

An *in situ* approach to characterizing photosynthetic gas exchange of rice panicle

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Methodology

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Abstract

Background: Photosynthesis of reproductive organs in C₃ cereals is generally regarded as important to crop yield. Whereas, photosynthetic characteristics of reproductive organs are much less understood as compared to leaf photosynthesis, mainly due to methodological limitations. To date, many indirect methods have been developed to study photosynthesis of reproductive organs and its contribution to grain yield, such as organ shading, application of herbicides and photosynthetic measurement of excised organs or tissues, which might be intrusive and cause biases. Thus, a robust and *in situ* approach needs to be developed.

Results: Here we report the development of a custom-built panicle photosynthesis chamber (P-chamber), which can be connected to standard infrared gas analyzers to study photosynthetic/respiratory rate of a rice panicle. With the P-chamber, we measured panicle photosynthetic characteristics of seven high-yielding elite *japonica*, *japonica-indica* hybrid and *indica* rice cultivars. Results show that, 1) rice panicle is photosynthetically active during grain filling, and there are substantial inter-cultivar variations in panicle photosynthetic and respiratory rates, no matter on a whole panicle basis, on an area basis or on a single spikelet basis; 2) among the seven testing cultivars, whole-panicle gross photosynthetic rates are 17 – 54 nmol s⁻¹ 5 days after heading under photon flux density (PFD) of 2000 μmol (photons) m⁻² s⁻¹, which represent some 20-38% of that of the corresponding flag leaves; 3) rice panicle photosynthesis has higher apparent CO₂ compensation point, light compensation point and apparent CO₂ saturation point, as compared to that of a typical leaf; 4) there is a strong and significant positive correlation between gross photosynthetic rate 5 days after heading on a single spikelet basis and grain setting rate at harvest (Pearson correlation coefficient $r = 0.93$, p -value < 0.0001).

Conclusions: Rice panicle gross photosynthesis is significant, has great natural variation, and plays an underappreciated role in grain yield formation. The P-Chamber can be used as a tool to study *in situ* photosynthetic characteristics of irregular non-foliar plant organs, such as ears, culms, leaf sheaths, fruits and branches, which is a relatively less explored area in current cereal breeding community.

Background

In cereal crops such as rice and wheat, there are two major sources of assimilates used for grain filling, i.e., the post-anthesis photosynthesis and the pre-anthesis assimilates stored in the stem. Under non-stress conditions, the post-anthesis photosynthesis contributes greater to grain yield than the pre-anthesis stored assimilates [1-4]. Leaves, especially flag leaves located at the upmost of stems, have been believed to be the dominant contributor of post-anthesis photosynthesis [5, 6]. Nevertheless, reproductive organs, including glumes, caryopsis pericarps, and awns, are green before mature and photosynthetically active [7, 8], suggesting a possibility of them being a major source of photosynthate for grain filling as well. In fact, reproductive organs have several potential advantages over leaves in terms of photosynthate production and transportation: 1) they generally locate at the upper canopy, where light levels are higher than those at the lower canopy; 2) they are physically closer to the

developing grains than the leaves, which shorten the path of assimilates transport to developing grains [9]; 3) they have large surface area, e.g. the ear surface areas were 1.1-5.5 folds greater than that of the corresponding flag leaves in wheat and barley [6]; 4) reproductive organs tend to senesce later than leaves, especially under adverse conditions such as drought [10, 11]. Due to these reasons, photosynthesis of reproductive organs gained more and more attraction in recent years (see reviews in [12-15]).

In wheat and barley, ear photosynthesis is a major source of photosynthate for grain filling; a large photosynthetic area and high photosynthetic activity from both the green glumes and awns are major factors underlying higher ear photosynthesis [6, 16, 17]. Using ear-shading and/or stem-shading methods, ear photosynthesis was estimated to contribute 10% to 65% of the grain yield in different wheat lines and under different growth conditions [16-20]. By analyzing the carbon isotope composition in different organs, it was estimated that ear photosynthesis contributed 47% to 100% of the grain yield in six high-yielding bread wheat lines under well-managed conditions [21, 22]. In a retrospective study, by using 15 winter wheat cultivars released from 1980 to 2012 in North China Plain, it was reported that the contribution of ear photosynthesis to grain yield doubled (from around 30% to around 60%) during breeding [18]. Although the exact proportion of ear photosynthesis to total grain yield varies, breeding cultivars with higher ear photosynthesis was recognized as a new mechanism to improve wheat yield potential [14, 15, 23].

In contrast to the significant importance of ear photosynthesis in wheat and barley, rice panicle has been traditionally regarded as a sink organ with negligible photosynthetic contribution to grain yield [24]. For example, Tsuno et al. (1975) estimated that net panicle photosynthesis is about 5% of that of an active photosynthesizing rice leaf [25]; Takeda and Maruta (1956) estimated that panicle photosynthesis contributes 8% of the rice grain yield [26]. Reflecting the notion of rice ear photosynthesis being of little importance to grain filling, some researchers recommended lowering spike position in a canopy as much as possible to increase canopy photosynthesis by minimizing shading leaves by panicles [27].

Is panicle photosynthesis in rice really of little importance for grain filling? More and more evidences are suggesting the opposite. For example, Imaizumi et al. (1990) found that whole-panicle gross amount of photosynthetically assimilated CO₂ was 30% of that in a flag leaf, and spikelets had a similar photosynthetic capability to a flag leaf on the basis of Chl [8]. It has been shown that rice panicle contains higher amount of chloroplast NAD(P)H dehydrogenase-like complex, and maintains a similar or even higher quantum yield of PSII than that of the flag leaf from heading to harvest [28]. There are also evidences showing a potential important role of rice panicle photosynthesis in maintaining grain filling under moderate stress conditions. For example, under moderate wetting and drying irrigation after anthesis, genes that function in photosynthesis and carbohydrate metabolism were upregulated in inferior spikelets, together with a simultaneous increase of grain filling rate and grain weight of the inferior spikelets [29].

One major factor attributing to the inconsistency regarding the importance of photosynthesis of reproductive organs in grain filling is to the lack of robust methods to measure photosynthesis in organs with a complex three-dimensional geometry [12, 22, 23]. Many indirect methods have been developed to study photosynthesis of reproductive organs [23], e.g. through ear (awn) removal/shading or application of herbicides [17, 27, 30], isotope tracing [21], gas exchange measurement of excised spikelets [6, 8] and chlorophyll fluorescence measurement for plants cultured in pots [28]. Among these methods, the shading treatments or applications of herbicides may affect other function of treated organs and/or trigger compensation effects of unaffected organs [20, 22, 31-35]. The stable isotope discrimination method calculated contributions of different organs to grain yield based on instant carbon isotope in water-soluble fraction in different tissues at mid-grain filling stage and carbon isotope in mature grains [12, 21, 22]. The accuracy of this isotope discrimination method might be affected by the dynamic environment (e.g. temperature and air humidity) and physiological processes other than photosynthesis (e.g. assimilates transport and ear night respiration) during the whole grain filling season [12, 21, 22]. Gas exchange measurement on excised ears or spikelets is an intrusive method and it can cause bias due to an alteration of ear natural architecture and posture [27].

Thus, the main objectives of this study are: 1) to develop a direct and non-intrusive method for gas exchange measurement of rice panicles; 2) to explore panicle photosynthetic characteristics and the inter-cultivar variations of photosynthetic gas exchange parameters; 3) and to study the relationship between panicle photosynthetic gas exchange parameters and grain yield related agronomic traits.

Results

Performance of the panicle photosynthesis P-chamber

The P-chamber is connected to a gas analyzer Li-6400 (Li-Cor Inc., Lincoln, Nebraska, USA) to measure gas exchange rate of a rice panicle (Fig. 1a, b). The panicle is illuminated with LED light source (Fig. 1c, d). The LED light source emits pure white light (light wavelength information is shown in Fig. 1f). The photon flux density (PFD) is in a range of $\pm 10\%$ of set value on 87% of the total 30×5 cm light field (Fig. 1e). At a room temperature of $25\text{ }^{\circ}\text{C}$ with a flow rate of $700\text{ }\mu\text{mol s}^{-1}$, temperature rise in the chamber at PFD of $2000\text{ }\mu\text{mol (photons) m}^{-2}\text{ s}^{-1}$ is $4.3 \pm 0.7\text{ }^{\circ}\text{C}$.

To test the leakage of the P-chamber, we recorded $\Delta[\text{CO}_2]$ (the difference between CO_2 concentration in the P-chamber and the set reference CO_2 concentration in the reference pipe) at different time after closure of the P-chamber. With an initial $\Delta[\text{CO}_2]$ value of around $500\text{ }\mu\text{mol mol}^{-1}$, the $\Delta[\text{CO}_2]$ values were stabilized within 8 min under different flow rates and different CO_2 concentrations (Fig. 1g). The final $\Delta[\text{CO}_2]$ values were $\leq 1.6\text{ }\mu\text{mol mol}^{-1}$ and $\leq 1.3\text{ }\mu\text{mol mol}^{-1}$ at a flow rate of 500 and $700\text{ }\mu\text{mol s}^{-1}$, respectively (Fig. 1g). During a typical measurement of panicle photosynthetic rate and respiratory rate, the light source was turned on before the rice panicle was enclosed, and the photosynthetic rate was recorded after photosynthetic rate reading was stabilized; then the light source was turned off, and the

respiratory rate was recorded after respiratory rate reading was stabilized (Fig. 1h). A comprehensive measurement for photosynthetic and respiratory rates of a rice panicle usually took 10 ~ 20 min after the panicle was enclosed in the P-chamber, depended on size and photosynthetic activity of the panicle.

Quantification of rice panicle photosynthesis

We first used the P-chamber to measure the *in situ* whole-panicle photosynthetic gas exchange parameters 5 days after heading for different rice cultivars when the panicles were still erect or semi-erect. Specifically, whole-panicle net photosynthetic rate (panicle A_{net}) and dark respiratory rate (panicle R_{d}) were measured; and whole-panicle gross photosynthetic rate (panicle A_{gross}) was calculated as a sum of panicle A_{net} and panicle R_{d} . In addition, average spikelet A_{gross} , A_{net} and R_{d} were calculated by dividing panicle A_{gross} , A_{net} and R_{d} by the spikelet number of a panicle.

Notably, substantial inter-cultivar variations in photosynthetic gas exchange parameters among rice cultivars were found, for both values of a whole panicle and values normalized by spikelet number. For example, the spikelet A_{net} and panicle A_{net} were 0.012 ~ 0.051 nmol s⁻¹ and 2.7 ~ 12.0 nmol s⁻¹ in 2015, and were -0.014 ~ 0.075 nmol s⁻¹ and -3.3 ~ 15.0 nmol s⁻¹ in 2016, respectively (Table 1). Panicle A_{gross} varied from 17.1 to 53.6 nmol s⁻¹ under PFD of 2000 μmol (photons) m⁻² s⁻¹ for the seven rice cultivars (Table 1). Moreover, *indica* type rice cultivars (YLY900, CY1000, SY63 and 9311) had an overall significantly higher A_{gross} , A_{net} and R_{d} than *japonica* and *japonica-indica* hybrid type rice cultivars (XS134, YY538 and YY17), both on whole panicle basis and on single spikelet basis (Fig. 2).

In 2016, we further measured area of panicles and area of the corresponding flag leaves, and then quantified A_{gross} , A_{net} and R_{d} on an area basis by dividing panicle A_{gross} , A_{net} and R_{d} by the area of a panicle (Fig. S1; see detailed procedure used to calculate panicle surface area in Materials and Methods). As a result, we found spikelets contributed the majority of the panicle area, whereas branches only accounted for 24 ~ 33% of the panicle area; moreover, total area of a panicle (single side) was 72 ~ 121 cm², which was 20 ~ 124% higher than that of a flag leaf (Table 2). On an area basis, A_{net} of panicles were very low, i.e., only -0.6 ~ 2.2 $\mu\text{mol m}^{-2} \text{s}^{-1}$, under a PFD of 2000 μmol (photons) m⁻² s⁻¹; whereas A_{gross} were 2.1 ~ 7.2 $\mu\text{mol m}^{-2} \text{s}^{-1}$, much higher than A_{net} , as a result of high R_{d} , i.e., 2.1 ~ 6.1 $\mu\text{mol m}^{-2} \text{s}^{-1}$ (Table 3).

Given these different representations of gas exchange rates, i.e., on a panicle basis, on a spikelet basis and on an area basis, what's the relations between them? Positive correlations between values on different bases were found for all the three parameters A_{gross} , A_{net} and R_{d} (the Pearson correlation coefficient $r=0.59 \sim 0.99$; Fig. 3a-i). Interestingly, we found that A_{net} per panicle was strongly positively correlated with A_{net} per unit panicle area (the Pearson correlation coefficient $r=0.97$; Fig. 3e), in contrast to that for A_{gross} and R_{d} (the Pearson correlation coefficient $r=0.77$ and 0.59, respectively; Fig. 3b, h). In addition, spikelet based A_{gross} , A_{net} and R_{d} were closely related to area based A_{gross} , A_{net} and R_{d} (the

Pearson correlation coefficient $r=0.97-0.99$; Fig. 3c, f, i), as a result of a tight relationship between panicle area and spikelet number (the Pearson correlation coefficient $r=0.95$; Fig. 4).

Characterization of rice panicle photosynthesis

To further characterize rice panicle photosynthesis, we measured dynamic changes of panicle A_{gross} and R_d for seven rice cultivars during grain filling in 2016 (Fig. 5). Firstly, panicle A_{gross} and R_d were highest at early grain filling stage, then slowly decreased over time. Notably, both A_{gross} and R_d could still be detected even 40 days after heading, i.e., panicle photosynthesis and respiration were active throughout the grain-filling season. It is worth mentioning here that the panicle A_{gross} should be interpreted as the maximal photosynthetic potential, as the panicle was held upright during measurement, which was not the natural state after early grain filling stage for some cultivars.

We further measured the photosynthetic light and CO_2 response patterns of a panicle using rice cultivar YY17. Although the shapes of both response curves are similar to that of a leaf, there are several differences. Firstly, the apparent CO_2 and light compensation points of a panicle were about $180 \mu\text{mol mol}^{-1}$ and $390 \mu\text{mol m}^{-2} \text{s}^{-1}$ (Fig. 6), much higher than those of a leaf, which are usually $<100 \mu\text{mol mol}^{-1}$ and $<50 \mu\text{mol m}^{-2} \text{s}^{-1}$, respectively. Secondly, the panicle photosynthetic rate was still not saturated at very high air CO_2 concentration ($1800 \mu\text{mol mol}^{-1}$; Fig. 6a), which also differed from that of a typical leaf.

Correlation between panicle photosynthetic gas exchange parameters and grain yield related agronomic traits

Finally, we studied the relationship between panicle photosynthetic gas-exchange parameters measured 5 days after heading (Table 1) and six grain yield related agronomic traits collected at harvest across two years, i.e., spikelet number per panicle, panicle length, spikelet density (defined as spikelet number per centimeter of panicle length), 1000-grain weight, grain setting rate and panicle weight (Table 4). Firstly, there were positive correlations between different panicle photosynthetic gas-exchange parameters (Fig. 7). For example, significant positive correlations were found between A_{gross} and A_{net} of a whole panicle (Pearson correlation coefficient $r = 0.87$, p -value < 0.001), between A_{gross} and R_d of a whole panicle ($r = 0.94$, p -value < 0.0001), between A_{gross} and A_{net} per spikelet ($r = 0.93$, p -value < 0.0001), and between A_{gross} and R_d per spikelet ($r = 0.97$, p -value < 0.0001). In contrast, both positive and negative correlations were found between grain yield related agronomic traits, e.g. significant negative correlations were found between 1000-grain weight and spikelet number per panicle, but significant positive correlations were found between 1000-grain weight and grain setting rate (Fig. 7). Notably, strong positive correlations were found between A_{gross} , A_{net} , R_d per spikelet and grain setting rate, 1000-grain weight. In particular, the Pearson correlation coefficient r is 0.93 (p -value < 0.0001) between A_{gross} per spikelet and grain setting rate (Fig. 7). Concurrently, we found that A_{gross} , A_{net} and R_d of a whole panicle had a moderate positive correlation with panicle dry weight at harvest ($r = 0.39 \sim 0.55$); whereas A_{gross} , A_{net} and R_d per spikelet were not correlated with panicle dry weight at harvest ($r = -0.18 \sim 0.12$). In addition, all the six panicle

photosynthetic gas exchange parameters were found to be positively correlated with panicle length ($r = 0.6 \sim 0.85$), whereas they were negatively correlated with spikelet density ($r = -0.82 \sim -0.28$). Finally, no correlations were found between A_{gross} , A_{net} , R_d of a whole panicle and spikelet number per panicle ($r = -0.2 \sim 0.1$).

Discussion

Reproductive organs of cereal crops have been shown photosynthetically active, and photosynthesis in ear of wheat and barley has attracted great research interest [12, 13] [14, 15]. Current high-yield rice cultivars usually have much bigger ears with many spikelets compared with wheat or barley. For example, the newly bred rice cultivar YLY900, which have reached record-high yield of 15 tons ha^{-1} [36], and another even higher yielded rice cultivar CY1000, have an extra-large panicle size of exceeding 350 and 400 spikelets per panicle, respectively (Table 4). As a result, the green area of a rice panicle can even be larger than a flag leaf (Table 2). Furthermore, considering that the panicles locate at the upper part of a canopy, they can intercept a considerably amount of solar radiation. All these suggest a great role of panicle photosynthesis in rice grain filling. However, compared to leaf photosynthesis, panicle photosynthesis is much less studied, largely due to the methodological limitations [12, 22, 23]. Therefore, new tools and methods are needed to enable better characterization of rice panicle photosynthesis and provide an accurate evaluation of the impact of panicle photosynthesis on rice grain yield.

Here we reported the development of a new panicle photosynthesis P-chamber, which can be connected to a standard infrared gas analyzer to enable direct measurement of gas exchange parameters. Compared to previously reported cuvettes for rice or wheat ear gas exchange measurement [8, 17, 22, 37], the P-chamber has several new features. First, it is made air-tight with very limited leakage under different CO_2 concentrations and flow rates, which enables accurate and fast measurement of gas exchange rate. Secondly, it has a large cuvette volume for gas exchange measurement of a whole rice panicle *in situ*. The large cuvette volume enables measurement of panicles of irregular shape. We expect that it can also be used to measure inflorescence of Arabidopsis, wheat, rape and soybean and other plant organs such as stem, fruit, and branch *in situ*. Moreover, the chamber is opaque and the PFD inside is uniformly distributed and can be precisely controlled, which enables studying response of photosynthesis to PFD.

Using the P-chamber, we studied quantitative characteristics of rice panicle photosynthesis. First of all, to facilitate comparison between tissues, gas exchange rates are normalized. There are three commonly adopted approaches to normalize gas exchange rates of foliar and non-foliar organs, i.e., 1) normalization based on organ number, which is widely used for cereal ears [25, 38]; 2) normalization based on area, which is commonly used for leaves [37, 39]; and 3) normalization based on chlorophyll content, which is applicable for both foliar and non-foliar organs [8, 40]. For ears of cereal crops, it is labor-saving and technically easier to measure gas exchange rate on a whole ear basis or on a spikelet basis rather than on an area basis or chlorophyll basis. However, what's the relationship between gas exchange rates normalized on different bases? Here we show that gas exchange rate on a spikelet basis is highly correlated to gas exchange rate on an area basis (Pearson correlation coefficient $r=0.97-0.99$;

Fig. 3). This result indicates that we may use spikelet-based spike photosynthesis rate as a surrogate for the area-based spike photosynthesis rate, as the former one is non-intrusive and technically easier. For the seven rice cultivars used in this study, we have the following equation to convert spikelet number to panicle area (Fig. 4):

$$\text{Panicle area} = 0.266 * \text{Spikelet number} + 9.84 \text{ Eqn. 1}$$

Strikingly, A_{net} on a panicle basis is also highly correlated to A_{net} on an area basis for the seven rice cultivars used in this study (Pearson correlation coefficient $r=0.97$; Fig. 3). This result indicates that panicle A_{net} is largely determined by A_{net} per unit area rather than its spikelet number or green surface area.

Secondly, we found that as a result of high respiratory rate and large panicle area, A_{gross} of a whole panicle 5 days after heading could be as high as 17 to 54 nmol s^{-1} under a PFD of 2000 $\mu\text{mol (photons) m}^{-2} \text{s}^{-1}$. With an assumed typical photosynthetic rate of 25 $\mu\text{mol m}^{-2} \text{s}^{-1}$ under a PFD of 2000 $\mu\text{mol (photons) m}^{-2} \text{s}^{-1}$ [41-43], the A_{gross} of whole flag blades for the seven rice cultivars were 82 to 220 nmol s^{-1} . This estimation resulted in the ratio between panicle A_{gross} and flag leaf A_{gross} to be 20 - 38% (Table 5). This result is comparable to a previous report, where A_{gross} of a panicle is 30% of a flag leaf [8]. Concurrently, though panicle photosynthetic activity gradually decreased during grain filling, it still maintained CO_2 uptake activity even 40 days after heading, especially for the newly released super high yield hybrid rice cultivars, YLY900 and CY1000 (Fig. 5a). Maintaining photosynthetic activity of a panicle till the end of rice grain filling has been reported earlier as well [28, 40]. The contribution of panicle photosynthesis to rice grain yield cannot be precisely calculated in the current study due to the lack of the light environments for panicle throughout the grain filling season. However, the contribution of panicle photosynthesis to grain filling can be estimated based on measured rates of respiration. Specifically, if we assume a constant respiratory rate during the daytime, i.e., from 6:00 AM to 18:00 PM, the total daytime-respired CO_2 during the whole grain filling season for the seven cultivars is 21-40 mmol (an integration of the Tot. R_d in Fig. 2), which is equivalent to 0.6-1.2 g CH_2O and accounts for 12.1 - 17.0% of the panicle weight at harvest (Table 4). Namely, by refixation of daytime-respired CO_2 , panicle photosynthesis can apparently contribute 12 - 17% of the final grain yield. Considering that during most of the grain filling season, panicle maintains a positive carbon uptake (Fig. 5), we conclude that the apparent contribution of panicle photosynthesis to grain filling will be higher than 12%. Combination of the photosynthetic light response curve measured by P-chamber with an elaborate three-dimensional modeling of rice panicles and leaves [44], light environments within a canopy can be calculated and the daily and season-long photosynthetic contribution of a panicle can be precisely quantified.

Thirdly, there are substantial inter-cultivar variations in panicle photosynthetic and respiratory rates 5 days after heading, both on a whole panicle basis and on a spikelet basis (Table 1). *Indica* cultivars have ~402% and ~693% higher panicle net photosynthetic rates per panicle and per spikelet than *japonica* and *japonica-indica* hybrid cultivars, respectively; meanwhile, *indica* cultivars have ~68% and ~83% higher

panicle respiratory rates per panicle and per spikelet, respectively (Fig. 2). Overall, *indica* cultivars show ~94% and ~122% increase in panicle gross photosynthetic rates on the panicle basis and on the spikelet basis, respectively (Fig. 2). One of the factors contributing to these differences might be panicle morphology. For example, the high-yielding *japonica* rice cultivars usually have erect and densely-packed panicles [45], while the *indica* rice cultivars typically have drooping and loosely-packed panicles [46]. Indeed, highly negative correlations between spikelet density and respiratory rate per spikelet (Pearson correlation coefficient $r=-0.82$; Fig. 7), and between spikelet density and net photosynthetic rate per spikelet were found in this study (Pearson correlation coefficient $r=-0.66$; Fig. 7). The mechanisms by how spikelet density could influence both panicle photosynthesis and respiration are not clear. Inter-subspecies differences in photosynthetic apparatus and metabolism may also exist, which need to be further studied. Similar to this, there is a significant difference between leaf photosynthetic rates between *indica* and *japonica* cultivars, the molecular mechanism of which also awaits clarification [47, 48].

Fourthly, rice panicles show differences in photosynthetic response pattern to light intensities and CO₂ concentrations compared to the leaves. Specifically, rice panicle has high photosynthetic light compensation point, and has high apparent CO₂ compensation and saturation points. Consistent with our finding here in rice, wheat ear was also found to have higher CO₂ compensation and saturation points than that of a flag leaf, which might be due to its high respiratory rate and different stomatal conductance response pattern [37].

Correlation study between panicle photosynthetic gas exchange parameters and grain yield related agronomic traits further shows a significant role of panicle photosynthesis in rice grain filling. Specifically, we found spikelet gross photosynthetic rates at early grain filling stage were strongly and significantly correlated with grain setting rate ($r = 0.93$) and 1000-grain weight ($r = 0.84$; Fig. 7). These results indicate that activity of the spikelet as a source and as a sink may be closely linked. It is likely that the activity of the spikelet as a photosynthetic source promotes the activity of the grain as a sink. The detailed mechanism of this “source-sink” correlation is still not clear. Earlier studies on wheat offer a few possibilities: 1) panicle photosynthesis may supply oxygen to the hypoxic regions deep within the developing seed to increase its metabolic activity [49]; 2) the transpiration accompanying panicle photosynthesis not only supports the transfer of nutrients and signal molecules from xylem to spikelets, but also regulates panicle temperature to avoid over-heating [50, 51]; 3) panicle photosynthesis can generate assimilatory power, i.e., ATP and reduced ferredoxin, which are required to support glume nitrogen assimilation, a major source of nitrogen in wheat grain [52].

Finally, the method of directly measuring reproductive organ gas exchange rate has a range of potential applications. Firstly, it can help accurately simulate photosynthesis from non-foliar organs and evaluate their role in canopy photosynthesis based on three-dimensional canopy photosynthesis modeling [44, 53]. Secondly, as large natural variations of panicle photosynthetic and respiratory rates exist between cultivars, the genetic basis can be explored by measuring panicle gas exchange parameters in rice genetic populations. Thirdly, the diurnal and seasonal panicle respiratory rate change can be measured to

trace grain filling dynamics to support identification of new options to improve panicle grain filling patterns [43, 54].

Conclusions

Here we report the design and application of a new device, the P-chamber. The features of large cuvette volume, air tight, and programmable light intensity inside the P-chamber enable *in situ* characterization of photosynthetic gas exchange of a whole plant organ. With the P-chamber, for the first time, we characterized whole-panicle photosynthetic light and CO₂ response patterns, quantified variations of panicle photosynthesis between cultivars and identified correlations between panicle photosynthetic gas exchange parameters and agronomic traits in rice. Further application of the P-chamber will facilitate study on photosynthetic characteristics and contribution of irregular photosynthetic organs, especially the non-foliar photosynthetic organs, e.g. inflorescence of cereals and other plant organs such as stem and branch.

Materials And Methods

Plant materials and experimental design

Rice plants were grown in a paddy field at Songjiang breeding station of the Institutes of Plant Physiology and Ecology, Chinese Academy of Sciences, Shanghai, China (30°56' 44" N, 121°8' 1" E) in 2015 and 2016. Seeds were sown on seedbeds after germination on 1 June 2015 and 5 June 2016, and seedlings were transplanted to the field on 26 June 2015 and 29 June 2016. In both years, one plant was transplanted into a hill with a spacing of 20 cm between hills and 20 cm between rows. The cultivars were planted in plots. Plots were arranged in blocks with five replicates, each containing 56 (7 × 8) plants. Basal fertilizer was applied at a rate of 120 kg N ha⁻¹, 65 kg P ha⁻¹ and 65 kg K ha⁻¹. Additional N fertilizer was top-dressed at a rate of 80 kg N ha⁻¹ 3 weeks after transplanting. Weeds, pests and diseases were controlled following common agronomic practice in the region. In 2015, five rice cultivars were grown for experiments: two recently bred *indica* super high yield hybrid cultivars Y-Liang-You 900 (YLY900; <http://www.ricedata.cn/variety/varis/614537.htm>) and Chao-You 1000 (CY1000; <http://www.ricedata.cn/variety/varis/616642.htm>), one widely grown traditional *indica* hybrid cultivar Shan-You 63 (SY63; <http://www.ricedata.cn/variety/varis/601174.htm>), one *indica* inbred cultivar Yang-Dao 6 (9311; <http://www.ricedata.cn/variety/varis/600611.htm>) and one elite *japonica* inbred cultivar Xiu-Shui 134 (XS134; <http://www.ricedata.cn/variety/varis/603176.htm>). In 2016, in spite of the above mentioned 5 cultivars, two additional *japonica-indica* hybrid cultivars, Yong-You 538 (YY538; <http://www.ricedata.cn/variety/varis/613563.htm>) and Yong-You 17 (YY17; <http://www.ricedata.cn/variety/varis/612347.htm>), were grown for experiments. The heading date and harvest date of main stems for these rice cultivars in two years were given in Table S1.

Construction of the custom-designed panicle chamber

To enable direct and non-intrusive measurement of panicle photosynthesis, we built a panicle photosynthesis measurement chamber (P-chamber), which can be used together with infrared gas analyzers to measure photosynthetic and respiratory rates of a whole rice panicle. The chamber body is made of aluminum, with cooling fins and cooling fans on both sides (Fig. 1c). The chamber body has a dimension of 30 × 5 × 5 cm. The air inside the chamber was mixed with two air-mixing fans inside the chamber (F2008ES-05WAV, SHICOH IC FAN, Japan), light source and a PFD control system (Fig. 1c, d). The inner space is uniformly illuminated with 192 light-emitting diodes (LEDs; CREE XML T6, Guangdong Benbon Electrical Co., Ltd) on an 30 × 5 cm Aluminum substrate (Dongxing electronic technology Co., Ltd) for which the PFD can be precisely controlled to span from 0 to 2000 $\mu\text{mol (photons) m}^{-2} \text{s}^{-1}$ (Fig. 1d). The PFDs in the P-chamber under different input voltages on LEDs were measured with an irradiance sensor (LightScout Light Sensor Reader, LightScout, Spectrum Technologies, Inc. USA). During measurement, the P-chamber is connected to a standard Infra-red gas analyzer Li-6400, and the rice panicle is held upright and fully enclosed in the chamber (Fig. 1a, b). Flow rate and CO_2 concentration of influx air are controlled by LI-6400, and data are recorded and stored in LI-6400. Temperature inside the P-chamber was not controlled during measurement. The ambient air temperature for each measurement was recorded (see in Table S2 and Table S3).

Measurement of panicle photosynthetic gas exchange parameters

Twenty panicles were randomly selected and labeled at heading for each rice cultivar, which were used to measure photosynthetic gas exchange parameters and yield related agronomic traits. Three and five panicles among these twenty were used to measure whole-panicle net photosynthetic rate and dark respiratory rate in 2015 and 2016, respectively. During each measurement, the CO_2 concentration was set to 400 $\mu\text{mol mol}^{-1}$, the air flow rate was set to 700 $\mu\text{mol s}^{-1}$. For net photosynthetic rate measurement, the PFD was set to 2000 $\mu\text{mol (photons) m}^{-2} \text{s}^{-1}$; for dark respiratory rate measurement, the PFD was set to 0 $\mu\text{mol m}^{-2} \text{s}^{-1}$. Each time when the gas exchange rate readings were stabilized, data were logged after matching the two infrared gas analyzers (IRGAs).

The rice cultivar YY17 was used for photosynthetic light and CO_2 response curve measurements 10 days after heading in 2016. For photosynthetic light response curve measurement, the CO_2 concentration was set to 400 $\mu\text{mol mol}^{-1}$, the flow rate was set to 700 $\mu\text{mol s}^{-1}$; and the light levels were manually changed in the following sequence: 2000, 1500, 1000, 700, 500, 300, 200, 100, 50, 25 and 0 $\mu\text{mol (photons) m}^{-2} \text{s}^{-1}$ during measurement. Panicles were maintained in the cuvette at each light level for about 4 min before photosynthetic rate was stabilized and recorded. For photosynthetic CO_2 response curve measurement, the PFD was set to 2000 $\mu\text{mol (photons) m}^{-2} \text{s}^{-1}$, the flow rate was set to 700 $\mu\text{mol s}^{-1}$, and the reference CO_2 concentration was set to 400 $\mu\text{mol mol}^{-1}$. The panicles were maintained under the above described condition for 15 min. Then, the CO_2 concentrations were manually changed in the following sequence: 400, 300, 150, 100 and 50 $\mu\text{mol mol}^{-1}$ during measurement. Panicles were maintained in the cuvette at each CO_2 concentration for about 4 min before photosynthetic rate was stabilized and recorded. Next, the

CO₂ concentration was set back to 400 μmol m⁻² s⁻¹ and maintained at that concentration for 15 min. Finally, the CO₂ concentrations were manually changed in the following sequence: 600, 900, 1200, 1500 and 1800 μmol mol⁻¹. Again, panicles were maintained in the cuvette at each CO₂ concentration for about 4 min before photosynthetic rate was stabilized and recorded.

Calculation of panicle area and flag area

Areas of spikelets and branches in a panicle were calculated to obtain the total area of a panicle. Specifically, the non-degraded green spikelets were firstly detached from the panicle 5 days after heading; then, these spikelets were scanned with an Epson Perfection V300 Photo scanner (Epson, Tokyo, Japan; Fig. S1a). The areas of spikelets in the scanned images were obtained by image processing using ImageJ (US National Institutes of Health, Bethesda, USA). The flag leaf area was obtained in the same way (Fig. S1c).

The spikelet-less panicles were then spread out, scanned with an Epson Perfection V300 Photo scanner as well (Epson, Tokyo, Japan; Fig. S1b). The projected area of branches was obtained through image processing using ImageJ (US National Institutes of Health, Bethesda, USA). Branches were assumed as cylinders, and single-sided surface area of all branches on a panicle S was then calculated as:

$$S = s_{bp} * \pi / 2 \text{ Eqn. 1}$$

in which s_{bp} is the projected area of branches. Single-sided surface area of a whole panicle was calculated as a sum of the area of branches and the area of all spikelets.

Agronomic traits measurement

Five of the twenty labeled panicles were sampled and stored in envelopes individually for each cultivar at harvest. The panicle length was measured, and then the panicles were put in an oven for drying. The temperature of the oven was maintained at 110 °C for 1 hour, and then switched to 70 °C and maintained for 3 days until samples were completely dried. Dry weight of each panicle was measured with an electronic balance AL104 (JB / T, METTLER TOLEDO, USA). The panicles were then threshed to determine the total spikelet number and filled grain number. The weight of all filled grains were measured to determine the 1000-grain weight.

Data Analysis

Statistical analyses were carried out using R-project (version 3.6.2). One-way analysis of variance (ANOVA) and the Scheffe post-hoc test were used to compare data at a level of 5%. The Pearson correlation coefficients between panicle photosynthetic gas exchange parameters and grain yield related agronomic traits were calculated using the R package (Corrplot; version 0.84).

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Availability of data and material

All data generated or analyzed during this study are included in the article and in Additional file 1.

Competing interests

The authors declare that they have no conflict of interest.

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Authors' contributions

XGZ and TGC conceived the research plan; TGC and HLZ managed the experiments; TGC analyzed the data; TGC and XGZ drafted the article; QFS, CX, SC and MQ provided technical assistance and constructive suggestions for the P-chamber building, field experiment and manuscript improvement; all authors discussed the results and contributed to the final manuscript.

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Abbreviations

9311, an *indica* inbred cultivar Yang-Dao 6; A_{gross} per unit area, gross panicle photosynthetic rate on an area basis ($\mu\text{mol m}^{-2} \text{s}^{-1}$); A_{net} per unit area, net panicle photosynthetic rate on an area basis ($\mu\text{mol m}^{-2} \text{s}^{-1}$); C_a , ambient air CO_2 concentration ($\mu\text{mol mol}^{-1}$); CY1000, an *indica* super high yield hybrid rice cultivar Chao-You 1000; Panicle A_{net} , net photosynthetic rate of a whole panicle (nmol s^{-1}); Panicle A_{gross} , gross photosynthetic rate of a whole panicle (nmol s^{-1}); Panicle R_p , respiratory rate of a whole panicle (nmol s^{-1}); P-chamber, the custom-built chamber for panicle photosynthetic gas exchange rate measurements; PFD, photon flux density ($\mu\text{mol m}^{-2} \text{s}^{-1}$); R_d per unit area, panicle respiratory rate on an area basis ($\mu\text{mol m}^{-2} \text{s}^{-1}$); Spikelet A_{net} , net panicle photosynthetic rate on a spikelet basis (nmol s^{-1}); Spikelet A_{gross} , gross

panicle photosynthetic rate on a spikelet basis (nmol s^{-1}); Spikelet R_d , panicle respiratory rate on a spikelet basis (nmol s^{-1}); SY 63, a traditional *indica* hybrid cultivar Shan-You 63; XS134, an elite *japonica* inbred rice cultivar Xiu-Shui 134; YLY900, an *indica* super high yield hybrid rice cultivar Y-Liang-You 900; YY17, a *japonica-indica* hybrid rice cultivar Yong-You 17; YY538, a *japonica-indica* hybrid rice cultivar Yong-You 538.

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Tables

Table 1. Panicle photosynthetic gas exchange parameters of the rice cultivars measured 5 days after heading. Data presented are mean values with s.d. (n = 3 in year 2015; n = 5 in year 2016). Data followed by different letters are statistically different at *p*-value < 0.05.

Year	Cultivar	Panicle A_{net} (nmol s^{-1})	Panicle A_{gross} (nmol s^{-1})	Panicle R_d (nmol s^{-1})	Spikelet A_{net} (nmol s^{-1})	Spikelet A_{gross} (nmol s^{-1})	Spikelet R_d (nmol s^{-1})
2015	YLY900	12±2.4a	49±2.8a	37±1.5a	0.035±0.003a	0.147±0.011b	0.111±0.013a
	CY1000	7.2±1.3b	43.6±7a	36.4±7.3a	0.015±0.004b	0.091±0.005c	0.075±0.005b
	SY63	11.8±2.2a	42.8±2.7a	31±2.8a	0.051±0.012a	0.185±0.014a	0.133±0.007a
	9311	10.9±1.4ab	46±3.4a	35±3.9a	0.044±0.009a	0.182±0.007ab	0.138±0.003a
	XS134	2.7±1.1c	18.1±3.4b	15.4±2.5b	0.012±0.005b	0.078±0.022c	0.067±0.017b
2016	YLY900	7.3±2.4b	47.5±4.1a	40.2±5.6a	0.021±0.007bc	0.136±0.011b	0.115±0.014b
	CY1000	6.9±2.2b	44.3±4ab	37.4±4.6ab	0.017±0.006bc	0.106±0.013bc	0.089±0.01bc
	SY63	15±3a	49.6±4.3a	34.6±5.3ab	0.075±0.016a	0.248±0.021a	0.172±0.023a
	9311	11.9±2.6ab	53.6±3.9a	45.5±4.5a	0.052±0.009a	0.243±0.005a	0.205±0.021a
	XS134	-3.3±3.4c	17.1±4.4d	20.3±6.4c	-0.014±0.014d	0.075±0.018cd	0.089±0.026bc
	YY538	-0.4±2.4c	25.8±4.7cd	26.2±4.1bc	-0.001±0.006cd	0.06±0.009d	0.062±0.011c
	YY17	9.3±3.1ab	36±5.8bc	26.8±5.6bc	0.024±0.009b	0.091±0.01cd	0.068±0.01c

Table 2. The area of a flag leaf and the areas of all spikelets and branches in a panicle for different rice cultivars 5 days after heading (n=5). Data measured in 2016. Data presented are mean values with s.d. (n = 5). Data followed by different letters are statistically different at p -value < 0.05. Note that areas presented are single side based.

Name	Spikelets area (cm^2)	Branches area (cm^2)	Panicle area (cm^2)	Branches : Panicle area ratio	Flag leaf area (cm^2)
YLY900	79.7±6.5a	34.6±2.8a	115.5±11.7ab	0.3±0.01b	73.8±11b
CY1000	74.7±7.9a	28.4±1.4b	120.7±13a	0.28±0.02cd	88.1±11.3a
SY63	47±4.1bc	23±1.9cd	74.3±7.2d	0.33±0a	61.9±10.9cd
9311	54.2±4.2b	21.2±1.6d	71.6±6.8d	0.28±0bc	56.2±12.1de
XS134	39±3.5c	12.6±1.4e	56.3±8.1e	0.24±0.03d	32.6±4.4f
YY538	69.5±3.7a	26.8±1.6bc	109±9.1bc	0.28±0.02bcd	48.7±8e
YY17	72.5±6.7a	25.6±3.1bcd	106.3±8.7c	0.26±0.01cd	72.3±8.4bc

Table 3. Panicle photosynthetic gas exchange parameters on an area basis for different rice cultivars 5 days after heading (n=5). Data were measured in 2016. Data presented are mean values with s.d. (n = 5). Data followed by different letters are statistically different at p -value < 0.05.

Name	A_{net} per unit area ($\mu\text{mol m}^{-2} \text{s}^{-1}$)	A_{gross} per unit area ($\mu\text{mol m}^{-2} \text{s}^{-1}$)	R_d per unit area ($\mu\text{mol m}^{-2} \text{s}^{-1}$)
YLY900	0.7±0.2cd	4.2±0.4b	3.6±0.4bc
CY1000	0.6±0.2cd	3.6±0.4bc	3±0.3bcd
SY63	2.2±0.5a	7.2±0.6a	5±0.7ab
9311	1.6±0.3ab	7.2±0.1a	6.1±0.6a
XS134	-0.6±0.6e	3±0.7cd	3.6±1.1bc
YY538	0±0.2de	2.1±0.3d	2.1±0.4d
YY17	0.9±0.3bc	3.5±0.4bc	2.6±0.4cd

Table 4. Grain yield related agronomic traits of the rice cultivars at harvest. Data presented are mean values with s.d. (n = 5). Data followed by different letters are statistically different at p -value < 0.05.

Year	Cultivar	Spikelet number	Panicle length (cm)	Spikelet density	1000-grain weight (g)	Grain setting rate	Panicle weight (g)
2015	YLY900	356±24b	28.1±1.5a	13.1±1.4b	22.9±0.7d	0.84±0.07bc	7.4±1b
	CY1000	444±30a	21.8±1.1c	21±2.4a	21.6±0.7d	0.8±0.05c	8.7±0.5a
	SY63	231±21c	27.6±0.9a	8.4±0.6c	28.2±1.3b	0.92±0.04ab	6.4±0.6bc
	9311	234±40c	24.5±1b	9.5±1.3c	30.7±0.4a	0.97±0.02a	7.2±1.1b
	XS134	253±14c	17.1±0.7d	14.8±0.9b	25.2±0.8c	0.74±0.08c	5.7±0.4c
2016	YLY900	359±36b	32.7±0.6a	10.9±0.9d	22.2±4c	0.83±0.09bc	7±0.9ab
	CY1000	410±44a	26.5±1c	13.8±2.3bc	21.8±0.5c	0.81±0.03c	8.1±0.5a
	SY63	216±21c	30.5±1.2b	6.7±0.5e	29.6±1.1ab	0.96±0.01ab	6.4±0.4b
	9311	211±20c	27.2±1.3c	8.2±0.3e	30.8±1.6a	0.96±0.02a	6.3±0.7bc
	XS134	222±32c	18.2±1.5e	11.2±1.1cd	24.1±0.3bc	0.79±0.05c	4.7±0.3c
	YY538	420±35a	21.6±0.9d	17.2±0.7a	20.5±1.3c	0.77±0.06c	6.9±0.7ab
	YY17	403±33a	23.9±1.1d	15.5±0.9ab	22.3±1c	0.84±0.08abc	7.4±0.6ab

Table 5. Estimated ratios of light saturated gross photosynthetic rates between a panicle and a flag leaf 5 days after heading in seven rice cultivars. Data measured in 2016 (n=5).

Name	A_{gross} of a flag leaf (nmol s^{-1})	A_{gross} of a panicle (nmol s^{-1})	P/FL A_{gross}
YLY900	185	48	0.26
CY1000	220	44	0.2
SY63	155	50	0.32
9311	141	54	0.38
XS134	82	17	0.21
YY538	122	26	0.21
YY17	181	36	0.2

Notes: A_{gross} of a flag leaf, estimated light saturated gross photosynthetic rate of a whole flag leaf by assuming a constant light saturated photosynthetic rate of $25 \mu\text{mol m}^{-2} \text{s}^{-1}$ (the leaf area data was taken from Table 2); A_{gross} of a panicle, light saturated gross photosynthetic rate of a whole panicle (data was taken from Table 1); P/FL A_{gross} , the ratio between light saturated gross photosynthetic rate of a panicle and that of a flag leaf.

Figures

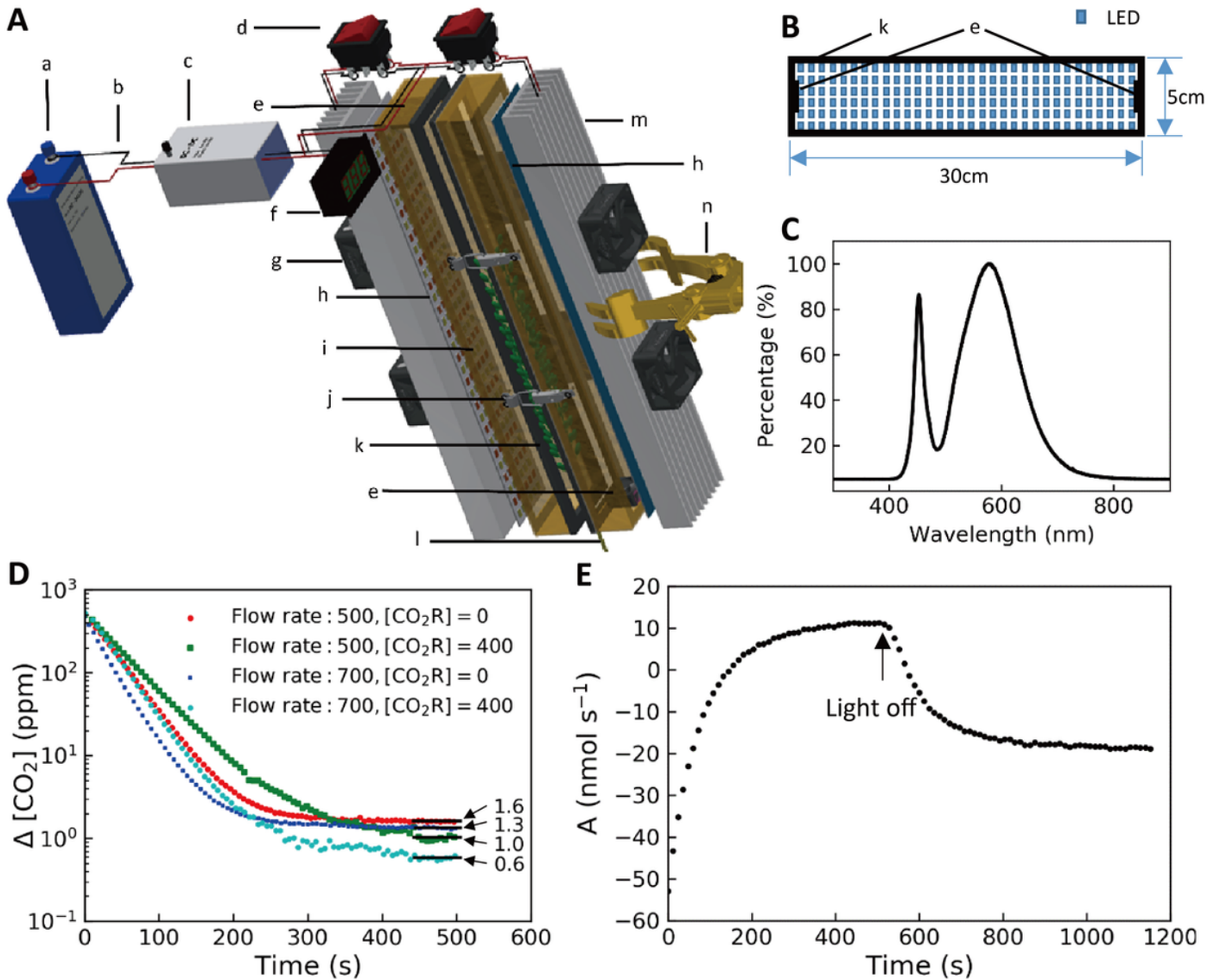


Figure 1.

Figure 1

A custom-built panicle photosynthesis chamber (P-chamber) for panicle gas exchange measurements. a, An illustration of how the P-chamber works together with the gas analyzer Li-6400 to measure the photosynthetic and respiratory rates of a rice panicle. b, The setting of a P-chamber in a paddy field. c, An

illustration of design of the P-chamber. d, A top-down view of the inner part of the P-chamber. e, The homogeneity of PFD in the chamber. f, Spectrum of the LED light source. g, The difference between the CO₂ concentration in the P-chamber and the set CO₂ concentration ([CO₂R]) in the reference pipe at different time after closure of the P-chamber. h, Recorded photosynthetic gas exchange rate of a rice panicle under a photon flux density (PFD) of 2000 μmol (photons) m⁻² s⁻¹ followed by dark treatment. Letters in panel (a) and (b): a. Battery; b. Wires; c. Voltage adaptor; d. Switch; e. Mixing fan (inside the chamber body); f. Digital displayed voltmeter; g. Cooling fan; h. LED array; i. Chamber body; j. Hasp; k. Sealing strip; l. Rice panicle; m. Cooling fin; n. Clamp. The horizontal lines in panel (g) represent mean values of the last 10 data points. The y-axis in panel (g) is log-scaled.

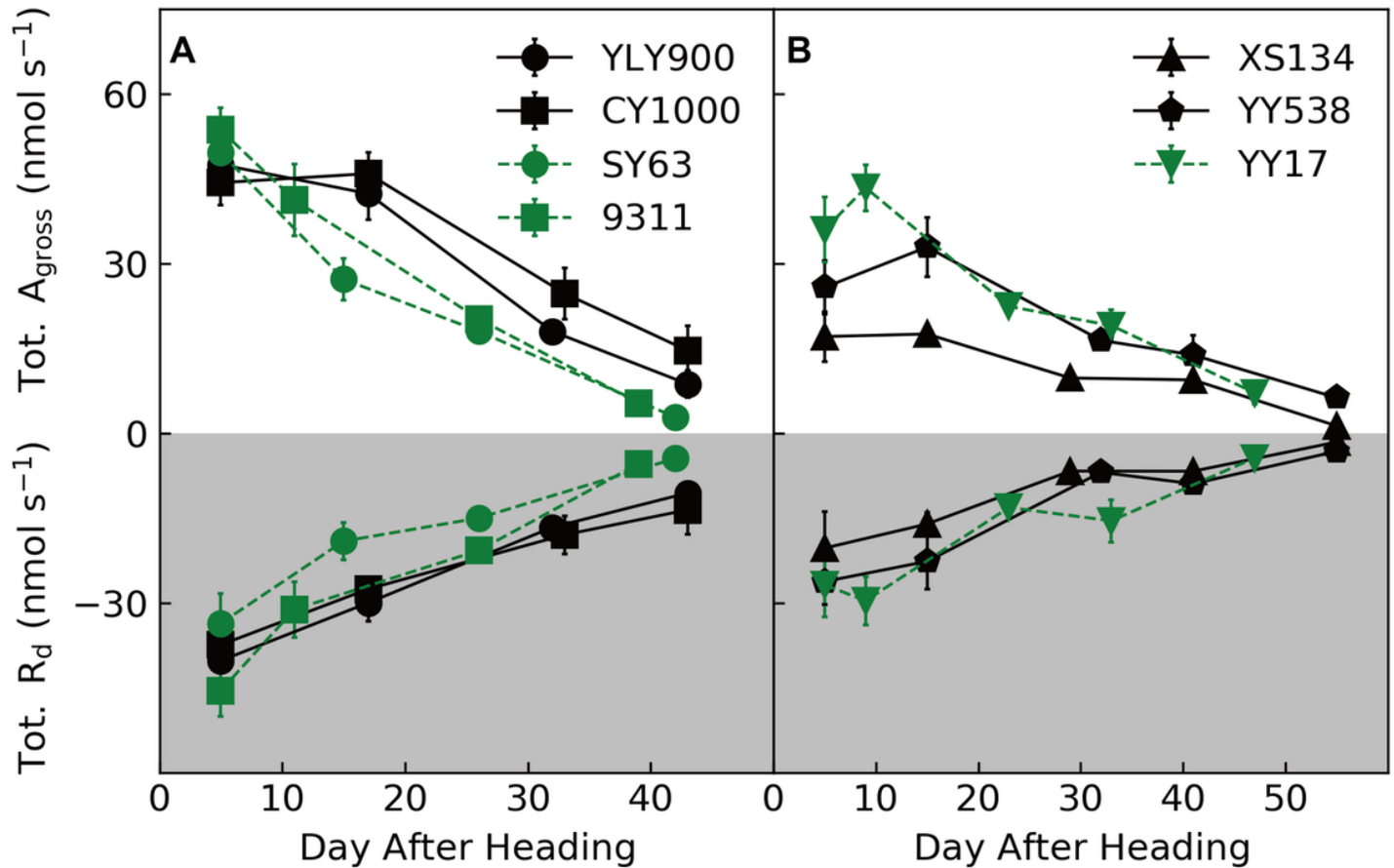


Figure 2.

Figure 2

Comparison of the gas exchange rates between indica and japonica rice cultivars 5 days after heading. a, Gross photosynthetic, net photosynthetic and respiratory rates of a whole rice panicle. b, Average gross photosynthetic, net photosynthetic and respiratory rates of a spikelet. Data of indica group were combined from rice cultivars YLY900, CY1000, SY63 and 9311; Data of japonica group were combined from rice cultivars XS134, YY538 and YY17. Data were combined from 2015 and 2016 experiments. Statistically different p-values were calculated from Student's t-test.

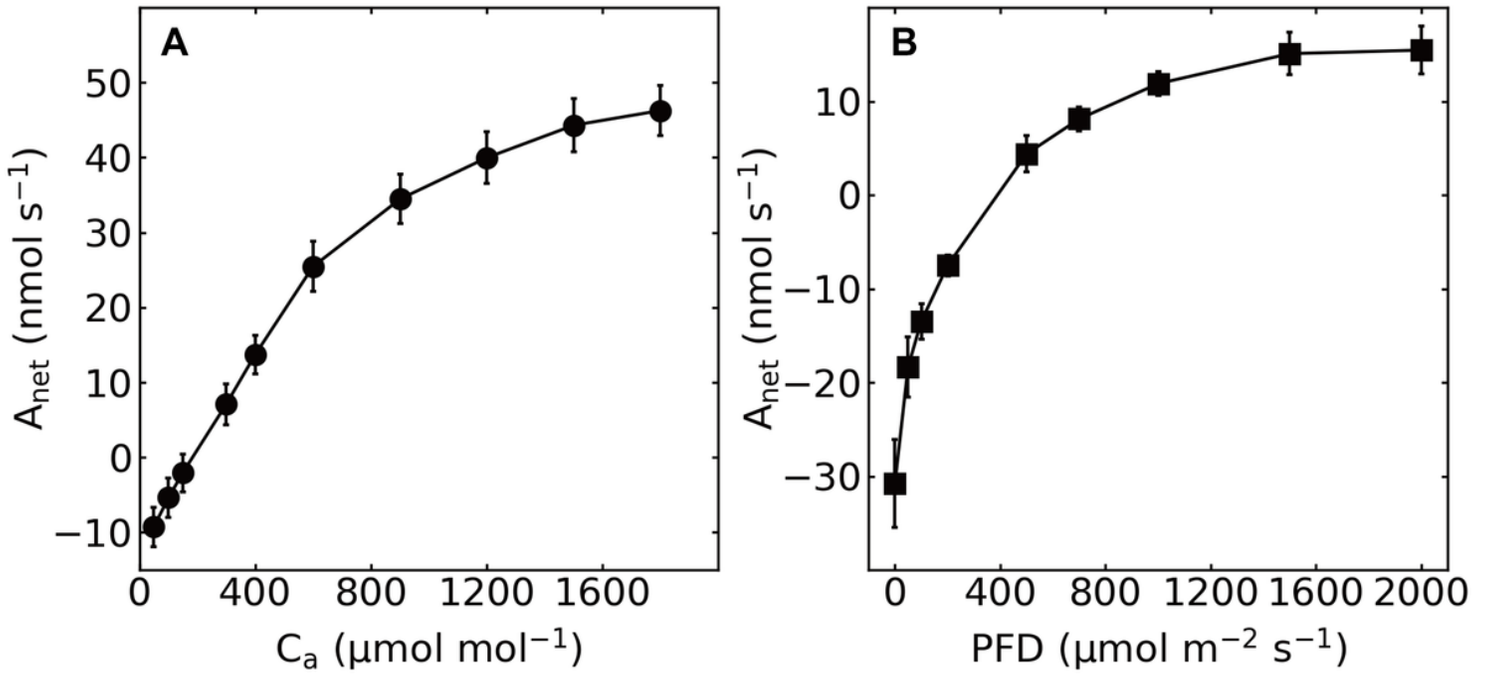


Figure 3.

Figure 3

Correlations between panicle photosynthetic gas exchange parameters normalized on a panicle basis, on a spikelet basis and on an area basis. a-c, Correlations between A_{gross} of panicles normalized on different bases. d-f, Correlations between A_{net} of panicles normalized on different bases. g-i, Correlations between R_d of panicles normalized on different bases. Data measured in 2016 ($n = 35$).

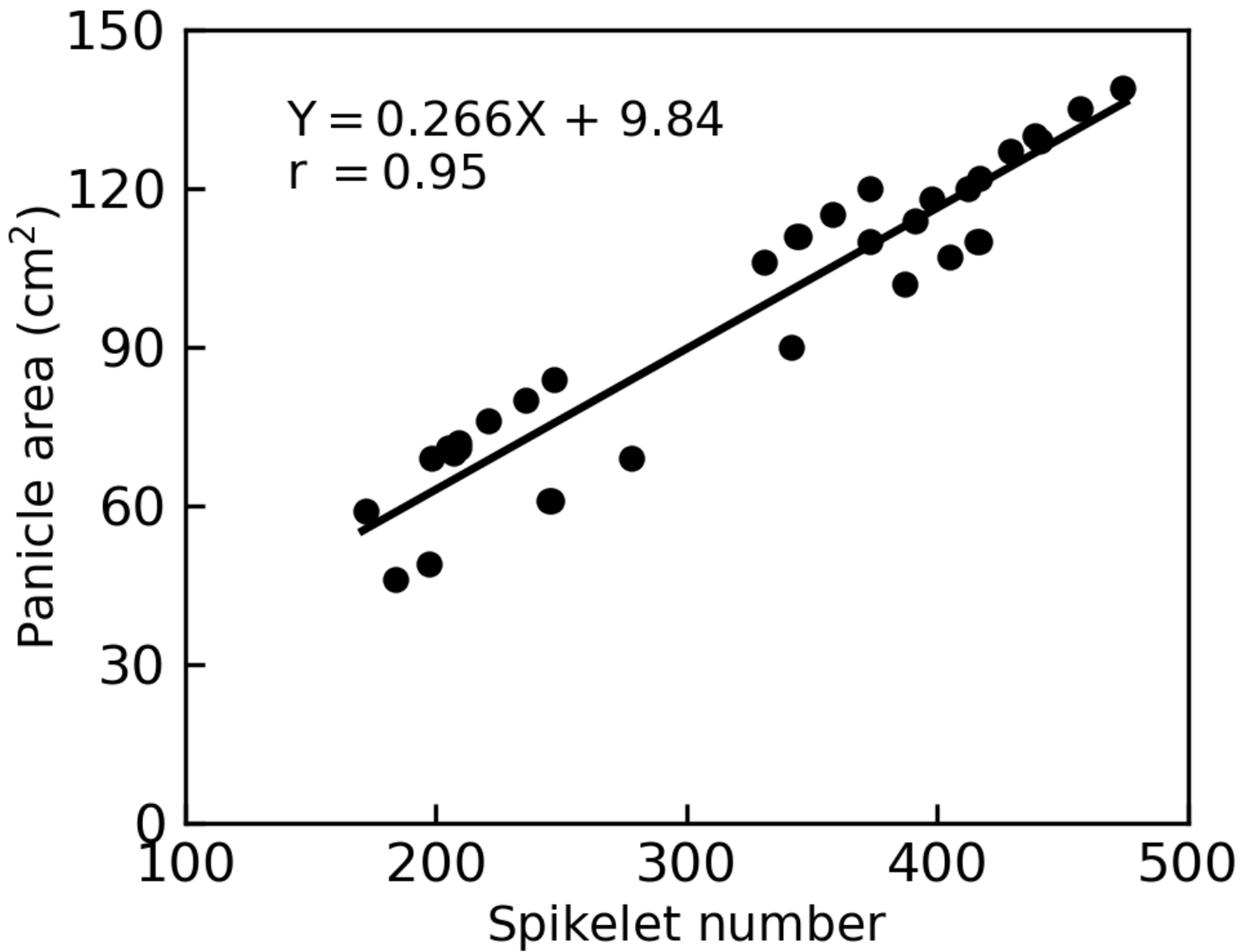


Figure 4

Correlation between panicle area and spikelet number. Data measured in 2016 (n = 35).

Supplementary Files

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- [supplementalinformation.docx](#)