

SI ADVANCES IN PHOTOSYNTHESIS

On the road to C₄ rice: advances and perspectivesMaria Ermakova[†] , Florence R. Danila[†] , Robert T. Furbank  and Susanne von Caemmerer* *Australian Research Council Centre of Excellence for Translational Photosynthesis, Division of Plant Science, Research School of Biology, The Australian National University, Acton, ACT 2601, Australia*

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SUMMARY

The international C₄ rice consortium aims to introduce into rice a high capacity photosynthetic mechanism, the C₄ pathway, to increase yield. The C₄ pathway is characterised by a complex combination of biochemical and anatomical specialisation that ensures high CO₂ partial pressure at RuBisCO sites in bundle sheath (BS) cells. Here we report an update of the progress of the C₄ rice project. Since its inception in 2008 there has been an exponential growth in synthetic biology and molecular tools. Golden Gate cloning and synthetic promoter systems have facilitated gene building block approaches allowing multiple enzymes and metabolite transporters to be assembled and expressed from single gene constructs. Photosynthetic functionalisation of the BS in rice remains an important step and there has been some success overexpressing transcription factors in the cytokinin signalling network which influence chloroplast volume. The C₄ rice project has rejuvenated the research interest in C₄ photosynthesis. Comparative anatomical studies now point to critical features essential for the design. So far little attention has been paid to the energetics. C₄ photosynthesis has a greater ATP requirement, which is met by increased cyclic electron transport in BS cells. We hypothesise that changes in energy status may drive this increased capacity for cyclic electron flow without the need for further modification. Although increasing vein density will ultimately be necessary for high efficiency C₄ rice, our modelling shows that small amounts of C₄ photosynthesis introduced around existing veins could already provide benefits of increased photosynthesis on the road to C₄ rice.

Keywords: C₄ photosynthesis, rice, metabolic engineering, bundle sheath cells, plasmodesmata, photosynthetic electron transfer.

INTRODUCTION

The C₄ photosynthetic pathway is characterised by a complex combination of biochemical and anatomical specialisation, which provides an elevation of the CO₂ partial pressure at the site of ribulose biphosphate carboxylase oxygenase (RuBisCO) in leaf bundle sheath (BS) cells. CO₂ is initially assimilated into C₄ acids by phosphoenolpyruvate (PEP) carboxylase in mesophyll (M) cells. These acids then diffuse to and are decarboxylated in BS cells in which CO₂ is concentrated. This spatial separation of initial CO₂ fixation in M cells and subsequent refixation by RuBisCO in BS cells is mostly associated with Kranz anatomy, but can also occur in single cells (Bowes and Salvucci, 1984; Voznesenskaya *et al.*, 2001). C₄ photosynthesis has evolved independently more than 60 times, providing one of the

most widespread and effective solutions for overcoming the catalytic inefficiency of RuBisCO (Sage *et al.*, 2012; Christin and Osborne, 2013). Because of the superior nitrogen and water use efficiency of C₄ plants, transfer of C₄ photosynthetic traits has been an early strategy for improving C₃ photosynthesis (Sage, 2004; Miyao *et al.*, 2011). Initial attempts were made to generate hybrids between closely related C₃ and C₄ plants by conventional crossing (reviewed in Brown and Bouton, 1993). With the advent of transformation technology there were numerous attempts to introduce C₄ photosynthetic genes into C₃ species (reviewed in Matsuoka *et al.*, 2001; Hausler *et al.*, 2002; Miyao, 2003). These all sought to insert a C₄ like pathway into M cells of C₃ species following the blue print of the aquatic

single cell organism *Hydrilla verticillata* (Bowes and Salvucci, 1984; von Caemmerer *et al.*, 2014). Four enzymes of the C₄ photosynthetic pathway were successfully introduced into rice by Miyao and collaborators and although no photosynthetic gains were observed, Miyao *et al.* (2011) have elegantly summarised the extensive contribution this endeavour made to our understanding of gene expression and regulation of C₄ genes expressed in rice. Current progress on building a two cell C₄ pathway in rice are built on these earlier insights.

The current C₄ rice project (<https://c4rice.com/>) which aims to introduce Kranz anatomy into rice was first conceived by John Sheehy (Mitchell and Sheehy, 2006) who invited a group of C₄ photosynthesis experts to the international rice research institute (IRRI) in the Philippines to discuss the potential of C₄ rice. Engineering the C₄ pathway into a C₃ plant requires manipulation of both anatomical and biochemical traits and progress of this research has been reviewed a number of times (Hibberd *et al.*, 2008; Hibberd and Covshoff, 2010; Langdale, 2011; Sedelnikova *et al.*, 2018). To introduce Kranz anatomy into rice requires a change of vein spacing patterns so that veins are closer together in the leaf and BS cells need to be 'functionalised' for increased photosynthetic capacity, including increased chloroplast content. At this point while there are established candidates for the genes and transcription factors that potentially control vein spacing, the complete transcriptional network remains to be elucidated (for review see Sedelnikova *et al.*, 2018). However, some progress has been made in photosynthetic functionalisation of the rice BS (Wang *et al.*, 2017b) While insertion of C₄ biochemistry in rice faces challenges of engineering high level and cell-specific gene expression (Hibberd and Covshoff, 2010), genes encoding the C₄ pathway enzymes and most of the metabolite transporters have now been identified. Here we review current progress and how they have been enabled by technological advances in cloning techniques and highlight future challenges.

BUILDING THE BIOCHEMISTRY FOR C₄ RICE

In 2008, the year the C₄ rice consortium commenced, a remarkable amount of information in regard to the genes encoding the key proteins in the C₄ pathway was already known. The cDNA sequences for maize phosphoenolpyruvate carboxylase (PEPC), pyruvate orthophosphate dikinase (PPDK), and NADP-malate dehydrogenase (MDH) had all been reported and expressed in the M of cells of rice (reviewed in Miyao *et al.*, 2011). The cDNA sequence for NADP-malic enzyme (NADP-ME) had also been reported for both the rice endogenous gene and the maize gene (Drincovich *et al.*, 2001). The genes encoding carbonic anhydrase (CA) in maize had also been cloned by this time, but the precise identity of the gene product located in the cytosol of the M cells in C₄ leaves was not definitively

proven (reviewed in DiMario *et al.*, 2016). Therefore, the key genes encoding the entire biochemical pathway of the NADP-ME type C₄ mechanism as shown in Figure 1 were all available at the commencement of the C₄ rice project.

What set the current C₄ rice project apart from previous activities was the desire to engineer a full Kranz two cell-type mechanism in rice (von Caemmerer *et al.*, 2012). This approach requires not only the cDNA sequences for the relevant photosynthetic proteins but high level expression with suitable promoters in the appropriate cell type of rice. For M cell-specific expression, the promoters of genes from C₄ species encoding PEPC, PPDK, and aspartate aminotransferase (AspAT) had all been tested in C₃ species and lead to M-specific accumulation of the β glucuronidase (GUS) reporter protein (Hibberd and Covshoff, 2010). The PEPC promoter from maize had been the most extensively tested in rice and various truncated versions have been shown to produce M-specific expression of the reporter gene GUS in rice (Hibberd and Covshoff, 2010).

High level BS cell-specific expression of proteins such as NADP-ME has proven to be the greatest challenge for establishing a C₄ metabolic pathway in rice. While there are potential anatomical constraints in regard to the photosynthetic competence of the rice BS compartment (Wang *et al.*, 2017b), the paucity of promoters available to drive BS expression in a C₃ plant has been a major obstacle. At the commencement of the C₄ rice project, the promoters of the genes encoding phosphoenolpyruvate carboxykinase (PEPCK), NADP-ME, AspAT, small subunit (Engelmann *et al.*) of RuBisCO, and the P subunit of glycine decarboxylase (GDCP) from C₄ species had been tested in C₃ plants (Chen *et al.*, 2001; Nomura *et al.*, 2005a; Nomura *et al.*, 2005b; Engelmann *et al.*, 2008). Of these, a version of the *Zoysia japonica* PEPCK promoter (Nomura *et al.*, 2005a) resulted in BS-specific GUS accumulation in rice. The *Flaveria trinervia* GDCP promoter was shown to be BS/vascular-specific in Arabidopsis and the promoter of this gene from *Flaveria anomala* (Chen *et al.*, 2001) was vascular-specific in rice. Both the PEPCK and GDCP promoters have subsequently been utilised in the C₄ rice strategy although promoter strength has been an ongoing issue for engineering efforts and it is notable that in the single cell C₄ project (reviewed in Miyao *et al.*, 2011), genomic clones often but not always gave superior expression levels compared with cDNAs driven by their own promoters.

Stacking of genes in transgenic rice in the initial phases of this project required the crossing of homozygous lines harbouring single gene constructs to build a rice plant expressing a complete set of the genes encoding the major enzymes of the C₄ pathway. Such a crossing strategy was a hugely time consuming effort as expression levels and cell-specific expression of the recombinant protein must be checked for each line and a crossing donor identified for each transgene. With segregation of the transgenes

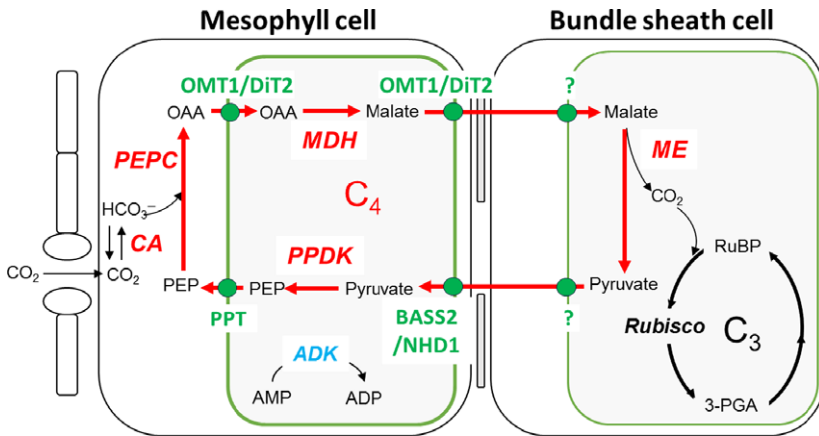


Figure 1. Enzymes and transporters included in the construction of C_4 rice. These include carbonic anhydrase (CA), PEP carboxylase (PEPC), malate dehydrogenase (MDH), NADP-malic enzyme (ME), and pyruvate Pi dikinase (PPDK). Adenylate kinase (ADK) has been added to ensure AMP is converted to ADP. The required transporters are oxoglutarate/malate transporter (OMT1), PEP/phosphate translocator (PPT), dicarboxylate transporter (DiT2), pyruvate/sodium symporter (BASS2), and sodium/proton antiporter (NHD1). Malate importer and pyruvate exporter in bundle sheath chloroplasts are not yet confirmed. Other abbreviations used are OAA for oxaloacetate and PEP for phosphoenolpyruvate.

inserted at different loci in the stacked lines, many hundreds of individuals had to be genotyped at each cross to obtain a single line harbouring the genes encoding the five key photosynthetic enzymes shown in Figure 1. Indeed, this crossing strategy has taken almost 6 years to achieve in *indica* rice in the current project. However, five genes are not enough for creating C_4 rice and the biochemical pathway is being complemented by a suite of membrane transporters ensuring fast transport of metabolites between cell compartments. Most of the transporters have now been identified and are listed in Figure 1, however there is still some uncertainty about malate import to the BS chloroplast and the export of pyruvate following malate decarboxylation. A recent review of transporters was given by Schuler *et al.* (2016). Physiological gas exchange techniques exist to allow for the identification of a complete, functioning C_4 pathway. These include measurements of CO_2 compensation points, reduced oxygen sensitivity of CO_2 assimilation and reduced carbon isotope discrimination (Furbank *et al.*, 2009). In the meantime new ^{13}C pulse chase labelling techniques have been developed to analyse the paths of carbon during C_3 and C_4 photosynthesis (Arrivault *et al.*, 2009) and confirmed that malate is in fact being formed in our current five gene rice transgenics.

SYNTHETIC BIOLOGY ACCELERATES THE PACE

A consensus definition drafted by a group of European experts more than a decade ago defined synthetic biology as follows: 'Synthetic biology is the engineering of biology: the synthesis of complex, biologically based (or inspired) systems, which display functions that do not exist in nature. This engineering perspective may be applied at all levels of the hierarchy of biological structures—from individual molecules to whole cells, tissues and organisms. In essence, synthetic biology will enable the design of 'biological systems' in a rational and systematic way' (Serrano, 2007). Synthetic biology has evolved and adopted many of the commonly used terms in mainstream engineering such as 'switch', 'rewire', 'design, test and redesign cycle'. Although in the

creation of C_4 rice, in which a template or design already existing in nature is being used, the installation of up to 20 genes to completely 'rewire' rice metabolism and anatomy surely fits the definition above. A major limitation in the synthetic biology approach however is the cycle time for the design, test and redesign in crops such as rice.

The last 5 years have seen an exponential growth in synthetic biology tools and the cost of gene synthesis has plummeted. This has enabled the C_4 rice consortium to adopt a more rapid cycle of design, test and prototype coupled to the adoption of a rapid *Agrobacterium*-based rice transformation system in the *japonica* rice variety 'Kitaake'. Kitaake is fast flowering, day neutral, small in stature and an established model for functional genomics studies (Li *et al.*, 2017). The obstacle of genetic transformation with a single gene construct at a time and crossing has largely been solved by gene synthesis and Golden Gate cloning or similar 'gene building block' approaches (Engler *et al.*, 2014). Gene synthesis allows the 'domestication' of coding sequences to remove or insert rare Type IIS restriction enzyme recognition sites while leaving the amino acid sequence unchanged, thus enabling the assembly of gene modules which can be pasted together, often in a 'one pot cloning' approach. Assembly of these modules into a T-DNA suitable for *Agrobacterium* transformation is therefore greatly accelerated over traditional restriction/ligation approaches (Andreou and Nakayama, 2018). In principle, assembling all the metabolic and transporter components of Figure 1 for rice transformation on a single construct should be readily achievable and this work is currently underway. This approach has so far enabled a 6-year crossing strategy to be reduced to a 6-month single transformation experiment in rice.

In a large multigene overexpression construct, it is not desirable to reuse 'parts' multiple times due to the possibility of recombination deletion, post-transcriptional gene silencing or inactivation at the promoter level via methylation (Wassenegger, 2002). Epigenetic promoter silencing is a poorly understood process and can be a major challenge

for metabolic engineering. This also presents a challenge for the design of the gene constructs described above for C₄ rice. For example, expression of just CA, PEPC, MDH and PPDK would ideally require four heterologous M-specific promoter sequences. While this may be possible for the M compartment, it is not for the BS compartment (see above).

Synthetic biology has also provided a potential solution to this paucity of promoters in rice. Brückner *et al.* (2015) described a system compatible with the Golden Gate cloning which utilises multiple promoters (Synthetic TALE Activated Promoters or STAPs) designed to be orthogonal to the genome of the plant to be transformed, which can be activated by a single Transcription Activator-Like Effector (TALE). This approach provides the opportunity to build multiple transcriptional units driven by different promoters on the same gene construct, *trans*-activated by a single transcription factor.

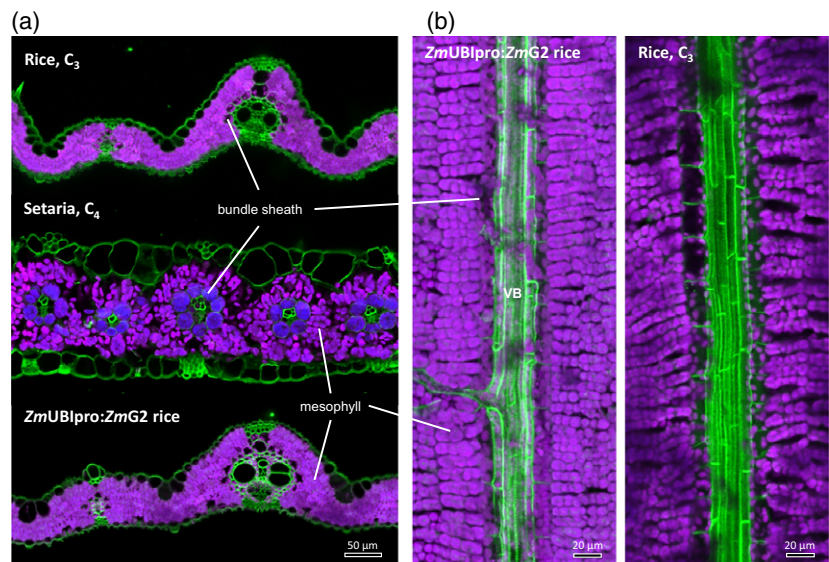
FUNCTIONALISATION OF THE BUNDLE SHEATH: AN ANATOMICAL ROADBLOCK?

Figure 2(a) shows fresh transverse sections of a rice (C₃) and *Setaria viridis* (C₄) leaf imaged with the laser confocal microscope using chlorophyll fluorescence overlaid with cell wall fluorescence. This enables visualisation of the chloroplast contents of BS cells in each species. This image clearly indicates that the BS compartment of the C₄ leaf is tightly packed with chloroplasts, whereas the rice BS compartment is only sparsely populated by chloroplasts and the cells are highly vacuolated. This has previously been pointed out and ‘photosynthetic functionalisation’ of the BS has been proposed as an early step in evolution of C₄ photosynthesis, probably occurring at the C₂ or proto-Kranz stage (Sage, 2004; Wang *et al.*, 2017b). The underlying mechanisms responsible for the

proliferation of chloroplasts in BS cells of C₄ leaves are largely unknown. However, recently a transgenic approach has been used in rice to ‘recreate’ this key step in evolution by overexpressing the transcription factors GOLDEN2 (G2) or GOLDEN2-LIKE (GLK) (Wang *et al.*, 2017b) which had been implicated as important in BS cell differentiation in terrestrial plants including *Zea mays* and more recently in rice (Wang *et al.*, 2013 and references therein). These transcription factors are thought to act in the cytokinin signalling pathway (Wang *et al.*, 2013). Overexpression of these *Z. mays* transcription factors in rice indeed increased the proportion of vascular cell area occupied by chloroplasts, and the mitochondrial population in rice BS cells, and the plasmodesmatal connectivity at the BS/M cell interface (Wang *et al.*, 2017b). The BS chloroplast abundance data from this work are summarised in Figure 3. It is evident from these quantitative data that overexpression of this class of transcription factor can increase BS chloroplast content as a proportion of total leaf chloroplasts by approximately five-fold (in the case of G2 expressed from the ubiquitin promoter of *Z. mays*: UBI-G2). However, another three-fold increase is required to reach levels equivalent to C₂ leaves and potentially another six-fold to reach C₄ levels of chloroplast area in the BS. This can be visualised in Figure 2 which shows a representative image of a leaf transverse (a) and paradermal (b) sections of the UBI-G2 lines (Wang *et al.*, 2017b) compared with rice and *Setaria* leaves. Comparison of rice and C₄ leaf chloroplast distribution is complicated, however, by the change in vein spacing and reduction in M chloroplast numbers seen in C₄ leaves relative to C₃ (Stata *et al.*, 2016).

Alternative approaches are also under investigation for increasing chloroplast abundance/ volume in the BS compartment of rice (Wang *et al.*, 2017a). The transcription factor CYTOKININ RESPONSIVE GATA FACTOR 1 (CGA1) has

Figure 2. Confocal micrographs of fresh hand-cut transverse (a) and paradermal (b) leaf sections of *Oryza sativa* (rice), *Setaria viridis* (*Setaria*), and a rice line with constitutively expressed GOLDEN2 transcription factor from *Zea mays* (*ZmUBIpro:ZmG2* rice). Excitation wavelength at 633 nm and dual emission wavelengths at 650–720 nm (magenta for Photosystem II) and 720–800 nm (blue for Photosystem I) were used to visualise chloroplasts. Note that bundle sheath cells of *Setaria* leaf have chloroplasts appearing pseudo-blue due to reduced Photosystem II content. Cell wall (green) was visualised at an excitation wavelength of 405 nm and emission wavelength of 420–480 nm. VB, vascular bundle.



also been shown to regulate leaf chloroplast abundance via the cytokinin signalling network (Chiang *et al.*, 2012; Hudson *et al.*, 2013; Wang *et al.*, 2017a) and overexpression of CGA1 in rice resulted in a 30% increase in flag leaf chlorophyll and close to a doubling of chloroplast numbers on a fresh weight basis (Hudson *et al.*, 2013). It has been proposed that regulation of the FtsZ chloroplast division gene by CGA1 is responsible for this phenotype, presenting opportunities for regulating levels of this protein directly or other genes in this pathway controlling chloroplast division (Hudson *et al.*, 2013; Wang *et al.*, 2017a).

It may be that part of the difficulty obtaining high level expression of chloroplast proteins in rice BS cells is the lack of sufficient chloroplast volume to house the recombinant proteins targeted to this compartment. The developmental programme of BS cells in a C₃ grass may reflect a more 'parenchyma-like' role in temporary sugar storage

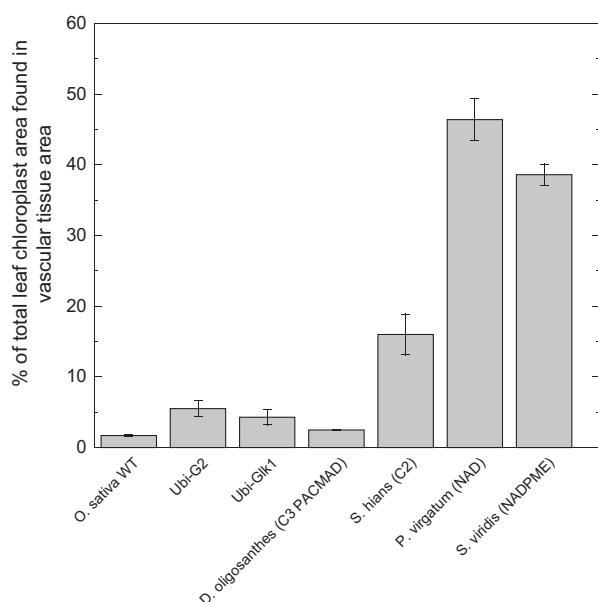


Figure 3. Per cent of total chloroplast area found in vascular tissue area. The data are taken from Table S5 Wang *et al.* (2017b). Ubi-G2 and Ubi-Glk1 are *Oryza sativa* plants in which the transcription factor GOLDEN2 (G2) or GOLDEN2-LIKE1 (GLK1) has been overexpressed from the ubiquitin promoter of *Zea mays* (Wang *et al.*, 2017b). Other monocots shown are *Dichanthelium oligosanthos*, a C₃ from the PACMAD clade; *Steinichisma hians*, which operates the C₂ photosynthetic pathway; *Panicum virgatum*, a C₄ species with the NAD-ME decarboxylation type; and *Setaria viridis*, a C₄ species with the NADP-ME decarboxylation type.

and the sugar status of the BS cells may not be conducive to photosynthetic functionalisation. The high level expression of sugar effluxers such as the SWEET13 gene family in the BS of C₄ plants has been proposed as evidence that sugar status in the BS or C₄ grasses may be quite different from that in C₃ grasses (Emms *et al.*, 2016). In addition, partitioning of starch almost exclusively into the BS of C₄ grasses is a curious and potentially relevant observation (Lunn and Furbank, 1999).

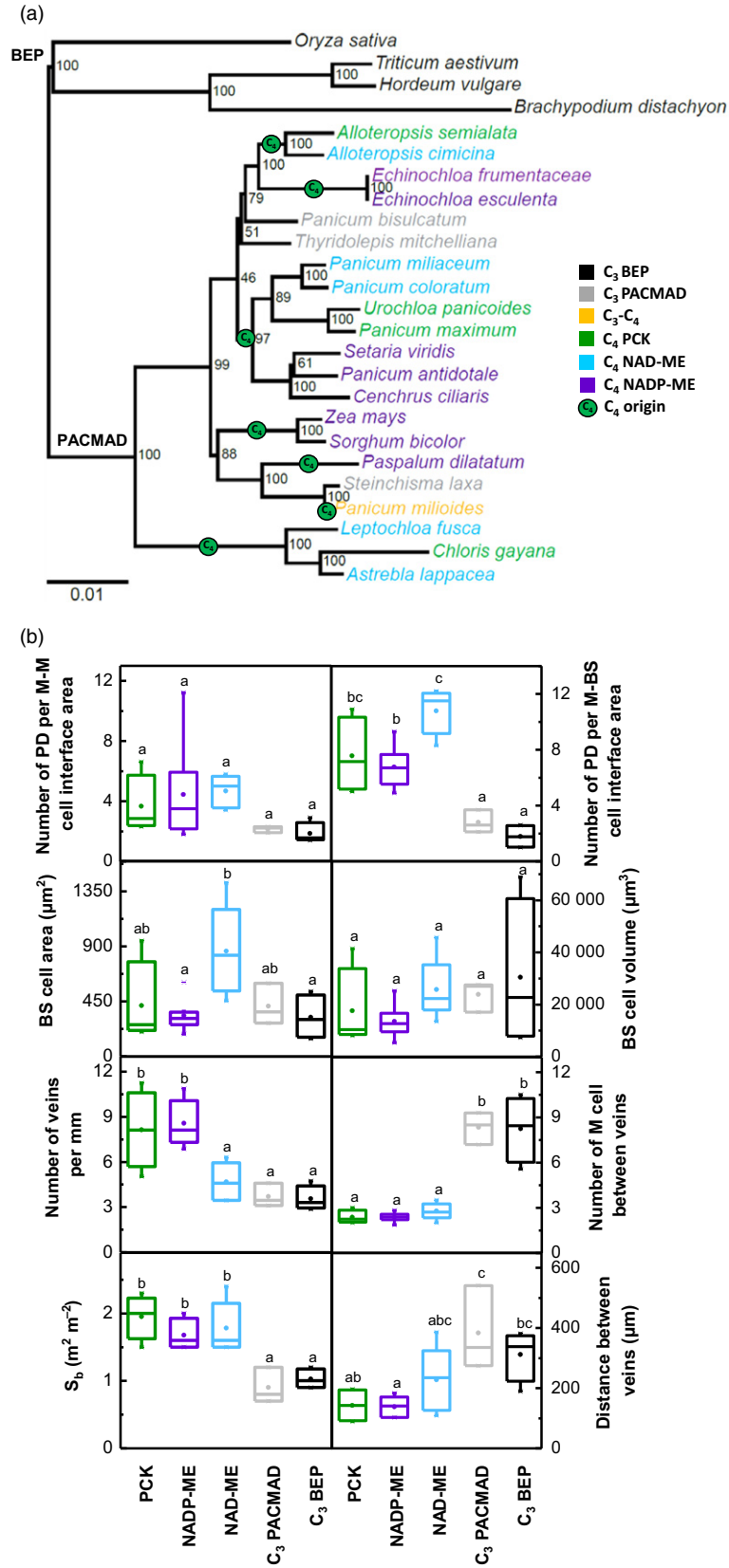
WHAT CAN WE LEARN FROM COMPARATIVE LEAF ANATOMY BETWEEN C₃ AND C₄ GRASS SPECIES?

It has previously been proposed that an early step in C₄ evolution was 'inflation' of BS cell size (Sage, 2004; Sage *et al.*, 2012; Christin *et al.*, 2013). However, our results from anatomical measurements performed in 25 grass species, representing different photosynthetic types and seven independent C₄ evolutionary origins (Figure 4a), reveal that C₄ leaves do not necessarily have larger BS cells, nor do they always have shorter interveinal distances than C₃ leaves (Figure 4b). Rather, a C₄ leaf has greater BS surface area per leaf area (S_b), fewer interveinal M cells than a C₃ leaf and most notably, more plasmodesmata (PD) at the BS/M cell interfaces (Figure 4; Danila *et al.*, 2016; Danila *et al.*, 2018). For a functional C₄ rice, these findings mean that increasing PD connections between M and BS, increasing S_b, and reducing M cells between veins of the rice leaves may be essential for the operation of an efficient photosynthetic pathway.

It has long been thought that a key feature of C₄ leaf anatomy was increased abundance of the symplastic nanochannels (PDs) that facilitate the rapid exchange of metabolites between M and BS cells during C₄ photosynthesis (Hatch and Osmond, 1976). Recent work, however, has provided quantitative data on this parameter (Danila *et al.*, 2016; Danila *et al.*, 2018); Figure 4). Enhancement of the PD connections between M and BS of rice to levels observed in closely related C₄ species would require increases of at least five-fold (Danila *et al.*, 2018). While there are few genes known which control PD development and proliferation, it was recently observed that when maize GLK genes were constitutively expressed in rice, increased organelle volume was accompanied by increased M-BS PD density; although the enhancement is only double that of wild type (Wang *et al.*, 2017b). Coordination of chloroplast

Figure 4. (a) Phylogenetic tree of the C₃ and C₄ grass species examined generated using sequences from *ndhF* and *rbcL* chloroplast genes. Species names are colour-coded according to photosynthetic types. The seven independent evolutionary origins of C₄ photosynthesis according to (GPWGII, 2012) are indicated with green circles at the midpoint of the branches. Note that *Panicum milioides*, a.k.a. *Steinichisma hians*, although technically a C₃-C₄ intermediate (Duvall *et al.*, 2003) was categorised as C₄ in GPWGII (2012). Numerical value at internal nodes is the percentage of non-parametric bootstrap replicates that support the bipartition. Scale bar indicates amino acid substitutions per site.

(b) Distribution of leaf trait values among photosynthetic types in (a) excluding the C₃-C₄ intermediate type, in which only one species was measured. The distribution of eight variables is summarised by boxplots. Box and whiskers represent the 25 to 75 percentile, and the minimum and maximum distribution. Means are denoted by (*). Letters show the statistical ranking using a post-hoc Tukey test among photosynthetic types (different letters indicate differences at P-value < 0.05). BS, bundle sheath; M, mesophyll; PD, plasmodesmata; S_b, bundle sheath surface area per unit leaf area.



and PD development has been suggested previously (Brunkard *et al.*, 2013) and cytokinin has also been implicated in the proliferation of PDs in the shoot apical meristem (Ormenese *et al.*, 2006). These findings offer some hope that a single gene or at least an established transcriptional network could provide a master switch for C₄ BS anatomy.

The BS surface area per leaf area (S_b) is a physiological parameter which has proven to be an important feature of modelling the M-BS interface, including to estimate BS conductance to CO₂ diffusion (first estimates ranged between 0.6 m² m⁻² and 3.1 m² m⁻² (Apel and Peisker, 1978; Brown and Byrd, 1993)). S_b was obtained by dividing measurements of BS tissue perimeter from micrographs by interveinal distance (Pengelly *et al.*, 2010). Increasing S_b can be achieved in multiple ways; modification of BS cell size, vasculature size, and the distance between BS. Because BS cell size does not differ substantially between C₃ and C₄ grass species (Figure 4, Danila *et al.*, 2018), reducing the interveinal distance appears to be the most logical path to increase S_b in rice. Ideally, this would mean upregulation of gene(s) that would promote insertion of additional veins between existing veins of rice, thus reducing interveinal distance and, at the same time, decreasing the number of M cells between veins (Sedelnikova *et al.*, 2018; Hughes *et al.*, 2019).

PAYING THE ENERGY COST OF C₄ PHOTOSYNTHESIS

The energy cost of C₄ photosynthesis is significantly higher compared with C₃ as it requires a minimum of two more ATP molecules per one CO₂ fixed (Furbank *et al.*, 1990). While high conductance of the M-BS interface to metabolites is essential for the operation of C₄ photosynthesis, it also allows a proportion of the CO₂ concentrated in the BS to escape back to M (called CO₂ leakage; von Caemmerer and Furbank, 2003). This CO₂ is either refixed or lost to the intercellular spaces in the M, which increases the cost of C₄ photosynthesis (Furbank *et al.*, 1990; von Caemmerer and Furbank, 2003). To sustain higher energy requirements, C₄ plants adapt the photosynthetic electron transfer chains of M and BS cells depending on specific needs of the C₄ subtype they belong to (Munekage and Taniguchi, 2016). As the efforts of the C₄ rice project have targeted NADP-ME as the decarboxylating enzyme, we focus here on specific energy requirements of the NADP-ME subtype of C₄ photosynthesis.

An early observation on the anatomy of tropical grasses which predated the discovery of C₄ photosynthesis was that chloroplasts in the two cell types of leaves with Kranz anatomy are dimorphic, with the BS chloroplasts often lacking grana stacks (Rhoades and Carvalho, 1944). Subsequently, it was discovered that these grasses such as *Z. mays* were in fact C₄ plants and specifically used the NADP-ME pathway of C₄ photosynthesis (Edwards *et al.*,

1971; Hatch and Kagawa, 1976). As NADP-ME plants primarily use malate as a C₄ acid diffusing from M to BS cells, NADPH produced in the M chloroplasts is consumed for malate synthesis but it is then produced in the BS upon malate decarboxylation to pyruvate (Figure 1). The net transfer of NADPH from M to BS cells means that there is a reduced requirement for linear electron flow to produce NADPH in the BS chloroplast, which have reduced Photosystem II (PSII) content but have highly developed cyclic electron flow (CEF) machinery around Photosystem I (PSI) (Figure 5). Therefore, the observation that BS chloroplasts of NADP-ME plants are mostly agranal with little or no grana thylakoids functionally reflects the reduced PSII, which would normally be located in the granal stacks (see Munekage and Taniguchi, 2016). However, grana content in BS chloroplasts of NADP-ME plants is rather variable between species (Ueno *et al.*, 2005) and also within species in response to environmental conditions (Omoto *et al.*, 2009; Danila *et al.*, 2019) and depending on leaf age (Andersen *et al.*, 1972). It has also been shown that aspartate can be used as a transported C₄ acid in many NADP-ME type C₄ plants, providing some flexibility in the amount of reducing power transferred to the BS from the M and suggesting that PSII content in the BS cells might in fact respond to the NADPH/NADP⁺ ratio or the redox state of the BS cells (see Furbank, 2011). This is supported by over-expression of maize NADP-ME in rice M cells which resulted in agranal chloroplasts, conceivably, due to the increased NADPH/NADP⁺ ratio depleting PSI of electron acceptor and causing over-reduction of the electron transfer chain (Takeuchi *et al.*, 2000). While prolonged reduction of PSII acceptors promotes the formation of reactive oxygen species and causes damage and degradation of PSII (Vass, 2012) and thus also grana, there might be also regulatory mechanisms preventing transcription and *de novo* assembly of PSII polypeptides and grana formation in over-reduced conditions (Pfannschmidt *et al.*, 1999).

It has been proposed that elevated CEF in BS chloroplasts allows NADP-ME plants to accommodate the extra costs of C₄ photosynthesis by producing ATP without affecting NADPH/NADP⁺ ratio (Furbank *et al.*, 1990). The chloroplast NADPH dehydrogenase-like complex and PROTON GRADIENT REGULATION 5 (PGR5) mediate two different CEF routes (Takabayashi *et al.*, 2005; Munekage, 2016). Interestingly, in C₃ plants, CEF is promoted in conditions causing over-reduction of the electron transport chain (Suorsa, 2015) and therefore CEF might be naturally upregulated in BS cells of C₄ rice in response to high NADPH/NADP⁺ ratio.

At present, it is unknown whether alteration of BS chloroplast electron transport components is a strict requirement for C₄ rice or only 'fine tuning'. If the required regulatory mechanisms already exist in rice, BS chloroplasts could conceivably adjust PSII and grana content

Figure 5. Transmission electron micrographs of the bundle sheath chloroplasts from *Oryza sativa* (rice) and *Setaria viridis* (Setaria) and schematic representation of their electron flow pathways. LEF, linear electron flow, is supported by Photosystem II, Cytochrome *b₆ f* complex and Photosystem I and is dominant in rice bundle sheath chloroplasts. CEF, cyclic electron flow, does not require Photosystem II and is dominant in Setaria bundle sheath chloroplasts. G, thylakoid grana stacks, are abundant in rice but not in Setaria because bundle sheath chloroplasts of NADP-ME C₄ plants require less Photosystem II that is typically localised to grana. NADPH/NADP⁺ balance in chloroplasts might be a factor defining the ratio between LEF and CEF.

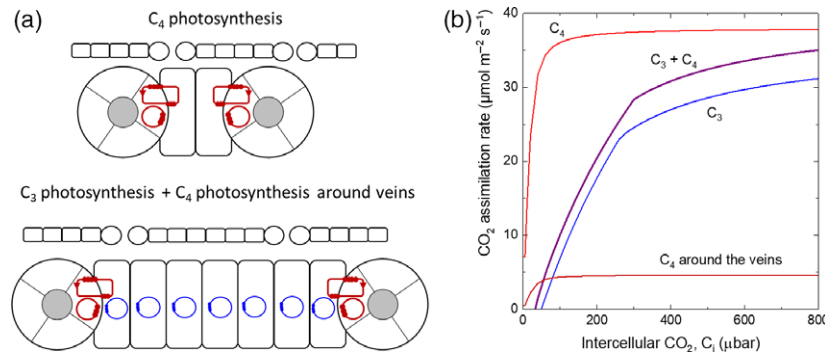
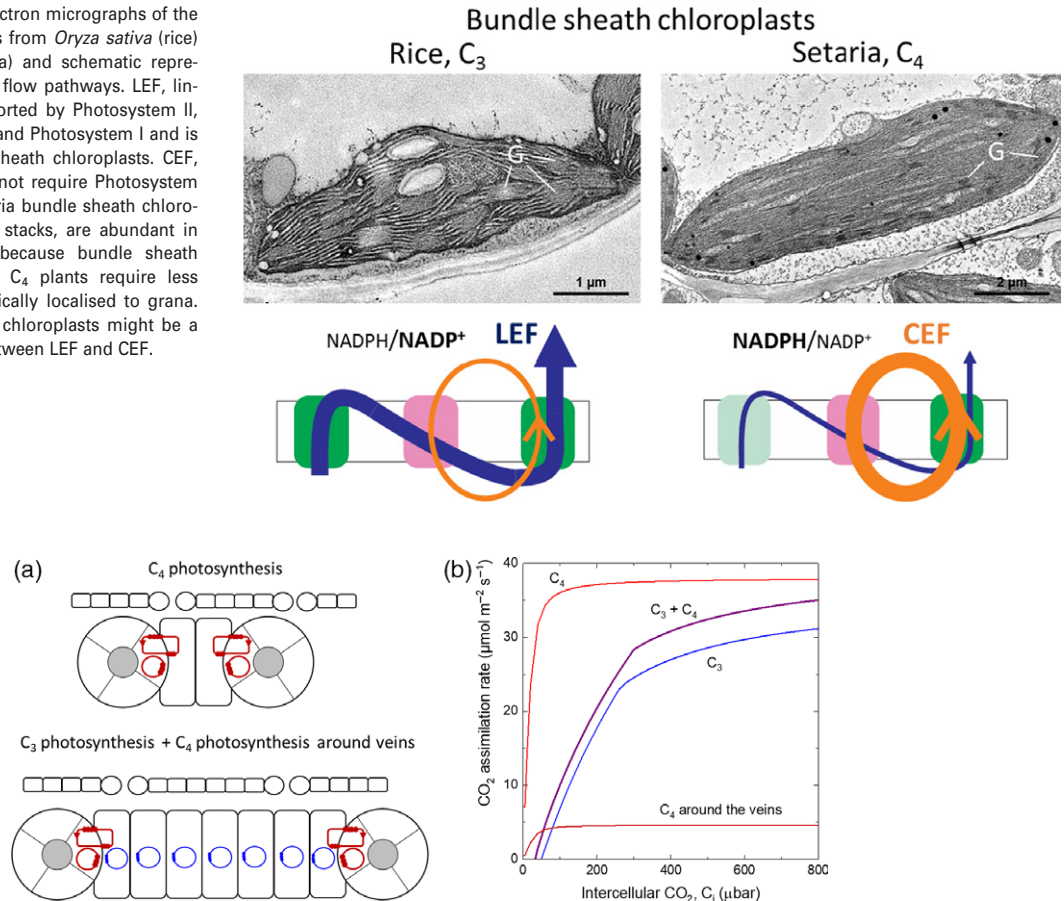


Figure 6. (a) Diagram comparing two cell-type C₄ photosynthesis and a rice leaf with low capacity C₄ photosynthesis occurring around the veins. In the latter, C₃ photosynthesis occurs in all mesophyll cells (rectangular cells) and in bundle sheath cells (cells inside circles), in addition to C₄ photosynthesis occurring only in mesophyll cells adjacent to bundle sheath cells. (b) Modelled CO₂ assimilation rates of a rice leaf with low capacity C₄ photosynthesis occurring around the veins (purple line). Also shown are rice leaf with C₃ photosynthesis (blue line), a standard high capacity C₄ photosynthesis (red line), and the C₄ photosynthesis around the vein contributing to the enhanced photosynthesis rate of the purple line (dark red line). C₃ leaf photosynthesis has been modelled with RuBisCO maximal activity, $V_{cmax} = 90 \mu\text{mol m}^{-2} \text{sec}^{-1}$ and electron transport rate, $J = 150 \mu\text{mol m}^{-2} \text{sec}^{-1}$. The standard C₄ photosynthesis was modelled with $V_{cmax} = 40 \mu\text{mol m}^{-2} \text{sec}^{-1}$ and PEP carboxylase maximal activity, $V_{pmax} = 300 \mu\text{mol m}^{-2} \text{sec}^{-1}$. The low capacity C₄ photosynthesis around the veins was modelled with $V_{cmax} = 5 \mu\text{mol m}^{-2} \text{sec}^{-1}$ and $V_{pmax} = 25 \mu\text{mol m}^{-2} \text{sec}^{-1}$. In the combined C₃ and C₄ photosynthesis, the total V_{cmax} was maintained at $90 \mu\text{mol m}^{-2} \text{sec}^{-1}$. C₃ RuBisCO kinetic constants by von Caemmerer (2000) were used together with the Michaels constant for CO₂ of PEPC of 154 μbar (Boyd *et al.*, 2015).

according to the redox state of cells in C₄ rice (Figure 5). Consequently, just the right amount of PSII will be fully assembled in BS to donate electrons for the CEF and compensate for the shortage of NADPH via linear electron transfer. However, the desired adaptation of electron transport requires a strictly coordinated expression and activity of NADP-ME in rice BS cells and efficient NADPH oxidation by the Calvin cycle to maintain an appropriate NADPH/NADP⁺ balance. It is worth mentioning that the flexibility of PGA reduction between M and BS cells in C₄ plants also contributes to the maintenance of NADPH/NADP⁺ balance in BS cells, however, it is not clear whether this pathway will be immediately available in C₄ rice. The composition of thylakoid protein complexes and energy requirements are similar between C₃ and C₄ M cells (Munekage and Taniuchi, 2016), but rice BS chloroplasts will require some

reorganisation of thylakoid complexes and increased abundance of the NADPH dehydrogenase-like complex (Majeran *et al.*, 2008; Hernández-Prieto *et al.*, 2019). This fine tuning will be necessary to run an efficient C₄ pathway as our recent research shows that the rate of C₄ photosynthesis is strongly dependent on the electron transport capacity of both M and BS cells (Ermakova *et al.*, 2019).

PARTIAL C₄: MODELLING MIXED C₃ AND C₄ PHOTOSYNTHESIS ON THE PATH TO C₄ RICE

The concept of C₄ rice has excited photosynthetic modellers and there are a number of photosynthetic models that have tried to evaluate the efficacy of introducing C₄ photosynthesis into rice (Bellasio, 2016; Yin and Struik, 2017; Wang *et al.*, 2017c; Bellasio and Farquhar, 2019). Each of these models has a different focus. Wang *et al.*

(2017c) developed a 3D reaction diffusion model of BS and connected M cells with anatomy based on a C₃ rice leaf in which C₄ photosynthesis was integrated with existing C₃ photosynthesis. They concluded that the C₄ cycle can operate adjacent to C₃ photosynthesis in rice M cells, but that the energy partitioning between the C₃ and C₄ cycle is an important consideration. In their current model every M cell is adjacent to a BS cell so it is built on a rice leaf with altered vein spacing. Bellasio's models consider C₄ photosynthesis as an addition to C₂ photosynthesis and also provides a valuable discussion on energy partitioning. Yin and Struik (2017) consider C₄ photosynthesis in rice at the canopy and crop level in different environments.

It is common to compare rice with *Z. mays* but it is informative to widen this comparison. The analysis of 25 monocot C₃ and C₄ leaves shows that all C₄ leaves have closer vein spacing than C₃ species with at most 2–3 M cells between BS (Figure 4). However there are examples in which C₄ photosynthesis is naturally supported around widely spaced veins such as in maize husk tissue, albeit at lower rates with little photosynthetic activity in the interveinal M cells (Langdale *et al.*, 1988; Pengelly *et al.*, 2011). Here, in the context of the introduction of C₄ metabolism into rice without altered vein spacing, we have asked the simple question: can we detect a small amount of C₄ photosynthesis introduced around existing veins using gas exchange techniques? The modelling here uses the Farquhar *et al.* (1980) C₃ model of photosynthesis combined with the enzyme limited C₄ model of photosynthesis described in von Caemmerer (2000) with C₃ kinetic constants for RuBisCO. Rice leaves have approximately 7 M cells between veins (Figure 4b; Chatterjee *et al.*, 2016). The modelled curves show that partitioning a small amount of RuBisCO to low capacity C₄ photosynthesis around veins (i.e. in 2 out of 7 M cells) lowers the compensation point and increase CO₂ assimilation rates at all CO₂ partial pressures (Figure 6). Hence this should be easily detected with gas exchange techniques. The modelling approach is very basic and has not considered the added energy requirements in M cells running both C₃ and C₄ photosynthesis given the low capacity C₄ photosynthetic rates considered here. Nevertheless it suggests that even this small addition, which would result in the small amount of RuBisCO in BS being more efficient, could have a physiological benefit.

CONCLUSIONS

Considerable progress has been made in building C₄ rice. The initial concept of assembling a tool box of components and a biochemical, anatomical and molecular blueprint based on evolution and 50 years of study has now become one of the largest synthetic biology projects of all time in plant biology. Discovering that our 'toolbox' was often lacking and that new knowledge and technologies

would continually reshape the strategy and redefine the blueprint should come as no surprise. 'Functionalisation' of the BS compartment is the immediate hurdle. For a fully functional C₄ cycle to operate in rice across two cell types, the BS must be functionally capable of housing the photosynthetic machinery necessary for C₄ acid decarboxylation and CO₂ refixation by RuBisCO. It appears that we have the biochemical and molecular-genetic components necessary for C₄ function; the means to prototype and fine tune them; a promise that C₄ photosynthesis around rice leaf veins may be possible and even beneficial, but there is still some distance yet to be travelled on the road to C₄ rice.

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AUTHOR CONTRIBUTIONS

All authors contributed to the writing of this review.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

DATA AVAILABILITY

The datasets generated in this paper are available from the corresponding author on request.

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