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RESEARCH PAPER

Variation between rice accessions in photosynthetic induction in flag leaves and underlying mechanisms

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Abstract

Several breeding initiatives have sought to improve flag leaf performance as its health and physiology are closely correlated to rice yield. Previous studies have described natural variation of photosynthesis for flag leaves; however, none has examined their performance under the non-steady-state conditions that prevail in crop fields. Photosynthetic induction is the transient response of photosynthesis to a change from low to high light. Rice flag leaf photosynthesis was measured in both steady- and non-steady-state conditions to characterize natural variation. Between the lowest and highest performing accession, there was a 152% difference for average CO₂ assimilation during induction (\bar{A}_{300}), a 77% difference for average intrinsic water use efficiency during induction (iWUE_{ava}), and a 185% difference for the speed of induction (IT₅₀), indicating plentiful variation. No significant correlation was found between steady- and nonsteady-state photosynthetic traits. Additionally, measures of neither steady-state nor non-steady-state photosynthesis of flag leaves correlated with the same measures of leaves in the vegetative growth stage, with the exception of iWUE_{ave}. Photosynthetic induction was measured at six [CO₂], to determine biochemical and diffusive limitations to photosynthesis in vivo. Photosynthetic induction in rice flag leaves was limited primarily by biochemistry.

Keywords: Atmospheric change, crop improvement, flag leaves, food security, natural variation, photosynthetic induction, rice, rice breeding, Rubisco activation, water use efficiency.

Introduction

on a mature flowering stem. The flag leaf has a higher

In cereals, the flag leaf is defined as the last leaf to emerge photosynthetic capacity relative to lower canopy leaves due to its position at the top of the canopy, which allows for greater

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interception of light (Adachi *et al.*, 2017). Additionally, high rice yields are closely correlated with the size and health of the flag leaf as they contribute some 50% of assimilates utilized for grain filling (Yoshida, 1981; Ishii, 1993; Nakano *et al.*, 1995; Li *et al.*, 1998). This large contribution of photosynthates is partially due to the proximity of the flag leaf to the grain, as the sink can more easily attract assimilates from closer sources (Sicher, 1993). The rest of the photosynthates used in grain filling are supplied by the leaf immediately below the flag leaf, and by remobilization of stored carbohydrates in leaf sheaths and older senescing leaves (Li *et al.*, 2017; Lin *et al.*, 2018). Agronomic strategies are typically aimed to protect the flag leaf, since its destruction during grain filling is associated with yield losses of up to 45% in rice (Abou–Khalifa *et al.*, 2008).

Consequently, great effort has been dedicated to understanding and optimizing the flag leaf to improve yields (Li et al., 1998). For example, flag leaf size and area can significantly influence grain yield (Li et al., 1998; Zhang et al., 2015). In rice, flag leaves with larger areas are significantly correlated with greater yields and have become a target for breeding programs seeking to achieve an ideal phenotype or ideotype (Zhang et al., 2015). Quantitative trait loci (QTLs) for larger flag leaves have been identified with the purpose of improving yield (Zhang et al., 2015). Additionally, rice yields have been increased by up-regulating NAL1, a gene that affects both flag leaf area and photosynthetic rates (Fabre et al., 2016). Increasing rice flag leaf photosynthesis, through either delaying senescence or a more acute leaf angle, has led to higher yields (Mantilla-Perez et al., 2017). Early flag leaf senescence significantly reduces seed-setting rate, 100-grain weight, and yield (Lin et al., 2018). Conversely, delaying senescence prolongs the period in which the flag leaf is photosynthetically active, resulting in more photosynthates that can fill grain (Ishii, 1993; Kobata et al., 2015; Leng et al., 2017). In previous studies, more vertical flag leaf angles resulted in 13% higher photosynthetic rates, reduced photoinhibition, delayed leaf senescence, and 15% higher yields (Chen et al., 2002; Mantilla-Perez et al., 2017).

Open-air elevation of $[CO_2]$ using free-air CO_2 enrichment technology is an artificial means of enhancing photosynthesis season-long in field-grown rice and observing the yield response. Modern high-yielding rice lines that have larger single-grain sizes and are able to produce larger panicles show a greater response to this elevation of photosynthesis, in contrast to older lines which appear partially sink limited. The fact that even under these circumstances only 70–80% of spikelets ripen—and less than in the older rice lines—suggests that these high-yielding lines are actually strongly source limited (Hasegawa *et al.*, 2013; Zhu *et al.*, 2015; Sakai *et al.*, 2019; Ainsworth and Long, 2020). These findings suggest that increasing the photosynthetic capacity of the flag leaf would be particularly valuable for these high-yielding cultivars.

Photosynthetic induction is the process by which leaves begin to assimilate CO_2 upon a transition from low light into high light, and is characterized by a lag in photosynthetic efficiency. Previously, examination of rice leaves in the vegetative growth stage found greater variation for photosynthetic traits during non-steady-state lighting compared with steady state (Acevedo-Siaca *et al.*, 2020). Despite the focus on flag leaf improvement, no studies have characterized the response of flag leaf photosynthesis in non-steady-state conditions or explored the possibility that there is variation in this character that could be utilized for improvement. Recently, it was shown that rice flag leaves are subject to different endogenous aging programs compared with other leaves (Lee *et al.*, 2017). Given its proximity to the major sink in the panicle, feedback would also be expected through non-structural carbohydrate-driven 'feast' and 'famine' gene expression response networks (Paul *et al.*, 2017). It is therefore likely that photosynthetic properties of flag leaves will differ from those of leaves formed in the vegetative growth stage.

Non-steady-state photosynthesis is important in a field agricultural setting where light is never constant. At the top of the canopy, flag leaves are subject to light fluctuations due to intermittent cloud cover and shadowing by other flag leaves or panicles as wind and sun angles change (Taylor and Long, 2017; Wang et al., 2020). In wheat, time taken for photosynthetic efficiency to recover from transient shadowing over the course of the day in the field was calculated to cost 21% of potential flag leaf assimilation (Taylor and Long, 2017). Improving the rate of recovery in this rarely measured photosynthetic parameter of flag leaves has the potential to increase yield. Limitations to the speed of photosynthetic induction on such shade to sun transitions are due to four main processes: the photoactivation of enzymes involved in the regeneration and production of ribulose-1,5-bisphosphate (RuBP); the buildup of intermediates of carbon metabolism; the activation of Rubisco by Rubisco activase; and stomatal opening (Pearcy et al., 1994; Mott and Woodrow, 2000; Slattery et al., 2018).

This study analyzed the performance of flag leaf photosynthesis in both steady-state and non-steady-state conditions. The objectives were limited to rice flag leaves and aimed to (i) determine the extent of variation between accessions in nonsteady-state and steady-state photosynthetic parameters relating to productivity and resource use efficiency across rice accessions; (ii) examine the response of photosynthetic induction of rice flag leaves at different $[CO_2]$ to understand limitations to induction; and (iii) compare the response of photosynthetic induction in rice flag leaves between accessions and with induction in leaves in the vegetative growth stage of the same accessions as reported in Acevedo-Siaca *et al.* (2020).

Materials and methods

Growing conditions and germplasm

Six accessions were selected from the 3000 Rice Genome Project (3K RGP) held at the Germplasm Resources Center at the International Rice Research Institute (IRRI) in Los Baños, Philippines. Accessions were selected on nucleotide mismatches for the gene encoding Rubisco activase, which plays a central role in photosynthetic induction. Seeds were maintained at 50 °C for 1 week to break dormancy and then

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sown into soil from the IRRI Upland Farm in small pots (4.5 cm diameter×12 cm) and fertilized using 0.4 g^{-1} Osmocote Plus 15-9-12 (The Scotts Company Ltd, Thorne, UK). Seedlings were transferred to larger individual pots (21.5 cm diameter×21.5 cm, 6 liters) after the emergence of the second leaf. These were then placed in a screen house, a type of greenhouse with a glass roof and screen-meshed walls, with no additional lighting or temperature control at IRRI, during the Philippines dry season from March to May 2017. Each pot was kept flooded using a drip irrigation system to simulate paddy conditions (Supplementary Fig. S1).

Gas exchange measurements

Photosynthetic induction

After anthesis, the flag leaf of the main stem was placed in the cuvette of an open gas exchange system (LI-6400XT, LI-COR, Lincoln, NE, USA). Light was provided by an integrated LED head (2×3 LED, LI-6400-02B). Within the cuvette, air temperature was 28 °C to approximate ideal growing conditions, flow rate was 400 μ mol s⁻¹, and [CO₂] was maintained at 400 μ mol mol⁻¹, and water vapor pressure deficit (VPD) at 1.3–1.7 kPa. Prior to measuring photosynthetic induction, rice plants were dark adapted for at least 1 h.

For induction, leaves were first allowed to reach a steady state in low light with a photosynthetically photon flux density (PPFD) of 50 µmol $m^{-2} s^{-1}$ ('shade') for 300 s followed by 720 s at 1700 µmol $m^{-2} s^{-1}$ ('sun'). Gas exchange measures were recorded every 10 s for the duration of the experiment. Measurements were repeated for all six accessions (*n*=8 plants per accession) over 4 d to minimize any age effects. Plants were selected by a randomized design and measured from 08.00 h to 12.00 h, to avoid confounding accessions with any diurnal influences. Net CO₂ uptake (*A*), stomatal conductance (*g*_s), intercellular CO₂ concentration (*C*_i), transpiration (*E*), and intrinsic water use efficiency (iWUE) were calculated following the equations of Farquhar *et al.* (1981). For a summary of all traits measured, see Table 1.

Two accessions, previously reported to show very different rates of induction for their leaves during the vegetative phase of growth, IR64-21 and AUS 278, were selected for further analysis of flag leaf induction at different [CO₂] (Acevedo-Siaca et al., 2020). Induction was measured following the protocol described above for induction, but at a cuvette [CO₂] of either 100, 200, 300, 400, 600, or 800 µmol mol⁻¹ during induction. The order of cuvette [CO₂] treatments for each individual leaf was randomized to avoid confounding [CO₂] with time. Leaves were dark adapted for a minimum of 1 h between measurements at the different [CO₂] at which they would later be measured. To determine limitations through induction, A was plotted against C_i for different time points, following the procedure of Soleh et al. (2016) and of Acevedo-Siaca et al. (2020). Five time points were selected for further analysis: 60, 180, 300, 360, and 700 s from the initiation of induction by transfer from darkness to high light (1700 µmol m⁻² s⁻¹). This allowed determination, at each time point, as to whether photosynthesis within the mesophyll was limited by the apparent maximum rates of RuBP regeneration (J_{max}) or RuBP carboxylation by Rubisco $(V_{c,max})$.

Steady-state measurements

For steady-state measurements, flag leaves were allowed to reach constant rates of A and stomatal conductance (g_s) at 1700 µmol m⁻² s⁻¹ PPFD. Cuvette conditions for steady-state measurements were as described above for photosynthetic induction.

Calculations

The rate constant of Rubisco activation, time to activation of

photosynthesis, and forgone assimilation

The rate constant of Rubisco activation $(1/\tau)$ was calculated by fitting the slope of the linear phase of the natural log of corrected photosynthetic induction $[\ln(A_t \star - A \star)]$ as described in Woodrow and Mott (1989).

Light condition	Trait	Description	Unit
Steady-state	A _{sat}	Leaf net CO ₂ uptake in saturating light	µmol m ^{−2} s ^{−1}
	$g_{ m s}$	Stomatal conductance	mol m ⁻² s ⁻¹
	C_{i}	Intercellular CO ₂ concentration	µmol mol⁻¹
	iWUE	Intrinsic water-use efficiency (iWUE=A/ $g_{\rm s}$)	μ mol CO ₂ mol H ₂ O ⁻¹
	V _{c, max}	Maximum rate of carboxylation	μ mol m ⁻² s ⁻¹
	J _{max}	Maximum rate of electron transport	μ mol m ⁻² s ⁻¹
	CE	Carboxylation efficiency	mol m ⁻² s ⁻¹
	Г	Compensation point	µmol m ⁻² s ⁻¹
	Φ	Quantum yield	Unitless (0–1)
Non-steady-state	Ā ₃₀₀	Average A during first 300 s of induction	μ mol m ⁻² s ⁻¹
	$\overline{g_s}$	Average g_s during first 300 s of induction	mol m ⁻² s ⁻¹
	$\overline{C_i}$	Average C_i during first 300 s of induction	µmol mol ⁻¹
	iWUE	Average iWUE during first 300 s of induction	μ mol CO ₂ mol H ₂ O ⁻¹
	IT_{50}	Time to 50% induction	Seconds
	IT ₉₀	Time to 90% induction	Seconds
	A_{Max}	Maximum A during induction	μ mol m ⁻² s ⁻¹
	A ₃₀₀	A at the end of 300 s	μ mol m ⁻² s ⁻¹
	A*	A corrected for stomatal limitation	μ mol m ⁻² s ⁻¹
	1/ τ	Rate constant of Rubisco activation	Seconds
	τ	Time to activation of photosynthesis	Seconds
	C Loss	Forgone assimilation	µmol m ^{−2}

Table 1. A summary of all traits measured and mentioned in the text

Units and light conditions are included.

 A^{\star} is photosynthesis at a point in time during induction corrected to a C_i value of 300 µmol mol⁻¹ to remove limitation from stomata. A_i^{\star} is the corrected value for photosynthesis at a C_i of 300 µmol mol⁻¹ at the end of the induction. The correction to C_i was made following the methods of Soleh *et al.* (2016):

$$A^* = A \times \frac{300}{C_i} \tag{1}$$

The time required to complete the activation of photosynthesis (τ) was calculated by taking the reciprocal of the rate constant of Rubisco activation (Woodrow and Mott, 1989).

The integrated amount of CO_2 uptake foregone due to the lower rates through induction compared with steady-state ($C Loss_t$) was calculated as in Acevedo-Siaca *et al.* (2020):

$$C \ Loss_t = \left(A - \overline{A}_t\right) \times t \tag{2}$$

where A is the steady-state rate of uptake and \bar{A}_t the average rate across the measured time period from the start of the induction (*t*), 300 s and 700 s, respectively.

Statistical analyses

All statistical analyses and model fitting used R (version 3.5.2, R-project) (R Core Team, 2020). Normal distribution and homogeneity of variances were tested by the Shapiro–Wilk test and Brown–Forsythe test, respectively. Assumptions were met and ANOVA was performed followed by Tukey's mean discrimination analysis, using the R-Project: 'agricolae:

Statistical Procedures for Agricultural Research' package. Pearson correlation coefficients between different photosynthetic measures were calculated using accession mean values (R; 'corrplot' and 'Hmisc').

Results

Characterizing photosynthesis in flag leaves

Significant variation was found across accessions for photosynthetic traits in both non-steady-state and steady-state conditions (Figs 1, 2). Significant differences were seen in steady-state parameters such as CO₂ uptake in saturating light (A_{sat} , P<0.0001), stomatal conductance (g_s , P=0.003), intercellular CO₂ concentration (C_i , P=0.03), and intrinsic water use efficiency (iWUE, P=0.006) (Fig. 2). Between the highest and lowest performing accessions, during induction there was a 152% difference for average CO₂ assimilation (\bar{A}_{300}) (M 102, 4.1 µmol m⁻² s⁻¹ versus IR64-21, 10.4 µmol m⁻² s⁻¹), 77% difference for average iWUE (iWUE_{avg}) (Fei Zhao 12, 36.2 µmol CO₂ mol H₂O⁻¹; AUS 278, 64.1 µmol CO₂ mol H₂O⁻¹), and a 185% difference for the time to 50% induction (IT₅₀) (Fei Zhao 12, 34.3 s; M 102, 98.2 s) (Fig. 3). Significant differences were also found among accessions for C Loss₃₀₀ and C Loss₇₀₀ (Fig. 4).



Fig. 1. The response of flag leaves from six rice accessions during photosynthetic induction. (A) Net leaf CO₂ assimilation (A), (B) stomatal conductance (g_s) , (C) intercellular CO₂ concentration (C_i), and (D) intrinsic water use efficiency (iWUE=A/g_s) with time (t) of induction upon change at 0 s from low light to high light (50 µmol m⁻² s⁻¹ to 1700 µmol m⁻² s⁻¹). The measurement was taken at an ambient [CO₂] of 400 µmol mol⁻¹. Two accessions, AUS 278 (red) and IR64-21 (black), were selected for further study at varied [CO₂]. The other four accessions are Dechangbyeo, Fei Zhao 12, M 102, and Malogbana. Each point is the mean ±SE) of eight plants (n=8).



Fig. 2. Mean and variation for flag leaf steady-state photosynthetic performance in six rice accessions. (A) Leaf CO_2 assimilation (A), (B) stomatal conductance (g_s), (C) intrinsic water use efficiency (iWUE=A/ g_s), and (D) intercellular CO_2 concentration (C_i). Accessions are ordered by median performance. Letters are indicative of a significant difference between accessions.

Understanding biochemical limitations to photosynthetic induction in flag leaves

Photosynthetic induction in flag leaves was limited primarily by biochemistry, rather than stomata, during the first 300 s of induction (Fig. 5). The rate of photosynthesis corrected to remove stomatal limitation (A^*) did not differ significantly from the uncorrected A values, suggesting that biochemical limitation dominated for these accessions (Fig. 5). This was also evident by the fact that C_i was higher than at steady state through most of the induction (Fig. 1C). A^* showed only small differences from the corresponding A (~3–4%), except for AUS 278 which had an average difference of ~13% between A and A^* (Fig. 5).

Photosynthetic induction was measured at six different $[CO_2]$ values in two selected accessions, AUS 278 and IR64-21. As expected, as $[CO_2]$ increased, *A* increased in both accessions (Supplementary Fig. S3). Also as expected, stomatal conductance decreased with increased $[CO_2]$ (Supplementary Fig. S3). As time after the beginning of induction increased, the amount of CO_2 assimilated increased as well (Fig. 6). This response to increasing CO_2 assimilation over time was seen in both selected accessions (Fig. 6).

Utilizing these data, the operating point at an ambient $[CO_2]$ (C_a) of 400 µmol mol⁻¹ was calculated for each curve.

The operating point fell on the initial slope of the A/C_i curve, indicating limitation by Rubisco for both accessions at all time points (Fig. 6). Photosynthesis was corrected for stomatal limitation at the different [CO₂] (Supplementary Figs S4, S5) to calculate the rate constant for Rubisco activation, τ , and forgone assimilation (*F*). The rate constant for Rubisco activation and τ was not significantly influenced by [CO₂] (Supplementary Figs S6, S7) while *C* Loss₃₀₀ increased significantly with [CO₂] (Supplementary Fig. S7).

Finally, the photosynthetic response curves to PPFD, measured at low $[CO_2]$ ($\leq 300 \ \mu mol \ mol^{-1}$), were used to infer the response of $V_{c,max}$ to a PPFD *in vivo*. These results indicated a significant difference between AUS 278 and IR64-21, with AUS 278 being more strongly limited by the rate of carboxylation (Supplementary Fig. S8). This contrasts with induction at ambient $[CO_2]$, where AUS 278 and IR64-21 did not vary significantly in photosynthetic induction traits \bar{A}_{300} , $g_{s avg}$, $C_{i avg}$, iWUE_{avg}, IT₅₀, and IT₉₀ (Fig. 3).

Comparing steady- and non-steady-state photosynthetic performance

The photosynthetic performance of accessions in non-steadystate and steady-state conditions was compared using Pearson's



Fig. 3. Mean and variation for flag leaf non-steady-state photosynthetic performance in six rice accessions. (A) CO₂ assimilation during the first 300 s of induction (\tilde{A}_{300}), (B) average stomatal conductance during the first 300 s of induction ($g_{s avg}$), (C) average intrinsic water use efficiency (iWUE_{avg}= $\tilde{A}_{300}/g_{s avg}$), (D) average intercellular CO₂ concentration ($C_{i avg}$), (E and F) time at which *A* reached 50% and 90% of A_{300} (IT₅₀ and IT₉₀, respectively). Accessions are ordered by median performance. Letters are indicative of a significant difference between accessions.

correlation coefficient. Significant (P<0.05) correlations were found between the different photosynthetic traits measured in steady-state and between the different traits measured in non-steady-state conditions (Fig. 7). However, there were no significant correlations between traits measured at steady state and their corresponding traits measured at non-steady state (Fig. 7). Additionally, as was previously found in vegetativephase leaves, there was a significant correlation (P<0.05) between \bar{A}_{300} and time at which A reached 50% induction (IT₅₀) (Fig. 7).

Comparing photosynthetic induction performance between flag leaves and vegetative-phase leaves

The photosynthetic traits measured here for flag leaves were compared with those made for leaves in the vegetative growth stage of the same accessions in a previous study (Acevedo-Siaca *et al.*, 2020). Significant correlations were found between steady-state C_i in leaves in the vegetative growth stage and iWUE in flag leaves, C_i in leaves in the vegetative growth stage and g_s in flag leaves, and iWUE in leaves in the vegetative growth phase and flag leaves (Fig. 8A). In non-steady-state conditions,



(a) p = 0.0002

Fig. 4. The integra1 of CO_2 uptake forgone due to the lower than steadystate rates through the first (A) 300 s and (B) 700 s of induction compared with steady state ($C \text{ Loss}_{300}$ and $C \text{ Loss}_{700}$, respectively).

significant correlations were found between $C_{i avg}$ in leaves in the vegetative growth stage and both $C_{i avg}$ and $iWUE_{avg}$ in flag leaves (Fig. 8). However, there was no significant correlation between any measure of CO₂ assimilation between flag leaves and leaves during the vegetative growth phase.

Discussion

Currently, no study has aimed to characterize the photosynthetic induction response in rice flag leaves. With respect to the objectives as outlined in the Introduction, the following key answers were obtained. (i) There were substantial differences among the six accessions of ~150% between leaf CO_2 uptake over the course of induction, with smaller differences in the light-saturated steady state (Figs 2, 3). This suggests significant scope for exploiting germplasm to increase rice photosynthesis in this key leaf for grain filling. (ii) Analysis of the responses of CO₂ uptake to intercellular CO₂ concentration showed that the *in vivo* activity of Rubisco $(V_{c,max})$ limited photosynthesis throughout induction (Fig. 6), suggesting that breeding or bioengineering increased Rubisco content and activity as a means to increase rice productivity. (iii) When the results obtained here for flag leaves were compared with those for the same accessions in the earlier study of leaves in the vegetative growth stage (Acevedo-Siaca et al., 2020), there was little correspondence between photosynthetic parameters for the two growth stages, with the exception of iWUE (Fig. 8). This suggests that breeding efforts to improve crop photosynthesis would need to address selection at both growth stages, although improved WUE could be selected at the vegetative stage alone, making this more tractable. The following discussion places these new findings in the context of prior studies and potential for increasing rice photosynthetic capacity, efficiency of resource use, and productivity.

Significant variation for photosynthetic traits in rice flag leaves

Previously, it was suggested that natural variation for photosynthesis was more prevalent in the flag leaves than for leaves in the vegetative growth stage of cereal crops, a finding confirmed here for rice (Dunstone et al., 1973; Bansal et al., 1993). Previous studies have focused on either steady-state or more recently non-steady-state photosynthesis, and demonstrated diversity within crop germplams (Flood et al., 2011; Gu et al., 2012; Driever et al., 2014; Soleh et al., 2016, 2017; Acevedo-Siaca et al., 2020; De Souza et al., 2020; Taniyoshi et al., 2020). Here, we found that significant natural variation exists for photosynthetic traits in both steadyand non-steady-state conditions (A 6.4–25.9 μ mol m⁻² s⁻¹; \bar{A}_{300} 2.3–16.7 µmol m⁻² s⁻¹) in the flag leaf. Indeed, 43, 12, and 68% more variation was found here between flag leaves when compared with leaves in the vegetative growth stage of the same accessions (Acevedo-Siaca *et al.*, 2020) for A_{300} , iWUE_{avg}, and IT₅₀, respectively. This is significant given the key role of the flag leaf.

Further evidence for the lack of correlation between steady- and non-steady-state measurements

Consistent with studies of soybean, cowpea, and leaves in the vegetative growth stage of rice, no significant correlations were found between parameters of photosynthetic CO_2 uptake in steady- and non-steady-state light in rice flag leaves (Soleh *et al.*, 2016; Acevedo-Siaca *et al.*, 2020; De Souza *et al.*, 2020). These studies, together with the present study, provide compelling evidence for a reconsideration of when during crop



Fig. 5. The response of uncorrected leaf CO₂ assimilation (A; filled circles) and the response of leaf CO₂ assimilation corrected for stomatal limitation (A^* ; open circles) over time in flag leaves of six rice accessions. The first line at 100 s indicates the mean time for the activation of Rubisco (τ) per accession. Each point represents the mean of at least six plants ±SE (n=6–8).

development we measure and how we measure photosynthesis when considering crop improvement. In particular, clear evidence that steady-state measurements do not indicate photosynthetic efficiency under the non-steady-state fluctuating light conditions that can be dominant in the field should be noted. However, the greater between-accession variability of non-steady-state photosynthesis highlights a greater opportunity for increasing net crop CO_2 uptake that could meet the apparent strong source limitation of modern high-yielding cultivars (Hasegawa *et al.*, 2013; Zhu *et al.*, 2015; Sakai *et al.*, 2019). It is possible that a balance for the distribution of resources between photosynthetic proteins underlies the differences between steady-state and non-steady-state photosynthesis. Previous studies have suggested a trade-off between maximum rates of photosynthesis in steady-state saturating light versus the speed of induction due to limited amounts of nitrogen invested and divided between Rubisco and Rubisco activase content (Woodrow and Mott, 1989; Mott and Woodrow, 2000). It is hypothesized that plants grown in fluctuating light environments that do not experience steady-state conditions frequently would benefit from having higher Rubisco activase



Fig. 6. The responses of leaf CO₂ assimilation (*A*) to intercellular [CO₂] (*C*₁) at different points in time after the beginning of photosynthetic induction for IR64-21 and AUS 278. Times after induction were: 60 s (filled circles), 180 s (open squares), 300 s (open circles), 360 s (filled squares), and 700 s (filled triangles) from the start of induction. The operating point of each curve at 400 µmol mol⁻¹ atmospheric [CO₂] (*C*₂) is indicated with a black arrow. Each point is the mean (±SE) of four plants of each rice accession.

content to be able to respond more quickly to changes in light (Mott and Woodrow, 2000; Yamori *et al.*, 2012; Carmo-Silva and Salvucci, 2013; Kaiser *et al.*, 2016). However, leaves that experience fewer sunflecks and have more exposure to direct sunlight would benefit from investing a higher proportion of resources in Rubisco to sustain higher photosynthetic rates at steady state (Mott and Woodrow, 2000). Such a trade-off is supported by the finding that antisense down-regulation of Rubisco activase increased Rubisco content in rice leaves (Jin *et al.*, 2006). This trade-off between Rubisco and Rubisco activase content could help to partially explain the lack of correlation between steady- and non-steady-state photosynthesis, as an increase in the protein that helps the leaf excel in induction acts to the detriment of the protein needed at steady state. The trade-off between steady- and non-steady-state photosynthesis was shown clearly in IR64-21, which had the highest steady-state A_{sat} yet was among the slowest to reach 90% of its steady-state level (IT_{90 A}) during induction (Figs 2, 3).

Photosynthetic induction is primarily limited by biochemistry in flag leaves

At ambient $[CO_2]$, rice flag leaves are predominantly limited by biochemistry, specifically the maximum activity of Rubisco (Figs 5, 6). Differences between A and A^{*} at ambient $[CO_2]$ were generally small, indicating little limitation by stomata (Fig. 5). This is likely to be due to the shape and small size of rice stomata allowing fast responses (McAusland *et al.*, 2016), but also the evolutionary history of rice, which was domesticated from emergent aquatic progenitors and then bred in paddy conditions where water would not be limiting to the plant (Nay-Htoon *et al.*, 2018).

Calculation of the operating point at ambient [CO₂] $(C_a=400 \text{ }\mu\text{mol mol}^{-1})$ in AUS 278 and IR64-21 showed that photosynthesis during induction was predominantly limited by Rubisco, and not affected by either the capacity for regeneration of RuBP or triose phosphate utilization, throughout the induction and into steady state. The operating point is on the initial slope of the A/C_i curve, throughout (Fig. 6). This parallels the previous findings for rice leaves in the vegetative growth stage (Acevedo-Siaca et al., 2020) and suggests that the predominant limitation to photosynthetic CO₂ assimilation at steady state and non-steady state throughout the life cycle of rice is consistently due to the in vivo capacity and, presumably, amount of Rubisco. This is also consistent with the recent observation that transgenic up-regulation of Rubisco in rice significantly increases paddy yield (Long, 2020; Yoon et al., 2020). One caveat is that this analysis is based on C_i and not the CO_2 concentration at Rubisco (C_c). Ease of movement of CO₂ from the intercellular space to Rubisco is governed by mesophyll conductance (g_m) , which was not measured here. So, it is possible that activation of $g_{\rm m}$, as well as Rubisco, could be a limiting factor. However, prior work with other species has suggested that activation of g_m is likely to be faster than activation of Rubisco (Deans et al., 2019).

Photosynthetic performance shows little correlation between flag and vegetative-phase leaves

The only significant correlations found between flag and leaves in the vegetative growth stage were for C_i and iWUE (Fig. 8). These results suggest that iWUE is not be affected by rice developmental stage and could be consistent throughout the lifetime of these rice plants. Water availability is the biggest limitation to agricultural production worldwide and is expected to pose an even greater limitation with climate change and population growth (Wassmann *et al.*, 2009; Ort and Long, 2014; Oladosu *et al.*, 2019). If iWUE is consistent throughout the life cycle of rice, it could allow breeders to screen for high iWUE



Fig. 7. Pearson correlation (*R*) of all measured dynamic and steady-state (filled tiangles) photosynthetic traits measured in rice flag leaves. Negative correlations are in blue, positive correlations are in red. Traits at steady-state are: intrinsic water-use efficiency (iWUE=A/g_s), transpiration (*E*), intercellular CO₂ concentration (*C*_i), stomatal conductance (*g*_s), and net CO₂ assimilation in saturating light (*A*_{sat}). Traits at non-steady state over the first 300 s of induction are: the time at which *A* reached 50% and 90% of *A*₃₀₀ (IT₅₀ and IT₉₀, respectively), average *C*_i during first 300 s of induction (*C*_{i avg}), average intrinsic water use efficiency (iWUE_{avg}= $\bar{A}_{300}/g_{s avg}$), average *g*_s, the maximum *A* during induction (*A*_{Max}), *A* at the end of this period (*A*₃₀₀), and the average *A* (\bar{A}_{300}). A significant *R* value is marked by a black line on the scale (0.8).

early in development, saving time and resources in selecting germplasm in breeding more water use-efficient plants.

Otherwise, no significant correlations were found between the photosynthetic performance of flag leaves and leaves in the vegetative growth stage in both steady- and non-steadystate conditions (Fig. 7). For example, elite cultivar IR64-21 was outperformed by other accessions in measurements of photosynthesis in leaves in the vegetative growth stage during steady- and non-steady-state conditions (Acevedo-Siaca et al., 2020). However, IR64-21 flag leaves had the highest photosynthetic rates during photosynthetic induction and at steady state, significantly outperforming the other accessions (Figs 1, 2; Supplementary Fig. S3). This suggests that increased flag leaf CO2 uptake may have been inadvertently improved through conventional breeding selection. This corresponds with other evidence that flag leaf photosynthesis has been improved unintentionally through breeding for higher yield potential. Newer and higher yielding rice varieties have flag leaves with higher photosynthetic rates per unit area than older varieties (Ishii, 1993). Additionally, flag leaves in Oryza sativa were found to maintain higher photosynthetic rates for longer relative to wild Oryza species (Ishii, 1993), which is curiously in contrast to wheat (Dunstone et al. 1973). These studies suggest that deliberate breeding for increased flag leaf photosynthesis might be a fertile avenue for a further increase in rice yield potential.

However, while an increased emphasis on flag leaf photosynthesis can lead to higher yields (Fabre *et al.*, 2016), there should still be a focus on improving photosynthetic efficiency throughout the life cycle. Photosynthesis in leaves during the vegetative stage is important in establishing the plant and developing a robust root system and tillers capable of becoming reproductive. Improved photosynthesis in leaves in the vegetative growth stage results in increases in non-storage



Fig. 8. Pearson correlation analysis between photosynthetic traits measured in flag leaves for the six cultivars measured here with the values obtained on leaves in the vegetative growth stage for the same cultivars in a previous study (Acevedo-Siaca *et al.* 2020). (A) Non-steady state; (B) steady state. Traits are as defined in Fig. 3.

carbohydrates in leaves and stems, which can subsequently be remobilized, for grain filling. The expected ideotype would therefore be an accession which shows high capacity and efficiency during both growth phases.

Non-steady-state photosynthesis—a practical target for breeding?

Non-steady-state photosynthesis has received little attention in breeding, largely because the gas exchange methods required to effectively phenotype traits are not practical for large-scale testing. However, it was recently shown in wheat that large-scale screening of non-steady-state photosynthesis could be achieved effectively with excised leaves using modulated chlorophyll fluorescence imaging (McAusland et al., 2019). This high-throughput technique could make selection and breeding for improved efficiency under nonsteady-state conditions practical, and probably more effective than selecting for improvement under steady-state conditions. Additionally, these methods could allow for the improvement of iWUE under non-steady-state conditions. As noted above, current methods of breeding paddy rice for yield may have lowered WUE. This study has revealed considerable variation-even within the limited germplasm examined-and a correlation in WUE between vegetative and reproductive growth. This finding suggests that selection of improved WUE could be achieved by screening during early growth. Furthermore, the development of integrated thermal and modulated fluorescence imaging of instantaneous WUE would now allow high-throughput phenotyping of this trait (McAusland et al., 2013). Accelerated breeding of rice lines requiring less water would help address the rising pressures on water supplies in many paddy rice-growing regions (Schyns et al., 2019). These benefits that go beyond increased yield are imperative for creating more sustainable agricultural systems that utilize this planet's limited resources with more discretion, especially in the face of global climate change and a growing human population.

Supplementary data

The following supplementary data are available at JXB online.

Fig. S1. Growing conditions at the IRRI used in the study.

Fig. S2. Corrected photosynthesis for time to activation of photosynthesis (τ).

Fig. S3. Induction of leaf CO_2 uptake (*A*) and stomatal conductance (g_s) at six [CO₂] in IR64-21 and AUS 278.

Fig. S4. Leaf CO₂ uptake (A) and leaf CO₂ uptake corrected for stomatal limitation (A^*) in AUS 278.

Fig. S5. Leaf CO_2 uptake (*A*) and leaf CO_2 uptake corrected for stomatal limitation (*A**) in IR64–21.

Fig. S6. Corrected photosynthesis for time to activation of photosynthesis (τ) in AUS 278 and IR64-21.

Fig. S7. Rate constant of Rubisco activation $(1/\tau)$, time to activation of photosynthesis (τ), and forgone assimilation in six $[CO_2]$ in AUS 278 and IR64-21.

Fig. S8. $V_{c,max}$ during photosynthetic induction in AUS 278 and IR64-21.

Table S1. Description of accessions used in this study.

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Author contributions

LGA and SPL planned the research. SPL and WPQ supervised the project. LGA and RC conducted the experimental work, LGA analyzed the data. LGA and SPL wrote the manuscript with the input of all the other authors.

Data availability

The data that support the findings of this study are openly available in the University of Illinois Data Bank at https://doi.org/10.13012/ B2IDB-3596430_V1 (Acevedo-Siaca, Liana; Long, Stephen (2020): Photosynthetic induction of rice flag leaves. University of Illinois at Urbana-Champaign.)

References

Abou-Khalifa ABA, Misra N, El-Azeem A, Salem KM. 2008. Effect of leaf cutting on physiological traits and yield of two rice cultivars. African Journal of Plant Science **2**, 147–150.

Acevedo-Siaca LG, Coe R, Wang Y, Kromdijk J, Quick WP, Long SP. 2020. Variation in photosynthetic induction between rice accessions and its potential for improving productivity. New Phytologist **227**, 1097–1108.

Adachi S, Yoshikawa K, Yamanouchi U, Tanabata T, Sun J, Ookawa T, Yamamoto T, Sage RF, Hirasawa T, Yonemaru J. 2017. Fine mapping of *Carbon Assimilation Rate 8*, a quantitative trait locus for flag leaf nitrogen content, stomatal conductance and photosynthesis in rice. Frontiers in Plant Science **8**, 60.

Ainsworth EA, Long SP. 2020. 30 years of free-air carbon dioxide enrichment (FACE): What have we learned about future crop productivity and its potential for adaptation? Global Change Biology doi: 10.1111/gcb.15375.

Bansal KC, Uprety DC, Abrol YP. 1993. Genetic variation in photosynthetic characteristics in wheat: causes and consequences. In: Abrol YP, Mohanty P, Govindjee, eds. Photosynthesis: photoreactions to plant productivity. Dordrecht: Springer Science + Business Media, 527–547.

Carmo-Silva E, Salvucci ME. 2013. The regulatory properties of rubisco activase differ among species and affect photosynthetic induction during light transitions. Plant Physiology **161**, 1645–1655.

Chen B, Zhang Y, Li X, Jiao D. 2002. Photosynthetic characteristics and assimilate distribution in super hybrid rice Liangyoupeijiu at late growth stage. Acta Agronomica Sinica 28, 777–782. [In Chinese, with an English abstract].

Deans RM, Farquhar GD, Busch FA. 2019. Estimating stomatal and biochemical limitations during photosynthetic induction. Plant, Cell & Environment **42**, 3227–3240.

De Souza AP, Wang Y, Orr DJ, Carmo-Silva E, Long SP. 2020. Photosynthesis across African cassava germplasm is limited by Rubisco and mesophyll conductance at steady state, but by stomatal conductance in fluctuating light. New Phytologist **225**, 2498–2512.

Driever SM, Lawson T, Andralojc PJ, Raines CA, Parry MA. 2014. Natural variation in photosynthetic capacity, growth, and yield in 64 fieldgrown wheat genotypes. Journal of Experimental Botany **65**, 4959–4973.

Dunstone RL, Gifford RM, Evans LT. 1973. Photosynthetic characteristics of modem and primitive wheat species in relation to ontogeny and adaptation to light. Australian Journal of Biological Sciences **26**, 295–307.

Fabre D, Adriani DE, Dingkuhn M, Ishimaru T, Punzalan B, Lafarge T, Clément-Vidal A, Luquet D. 2016. The qTSN4 effect on flag leaf size, photosynthesis and panicle size, benefits to plant grain production in rice, depending on light availability. Frontiers in Plant Science 7, 623.

Farquhar GE, von Caemmerer S, Berry JA. 1981. A biochemical model of photosynthetic CO_2 assimilation in leaves of C_3 species. Planta 149, 78–90.

Flood PJ, Harbinson J, Aarts MG. 2011. Natural genetic variation in plant photosynthesis. Trends in Plant Science **16**, 327–335.

Gu J, Yin X, Stomph TJ, Wang H, Struik PC. 2012. Physiological basis of genetic variation in leaf photosynthesis among rice (*Oryza sativa* L.) introgression lines under drought and well-watered conditions. Journal of Experimental Botany **63**, 5137–5153.

Hasegawa T, Sakai H, Tokida T, et al. 2013. Rice cultivar responses to elevated CO_2 at two free-air CO_2 enrichment (FACE) sites in Japan. Functional Plant Biology **40**, 148–159.

Ishii R. 1993. Leaf photosynthesis in rice in relation to grain yield. In: Abrol YP, Mohanty P, Govindjee, eds. Photosynthesis: photoreactions to plant productivity. Dordrecht: Springer Science + Business Media, 561–570.

Jin SH, Hong J, Li XQ, Jiang DA. 2006. Antisense inhibition of Rubisco activase increases Rubisco content and alters the proportion of Rubisco activase in stroma and thylakoids in chloroplasts of rice leaves. Annals of Botany 97, 739–744.

Kaiser E, Morales A, Harbinson J, Heuvelink E, Prinzenberg AE, Marcelis LFM. 2016. Metabolic and diffusional limitations of photosynthesis in fluctuating irradiance in *Arabidopsis thaliana*. Scientific Reports 6, 31252.

Kobata T, Shinonaga M, Yoshida H, Tomisaka K, Akai K. 2015. Staygreen trait assessment using the leaf incubation method to examine the maintenance of assimilation rates under high temperature conditions during the grain-filling period in rice. Plant Production Science **18**, 254–266.

Lee S, Jeong H, Lee S, et al. 2017. Molecular bases for differential aging programs between flag and second leaves during grain-filling in rice. Scientific Reports **7**, 9792.

Leng Y, Ye G, Zeng D. 2017. Genetic dissection of leaf senescence in rice. International Journal of Molecular Breeding 18, 2686.

Li Z, Pinson SRM, Stansel JW, Paterson AH. 1998. Genetic dissection of the source–sink relationship affecting fecundity and yield in rice (*Oryza sativa* L.). Molecular Breeding **4**, 419–426.

Li Z, Wang F, Lin W, Zhao Q, Liu J, Cheng F. 2017. Carbon reserve and remobilization in leaf sheaths during the grain-filling stage in response to leaf early senescence. Acta Physiologiae Plantarum **38**, 10.

Lin W, Guo X, Pan X, Li Z. 2018. Chlorophyll composition, chlorophyll fluorescence, and grain yield change in es/ mutant rice. International Journal of Molecular Sciences **19**, 2945.

Long SP. 2020. Photosynthesis engineered to increase rice yield. Nature Food 1, 105–105.

Mantilla-Perez MB, Salas Fernandez MG. 2017. Differential manipulation of leaf angle throughout the canopy: current status and prospects. Journal of Experimental Botany **68**, 5699–5717.

McAusland L, Atkinson JA, Lawson T, Murchie EH. 2019. High throughput procedure utilising chlorophyll fluorescence imaging to phenotype dynamic photosynthesis and photoprotection in leaves under controlled gaseous conditions. Plant Methods **15**, 109.

McAusland L, Davey PA, Kanwal N, Baker NR, Lawson T. 2013. A novel system for spatial and temporal imaging of intrinsic plant water use efficiency. Journal of Experimental Botany **64**, 4993–5007.

McAusland L, Vialet-Chabrand S, Davey PA, Baker NR, Brendel O, Lawson T. 2016. Effects of kinetics of light-induced stomatal responses on photosynthesis and water-use efficiency. New Phytologist **211**, 1209–1220.

Mott KA, Woodrow IE. 2000. Modelling the role of Rubisco activase in limiting non-steady-state photosynthesis. Journal of Experimental Botany **51**, 399–406.

Nakano H, Makino A, Mae T. 1995. Effects of panicle removal on the photosynthetic characteristics of the flag leaf of rice plants during the ripening stage. Plant & Cell Physiology **36**, 653–659.

Nay-Htoon B, Xue W, Lindner S, Cuntz M, Ko J, Tenhunen J, Werner C, Dubbert M. 2018. Quantifying differences in water and carbon cycling between paddy and rainfed rice (*Oryza sativa* L.) by flux partitioning. PLoS One **13**, e0195238.

Oladosu Y, Rafii MY, Samuel C, Fatai A, Magaji U, Kareem I, Kamarudin ZS, Muhammad I, Kolapo K. 2019. Drought resistance in rice from conventional to molecular breeding: a review. International Journal of Molecular Sciences **20**, 3519.

Ort DR, Long SP. 2014. Limits on yields in the corn belt. Science 344, 484–485.

Paul MJ, Oszvald M, Jesus C, Rajulu C, Griffiths CA. 2017. Increasing crop yield and resilience with trehalose 6-phosphate: targeting a feast-famine mechanism in cereals for better source-sink optimization. Journal of Experimental Botany **68**, 4455–4462.

Pearcy RW, Chazdon RL, Gross LJ, Mott KA. 1994. Photosynthetic utilization of sunflecks: a temporally patchy resource on a timescale of seconds to minutes. In: Caldwell MM, Pearcy RW, eds. Exploration of environmental heterogeneity by plants. San Diego: Academic Press, 175–208.

R Core Team. 2020. R: a language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing.

Sakai H, Tokida T, Usui Y, Nakamura H, Hasegawa T. 2019. Yield responses to elevated CO₂ concentration among Japanese rice cultivars released since 1882. Plant Production Science **22**, 352–366.

Schyns JF, Hoekstra AY, Booij MJ, Hogeboom RJ, Mekonnen MM. 2019. Limits to the world's green water resources for food, feed, fiber, timber, and bioenergy. Proceedings of the National Academy of Sciences, USA **116**, 4893–4898.

Sicher, R. 1993. Assimilate partitioning within leaves of small grain cereals. In: Abrol YP, Mohanty P, Govindjee, eds. Photosynthesis: photoreactions to plant productivity. Dordrecht: Springer Science + Business Media, 351–360.

Slattery RA, Walker BJ, Weber APM, Ort DR. 2018. The impacts of fluctuating light on crop performance. Plant Physiology **176**, 990–1003.

Soleh MA, Tanaka Y, Kim SY, Huber SC, Sakoda K, Shiraiwa T. 2017. Identification of large variation in the photosynthetic induction response among 37 soybean [*Glycine max* (L.) Merr.] genotypes that is not correlated with steady-state photosynthetic capacity. Photosynthesis Research **131**, 305–315.

Soleh MA, Tanaka Y, Nomoto Y, Iwahashi Y, Nakashima K, Fukuda Y, Long SP, Shiraiwa T. 2016. Factors underlying genotypic differences in the induction of photosynthesis in soybean [*Glycine max* (L.) Merr.]. Plant, Cell, & Environment **39**, 685–693.

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Taniyoshi K, Tanaka Y, Shiraiwa T. 2020. Genetic variation in the photosynthetic induction response in rice (*Oryza sativa* L.). Plant Production Science **23**, 513–521.

Taylor SH, Long SP. 2017. Slow induction of photosynthesis on shade to sun transitions in wheat may cost at least 21% of productivity. Philosophical Transactions of the Royal Society B: Biological Sciences **372**, 20160543.

Wang Y, Burgess SJ, de Becker EM, Long SP. 2020. Photosynthesis in the fleeting shadows: an overlooked opportunity for increasing crop productivity? The Plant Journal **101**, 874–884.

Wassmann R, Jagadish SVK, Sumfleth K, Pathak H, Howell G, Ismail A, Serraj R, Redona E, Singh RK, Heuer S. 2009. Regional vulnerability of climate change impacts on asian rice production and scope for adaptation. Advances in Agronomy **102**, 91–133.

Woodrow IE, Mott KA. 1989. Rate limitation of non-steady-state photosynthesis by ribulose-1,5-bisphosphate carboxylase in spinach. Functional Plant Biology **16**, 487–500. **Yamori W, Masumoto C, Fukayama H, Makino A.** 2012. Rubisco activase is a key regulator of non-steady-state photosynthesis at any leaf temperature and, to a lesser extent, of steady-state photosynthesis at high temperature. The Plant Journal **71**, 871–880.

Yoon DK, Ishiyama K, Suganami M, et al. 2020. Transgenic rice overproducing Rubisco exhibits high yields with high nitrogen use efficiency in a paddy field. Nature Food **1**, 134–139.

Yoshida, S. 1981. Physiological analysis of rice yield. In: Fundamentals of rice crop science. Makati City, Philippines: International Rice Research Institute, 231–251.

Zhang B, Ye W, Re D, et al. 2015. Genetic analysis of flag leaf size and candidate genes determination of a major QTL for flag leaf width in rice. Rice 8, 2.

Zhu C, Xu X, Wang D, Zhu J, Liu G. 2015. An indica rice genotype showed a similar yield enhancement to that of hybrid rice under free air carbon dioxide enrichment. Scientific Reports 5, 12719.