [8].

Advances in observing solar-induced chlorophyll fluorescence (SIF) from space [9,10] have documented promise for improved mapping of global terrestrial gross primary productivity (GPP) [11–21]. Numerous studies have already reported that SIF and GPP are well correlated for a variety of terrestrial ecosystems. For example, Guanter et al. [22] showed that space-borne SIF observations exhibited good performance in estimating crop photosynthesis across the US Corn Belt region. Yang et al. [23] showed that ground-based SIF measurements exhibit a strong linear relationship with GPP at a deciduous forest site. It should be noted that fluorescence emanates from the light reactions, whereas the assimilation of CO$_2$ is accomplished in the dark reactions of photosynthesis [7]. The difference in light and dark reactions naturally raises an important question: does ChlF respond to the efficiency of the dark reactions in photosynthesis? If the answer is no, it might suggest that the strong SIF–GPP correlations found in many previous studies are only because they have a shared driver, namely, absorbed photosynthetically active radiation (APAR). If the answer is yes, it indicates that SIF does provide direct insight into the processes occurring in the dark reactions, including gas exchange and carbon assimilation. This has important implications for better understanding the SIF–GPP relationship when APAR is not a major limiting factor. For instance, it can inform our interpretation of satellite SIF observations commonly acquired around noon (high light) when the dark reactions of photosynthesis on top of the canopy tend to become more prominent. This is also particularly relevant in the context of rising temperatures and atmospheric CO$_2$, as more climate extremes are likely to occur, but the variability of incident PAR will be relatively small [24].

Previous studies on this question have provided inconsistent results. Marrs et al. [25] found a significant decrease in photosynthetic CO$_2$ assimilation as stomata closed, but no clear trend in fluorescence measurements, suggesting that SIF did not directly track carbon assimilation. Helm et al. [26] also reported that water stress caused a clear and fast decline in both net photosynthesis and stomatal conductance but had a smaller effect on SIF emission. They concluded that SIF may not track changes in photosynthesis due to stomatal closure as long as the light reactions of photosynthesis remained largely unchanged. However, Gu et al. [27] observed saturation of, or even a decrease in, canopy SIF under high irradiance. The saturation of canopy SIF was probably due to their bi-hemispheric instrument setup and the sensor–canopy geometry, but it also implied that SIF might respond to changes in the dark reactions because the SIF yields decreased at a higher light intensity at the top of the canopy [28]. This discrepancy may be partially due to the designs of SIF-observing systems and protocols for measurements [29]. Compared with SIF variability driven by incident light, the magnitude of changes in SIF in response to changes in the dark reactions is so subtle that it may be unobservable or uninterpretable for some measuring systems [30]. In order to quantify possible changes in ChlF due to changes in the dark reactions, all of the involved processes in both the light and dark reactions of photosynthesis should be carefully observed. Specifically, one should simultaneously obtain not only high-quality measurements of passive ChlF radiance and CO$_2$ assimilation but also fluorescence parameters deriving from active pulse amplitude-modulated (PAM) ChlF yields under controlled conditions.

In this study, we developed a leaf-scale measurement system by integrating active (PAM fluorometer) and passive (spectrometer) instruments into a portable gas exchange system. We then used the system to carry out two controlled leaf-scale experiments under constant light conditions to answer the following questions: (1) How do measurements
of passive and active ChlF, the derived photosynthetic parameters, and gas exchange measurements respond to changing CO₂ concentrations and leaf temperatures? (2) How do these measurements/parameters vary after inducing stomatal closure? We show how leaf-scale energy partitioning changes due to changes in the dark reactions of photosynthesis and relate it to previous studies reporting seemingly inconsistent results. We also discuss the key requirements needed for an SIF observing system to accurately track changes in the dark reactions of photosynthesis under natural conditions.

2. Materials and Methods

2.1. Leaf-Scale Measurement System Setup

We developed a leaf-scale measurement system to concurrently measure CO₂ assimilation, active ChlF yields, and passive ChlF radiances from both adaxial and abaxial leaf sides, which was similar to the instrument in Magney et al. [31]. A gas exchange system was modified to integrate one external LED light source, one PAM fluorometer, and four spectrometers (Figure 1). The main components included: (1) an LI-6800 portable gas exchange system with a clear-top leaf chamber (LI-COR Biosciences, Lincoln, NE, USA), (2) a DUAL-PAM-100 portable fluorometer (Heinz Walz GmbH, Effeltrich, Germany) measuring active ChlF yields, (3) two QEpro spectrometers (Ocean Optics, Dunedin, FL, USA; 630–860 nm, 0.24 nm spectral resolution) measuring backward and forward ChlF emissions in the 640–850 nm region, and (4) a white LED fiber optic O-ring light (S5000, Nanjing Hecho Technology Co., Ltd., Nanjing, China) with a 625 nm short-pass filter (Edmunds Optics, Barrington, NJ, USA).

![Figure 1. Photograph of the practical synchronous measurement system. The picture demonstrates a custom-made leaf chamber system for concurrent fluorescence measurements on a leaf sample under various environmental factors. It shows the main components in this system, including an LED light source with a short-pass filter, a PAM fluorometer, an LI-6800 gas exchange system, and four spectrometers with two bundled fibers.](image)

The top of the leaf chamber was equipped with two vertical airtight slots (Figure 2). One was installed on the top of the leaf chamber, fitting the bundled fiber optic for two spectrometers (backward spectral radiance) and housing the fiber optic of the PAM fluorometer; the second slot was on the bottom of the leaf chamber to fit another bundled fiber optic for the two spectrometers (forward spectral radiance). The top and bottom fibers

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CO₂ concentrations were manually adjusted as follows (in ppm): 50, 100, 200, 300, 400, 900, 1200; for the temperature curve, the air CO₂ concentration was set at 400 ppm, and leaf temperatures were manually adjusted as follows (in °C): 15, 20, 25, 30, 35, 500, 700, 900, 1200; for the temperature curve, the air CO₂ concentration was set at 400 ppm, and leaf temperatures were manually adjusted as follows (in °C): 15, 20, 25, 30, 35, 40.

2.2. Experimental Design

The experiment was conducted at the State Key Laboratory of Soil Erosion and Dryland Farming on the Loess Plateau, Northwest A&F University, Yangling, China, from October to December in 2020. Both maize (*Zea mays* L.) and soybean (*Glycine max* Linn. Merr.) were grown in a climate chamber under the following conditions: 12 h light/12 h dark, 800 μmol m⁻² s⁻¹ irradiance, 25 °C, and 50% relative humidity.

2.2.1. Synchronous Measurements of CO₂ and Temperature Curve

Measurements were collected on attached and healthy leaves of maize and soybean. After a >1 h dark adaptation period, a saturation flash from the PAM fluorometer was used to determine minimal fluorescence (Fₐ) and maximal fluorescence in the dark (Fₘ) of dark-adapted leaves. One minute later, the dark respiration (Rₐ, μmol m⁻² s⁻¹) at the leaf temperature of 25 °C was recorded by the gas exchange system. Next, the leaves were equilibrated inside the synchronous chamber at a CO₂ concentration of 400 μmol mol⁻¹ and under an illumination of 800 μmol m⁻² s⁻¹ (soybean) and 1200 μmol m⁻² s⁻¹ (maize) for at least 30 min to reach a stable CO₂ assimilation. These two high light intensities were kept constant because the impact of the dark reaction aspects of photosynthesis becomes more marked at high light. For both species, the temperature and the CO₂ response curves were measured for each leaf and repeated for three individual leaves. During the measurements, the gas exchange system was set to a constant flow rate of 500 μmol s⁻¹ and a relative humidity of 50%. For the CO₂ curve, the leaf temperature was set at 25 °C, and air CO₂ concentrations were manually adjusted as follows (in ppm): 50, 100, 200, 300, 400, 500, 700, 900, 1200; for the temperature curve, the air CO₂ concentration was set at 400 ppm, and leaf temperatures were manually adjusted as follows (in °C): 15, 20, 25, 30, 35, 40.

Through the procedure of determining CO₂ and temperature response curves, spectral measurements (reflected radiance, transmitted radiance, backward–forward fluorescence spectral radiant energy flux) were constantly recorded at 1 s intervals. In this study, we focused on the total fluorescence radiant energy flux emitted from both abaxial and adaxial surfaces of a leaf at one oxygen absorption line frequently used in SIF-related applications, namely, 760 nm (*T*ChIF₇₆₀, mW m⁻² nm⁻¹ sr⁻¹). We also estimated the fluorescence quantum yield at 760 nm (ΦChIF₇₆₀) from leaf reflectance (R) and transmittance (T), which is described in Text S1. Note that *T*ChIF₇₆₀ comes from both upper and deeper leaf layers,
and thus it is also affected by the inner leaf structure and leaf pigment content [31]. To better interpret passive ChlF radiance, steady-state fluorescence emission (\(F_s\)) induced by the measuring beam of the PAM fluorometer was also included in this study. Compared with passive ChlF, \(F_s\) mainly comes from the upper leaf surface such that it is less likely to be reabsorbed/scattered. Maximal fluorescence emissions in the dark-adapted and light-adapted states (\(F_m\) and \(F'_m\)) from PAM were also collected for 45 s and were used to calculate the required photosynthetic parameters, including NPQ and the photochemical quantum yield of PSII (\(\Phi_{PSII}\)), which are expressed as [32,33]

\[
\text{NPQ} = \frac{F_m - F'_m}{F'_m} \quad (1)
\]

\[
\text{\(\Phi_{PSII}\)} = \frac{F'_m - F_s}{F'_m} \quad (2)
\]

The net \(\text{CO}_2\) assimilation rate (\(A_{\text{net}}, \mu\text{mol m}^{-2}\text{s}^{-1}\)) and stomatal conductance to water vapor (\(g_{sw}, \text{mol m}^{-2}\text{s}^{-1}\)) provided by the gas exchange system were automatically stored every 5 s. Leaf-scale gross \(\text{CO}_2\) assimilation (\(A_G, \mu\text{mol m}^{-2}\text{s}^{-1}\)) was obtained from gas exchange measurements:

\[
A_G = A_{\text{net}} + R_d \quad (3)
\]

where \(R_d\) is dark respiration (\(\mu\text{mol m}^{-2}\text{s}^{-1}\)) measured in a period of darkness. All measured variables and related equipment are listed in Table 1. For the temperature response curves, leaf respiration was estimated by using the global polynomial respiration-temperature model [34]. To ensure that the gas exchange and fluorescence conditions were stable, measurements for \(\text{CO}_2\) and temperature curves were performed after waiting at least 5 min between \(\text{CO}_2\) changes, and after waiting at least 10 min between temperature changes, respectively. We then extracted spectral radiance immediately before the saturation pulse as the steady-state spectral measurements.

### Table 1. Measured variables and related equipment used in this study.

| Measured Variables | Definitions | Related Equipment |
|--------------------|-------------|-------------------|
| \(A_{\text{net}}\)  | leaf net \(\text{CO}_2\) assimilation (\(\mu\text{mol m}^{-2}\text{s}^{-1}\)) | LI-6800 portable gas exchange system |
| \(R_d\)            | leaf dark respiration (\(\mu\text{mol m}^{-2}\text{s}^{-1}\)) | LI-6800 portable gas exchange system |
| \(A_G\)            | leaf-scale gross photosynthetic \(\text{CO}_2\) assimilation estimated from the gas exchange system (\(\mu\text{mol m}^{-2}\text{s}^{-1}\)) | LI-6800 portable gas exchange system |
| \(g_{sw}\)         | stomatal conductance to water vapor (\(\text{mol m}^{-2}\text{s}^{-1}\)) | LI-6800 portable gas exchange system |
| \(F_m\)            | maximum fluorescence yield induced by a saturation pulse in the dark-adapted state | DUAL-PAM-100 fluorometer |
| \(F'_m\)           | maximum fluorescence yield induced by a saturation pulse in the light-adapted state | DUAL-PAM-100 fluorometer |
| \(F_s\)            | steady-state fluorescence yield from PAM | DUAL-PAM-100 fluorometer |
| \(\Phi_{PSII}\)    | photochemical quantum yield of PSII | DUAL-PAM-100 fluorometer |
| NPQ                | nonphotochemical quenching | DUAL-PAM-100 fluorometer |
| TChlF\(_{760}\)    | the total chlorophyll fluorescence radiance emitted from both abaxial and adaxial surfaces of a leaf at 760 nm (\(\text{mW m}^{-2}\text{nm}^{-1}\text{sr}^{-1}\)) | QEpro spectrometer |
2.2.2. Synchronous Measurements after ABA Treatment

To determine the response of ChlF emission to inducing stomatal closure (i.e., the decrease in the dark reactions), we also obtained measurements of gas exchange, spectral ChlF, reflectance, transmittance, and PAM fluorescence before and after adding the plant hormone abscisic acid (ABA, Sigma-Aldrich, St. Louis, MO, USA; purity ≥ 99%). To do so, we followed the protocol proposed by Ceciliato et al. [35] to measure the time-resolved stomatal conductance response to ABA in intact leaves. Because this method is only suitable for leaves with long petioles (leaf stalks), we only included soybean leaves in this experiment. Petioles of soybean leaves were cut and immediately submerged in deionized water. Petioles were then cut a second time under water and transferred to a microcentrifuge tube with the petiole tips submerged in water [35]. Before the beginning of the experiments, leaves attached with a microcentrifuge tube were placed inside the modified leaf chamber and equilibrated for at least 45 min to reach a stable CO$_2$ assimilation and stomatal conductance. Concurrent measurements were recorded for at least 30 min at a steady-state level prior to the addition of ABA. Hereafter, ABA was added to the deionized water in the microcentrifuge tube, reaching a concentration of 10 µM. Concurrent measurements were then recorded for at least 30 min. All of the measurements for pre-treatment and post-treatment of ABA were conducted under the following conditions: the LED light source was set at 800 µmol m$^{-2}$ s$^{-1}$, leaf temperature was 25 °C, relative humidity in the chamber was 50%, and a constant flow rate of 500 µmol s$^{-1}$ was maintained.

3. Results
3.1. Effects of CO$_2$ Concentration on the Variations in Carbon Assimilation and Fluorescence Emission

For the CO$_2$ curves with constant irradiance, the proxies of photosynthetic capacity including $A_G$ and $\Phi_{PSII}$ for soybean (Figure 3a,e) and maize (Figure 3c,g) increased with increasing CO$_2$, until plateaus occurred at a CO$_2$ concentration of about 500 ppm. However, $g_{sw}$ curves declined continuously to nearly 0 in response to the increased CO$_2$ concentration (Figure 3b,d). Except for abrupt increases at the lowest CO$_2$ concentrations (<50 ppm) of maize, TChlF$_{760}$ and $F_s$ for both species responded similarly: they continuously increased until the external concentration was around 500 ppm CO$_2$ (Figure 3j–l). With the increases in both $A_G$ and TChlF$_{760}$ ($F_s$), steep decreases in NPQ were observed, as expected, but no adjustments in leaf-level energy partitioning were found when the CO$_2$ concentration exceeded 500 ppm (Figure 3f,h). Excluding measurements at low CO$_2$ concentrations (<50 ppm), TChlF$_{760}$ of soybean and maize was highly correlated with $A_G$, with $R^2 = 0.96$ and $R^2 = 0.87$, respectively (Table 2); $F_s$ also explained 94% and 95% of the variability in the $A_G$ measurements for soybean and maize, respectively (Table 2).
Figure 3. Variation in gross carbon assimilation ($A_G$, μmol m$^{-2}$ s$^{-1}$, a,c), stomatal conductance to water vapor ($g_{sw}$, mol m$^{-2}$ s$^{-1}$, b,d), quantum efficiency of photosystem II ($\Phi_{PSII}$, e,g), nonphotochemical quenching (NPQ, f,h), total fluorescence radiance at 760 nm measured from both sides of leaves ($TChlF_{760}$, mW m$^{-2}$ nm$^{-1}$ sr$^{-1}$, i,j), and light-adapted steady-state PAM fluorescence ($F_s$, j,l) with CO2 concentrations (ppm) measured on three leaves of soybean (left two columns) and maize (right two columns) under irradiance of 800 (soybean) and 1200 (maize) μmol m$^{-2}$ s$^{-1}$. The three different colored lines represent three individual samples. $n_{CO2} = 8$ for individual leaves and 24 for a set. $n_{ABA} = 30$ for individual leaves and 90 for a set.

Table 2. Coefficients of determination ($R^2$) for the linear regressions between fluorescence radiance and photosynthetic capacity for the measurements driven by CO2 concentration (ppm), leaf temperature ($^\circ$C), and abscisic acid (ABA) treatments under constant irradiance of 800 μmol m$^{-2}$ s$^{-1}$. $A_G$ represents gross carbon assimilation (μmol m$^{-2}$ s$^{-1}$). $TChlF_{760}$ and $F_s$ represent the total emitted fluorescence radiance at 760 nm (mW m$^{-2}$ nm$^{-1}$ sr$^{-1}$) and light-adapted steady-state PAM fluorescence. Bold values indicate regression insignificance ($p \geq 0.05$). This table analyzes four groups of datasets: three individual samples (leaf numbers 1–3) and their average (Set). $n_{CO2} = 8$ for individual leaves and 24 for a set. $n_{ABA} = 30$ for individual leaves and 90 for a set.

| Species | Treatments | Fluorescence | $A_G$     |
|---------|------------|--------------|-----------|
|         |            |              | Leaf 1    | Leaf 2 | Leaf 3 | Set      |
| Soybean | CO2        | $TChlF_{760}$| 0.93      | 0.94  | 0.95  | 0.96     |
|         | Leaf       | $TChlF_{760}$| 0.93      | 0.94  | 0.91  | 0.94     |
|         | temperature| $F_s$        | 0.86      | 0.91  | 0.95  | 0.96     |
|         | ABA        | $TChlF_{760}$| 0.81      | 0.92  | 0.75  | 0.89     |
|         | CO2        | $F_s$        | 0.99      | 0.95  | 0.99  | 0.99     |
| Maize   | Temperature| $TChlF_{760}$| 0.86      | 0.76  | 0.73  | 0.93     |
|         |            | $F_s$        | 0.90      | 0.92  | 0.94  | 0.95     |
|         |            | $TChlF_{760}$| 0.83      | 0.83  | 0.92  | 0.87     |
|         |            | $F_s$        | 0.79      | 0.93  | 0.94  | 0.92     |

For the temperature response of CO2 assimilation, soybean $\Phi_{PSII}$ was likely symmetrical but more shallow, showing that the potential range of $T_{opt}$ for soybean $\Phi_{PSII}$ was likely
3.2. Effects of Temperature on the Variations in Carbon Assimilation and Fluorescence Emission

For the temperature response of CO₂ assimilation, soybean A₇ and gsₜ showed a bell-shaped response to leaf temperatures (Tleaf), with an optimum temperature (Topt) at around 25 °C (Figure 4a,b). The temperature response curves of soybean ΦPSII were also symmetrical but more shallow, showing that the potential range of Topt for soybean ΦPSII was likely broader (Figure 4e). Soybean NPQ changed with changing Tleaf in an opposite direction to A₇ (Figure 4f), exhibiting a bowl-like shape, and the net effect of A₇ and NPQ on fluorescence flux tended to diminish. As a result, TChlF₇₆₀ and Fs of soybean appeared to be sensitive to varying leaf temperatures (Tleaf < 30 °C) when the light intensity was fixed (Figure 4i,j). Maize showed higher Topt and CO₂ fixation rates at Topt: A₇ and ΦPSII of maize increased rapidly with increasing Tleaf below Topt, and they tended to stabilize or slightly decrease at high Tleaf (Figure 4c,g). The temperature dependency of maize NPQ also followed a bowl-shaped optimum curve, but with a more asymmetrical pattern than observed for soybean (Figure 4h). At variable Tleaf and a fixed light intensity, TChlF₇₆₀ accounted for more than 92% of the variability in A₇ (Table 2). By contrast, Fs showed an unstable predictive power, accounting for 89% of the variability in soybean A₇, compared with 71% for maize A₇ (Table 2). This variation was possibly due to the fluorescence measurements probing a small leaf area, while gas exchange techniques provide an integrated measurement over an entire leaf.

![Figure 4](image-url)

**Figure 4.** Variation in gross carbon assimilation (A₇, µmol m⁻² s⁻¹, (a,c)), stomatal conductance to water vapor (gsₜ, mol m⁻² s⁻¹, (b,d)), quantum efficiency of photosystem II (ΦPSII, (e,g)), nonphotochemical quenching (NPQ, (f,h)), total fluorescence radiance at 760 nm measured from both sides of leaves (TChlF₇₆₀, mW m⁻² nm⁻¹ sr⁻¹, (i,k)), and light-adapted steady-state PAM fluorescence (Fs, (j,l)) with leaf temperature (T, °C) measured on three leaves of soybean (left two columns) and maize (right two columns) under irradiance of 800 (soybean) and 1200 (maize) µmol m⁻² s⁻¹. The three different colored lines represent three individual samples. nₜ = 6 for each leaf.
3.3. Effects of Stomatal Closure on the Variations in Carbon Assimilation and Fluorescence Emission

An apparent decrease in both $A_G$ and $g_{sw}$ was observed within 30 min after ABA addition (Figure 5a–c), confirming that the ABA treatment effectively induced stomatal closure and inhibited leaf CO$_2$ assimilation. Moreover, in the context of the energy distribution in these treatment leaves, a similarly large reduction was also observed in $\Phi_{PSII}$ (Figure 5d–f), thereby revealing the decline in photochemical quenching and lower light use efficiency of PSII. As a compensating pathway for dissipating the extra energy that could not be used in photochemistry, NPQ showed a significant increase following treatment (Figure 5d–f). More importantly, the apparent decreases in both TChlF$_{760}$ and $F_s$ were consistently observed for all three ABA-treated leaves (Figure 5g–i): TChlF$_{760}$ and $F_s$ decreased by 15.4 ± 2.0% and 13.5 ± 3.0%, respectively, after the treatment. We found that ChlF emission was well correlated with CO$_2$ fixation rates during stomatal closure invoked by ABA: TChlF$_{760}$ explained 99% and 73% of the variability in $A_G$ after the treatment. These results confirm that far-red fluorescence radiance could respond to the ABA treatments, resulting in a reduction in the efficiency of dark reactions.

Figure 5. Variation in soybean gross carbon assimilation ($A_G$, µmol m$^{-2}$ s$^{-1}$, (a–e), left $y$-axis), stomatal conductance to water vapor ($g_{sw}$, mol m$^{-2}$ s$^{-1}$, (a–c), right $y$-axis), quantum efficiency of photosystem II ($\Phi_{PSII}$, (d–f), left $y$-axis), nonphotochemical quenching (NPQ, (d–f), right $y$-axis), total fluorescence radiance at 760 nm measured from both sides of leaves (TChlF$_{760}$, mW m$^{-2}$ nm$^{-1}$ sr$^{-1}$, (g–i), left $y$-axis), and light-adapted steady-state PAM fluorescence ($F_s$, (g–i), right $y$-axis) for pre-treatment and post-treatment of ABA under irradiance of 800 µmol m$^{-2}$ s$^{-1}$. The vertical green dashed line indicates the start of the ABA treatment. The three columns represent three individual soybean leaves. $n_{ABA} = 30$ for each leaf.
4. Discussion and Conclusions

Marrs et al. [25] showed that leaf-scale fluorescence signals (SIF and F_s) did not exhibit a clear trend after stomatal closure, suggesting that changes in SIF emissions only responded to changes in the light reactions of photosynthesis. They further hypothesized that the linearity between SIF and GPP observed in many previous studies resulted from a shared driver: APAR. However, our results show that both ChlF and F_s could track changes in the photosynthetic CO_2 rate due to variations in the CO_2 concentration and leaf temperature under constant light conditions. Further, we observed a significant reduction in both ChlF(F_s) and A_G after inducing stomatal closure. Our findings, in contrast, suggest that changes in SIF can not only track the impacts of stresses on the light reactions of photosynthesis but also offer relevant insight into carbon assimilation.

Under steady-state conditions, the efficiencies of the light and dark reactions would coordinate with each other to optimally match the surrounding environmental conditions and internal substrate availability [36]. Downregulation or upregulation of the photochemical quantum yield should accompany the inhibition or promotion of light reaction centers, resulting in variations in the fluorescence intensity and NPQ. However, changes in the SIF magnitude related to changes in the overall efficiency of the dark reactions should be limited. Our experiment showed that the variations in ChlF and F_s were less than 22% in response to the experimental treatments. It should also be noted that the light intensities were kept constant in this study. Further research is needed to quantify the contribution of the dark reactions to SIF emissions under other light levels.

Marrs et al. [25] used a two-spectrometer approach to retrieve SIF at O_2-A absorption, that is, two spectrometers were used to measure incoming solar irradiance (E_I) and outgoing radiance from the canopy (E_O). As described by Gu et al. [27], however, it is virtually impossible to perform a precise cross-calibration between the upwelling and downwelling channels, and therefore E_I and E_O measurements are not sufficiently compatible for retrieving SIF emissions. To analyze the SIF response to factors other than incident PAR, it is important to reduce (or normalize) the impact of the variations in incoming irradiance. Although Marrs et al. [25] tried to minimize the impact of varying PAR by performing the pre/post-treatment measurements at nearly the same time (but on a different date), their strategy may not effectively rule out uncertainties such as cloud passages that may result in fluctuating light conditions. Considering these issues in their experiment, it is unlikely that they could accurately identify the response of the dark reactions of photosynthesis from their measurements. Helm et al. [26] suggested that SIF was mainly controlled by the light reactions, and the fluorescence response occurred much later than the stomatal response. However, most of SIF and SIF yields in their study showed a large standard deviation of about 30% of their mean values, most likely caused by large differences among the samples. Thus, the simultaneous SIF response to stomatal closure might be overwhelmed by their large standard deviations.

Our experiments also showed that changes in Φ_{ChlF} and Φ_{PSII} are largely congruent under constant light, suggesting that both of them have a similar response to changes in the dark reactions. This appeared to be inconsistent with previous studies that reported the decoupling of SIF emissions and the carbon reactions of photosynthesis under certain conditions. For example, SIF emissions showed a rapid increase shortly after the application of a herbicide, resulting in a large reduction in CO_2 assimilation rates [37]. Liu et al. [38] showed that the GPP of C3 crops exhibited so-called ‘midday depression’, a stable or even decreasing trend in photosynthesis at high light, whereas an increase in the SIF curve was observed. Kim et al. [39] reported a strongly non-linear relationship between GPP and SIF in an evergreen needleleaf forest based on tower observations, where GPP exhibited a saturation pattern under high-light conditions, while SIF did not.

We propose that these findings are not contradictory to our results. In the application of the herbicide, the electron transport chain was blocked such that carbon assimilation decreased and SIF emissions were triggered as one of the de-excited paths to quickly dissipate absorbed excessive light energy, causing a divergence in the SIF–GPP relation-
ship over short time scales. In contrast, the electron transport was not inhibited in the ABA treatment, and the feedback between light and dark reactions still remained normal. Although Liu et al. [38] used one single spectrometer in their field experiment, avoiding wavelength mismatch between $E_I$ and $E_O$ [27], the uncertainties in their SIF observations due to the too long basic set sampling time (30 min) may very likely have caused the impact of the dark reactions to be indistinguishable. It is worth noting that Kim et al. [39] also reported that leaf-level $\Phi_{ChlF}$ exhibited a negative response to light intensity. This somewhat counterintuitive linkage between decreasing photosynthesis, increasing SIF, and decreasing SIF yield could be explained by the different sensitivity of $\Phi_{PSII}$ and $\Phi_{ChlF}$ to light levels: $\Phi_{PSII}$ shows a much steeper decline than $\Phi_{ChlF}$ in response to increasing stress levels such as light intensities [36]. As a consequence, the small decrease in $\Phi_{ChlF}$ may be overwhelmed by the large increase in PAR, with the net effect of an increase in SIF emissions. Taken together, a well-designed SIF-observing system should have the potential to detect the contributions of the dark reactions of photosynthesis under field conditions. A tower-based canopy SIF system called FAME, developed very recently by Gu et al. [27], significantly improved both the hardware and software designs, and their measurements showed that canopy-leaving directional SIF emissions saturated or even decreased at high light. In a recent high-temperature experiment, Kimm et al. [30] found that both light use efficiency and SIF yield decreased with increasing heat stress when PAR was not a limiting factor. These two studies highlighted the role of the dark reactions in the partitioning of absorbed solar radiation to the different pathways.

Our experiments confirm that SIF had the capacity to track changes in the dark reactions of photosynthesis—in other words, the gas exchange and carbon assimilation of plants. A well-designed measurement system with the ability to measure $E_I$ and $E_O$ almost simultaneously is needed to fully realize its potential. This study highlights the role of the dark reactions that have been ignored or misinterpreted in previous studies and will support our understanding of how elevated CO$_2$ concentrations and warming under future climate change influence photosynthesis activities.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/app112210821/s1, Text S1: Fluorescence quantum yield, Figure S1: Fluorescence yield at 760 nm ($\Phi_{ChlF,760}$) with CO$_2$ concentrations (ppm) measured on three leaves of soybean (left column) and maize (right column) under illumination of 800 (soybean) and 1200 (maize) $\mu$mol m$^{-2}$ s$^{-1}$, Figure S2: Fluorescence yield at 760 nm ($\Phi_{ChlF,760}$) with leaf temperature ($T, \degree C$) measured on three leaves of soybean (left column) and maize (right column) under illumination of 800 (soybean) and 1200 (maize) $\mu$mol m$^{-2}$ s$^{-1}$, Figure S3: Fluorescence yield at 760 nm ($\Phi_{ChlF,760}$) for pre-treatment and post-treatment of abscisic acid (ABA) under the illumination of 800 $\mu$mol m$^{-2}$ s$^{-1}$. The vertical green dashed line indicates the start of ABA treatment.

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