



# Phosphorylation of Phosphoenolpyruvate Carboxylase Is Essential for Maximal and Sustained Dark CO<sub>2</sub> Fixation and Core Circadian Clock Operation in the Obligate Crassulacean Acid Metabolism Species *Kalanchoë fedtschenkoi* <sup>CC-BY</sup>

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**Phosphoenolpyruvate carboxylase (PPC; EC 4.1.1.31) catalyzes primary nocturnal CO<sub>2</sub> fixation in Crassulacean acid metabolism (CAM) species. CAM PPC is regulated posttranslationally by a circadian clock-controlled protein kinase called phosphoenolpyruvate carboxylase kinase (PPCK). PPCK phosphorylates PPC during the dark period, reducing its sensitivity to feedback inhibition by malate and thus enhancing nocturnal CO<sub>2</sub> fixation to stored malate. Here, we report the generation and characterization of transgenic RNAi lines of the obligate CAM species *Kalanchoë fedtschenkoi* with reduced levels of *KfPPCK1* transcripts. Plants with reduced or no detectable dark phosphorylation of PPC displayed up to a 66% reduction in total dark period CO<sub>2</sub> fixation. These perturbations paralleled reduced malate accumulation at dawn and decreased nocturnal starch turnover. Loss of oscillations in the transcript abundance of *KfPPCK1* was accompanied by a loss of oscillations in the transcript abundance of many core circadian clock genes, suggesting that perturbing the only known link between CAM and the circadian clock feeds back to perturb the central circadian clock itself. This work shows that clock control of *KfPPCK1* prolongs the activity of PPC throughout the dark period in *K. fedtschenkoi*, optimizing CAM-associated dark CO<sub>2</sub> fixation, malate accumulation, CAM productivity, and core circadian clock robustness.**

## INTRODUCTION

Crassulacean acid metabolism (CAM) is one of several higher plant photosynthetic CO<sub>2</sub>-concentrating mechanisms. It is particularly noteworthy for its high water-use efficiency (WUE) relative to C<sub>3</sub> and C<sub>4</sub> photosynthesis (Borland et al., 2009; Cushman et al., 2015). The functional genomics and evolutionary biology of CAM species has recently received a dramatic upsurge in interest due to the potential of CAM crops, such as agaves and opuntias, as water-wise sources of biomass for biofuels, platform chemicals, and food (Cushman et al., 2015; Yang et al., 2015). In addition, efforts are underway to engineer CAM into C<sub>3</sub> species using synthetic biology approaches (Borland et al., 2014, 2015). CAM into C<sub>3</sub> aims to improve the WUE of key C<sub>3</sub> crops, thereby adapting them to the increased frequency and intensity of droughts and extreme high temperatures predicted under future climate change scenarios. The further exploitation of CAM in these ways will require detailed characterization of CAM gene function, especially with respect to their daily regulation by the circadian clock (Yang et al., 2015; Hartwell et al., 2016).

CAM is a classic example of convergent evolution that is known from 35 families of plants spanning monocots, dicots, and ferns (Smith and Winter, 1996; Holtum et al., 2007; Tsen and Holtum, 2012; Christin et al., 2014; Silvera et al., 2014). Based on our currently incomplete understanding of the molecular-genetic blueprint for CAM (Hartwell et al., 2016), C<sub>3</sub> species are believed to possess ancestral copies of all of the genes required for CAM (Cushman and Bohnert, 1999; Aubry et al., 2011; Ming et al., 2015). In principle, for CAM to evolve from the C<sub>3</sub> ancestral state in a leaf succulent CAM species such as *Kalanchoë fedtschenkoi*, the corresponding metabolic enzymes, metabolite transporters, and their cognate regulatory proteins must increase in abundance in leaf mesophyll cells and come under tight temporal control such that certain metabolic steps are confined to the dark period, and others are confined to the light (Cushman and Bohnert, 1999; Borland et al., 2009; Abraham et al., 2016; Brilhaus et al., 2016; Hartwell et al., 2016). However, only limited work has been reported testing this principle using a genetic approach, which represents a powerful method for in planta functional testing of candidate CAM genes when combined with detailed characterization of CAM-associated phenotypes in the resulting transgenic lines (Hartwell et al., 2016; Dever et al., 2015).

During the daily CAM cycle, CO<sub>2</sub> fixation occurs via two temporally separated steps. In the dark, atmospheric CO<sub>2</sub> enters through open stomata and is converted to HCO<sub>3</sub><sup>-</sup> either passively or via the reaction catalyzed by a β-carbonic anhydrase. Primary CO<sub>2</sub> fixation is catalyzed by phosphoenolpyruvate carboxylase (PPC), which converts PEP and HCO<sub>3</sub><sup>-</sup> to OAA and inorganic phosphate (Chollet et al., 1996). Malate dehydrogenase reduces OAA to malate, which is transported to and stored in the vacuole as malic acid

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(Borland et al., 2009). In the light, malate decarboxylation generates CO<sub>2</sub> that is refixed by Rubisco in the Calvin cycle, referred to as secondary CO<sub>2</sub> fixation. Both primary and secondary CO<sub>2</sub> fixation occurs in each leaf mesophyll cell; therefore, strict temporal regulation is required to prevent futile cycling between carboxylation and decarboxylation (Hartwell, 2005, 2006).

In the model obligate CAM species *K. fedtschenkoi* (Hartwell et al., 2016), both PPC abundance and specific activity are stable over the light/dark cycle (Nimmo et al., 1984; Dever et al., 2015), but its allosteric properties are subject to tight temporal control (Nimmo et al., 1986, 1987). Several primary metabolites have been reported to influence the activity of PPC in vitro, including malate (allosteric inhibitor) and glucose 6-phosphate (allosteric activator) (Doncaster and Leegood, 1987; Vidal and Chollet, 1997). For a CAM leaf, as the dark period progresses, the vacuolar storage capacity for malic acid reaches its limit, and malate begins to accumulate in the cytosol where it feeds back to inhibit PPC activity. In *K. fedtschenkoi*, the K<sub>i</sub> of PPC for malate was found to be increased by dark period phosphorylation of the enzyme on an N-terminal serine residue catalyzed by the activity of a specific, Ca<sup>2+</sup>-independent protein kinase, named PPC kinase (PPCK) (Carter et al., 1991; Hartwell et al., 1999; Nimmo, 2000, 2003). The circadian clock controls PPCK transcript levels and activity, such that they peak during the dark period (Hartwell et al., 1996, 1999; Taybi et al., 2000; Boxall et al., 2005; Theng et al., 2008). In the light, a protein phosphatase type 2A dephosphorylates PPC, leaving it up to 10 times more sensitive to malate inhibition (Carter et al., 1990). This phosphatase was found to be active throughout the light/dark cycle (Carter et al., 1991), supporting the proposal that the light/dark regulation of PPC is mainly a result of the circadian rhythm of PPCK levels (Hartwell et al., 1999), although this has yet to be demonstrated in planta.

PPCK remains the only well-defined, temporal/circadian control point for CAM (Hartwell, 2006, 2016). To further elucidate the role of PPC phosphorylation in the temporal control of CO<sub>2</sub> fixation associated with CAM, and the extent to which this phospho-regulation step helps to prevent metabolic futile cycling during CAM, we generated stable transformants of *K. fedtschenkoi* expressing RNA interference (RNAi) constructs that targeted the degradation of endogenous *KfPPCK1* transcripts. We recovered independent transgenic lines with reduced *KfPPCK1* transcripts and reduced phosphorylation of PPC in the dark and found that phosphorylation was vital for optimizing and sustaining dark CO<sub>2</sub> fixation associated with CAM. Furthermore, the loss of dark phosphorylation of PPC led to a loss of robust rhythmicity for certain components of the central molecular circadian clock and for the regulation of several distinct circadian clock outputs. Surprisingly, several key components of the central oscillator were induced in the *KfPPCK1* RNAi lines and these genes maintained robust transcript oscillations under constant light and temperature conditions.

## RESULTS

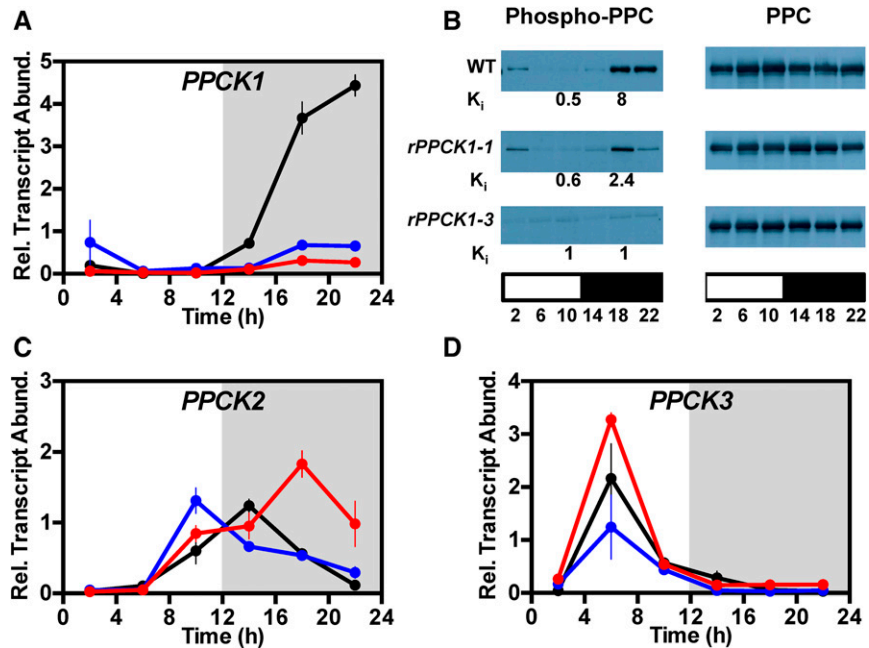
### Initial Screening and Characterization of *KfPPCK1* RNAi Lines of *K. fedtschenkoi*

As *K. fedtschenkoi* does not produce viable seed (Garcês et al., 2007; Hartwell et al., 2016), data are presented for primary

transformants that were propagated clonally via leaf plantlets and/or stem cuttings. Primary transformants were screened initially using high-throughput leaf disc tests for starch and acidity (Cushman et al., 2008; Dever et al., 2015). Leaf disc screening identified *KfPPCK1* RNAi lines because they had higher leaf pH at dawn than the wild type due to reduced malic acid accumulation during the dark period. Transgenic lines that acidified less during the dark period were screened for the steady state abundance of *KfPPCK1* transcript levels with RT-qPCR (Figure 1). Comparison of *KfPPCK1* transcript levels over a 12-h-light/12-h-dark cycle revealed that, relative to the wild type, the two best RNAi lines (*rPPCK1-1* and *rPPCK1-3*) had a large reduction in the transcript abundance of the target transcript (Figure 1A). While both lines displayed reduced levels of *KfPPCK1* transcripts relative to the wild type, line *rPPCK1-1* had a higher level of *KfPPCK1* transcripts in the middle of the dark period than line *rPPCK1-3*, and neither line displayed a complete loss of *KfPPCK1* transcript accumulation in the dark (Figure 1A). These results indicated that we had recovered independent lines with different degrees of silencing of the endogenous transcript level, with line *rPPCK1-3* showing the greatest reduction (Figure 1A).

### PPC Phosphorylation and the Apparent K<sub>i</sub> of PPC for Feedback Inhibition by L-Malate

In *K. fedtschenkoi*, PPC has been shown to be subject to a circadian rhythm of dark period phosphorylation/light period dephosphorylation as a result of rhythmic synthesis and degradation of PPCK (Hartwell et al., 1999). Immunoblotting using an anti-phospho-PPC antibody was used to investigate the phosphorylation state of PPC over a 12-h-light/12-h-dark cycle in CAM leaves of both the wild type and the *rPPCK1* lines (Figure 1B, left panel). The abundance of PPC itself was measured on replicate immunoblots to check for even protein loading of the blots (Figure 1B, right panel). In the wild-type CAM leaves, PPC was phosphorylated to a high level both at 18:00 and 22:00, 6 and 10 h into the 12-h dark period (Figure 1B, left panel). Wild-type PPC phosphorylation declined rapidly in the light, with a low level of phospho-PPC detected at 2 h into the 12-h light period, but virtually no phosphorylation detected at both 6 and 10 h into the light period (Figure 1B, left panel). The phosphorylation of PPC in the dark was reduced in both of the *rPPCK1* lines (Figure 1B, left panel). Relative to the wild type, line *rPPCK1-1* had a reduced peak of phospho-PPC in the middle of the dark period (18:00), but there was a dramatic reduction relative to the wild type at the end of the dark period (22:00) (Figure 1B, left panel). Line *rPPCK1-3* failed to achieve a detectable level of PPC phosphorylation above background at any of the sampled light/dark time points. Thus, *rPPCK1-1* had intermediate levels of PPC phosphorylation in the dark, and *rPPCK1-3* had the strongest phenotype, as it lacked detectable levels of dark period phosphorylation of PPC (Figure 1B, left panel). These perturbations of dark phosphorylation of PPC correlated well with the level of *KfPPCK1* transcripts in the dark in the two lines, with *rPPCK1-3* showing the greatest reduction in *KfPPCK1* transcript abundance (Figure 1A), consistent with the loss of PPC phosphorylation at the same time points (Figure 1B, left panel). The relative transcript abundance of two other detectable *KfPPCK* genes, namely, *KfPPCK2* and *KfPPCK3*, was higher in the *rPPCK1-3* lines relative to the wild type (Figures 1C and 1D).



**Figure 1.** Confirmation of Target Gene Silencing in Transgenic *K. fedtschenkoi* RNAi Lines *rPPCK1-1* and *rPPCK1-3*.

Downregulation of the target endogenous *KfPPCK1* gene was confirmed at the level of *KfPPCK1* transcript abundance (A), and target protein PPC phosphorylation and the apparent  $K_i$  of PPC for L-malate in rapidly desalted leaf extracts, measured as L-malate (mM) required for a 50% reduction in extractable PPC activity (B). Gene transcript abundance was measured using RT-qPCR for the wild type, *rPPCK1-1*, and *rPPCK1-3* for target genes: *KfPPCK1* (A), *KfPPCK2* (C), and *KfPPCK3* (D). Mature leaves (leaf pair 6) were sampled from three individual, clonal, biological replicate plants collected every 4 h across the 12-h-light/12-h-dark cycle. A thioesterase/thiol ester dehydrase-isomerase superfamily gene (*KfTED1*) was amplified from the same cDNAs as a reference gene. Gene transcript abundance data represent the mean of three technical replicates of each of three biological replicates and were normalized to reference gene *KfTED1*; error bars represent the SE of the mean calculated for each biological replicate. In all cases, plants were entrained under 12-h-light/12-h-dark cycles for 7 d prior to sampling. In (B), protein abundance (right panel) and the phosphorylation state of PPC (left panel) were determined by immunoblot analyses, and the apparent  $K_i$  of PPC for L-malate was measured using rapidly desalted leaf extracts ( $K_i$  values in mM below the corresponding left panel immunoblot time points). Total leaf protein (leaf pair 6) was isolated from leaves sampled every 4 h across the 12-h-light/12-h-dark cycle, separated using SDS-PAGE, and used for immunoblot analyses with antibodies raised to a phospho-PPC peptide (left panel). Sample loading was normalized according to total protein and confirmed using the immunoblot for total PPC protein, which is stable over the 12-h-light/12-h-dark cycle (right panel). The white bar below each panel represents the 12-h-light period, and the black bar below each panel represents the 12-h-dark period. For (A), (C), and (D), black data are for the wild type, blue for *rPPCK1-1*, and red for *rPPCK1-3*.

Interestingly, the increased transcript levels for *KfPPCK2* and *KfPPCK3* in the *rPPCK1-3* line (Figures 1C and 1D) did not rescue the dark period phosphorylation of PPC (Figure 1B, left panel). The fourth *PPCK* gene (*KfPPCK4*) in the *K. fedtschenkoi* genome could not be detected reliably using RT-qPCR, even after 40 cycles.

As the phosphorylation of PPC in the dark results in an increase in its apparent  $K_i$  for its feedback inhibitor L-malate (Nimmo et al., 1984, Carter et al., 1991), we also investigated the apparent  $K_i$  of PPC for L-malate using rapidly desalted extracts of LP6 sampled in the light at 10:00 (2 h before dusk) and in the dark at 18:00 (middle of the 12-h dark period) (Figure 1B,  $K_i$  values below corresponding time points on the left panel; Supplemental Figure 1). The wild type displayed the well-documented large increase in the apparent  $K_i$  of PPC for L-malate in the dark period, with the apparent  $K_i$  L-malate increasing from 0.5 mM for leaves sampled 2 h before the end of the light period up to 8 mM in the middle of the dark period, the latter coincident with high PPC phosphorylation detected at the same time point (Figure 1B, left panel; Supplemental Figure 1). Line *rPPCK1-1* displayed a large reduction in the light to dark

differential for the apparent  $K_i$  of PPC for L-malate, with the measured values being reduced to an average of 0.6 mM in the light samples, and 2.4 mM for the dark samples (Figure 1B; Supplemental Figure 1). This was consistent with the reduced level of PPC phosphorylation in *rPPCK1-1* in the middle of the dark period (Figure 1B, left panel). In line *rPPCK1-3*, which displayed the largest reduction in PPC phosphorylation in the dark period, the apparent  $K_i$  of PPC for L-malate was 1 mM in both the light and the dark, confirming that the lack of measurable PPC phosphorylation in the dark period in line *rPPCK1-3* resulted in a failure of the leaf mesophyll cells to convert the major, CAM-associated PPC isoform into the phosphorylated, malate-insensitive form that occurs in wild-type leaves in the dark period (Figure 1B, left panel; Supplemental Figure 1).

#### Gas Exchange Characteristics in Light/Dark Cycles

In *K. fedtschenkoi*, CAM is induced developmentally along the growing shoot, with the youngest leaves performing only daytime

CO<sub>2</sub> fixation via Rubisco and older leaves developing nocturnal CO<sub>2</sub> fixation associated with increasing CAM (Jones, 1975; Borland et al., 2009; Dever et al., 2015; Hartwell et al., 2016). Using a multi-cuvette, gas switching, infrared gas analyzer system (described in detail in Dever et al., 2015), we investigated the 24 h pattern of CO<sub>2</sub> exchange for detached, full-CAM leaves (leaf pair 6 [LP6]) from *rPPCK1-1*, *rPPCK1-3*, and the wild type that had been entrained in 12-h-light/12-h-dark cycles (Figure 2). Both the wild type and the transgenic lines with reduced *KfPPCK1* fixed the majority of their CO<sub>2</sub> at night (phase I of CAM: stomata open, primary atmospheric CO<sub>2</sub> fixation via PPC; Figure 2A) (Osmond, 1978). Lines *rPPCK1-1* and *rPPCK1-3* fixed only 53% and 34%, respectively, of wild-type CO<sub>2</sub> uptake in the dark (Figure 2A). *rPPCK1-1* and *rPPCK1-3* also fixed a small amount of CO<sub>2</sub> at dawn (phase II; early hours of the light period, stomata remain open briefly, CO<sub>2</sub> fixation via PPC and Rubisco), and this phase II peak of CO<sub>2</sub> fixation was greater in *rPPCK1-3* than in the wild type (Figure 2A). The wild type reached a maximum CO<sub>2</sub> fixation rate in the dark period of 4.4 μmol m<sup>-2</sup> s<sup>-1</sup>, whereas lines *rPPCK1-1* and *rPPCK1-3* reached a peak dark CO<sub>2</sub> fixation rate of only 2.4 μmol m<sup>-2</sup> s<sup>-1</sup>. It was noteworthy that the *rPPCK1-3* line that had the greatest reduction in dark phosphorylation of PPC (Figure 1B) reached its peak dark CO<sub>2</sub> fixation rate 1 h earlier in the dark period than either the wild type or *rPPCK1-1* line (Figure 2A). Furthermore, the CO<sub>2</sub> fixation of line *rPPCK1-3* began to decline after only 4.5 h of the dark period, to such an extent that, from 2 h before dawn, this line was respiring an increasing amount of CO<sub>2</sub> throughout the remainder of the dark period (Figure 2A). By contrast, the wild type and intermediate *rPPCK1-1* line continued net atmospheric CO<sub>2</sub> fixation for the whole of the dark period (Figure 2A).

Intact small plants (8-leaf-pair stage) with their root system in soil performed all four phases of CAM (labeled I–IV in Figure 2B). *rPPCK1-1* and *rPPCK1-3* fixed 72% and 62%, respectively, of wild-type light period CO<sub>2</sub> fixation. In the dark, *rPPCK1-1* fixed only 12%, and *rPPCK1-3* respired –76%, of wild-type CO<sub>2</sub> fixation (Figure 2B). Whole young plants of *rPPCK1-3* respired CO<sub>2</sub> throughout the dark period, but respiratory CO<sub>2</sub> loss from the plants declined from an initial peak straight after the lights went off to a minimum 4 h into the dark period, before rising gently throughout the remainder of the dark (Figure 2B). However, whole plants of all three lines showed a clear phase III of CAM in the light, evidenced by the pronounced dip in CO<sub>2</sub> fixation in the middle of the light period (Figure 2B).

### Malate, Starch, and Soluble Sugar Levels

Malate and starch levels reciprocate over the light/dark cycle in starch-accumulating CAM species such as *K. fedtschenkoi* (Borland et al., 2016). Starch breakdown in the dark generates the substrate for glycolysis, which in turn supplies PEP as a substrate for PPC to use in dark CO<sub>2</sub> fixation. As we had observed such large changes in the magnitude and duration of dark CO<sub>2</sub> fixation in the *rPPCK1* lines (Figures 2A and 2B), it was important to measure the levels and daily light/dark dynamics of the total leaf malate and starch pools.

The wild type, *rPPCK1-1*, and *rPPCK1-3* all accumulated malate in the dark (Figure 2C). Wild-type plants accumulated the most

malate by dawn (75.76 μmol gFW<sup>-1</sup> SE 4.65). Line *rPPCK1-1* accumulated 78% and *rPPCK1-3* accumulated 64%, of the wild-type level of malate by dawn (Figure 2C). At the end of the light period, ~90% of the dark accumulated malate had been decarboxylated in the wild-type and *rPPCK1* lines. The wild type had a Δ malate of 59.19 ± 8.82 μmol gFW<sup>-1</sup>, *rPPCK1-1* had a Δ malate of 47.16 ± 5.23 μmol gFW<sup>-1</sup>, and *PPCK1-3* had a Δ malate of 38.54 ± 4.89 μmol gFW<sup>-1</sup>.

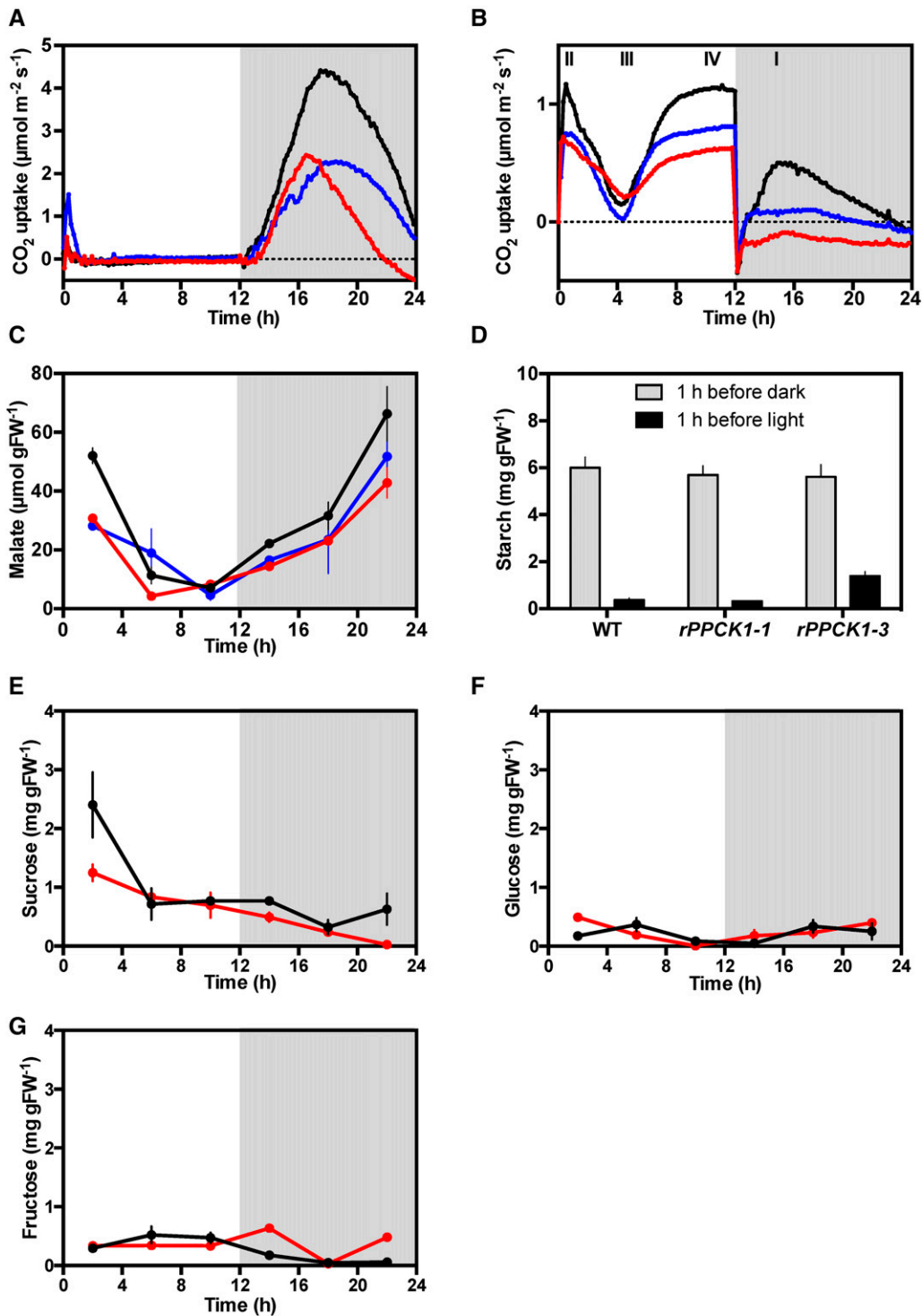
At the end of the light period, wild-type and *rPPCK1* CAM leaves (LP6) had synthesized similar amounts of starch (Figure 2D). Wild-type and *rPPCK1-1* leaves broke down 94% of the accumulated starch during the dark period, whereas *rPPCK1-3* only broke down 82% of its starch, consistent with the reduced CO<sub>2</sub> fixation and malate accumulation achieved by this line (Figures 2A and 2C). The wild type had a Δ starch of 5.63 ± 0.47 mg gFW<sup>-1</sup>, *PPCK1-1* had a Δ starch of 5.37 ± 0.38 mg gFW<sup>-1</sup>, and *PPCK1-3* had a Δ starch of 4.23 ± 0.67 mg gFW<sup>-1</sup>.

CAM leaves of the closely related obligate species *Kalanchoë daigremontiana* have been reported previously to display a characteristic postdawn peak of sucrose peaking at the end of phase II of CAM (Wild et al., 2010). At the beginning of the light period, wild-type *K. fedtschenkoi* had a peak of sucrose phased to 2 h after dawn that was reduced to around half the wild-type level in line *rPPCK1-3* (Figure 2E). Fructose accumulation peaked slightly later than the wild type in *rPPCK1-3* at 2 h into the dark period (Figure 2G), whereas the temporal pattern of glucose accumulation was similar between *rPPCK1-3* and the wild type (Figure 2F).

### Gas Exchange Characteristics under Constant Light and Temperature Free-Running Conditions

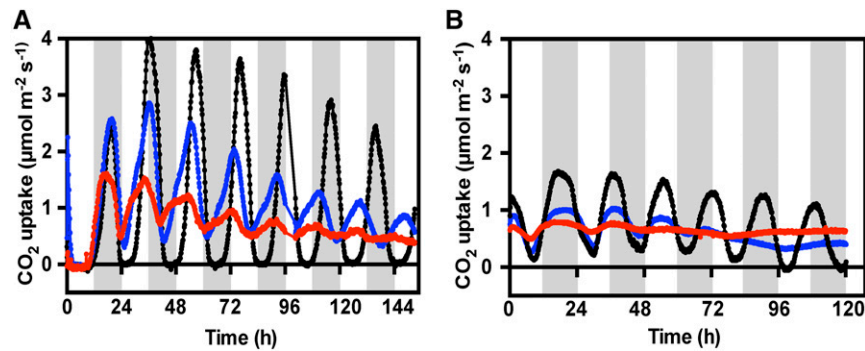
In constant light, free-running circadian conditions (LL; constant light 100 μmol m<sup>-2</sup> s<sup>-1</sup>, and temperature, 15°C), detached LP6 from wild-type plants displayed the characteristic, robust, and persistent CAM circadian rhythm, with oscillating peaks and troughs of CO<sub>2</sub> fixation (Figure 3) (Wilkins, 1992; Hartwell, 2006). *rPPCK1-1* and *rPPCK1-3* showed a dampening of the CO<sub>2</sub> fixation rhythm in constant conditions, with a dramatic reduction in rhythm amplitude (Figure 3A). The *rPPCK1-1* CO<sub>2</sub> fixation rhythm maintained clear oscillations throughout the 6.5-d experiment, but the period of the rhythm was shorter than the wild type such that, after 6.5 d, the wild-type CO<sub>2</sub> fixation trough or minimum coincided with the *rPPCK1-1* peak of CO<sub>2</sub> fixation (Figure 3A). The *rPPCK1-3* CO<sub>2</sub> fixation rhythm dampened more rapidly than *rPPCK1-1*, so much so that it was virtually arrhythmic after 6.5 d. The weak LL rhythm in line *rPPCK1-3* also had an even shorter period than *rPPCK1-1* (Figure 3A).

When whole young plants at the 8-leaf-pair stage were measured in the gas exchange system under LL free-running conditions, the CO<sub>2</sub> exchange rhythms in the *rPPCK1-1* and *rPPCK1-3* lines both collapsed to arrhythmia after 2 to 3 d (Figure 3B). The amplitude of the *rPPCK1-1* and *rPPCK1-3* CO<sub>2</sub> fixation rhythms was even lower for whole plants than for detached LP6 (compare Figure 3B with 3A). In particular, comparing the first peak and trough of its CO<sub>2</sub> fixation rhythm, *rPPCK1-3* achieved a maximum amplitude of ~0.25 μmol m<sup>-2</sup> s<sup>-1</sup>. This contrasted strongly with the amplitude of the wild-type rhythm for whole



**Figure 2.** Impact of Silencing *KPPCK1* on 24-h-Light/Dark Gas Exchange Profiles, and Malate, Starch, and Soluble Sugar Levels for the Wild Type, *rPPCK1-1*, and *rPPCK1-3* under Well-Watered Conditions.

- (A) Gas exchange profile for CAM leaves (leaf pair 6) using plants pre-entrained for 7 d under 12-h-light/12-h-dark cycles.
- (B) Gas exchange profile for well-watered whole young plants (8-leaf-pairs stage) using plants entrained for 7 d under 12-h-light/12-h-dark cycles.
- (C) Malate content was determined from leaf pair 6 samples collected every 4 h using plants entrained under 12-h-light/12-h-dark cycles using methanol extracts of leaves from the wild type, *rPPCK1-1*, and *rPPCK1-3*.
- (D) Starch content was determined from leaf pair 6 samples collected at 1 h before dawn and 1 h before dusk.



**Figure 3.** Effects of Silencing *KfPPCK1* on CAM CO<sub>2</sub> Exchange Rhythms Measured under LL Conditions.

**(A)** Gas exchange profile for CAM leaves (leaf pair 6) was measured using leaves entrained under a 12-h-light/12-h-dark cycle followed by release into constant LL conditions ( $100 \mu\text{mol m}^{-2} \text{s}^{-1}$  light at  $15^\circ\text{C}$ ).

**(B)** Gas exchange profile for well-watered whole young plants (8-leaf-pair stage) using plants entrained under 12-h-light/12-h-dark cycles followed by release into constant LL conditions ( $100 \mu\text{mol m}^{-2} \text{s}^{-1}$  light at  $15^\circ\text{C}$ ). Black data traces represent the wild type, blue data *rPPCK1-1*, and red data *rPPCK1-3*.

plants during the first peak and trough when the amplitude reached over  $1.5 \mu\text{mol m}^{-2} \text{s}^{-1}$  (Figure 3B).

### Phenotypic Characterization under Well-Watered and Drought-Stressed Conditions

CAM is widely recognized and characterized as a water-conserving adaptation to seasonally dry environments. It was therefore important to investigate the physiological responses and performance of the *rPPCK1* lines under drought-stress conditions. CO<sub>2</sub> exchange was measured for drought-stressed whole plants. Wild-type and *rPPCK1* plants were grown from plantlets under identical conditions to the LP6 stage. Water was then withheld for 25 d. Wild-type drought-stressed plants displayed strong CAM in light/dark cycles, with a large nocturnal phase I period of dark CO<sub>2</sub> fixation and a net loss of CO<sub>2</sub> in the light period (Figure 4). Phase II CO<sub>2</sub> uptake at dawn was present in both wild-type and RNAi lines, but phase IV CO<sub>2</sub> fixation at the end of the light period, which is characteristically observed in well-watered wild-type plants (Figure 2B), was absent in both the RNAi lines and the wild type.

As observed in well-watered conditions (Figure 3B), the LL circadian rhythm of CO<sub>2</sub> exchange persisted robustly in the wild-type drought-stressed plants, but collapsed rapidly toward arrhythmia in both *rPPCK1* lines (Figure 4A). In particular, after 2.5 d of LL, drought-stressed *rPPCK1-3* moved into net respiratory loss of CO<sub>2</sub>, and this continued throughout the remaining 4.5 d of the LL experimental run (Figure 4A). This contrasted strongly with the LL gas exchange for line *rPPCK1-3* under well-watered conditions,

where it maintained positive net CO<sub>2</sub> fixation throughout the LL experiment, both for detached LP6 and young whole plants (Figures 3A and 3B).

### Growth Performance under Well-Watered and Drought-Stressed Conditions

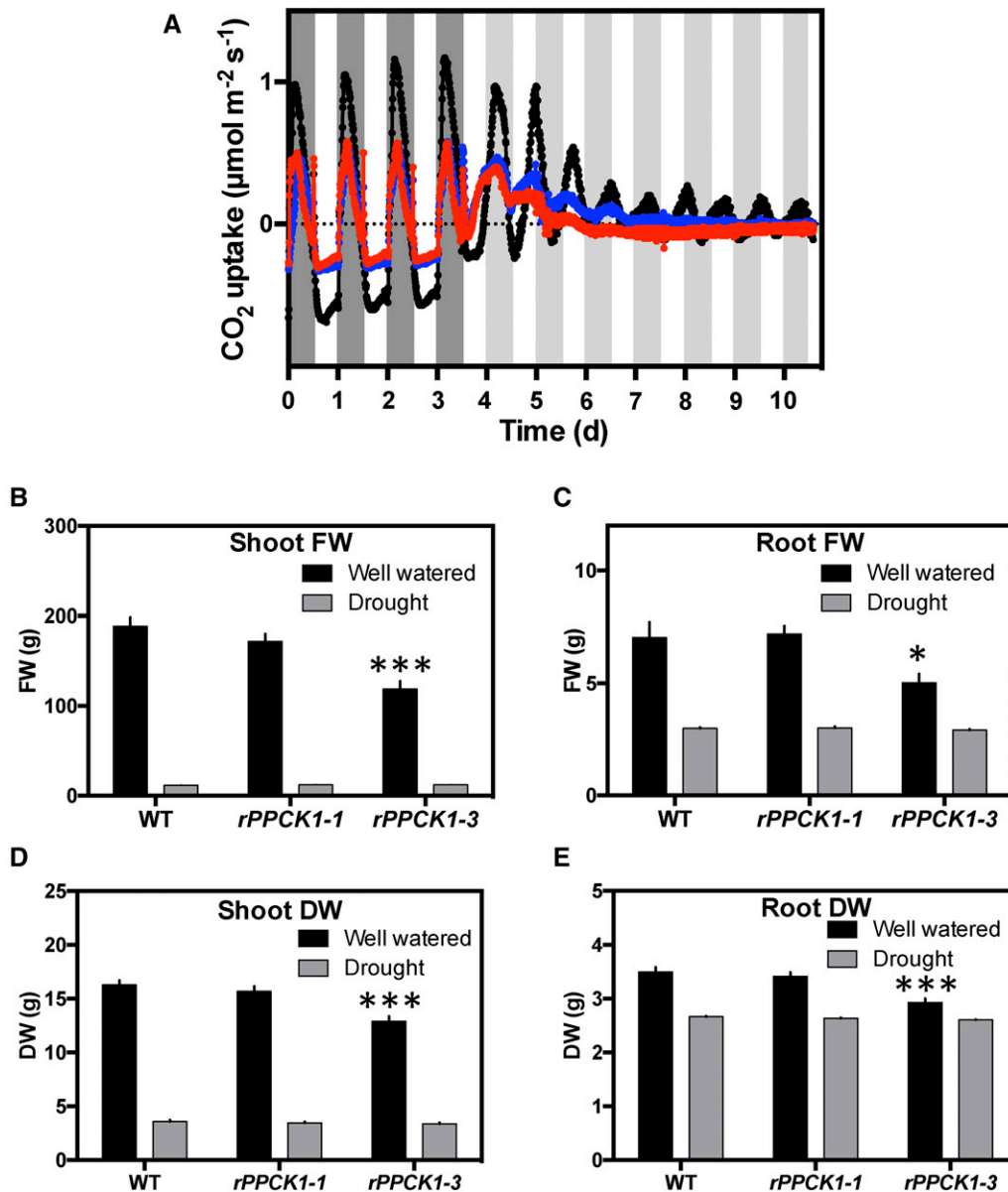
The high WUE of CAM plants allows them to grow productively in arid and semi-arid environments (Borland et al., 2009; Davis et al., 2017). Multiple clones of all lines were grown from plantlets under identical conditions to the LP6 stage. Half of the plants were maintained well-watered as controls, and half were subjected to drought stress for 138 d. Both sets of plants were maintained in the same growth cabinet such that the only environmental variable was water supply.

In well-watered conditions, all plants grew well. However, weighing aboveground shoots and belowground roots, both as fresh weight and dry weight, revealed that the average fresh and dry weight of the shoots and roots was significantly less than the wild type for line *rPPCK1-3* under well-watered conditions (Figures 4B to 4E). The intermediate line *rPPCK1-1* also had a lower average fresh and dry weight for its shoots and roots under well-watered conditions, although the reductions were not significantly different from the wild type based on Student's *t* tests (Figures 4B and 4D). The wet weight of the shoots of *rPPCK1-1* weighed 91% of the wild type, whereas the shoots of *rPPCK1-3* weighed 55% of the wild type ( $n = 7$ ). Differences in shoot dry weight were smaller, with *rPPCK1-1* shoots weighing 96% of the wild type, and *rPPCK1-3* weighing 76% of the wild

**Figure 2.** (continued).

**(E) to (G)** Soluble sugars levels were determined separately for sucrose, glucose, and fructose using three biological replicates of leaf pair 6 sampled every 4 h using plants entrained under 12-h-light/12-h-dark cycles.

In **(C) to (G)**, the error bars represent the SE of the mean calculated for the three biological replicates measured at each time point. Black data traces represent the wild type, blue data *rPPCK1-1*, and red data *rPPCK1-3*.



**Figure 4.** Impact of Loss of KfPPCK1 Activity on Gas Exchange Profiles and Vegetative Yield for Plants Subjected to Drought Stress.

**(A)** Drought-stressed whole plants (8-leaf-pair stage) were entrained under 12-h-light/12-h-dark cycles followed by release into constant LL conditions ( $100 \mu\text{mol m}^{-2} \text{s}^{-1}$  light at  $15^\circ\text{C}$ ). The black data trace represents the wild type, blue *rPPCK1-1*, and red *rPPCK1-3*.

**(B) to (E)** Fresh **(B)** and **(C)** and dry **(D)** and **(E)** weight of aboveground biomass (shoot; **(B)** and **(D)**) and belowground tissues (roots; **(C)** and **(E)**) at maturity (138 d under greenhouse conditions) for the wild type, *rPPCK1-1*, and *rPPCK1-3* under well-watered and drought-stressed conditions.  $n = 7$  plants; error bars represent se of the mean. Asterisks indicate significant difference from the wild type based on Student's  $t$  test: \* $P < 0.05$  and \*\*\* $P < 0.001$ .

type shoot dry weight ( $n = 7$ ). Belowground, root growth differences were less pronounced, although line *rPPCK1-3* achieved an average root wet weight that was only 71% of the wild type (Figure 4C).

The replicate plants subjected to 138 d of drought stress had fresh weights for both shoots and roots that were much lower than the fresh weights for the well-watered plants. For example, the

average fresh weight of wild-type shoots from well-watered plants was 188 g, whereas the average fresh weight for the drought-stressed wild type shoots was only 11 g. This demonstrated that the 138-d drought-stress treatment was a severe drought-stress regime, as might be experienced by *K. fedtschenkoi* during the dry season in its native Madagascar environment. After drought stress, the fresh and dry weights of both the shoots and roots of

the *rPPCK1* lines did not differ significantly from the wild-type values. The very low dry weights of the drought-stressed shoots of all lines relative to the dry weights achieved by all of the well-watered plants indicated that all lines performed very little shoot growth under such a long-term drought-stress regime (Figure 4D). However, the reduction in the dry weight of drought-stressed roots relative to the well-watered values was less dramatic (Figure 4E). Drought-stressed root dry weights were between 76 and 89% of the corresponding well-watered values, whereas drought-stressed shoots weighed less than 22 to 26% of the well-watered shoots for all lines. It is clear that all plants put more of their available fixed carbon into belowground root growth during the prolonged drought-stress treatment used in this experiment. Furthermore, the root fresh weights were almost the same as the root dry weights for the drought-stressed plants, suggesting that, by the end of the 138-d drought treatment, the roots of all lines were almost completely dry, even before being placed in a drying oven.

### Diurnal Regulation of the Transcript Abundance of Circadian Clock Genes

Having established that plants lacking *KfPPCK1* had lost the rhythm of CO<sub>2</sub> uptake in free-running conditions (Figures 3A and 3B), it was also important to investigate whether the regulation of genes in the core molecular circadian clock was affected under both light/dark cycles (Figure 5) and free-running LL conditions (Figure 6). *K. fedtschenkoii* has two copies of the circadian clock gene *TIMING OF CHLOROPHYLL A/B BINDING PROTEIN1* (*KfTOC1-1* and *KfTOC1-2*) and two *CIRCADIAN CLOCK-ASSOCIATED1/LATE ELONGATED HYPOCOTYL*-related genes (*KfCCA1-1* and *KfCCA1-2*). Both *KfTOC1* genes were upregulated under light/dark conditions in the *rPPCK1-3* line, with the phasing of the *KfTOC1-2* peak occurring 4 h earlier in *rPPCK1-3* (Figures 5A and 5B). In *rPPCK1-3*, both *KfCCA1-1* and *KfCCA1-2* were slightly upregulated at the time of their daily peak 2 h after dawn, whereas transcript levels of these genes in line *rPPCK1-1* were similar to or slightly lower than those in the wild type (Figures 5C and 5D). A *K. fedtschenkoii* *PSEUDO RESPONSE REGULATOR7* ortholog (*KfPRR7*) was markedly upregulated at 2 h after dawn in the *rPPCK1-3* line (Figure 5E), and a *KfPRR37* gene was also upregulated and peaked 4 h earlier than the wild type at 10:00 (2 h before dusk) in line *rPPCK1-3* (Figure 5F).

### Regulation of Circadian Clock Genes under Free-Running Conditions

The robust circadian rhythm of CAM CO<sub>2</sub> exchange in LL conditions for the wild type (Figures 3A and 3B) has been shown to correlate tightly with circadian clock control of PPCK and the phosphorylation state of PPC and its K<sub>i</sub> for malate (Hartwell, 2005, 2006; Dever et al., 2015). Since CO<sub>2</sub> exchange rhythms in LL conditions for *rPPCK1-1* and *rPPCK1-3* showed, at best, low-amplitude rhythmicity in well-watered plants (Figures 3A and 3B), it was important to also investigate the circadian regulation of the transcript abundance of *KfPPCK1* under LL conditions. In wild-type plants, *KfPPCK1* transcript levels

oscillated with a robust circadian rhythm throughout 3 d under LL (Figure 6A). The period of the rhythmic peaks and troughs matched the well-documented, temperature-dependent, short-period rhythm of gas exchange observed in *K. fedtschenkoii* (Anderson and Wilkins, 1989; Dever et al., 2015). Consistent with the RNAi-mediated silencing of the target gene, *KfPPCK1* transcript abundance was very low in *rPPCK1-3* in free-running conditions and did not oscillate (Figure 6A). The upregulation of *KfPPCK2* and *KfPPCK3* in plants lacking *KfPPCK1* under LD conditions (Figures 1C and 1D) led us to investigate whether these genes remained rhythmic in constant conditions in line *rPPCK1-3*. Both *KfPPCK2* and *KfPPCK3* transcript oscillations remained rhythmic and were dramatically upregulated at the start of the LL time course in *rPPCK1-3* (Figures 6B and 6C).

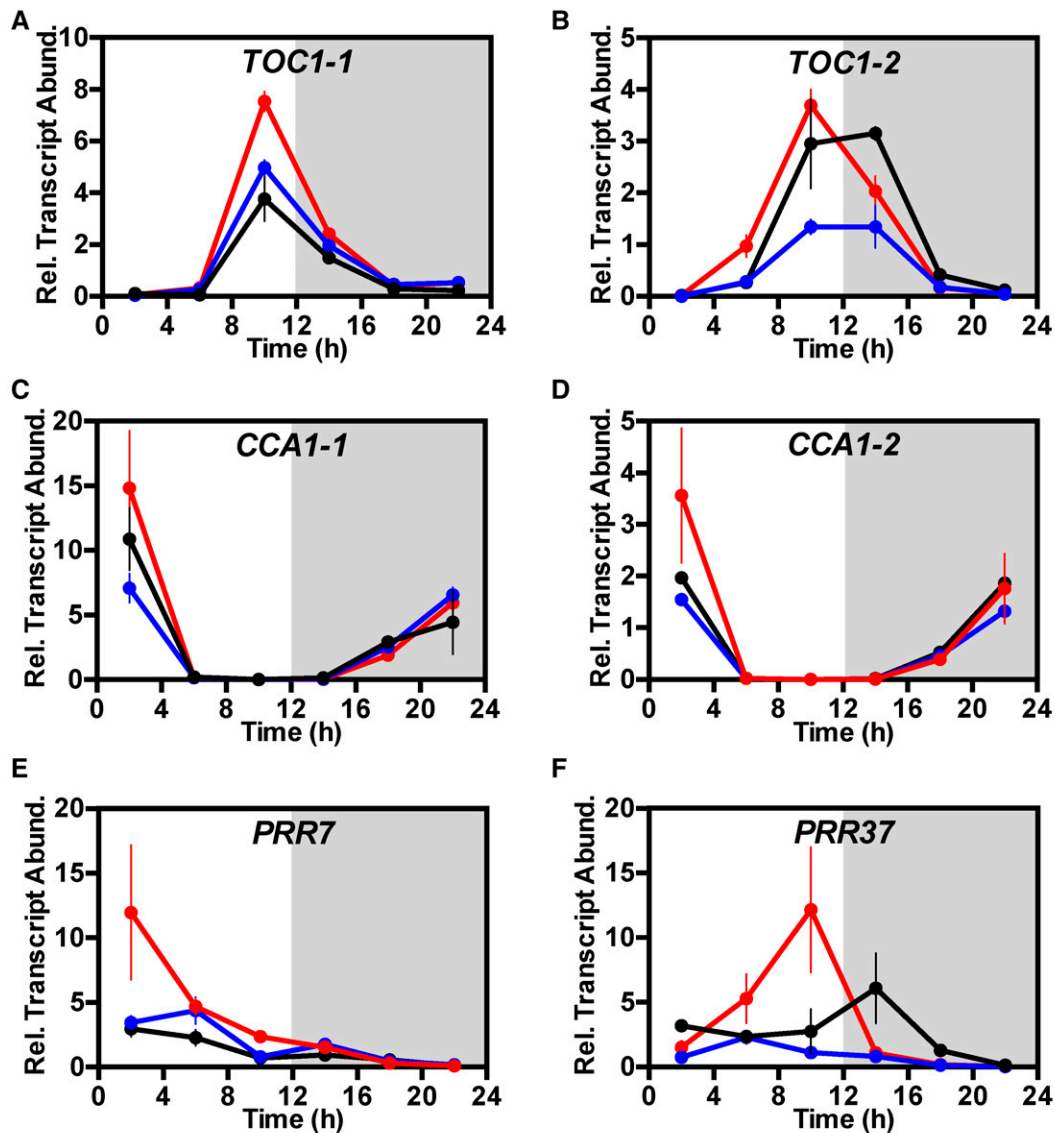
As these results revealed that certain circadian clock-controlled output genes were subject to robust oscillations under LL free-running conditions, it was also important to investigate the rhythmicity of core circadian oscillator genes. The rhythmicity of both *KfCCA1-1* and *KfCCA1-2* was dampened in *rPPCK1-3* (Figures 6D and 6E). Interestingly, the two *KfTOC1* genes were differentially regulated in *rPPCK1-3* under constant conditions with *KfTOC1-1* being upregulated and rhythmic, whereas *KfTOC1-2* was downregulated and its rhythm collapsed toward arrhythmia (Figures 6F and 6G). This was surprising as it indicated that the two *KfTOC1* genes may be involved in distinct circadian clocks having potentially divergent roles and responding to different entrainment signals, possibly in different tissues/cell types of the CAM leaf.

*KfPRR7*, *KfPRR37*, and *KfPRR9* all displayed dampening of their transcript rhythms toward arrhythmia in *rPPCK1-3* under constant conditions (Figures 6H to 6J). *K. fedtschenkoii* *FLAVIN BINDING KELCH-REPEAT F-BOX PROTEIN* (*KfFKF1*) and *Kf GIGANTEA* (*KfGI*) were both upregulated and rhythmic in LL (Figures 6K and 6L), whereas *Kf JUMONJIC-DOMAIN CONTAINING PROTEIN30* (*KfJM30*), *Kf NIGHT LIGHT-INDUCIBLE AND CLOCK-REGULATED3* (*KfLNK3-like*), and *Kf CYCLING DOF-TRANSCRIPTION FACTOR2* (*KfCDF2*) were all downregulated but rhythmic in *rPPCK1-3* (Figures 6M to 6O). *KfGI* was particularly noteworthy for the fact that the phasing of its transcript peaks and troughs shifted late relative to the wild type in line *rPPCK1-3* on the second and third 24-h period of the LL experimental run (Figure 6L), which contrasted strikingly with the earlier phase/ shorter period of the LL CO<sub>2</sub> fixation rhythm for *rPPCK1-3* (Figure 3A).

### Delayed Fluorescence Rhythms Revealed That a Further, Distinct Circadian Clock Output inside the Chloroplast Was Perturbed in the Absence of PPCK Activity

The level of delayed fluorescence (DF) has previously been reported to be under robust circadian control in *K. fedtschenkoii* and provides a simple high-throughput assay for measuring circadian period, robustness, and accuracy (Gould et al., 2009). DF was measured in the wild type, *rPPCK1-1*, and *rPPCK1-3* under LL free-running conditions (Figure 7), and the data were analyzed to calculate key circadian rhythm statistics using Biodare (Moore et al., 2014; Zielinski et al., 2014). The wild type had



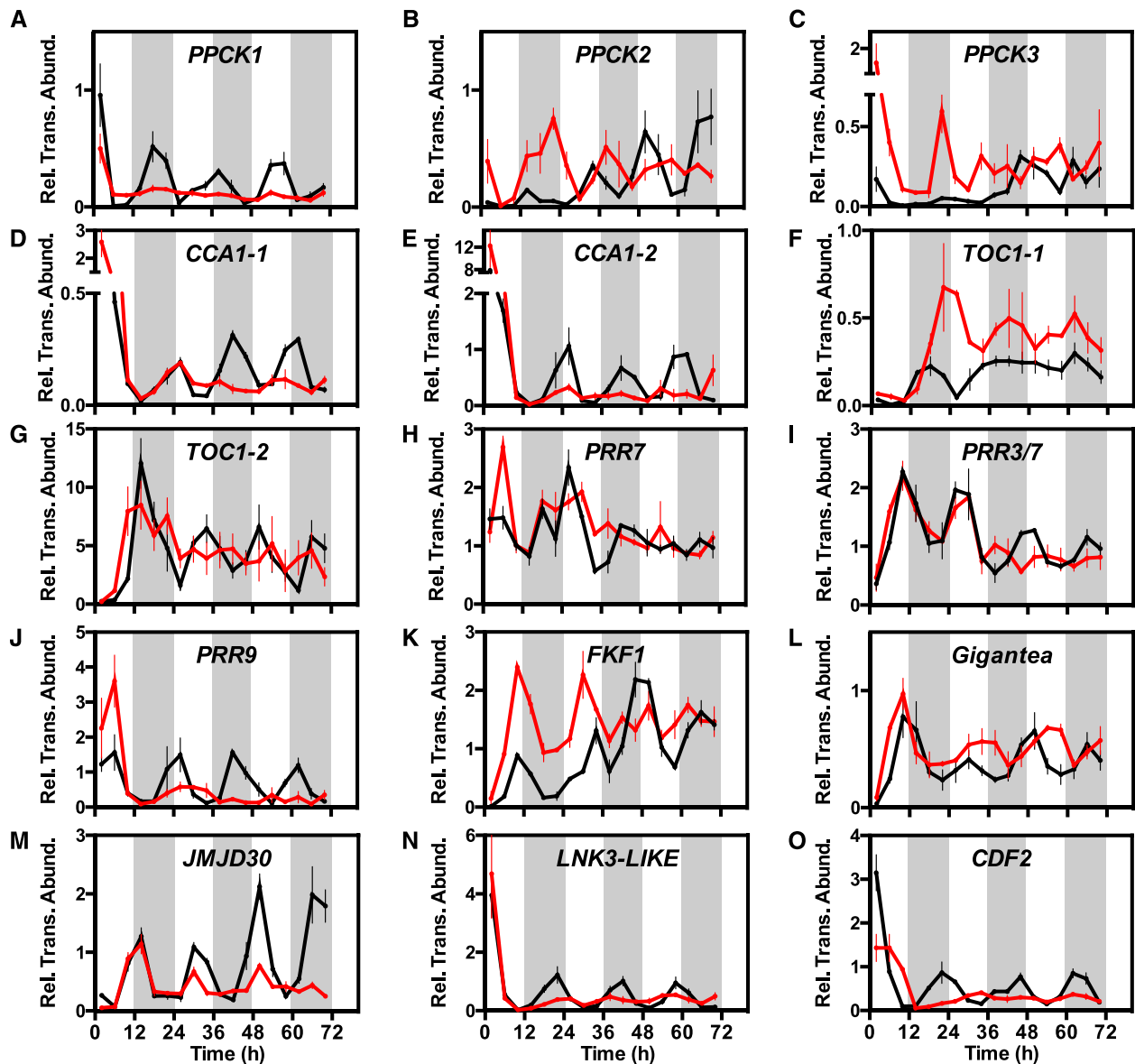


**Figure 5.** Impact of the Loss of KfPPCK1 Activity on the Light/Dark Regulation of the Transcript Abundance of Central Circadian Clock Genes *KfTOC1-1*, *KfTOC1-2*, *KfCCA1-1*, *KfCCA1-2*, *KfPRR7*, and *KfPRR37*.

Mature leaves (leaf pair 6) were sampled from three biological replicates every 4 h across the 12-h-light/12-h-dark cycle, RNA isolated, and used for real-time RT-qPCR. *TOC1-1* (A), *TOC1-2* (B), *CCA1-1* (C), *CCA1-2* (D), *PRR7* (E), and *PRR37* (F). A thioesterase/thiol ester dehydrase-isomerase superfamily (*KfTED1*) gene was amplified from the same cDNAs as a reference gene. Gene transcript abundance data represents the mean of three technical replicates for each of three biological replicates and was normalized to the reference gene (*KfTED1*); error bars represent the SE of the mean calculated for the three biological replicates. In all cases, plants were entrained under 12-h-light/12-h-dark cycles for 7 d prior to sampling. Black data are for the wild type, blue data *rPPCK1-1*, and red data *rPPCK1-3*.

a robust DF oscillation (Figure 7A). The wild-type relative amplitude error (RAE) plot (Figure 7B), and summary bar chart for the computed period length for the wild type (Figure 7C, top panel), demonstrated a short period (mean 17.8 h) from both fast Fourier transform (nonlinear least squares method) and spectral resampling analysis. The mean RAE plot revealed that the values were below the 0.6 cutoff (Figure 7C, bottom panel), supporting statistically the visibly robust and high-amplitude wild-type

short period DF rhythm (Figure 7A). Biodare analysis demonstrated that both *rPPCK1-1* (Figures 7D and 7E) and *rPPCK1-3* (Figures 7F and 7G) lost rhythm robustness through time, particularly in the later stages of the LL free-running conditions. Line *rPPCK1-1* had an intermediate effect with dampening of the rhythm occurring over several cycles (Figure 7D). The period length plot for *rPPCK1-1* suggested that its average free-running period was slightly longer than that of the wild type, but the larger



**Figure 6.** Impact of the Loss of *KfPPCK1* Activity on Circadian Clock-Controlled Gene Transcript Abundance during Constant Light and Temperature Free-Running Conditions.

(A) Circadian rhythm of *KfPPCK1* transcript abundance under constant LL conditions ( $100 \mu\text{mol m}^{-2} \text{s}^{-1}$  light at  $15^\circ\text{C}$ ) for the wild type (black line) and *rPPCK1-3* (red line).

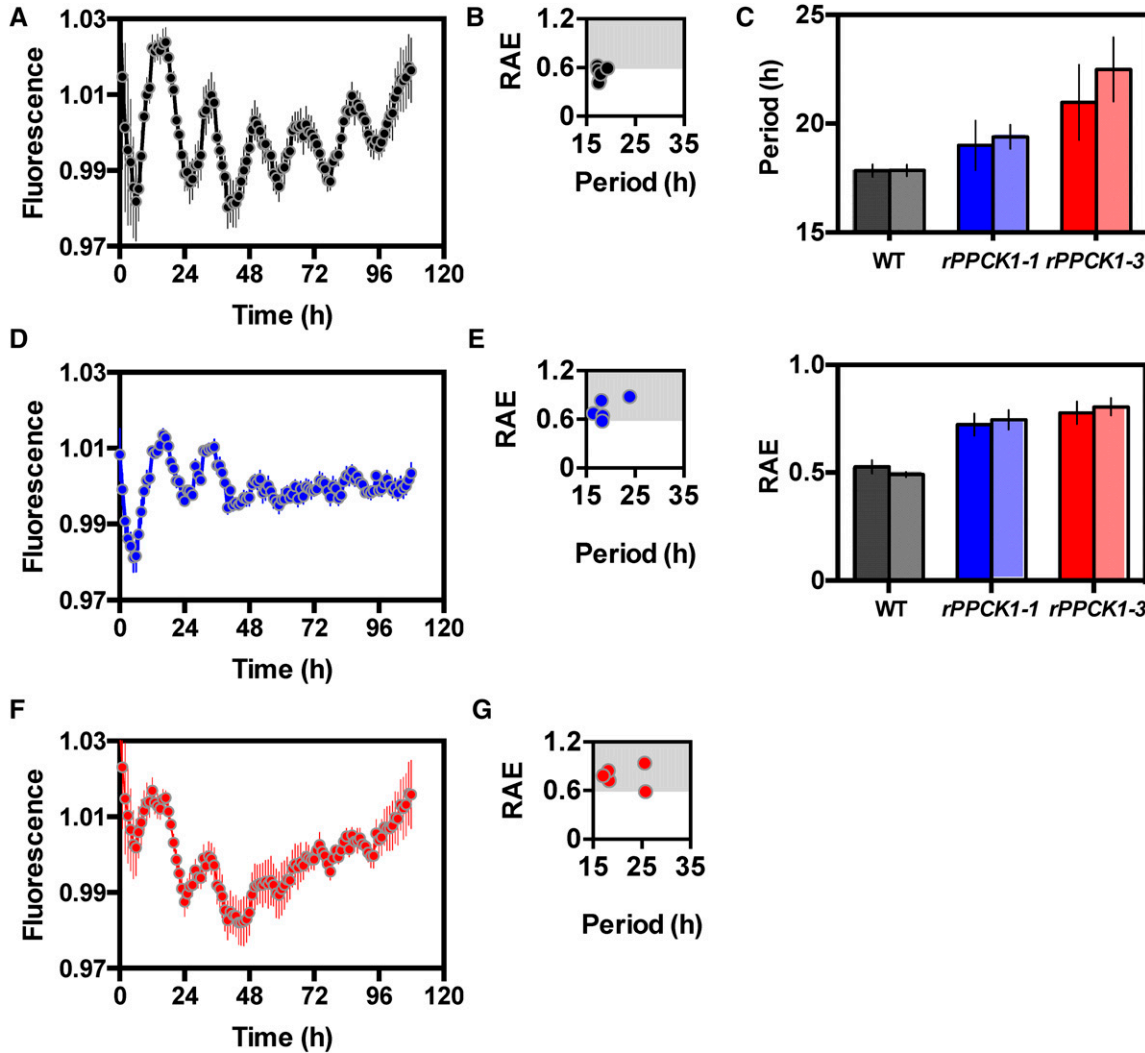
(B) to (O) *KfPPCK2* (B), *KfPPCK3* (C), *KfCCA1-1* (D), *KfCCA1-2* (E), *KfTOC1-1* (F), *KfTOC1-2* (G), *KfPRR7* (H), *KfPRR3/7* (I), *KfPRR9* (J), *KfFKF1* (K), *KfGIGANTEA* (L), *KfJMJD30* (M), *KfLNK3-like* (N), and *KfCDF2* (O). Mature leaves (leaf pair 6) were sampled from three biological replicates every 4 h under constant conditions ( $100 \mu\text{mol m}^{-2} \text{s}^{-1}$  light at  $15^\circ\text{C}$ ) for the wild type and *rPPCK1-3*. RNA was isolated and used for real-time RT-qPCR. A thioesterase/thiol ester dehydrase-isomerase superfamily gene (*KfTED1*) was amplified as a reference gene from the same cDNAs. Gene transcript abundance data represent the mean of three technical replicates for each of three biological replicates and were normalized to the reference gene (*KfTED1*); error bars represent the  $\pm$ SE of the mean calculated for the three biological replicates. In all cases, plants were entrained under 12-h-light/12-h-dark cycles prior to release into LL free-running conditions. Black data are for the wild type and red data *rPPCK1-3*.

error suggested this period estimate was not reliable (Figure 7C, top panel). The corresponding RAE values showed a much higher average for *rPPCK1-1* than the wild type due to its dampening, reflected in the wide spread of period lengths from

short to long (Figure 7E). The analysis outputs for the DF results from *rPPCK1-3* revealed a slightly more severe rhythm phenotype than *rPPCK1-1* (Figures 7F and 7G). Although the Biodare graph summarizing period length for DF rhythms in *rPPCK1-1*

and *rPPCK1-3* indicated longer DF period length for the transgenic lines relative to the wild type (Figure 7C, top panel), which was at odds with the shorter period for the gas exchange LL rhythms (Figures 3A and 3B), visual inspection of the DF rhythms revealed that there may have been a shorter period length for the very low amplitude DF rhythms in *rPPCK1-1* and

*rPPCK1-3* that were observed throughout the latter stages of the LL time course (Figures 7D and 7F). The RAE values above the 0.6 cutoff for *rPPCK1-1* and *rPPCK1-3* also indicated that the rhythms were not robust and thus that the period estimates in Figure 7C should not be overinterpreted due to their high associated RAE values.



**Figure 7.** DF Rhythms Collapsed toward Arrhythmia in Lines *rPPCK1-1* and *rPPCK1-3*.

Plants were entrained under 12-h-light/12-h-dark cycles before being transferred to constant red/ blue light under the CCD imaging camera ( $35 \mu\text{mol m}^{-2} \text{s}^{-1}$ ). DF was assayed with a 1-h time resolution for 108 h. The plots represent normalized averages for DF measured for six leaf discs sampled from three biological replicates of leaf pair 6 for each line. Error bars indicate  $\text{SE}$  of the mean calculated from three biological replicates. Black data are for the wild type, blue data *rPPCK1-1*, and red data *rPPCK1-3*.

(A) Wild-type DF rhythm under LL.

(B) RAE plot for wild-type DF rhythm.

(C) Mean period length (upper graph) and RAE plots (lower graph) for the wild type (black/gray), *rPPCK1-1* (blue/pale blue), and *rPPCK1-3* (red/pale red); the plotted values were calculated using the Biodare package for circadian rhythm analysis. In both graphs, the dark shade of the color represents data from fast Fourier transform-nonlinear least squares analysis and the paler shade represents data from spectral resampling analysis.

(D) *rPPCK1-1* DF rhythm.

(E) RAE plot for *rPPCK1-1*.

(F) *rPPCK1-3* DF rhythm.

(G) RAE plot for *rPPCK1-3*.

## DISCUSSION

### Physiological Consequences of Loss of Clock-Controlled PPC Phosphorylation in an Obligate CAM Plant

Previous work had shown that the *PPCK* gene responsible for light period phosphorylation and in vivo activation of photosynthetic PPC in the  $C_4$  species *Flaveria bidentis* was not essential for high  $C_4$  photosynthetic rates in this species (Furumoto et al., 2007). This work on a  $C_4$  species surprised the wider community because decades of prior work had argued that PPC phosphorylation by PPCK was vital for alleviating malate/aspartate inhibition of PPC in planta and thus for optimizing photosynthetic  $CO_2$  fixation in  $C_4$  and CAM species (Vidal and Chollet, 1997; Nimmo, 2003). These findings have become all the more surprising in the light of recent work on the evolution of the *PPCK* gene family in  $C_3$  and  $C_4$  species of *Flaveria* (Aldous et al., 2014). A specific, light-induced,  $C_4$ -recruited *PPCKA* gene was found in each  $C_4$  species of *Flaveria* and was found to have evolved rapidly since its recruitment to a  $C_4$ -specific function. All *Flaveria* species also possessed a *PPCKB* gene that displayed peak transcript abundance in the dark period and that was downregulated in the  $C_4$  species relative to the  $C_3$  species. Aldous et al. (2014) suggested that the  $C_4$ -recruited *ppcA* and *PPCKA* formed a coevolving substrate-kinase pair, and yet the findings of Furumoto et al. (2007) argued that *PPCKA* in *F. bidentis* was dispensable for optimal  $CO_2$  fixation via the  $C_4$  pathway. Why would there be strong evolutionary selection acting on the  $C_4$  *ppcA/PPCKA* substrate-kinase pair if the kinase and its phosphorylation of PPC were dispensable for optimal  $CO_2$  fixation via the  $C_4$  pathway? It was thus vital to investigate the physiological role, if any, of PPCK and PPC phosphorylation in relation to dark period  $CO_2$  fixation through to malate in a CAM plant.

Our findings revealed that, in the CAM species *K. fedtschenkoi*, phosphorylation of PPC during the dark period by the circadian clock-controlled *KfPPCK1* both maximized and sustained primary atmospheric  $CO_2$  fixation via PPC. Overall, our data showed that dark phosphorylation of PPC during CAM in *K. fedtschenkoi* is not a prerequisite for dark  $CO_2$  fixation, as 34% of dark  $CO_2$  fixation and 64% of nocturnal malate accumulation happened even in the absence of detectable levels of dark PPC phosphorylation in *rPPCK1-3* (Figure 2). However, *KfPPCK1* does play a critical role in fine-tuning the amount and duration of dark  $CO_2$  fixation, leading to greater malate accumulation and growth in the wild type relative to *rPPCK1-3*.

The high WUE of CAM plants is regarded as one of their key adaptive strengths and the main trait of interest for future exploitation in CAM crops and CAM biodesign for climate-resilient agriculture (Borland et al., 2009, 2014). It was thus important to compare the performance of the *rPPCK1* lines between well-watered and drought-stressed conditions. Gas exchange analysis of the *rPPCK1* lines subjected to drought stress demonstrated that both lines showed similar reductions in dark  $CO_2$  fixation relative to the wild type in comparison to the differences observed under well-watered conditions (Figure 4A). However, the *rPPCK1* lines did continue to fix atmospheric  $CO_2$  in the dark after drought stress, which contrasted strikingly with our previous findings for transgenic *K. fedtschenkoi* with reduced CAM and dark  $CO_2$  fixation due to reduced activity of NAD-ME or PPK, both of which

failed to fix  $CO_2$  for the majority of the light/dark cycle following drought stress (Dever et al., 2015).

Drought stress had a dramatic impact on both the wet and dry weight of plants for both the wild-type and the *rPPCK1* lines, but there was no significant difference between the fresh or dry weight of both the shoots and roots of wild-type and *rPPCK1* lines following drought stress (Figure 4). Thus, under our growth conditions, the increased dark  $CO_2$  fixation during drought stress achieved by the wild type relative to the *rPPCK1* lines (Figure 4A) did not lead to a measurable increase in yield for the wild type. Instead, all lines achieved little if any net  $CO_2$  fixation after prolonged drought stress, as their respiratory loss of  $CO_2$  in the light period largely balanced their fixation of  $CO_2$  in the dark period. This was reflected in the very low dry weight yields, which were >3-fold lower for drought-stressed shoots relative to the well-watered shoots (Figure 4D). It would be interesting to monitor the gas exchange of the wild-type and *rPPCK1* lines continuously throughout the drought-stress treatment, so that their gas exchange during the transition from fully hydrated to severely drought stressed could be investigated in detail. This may reveal whether or not the wild type is able to maintain net daily fixation of  $CO_2$  for longer into the drought-stress period. However, if this does occur, it was clearly not sufficient to generate a measurable difference in growth performance after the 138 d of drought treatment (Figure 4). It may be that all lines were locked down into a CAM-idling survival mode after 138 d of drought stress and that their wet and dry weights were indistinguishable because they had, by that time, respired away the small but significant differences in their shoot and root dry biomass that were evident under well-watered conditions.

### Molecular and Biochemical Phenotypes of *rPPCK1* Lines

Both *rPPCK1* lines possessed lower *KfPPCK1* transcript levels than the wild type throughout the dark period (Figure 1A). *rPPCK1-1* and *rPPCK1-3* had similar *KfPPCK1* transcript levels at both the beginning and end of the light period, but possessed a large difference in transcript abundance in the middle of the dark period, when line *rPPCK1-1* achieved a significantly higher transcript level than line *rPPCK1-3* (18:00, 6 h into the 12-h dark period; Figure 1A). This was consistent with the higher PPC phosphorylation and apparent  $K_m$  of PPC for L-malate detected in line *rPPCK1-1* relative to line *rPPCK1-3*, specifically in the middle of the dark period at 18:00 (Figure 1B), and the greater dark period  $CO_2$  fixation, malate accumulation, and starch turnover achieved by this line (Figures 2A to 2D). Overall, our biochemical measurements revealed a dosage-dependent impact of the level of *KfPPCK1* transcripts each night on the biochemical correlates of CAM.

### Impact of Silencing *KfPPCK1* on Circadian Rhythms of CAM $CO_2$ Fixation and the Operation of the Central Circadian Oscillator

Since the first discovery of PPCK activity and its circadian control in 1991 (Carter et al., 1991), through the subsequent cloning and characterization of the gene encoding this remarkable protein kinase in 1999 (Hartwell et al., 1999), there has been a long-held assumption that this circadian clock-controlled kinase is crucial

for the temporal coordination and optimization of nocturnal atmospheric CO<sub>2</sub> fixation in CAM species, and the associated persistent circadian rhythm of CO<sub>2</sub> exchange observed under constant conditions in CAM species such as *K. fedtschenkoi* and *K. daigremontiana* (Wilkins, 1992; Nimmo, 2003; Wyka et al., 2004; Hartwell, 2006). However, it has not been possible previously to perturb genetically the level of PPCK activity and dark period PPC phosphorylation in a CAM species; thus, it has not been possible to test the validity of this long held assumption about the pivotal role of PPCK in the circadian control of CAM.

By silencing the CAM-associated *KfPPCK1* gene with an RNAi construct, we have been able to test in planta the importance of *KfPPCK1* for the circadian rhythm of CO<sub>2</sub> exchange. The loss of dark period PPC phosphorylation in line *rPPCK1-3* led to a rapid decline in the circadian rhythm of CO<sub>2</sub> exchange when leaves or whole plants pre-entrained under 12-h-light/12-h-dark cycles were released into LL free-running conditions (Figures 3A, 3B and 4A). After 25 d of drought stress, the weak CO<sub>2</sub> exchange rhythms displayed by the well-watered *rPPCK1* lines (Figure 3B) collapsed to arrhythmia more rapidly (Figure 4A). In particular, line *rPPCK1-3* became arrhythmic and continued a steady rate of respiratory loss of CO<sub>2</sub> for the majority of the LL time course (Figure 4A), whereas the drought-stressed wild type continued with a persistent circadian rhythm of CO<sub>2</sub> fixation throughout the LL experiment (Figure 4A). Furthermore, silencing *KfPPCK1* also led to DF rhythms collapsing toward arrhythmia, revealing that a distinct circadian clock output that originates inside the chloroplast was also perturbed in the *rPPCK1* lines (Figure 7). These results revealed that wild-type levels of circadian clock controlled PPCK are essential for the amplitude, robustness, and persistence of the CO<sub>2</sub> fixation rhythm associated with CAM and the DF rhythms in the chloroplast in *K. fedtschenkoi*.

In light of the weakened rhythmicity of CO<sub>2</sub> exchange in the *rPPCK1* lines, it was important to investigate the robustness of transcript abundance oscillations for components of the central molecular oscillator that underpin plant circadian rhythms (Figures 5 and 6). Wild-type *K. fedtschenkoi* displayed robust oscillations in the transcript abundance of *KfCCA1-1*, *KfCCA1-2*, *KfTOC1-1*, and *KfTOC1-2*, but rhythms in the transcript abundance of *KfCCA1-1*, *KfCCA1-2*, and *KfTOC1-2* collapsed rapidly toward arrhythmia under LL in the *rPPCK1-3* line (Figures 6D, 6E, and 6G). This revealed that multiple components of the core molecular oscillator failed to maintain wild-type levels of robust rhythmicity in the absence of *KfPPCK1*. In contrast, *KfTOC1-1*, *KfPRR7*, *KfFKF1*, and *KfGI* were induced and displayed a rhythm with a greater amplitude than the wild type under LL in line *rPPCK1-3* (Figures 6F, 6H, 6K, and 6L), revealing that certain sub-components of the core molecular circadian oscillator continued to function with a robust circadian rhythm in the absence of *KfPPCK1* activity. These are paradigm-changing findings because *KfPPCK1* has previously been regarded as a circadian clock-controlled output pathway component that is vital for coupling the central clock to CAM. However, our current findings reveal that some components of the central clock itself stop working correctly in the absence of *KfPPCK1*. This suggests either that *KfPPCK1* is itself part of the central oscillator in *K. fedtschenkoi*, although this seems highly unlikely considering the known function of PPCK1 in PPC phosphorylation, or, more likely,

that the failure to phosphorylate PPC in the dark leads to metabolic changes that influence the operation of the core oscillator, perhaps via a mechanism not dissimilar to that proposed for sugars acting as signals from photosynthesis that can entrain the central clock in Arabidopsis (Haydon et al., 2013).

It will be important to investigate the wider metabolic and/or gene expression changes associated with the loss of PPCK and PPC phosphorylation in these lines. Such investigations should provide valuable insights into the types of signals that could link CAM-associated metabolites to the central clock during the daily operation of the CAM pathway. It is particularly noteworthy in this respect that we also saw a loss of central clock gene *TOC1* transcript oscillations in *K. fedtschenkoi* transgenic RNAi lines with reduced activity of NAD-ME (Dever et al., 2015). Those lines turned over only a small proportion of their malate each day and had a much greater reduction in starch accumulation and turnover than the *rPPCK1* lines reported here. This suggests that the metabolic and/or gene regulatory changes that led to dampening of oscillations for a subset of central clock genes in these two distinct sets of *K. fedtschenkoi* transgenic lines may be more complex than a direct impact of malate or carbohydrate levels on clock operation. However, when we examined sucrose, fructose, and glucose levels over the 12-h-light/12-h-dark cycle in the *rPPCK1* lines reported here, we discovered that line *rPPCK1-3* displayed an almost 50% reduction in the 2 h after dawn-phased peak of sucrose detected in the wild type (Figure 2E).

Taking into consideration the fact that sucrose has been demonstrated as a central clock-entraining metabolite in the C<sub>3</sub> species Arabidopsis (Haydon et al., 2013), our findings suggest that sucrose is a strong candidate for a metabolite linking the perturbations in CAM and core clock rhythms reported here for the *rPPCK1* lines (Figure 2E). It is particularly noteworthy in this context that *KfPRR7* and *KfPRR37* transcript levels were increased in *rPPCK1-3* at 2 and 10 h after dawn, respectively, relative to the wild type (Figures 5E and 5F). For *KfPRR7*, this change in transcript level was coincident with the decrease in the post-dawn sucrose peak (Figure 2E). Haydon et al. (2013) proposed that *PRR7* could be a key clock gene functioning in the entrainment of the central oscillator by sucrose signals. However, it is noteworthy that our *KfPRR7* ortholog displayed a transcript peak phased to 2 h after dawn, coincident with the daily sucrose peak, whereas the Arabidopsis *PRR7* transcript level peaked at 8 h into the 12-h light period and was also strongly correlated with the daily sucrose peak, which occurs in the latter part of the light period in Arabidopsis (Haydon et al., 2013). Transcript levels of our *KfPRR37* transcript did peak at an equivalent time (10:00, 2 h prior to dusk) to *PRR7* in Arabidopsis and reached its daily peak in *rPPCK1-3* some 4 h earlier than the wild type, which peaked at 14:00, 2 h into the dark (Figure 5F). These striking similarities in the timing and strong correlation of peak sucrose and peak *PRR7* transcript levels in both *K. fedtschenkoi* and Arabidopsis provide support for the proposal that sucrose is a key clock entraining signal in *K. fedtschenkoi* and that the decrease in the 2 h post-dawn sucrose peak in *rPPCK1-3* (Figure 2E) feeds back to perturb core clock gene regulation (Figure 6), potentially via *KfPRR7* and *KfPRR37* (Figures 5E and 5F).

Our data support the proposal that, in the CAM leaf, the circadian clock is set by both light and sucrose. Sucrose has been

demonstrated to be a product of CAM-derived starch degradation during the hours at the end of the phase I dark period, although sucrose was also shown to accumulate further during phase II after dawn as a direct result of Rubisco-mediated fixation of atmospheric CO<sub>2</sub>, as is classically observed in the late afternoon in C<sub>3</sub> plants such as *Arabidopsis* (Wild et al., 2010; Haydon et al., 2013). Both of the *KfCCA1* genes and one of the two *KfTOC1* genes were downregulated and displayed dampening rhythmicity in the *rPPCK1-3* plants under LL conditions (Figure 6), and this may be due to the *rPPCK1-3* plants being in a state of metabolic starvation after dawn. In C<sub>3</sub> leaves of *Arabidopsis* during starvation, *PRR7* represses *CCA1* and derepresses the *TOC1/GI* loop (Dalchau et al., 2011; Haydon et al., 2013, 2015). The *rPPCK1-3* transgenic lines of *K. fedtschenkoi* displayed derepression and rhythmicity for *KfPRR7*, *KfTOC1-1*, *KfGI*, and *KfFKF1*, supporting the proposal that the reduction in the post-dawn sucrose peak and associated increase in *KfPRR7* and *KfPRR37* may lead to the derepression of a *TOC1-1/GI* loop in *K. fedtschenkoi*. However, in *K. fedtschenkoi*, while *KfTOC1-1* was derepressed, which is similar to the *TOC1* regulation reported in *Arabidopsis*, *KfTOC1-2* was repressed, suggesting that in CAM leaves of *Kalanchoë* there are distinct suboscillators of the central circadian clock involving *KfTOC1-1* and *KfTOC1-2*. We cannot at this stage rule out the possibility that these striking differences in the regulation of core circadian clock genes in line *rPPCK1-3* were a result of differential regulation of the two *TOC1* genes occurring in different cell types of the leaf.

We have previously demonstrated the utility of *K. fedtschenkoi* as a model system for elucidating the in planta functioning of candidate core metabolic enzymes associated with CAM (Dever et al., 2015; Hartwell et al., 2016). Our findings here provide in planta confirmation of the importance of the regulatory protein kinase PPCK and its circadian rhythm for optimal and maximal dark CO<sub>2</sub> fixation. Plants lacking *KfPPCK1* also failed to maintain robust rhythmicity of the transcript oscillations of several core clock components, including both isogenes of *KfCCA1* and one isogene of *KfTOC1*. A key criterion for a gene to be classified as a core clock component is that misregulation of the gene should cause arrhythmia within the oscillator itself. Our data fulfill this criterion to some extent for *KfPPCK1*; thus, our data could be interpreted as evidence that *KfPPCK1* operates as a core component of the central circadian clock in *K. fedtschenkoi*. However, we do not favor that explanation. Instead, our data for daily sucrose oscillations and *KfPRR7*, *KfPRR37*, *KfTOC1-1*, and *KfGI* transcript regulation support the proposal that the pathway of metabolic regulation suggested to integrate photosynthetic sugar metabolism with the clock in *Arabidopsis* plays a role in the partial loss of rhythmicity observed in this *K. fedtschenkoi rPPCK1-3* line (Haydon et al., 2013). Alternatively, malate changes in the *rPPCK1-3* line may influence the clock, as malate has been demonstrated to function to influence gene regulation (Finkemeier et al., 2013).

This work extends the categories of CAM-associated genes for which valuable functional insights have been gained using transgenic *K. fedtschenkoi*, moving beyond core CAM enzymes to include the circadian clock-controlled posttranslational regulatory step catalyzed by PPCK1. Based on these findings, we predict that transgenic approaches applied in *K. fedtschenkoi* will also

prove useful for demonstrating the functional role of other proposed CAM regulators, such as PPK regulatory protein (Dever et al., 2015), or pyruvate dehydrogenase kinase (Thelen et al., 2000), or as yet unknown transcription factors and other regulatory proteins that likely function in the circadian clock output pathway that couples the core molecular oscillator to the regulation of CAM enzymes via proteins such as PPCK (Hartwell, 2006; Borland et al., 2009; Hartwell et al., 2016).

### Relevance of These Findings to Efforts to Engineer CAM into C<sub>3</sub> Species

Our findings are vitally important to current and ongoing efforts to engineer the CAM pathway into C<sub>3</sub> crops using plant synthetic biology approaches because they demonstrate that expressing high levels of PPC alone in leaf mesophyll cells will not be sufficient to achieve optimized operation of CAM dark CO<sub>2</sub> fixation. Expressing elevated levels of PPC may yield plants that can fix some atmospheric CO<sub>2</sub> in the dark and accumulate a certain level of malate by dawn, which would be a significant advance in and of itself. However, our data for the *KfPPCK1* RNAi lines demonstrate that in order for an engineered CAM system to work with the full efficiency demonstrated by existing CAM species, C<sub>3</sub> crops will need to be engineered to express both high levels of PPC in leaf mesophyll cells and a circadian clock-controlled PPCK that peaks in the dark period and achieves a 10-fold increase in the K<sub>i</sub> of the PPC for malate.

We propose that the PPCK and PPC used for this forward engineering work should ideally be a pre-evolved substrate-kinase pair, as has been suggested to have coevolved in the C<sub>4</sub> *Flaveria* system (Aldous et al., 2014). For example, *KfPPCK1* (GenBank KM078709) and *KfPPCK1* (GenBank AF162662) from *K. fedtschenkoi* would be an ideal substrate-kinase pair for the forward engineering of CAM into C<sub>3</sub> species, although care would need to be taken with promoter choice for the transgenes in order to ensure that the *KfPPCK1* gene was expressed to very high levels in the leaf mesophyll cells, and *KfPPCK1* was expressed under circadian clock control to peak in the middle of the dark in leaf mesophyll cells.

## METHODS

### Plant Materials

Unless otherwise stated, *Kalanchoë fedtschenkoi* Hamet et Perrier plants were propagated clonally from stem cuttings or leaf margin adventitious plantlets using the same clonal stock originally obtained from the Royal Botanic Gardens, Kew, by Malcolm Wilkins (Wilkins, 1959). Plants were grown and entrained to light/dark cycles as described by Dever et al. (2015).

### Time-Course Experiments

For light/dark time-course experiments, opposite pairs of LP6 (where leaf pair 1/LP1 are the youngest leaf pair flanking the shoot apical meristem and leaf pairs are counted down the stem) were collected every 4 h over a 12-h-light (450 μmoles m<sup>-2</sup> s<sup>-1</sup>, 25°C, 60% humidity)/12-h-dark (15°C, 70% humidity) cycle, starting 2 h (02:00) after the lights came on at 00:00 h using plants raised and entrained as described by Dever et al. (2015). For constant light, constant temperature, constant humidity (LL) free-running circadian time-course experiments, plants were entrained in 12-h-light/

12-h-dark conditions for 7 d and switched to LL after a dark period as described by Dever et al. (2015). For LL, the constant conditions were as follows: light  $100 \mu\text{mol m}^{-2} \text{s}^{-1}$ , temperature  $15^\circ\text{C}$ , and humidity 70%. LP6 were sampled every 4 h from three individual (clonal) plants, starting at 02:00 (2 h after lights on). All sampling involved immediate freezing of leaves in liquid nitrogen. Samples were stored at  $-80^\circ\text{C}$  until use.

### Generation of Transgenic *K. fedtschenkoi* Lines

An intron containing hairpin RNAi construct was designed to target the silencing of the CAM-induced and clock-controlled *KfPPCK1* gene (GenBank accession AF162661) (Hartwell et al., 1999). A 359-bp fragment was amplified from CAM leaf cDNA using high fidelity PCR with KOD Hot Start DNA Polymerase (Merck) and the primers *KfPPCK1* RNAi F 5'-CACCGCGCAGGACATATTTGAGG-3' and *KfPPCK1* RNAi R 5'-GGCCGAATCCCTCACTCTGCTTC-3'. The amplified fragment spanned the 3' end of the coding sequence and extended into the 3' untranslated region to ensure specificity of the silencing to a single member of the *K. fedtschenkoi* PPCK gene family. Alignment of the 359-bp region with the homologous region from the three other PPCK genes in the *K. fedtschenkoi* genome demonstrated that none of the other PPCK genes shared any 21 nucleotide stretches that were an exact match for *KfPPCK1*. PCR products were cloned into the pENTR/D Gateway-compatible entry vector via directional TOPO cloning (Life Technologies) and recombined into the intron containing hairpin RNAi binary vector pK7GWIWG2(II) (Karimi et al., 2002) using LR Clonase II enzyme mix (Life Technologies). Constructs were confirmed by DNA sequencing and introduced into *Agrobacterium tumefaciens* strain GV3101 using the freeze-thaw method (Höfgen and Willmitzer, 1988).

*Agrobacterium*-mediated stable transformation of *K. fedtschenkoi* was achieved using the tissue culture based method described by Dever et al. (2015).

### High-Throughput Leaf Acidity and Starch Content Screens

Leaf acidity (as a proxy for leaf malate content) and leaf starch content were screened with leaf disc stains using chlorophenol red and iodine solution, respectively, at both dawn and dusk, as described by Cushman et al. (2008). For each transgenic line, leaf discs were sampled in triplicate at 1 h before dawn and 1 h before dusk and stained in a 96-well plate format.

### Net CO<sub>2</sub> Exchange

Gas exchange measurements were performed using a six-channel, custom-built infrared gas analyzer system (PP Systems), which allowed the individual environmental control (CO<sub>2</sub>/H<sub>2</sub>O) and measurement of rates of CO<sub>2</sub> uptake for each of six gas exchange cuvettes with measurements collected every 10 min (described in full in Dever et al., 2015). All experiments were repeated at least three times using three separate individual young plants (eight leaf pairs), or detached LP6 from three separate clonal plants, and representative gas exchange traces are shown. Wild-type and *rPPCK1* lines were compared in neighboring gas exchange cuvettes during each run, such that the data are directly comparable between each line. For example, many runs included two cuvettes containing the wild type, two cuvettes containing *rPPCK1-1*, and two with *rPPCK1-3*.

### Drought-Stress Experiments

For the drought-treatment gas exchange experiments, clonal leaf margin plantlets from wild-type and the *rPPCK1* lines were grown to the 8-leaf-pair stage using 150 mL of our standard compost/perlite mix (Dever et al., 2015) and maintained well-watered throughout their development. Water was withheld for 25 d under our standard 12:12 LD growth conditions according to Dever et al. (2015).

### Leaf Malate, Starch, and Sucrose Content

Leaf pair 6 (LP6; full CAM in the wild type) from mature plants was sampled into liquid nitrogen at the indicated times and stored at  $-80^\circ\text{C}$  until use. The frozen leaf samples were prepared and assayed for malate and starch as described by Dever et al. (2015) using the standard enzyme-linked spectrophotometric methods for assaying malate (Möllering, 1985) and starch (Smith and Zeeman, 2006).

The concentrations of sucrose, D-fructose, and D-glucose were determined using enzyme-linked spectrophotometric assays according to the manufacturer's instructions (K-SUFRG sucrose/D-fructose/D-glucose assay kit; Megazyme International). This involved the stepwise enzymatic conversion of each sugar to glucose-6-phosphate (G6P) and subsequent quantification by measuring NADPH production at 340 nm by following oxidation of the G6P in the presence of NADP<sup>+</sup> and G6P-dehydrogenase.

### Total RNA Isolation and RT-qPCR

Total RNA was isolated from 100 mg of frozen, ground leaf tissue using the Qiagen RNeasy kit following the manufacturer's protocol with the addition of  $13.5 \mu\text{L } 50 \text{ mg mL}^{-1}$  PEG 20,000 to the 450  $\mu\text{L}$  RLC buffer used for each extraction. cDNA was synthesized from the total RNA using the Qiagen Quantitect RT kit according to the manufacturer's instructions (Qiagen). The resulting cDNA was diluted 1:4 with molecular biology grade water prior to use in RT-qPCR. Transcript levels were determined using three technical replicates and three biological replicates with SensiFAST SYBR No Rox kit (Bioline) in an Agilent MX3005P qPCR system cycler using the following program:  $95^\circ\text{C}$  for 2 min, 40 cycles of  $95^\circ\text{C}$  for 5 s,  $60^\circ\text{C}$  for 10 s, and  $72^\circ\text{C}$  for 10 s, and primers designed with Primer3. Primer specificity was checked using BLAST searches, and primer efficiency was checked using a melting curve profile standard curve. In every PCR plate an interplate calibrator (made from a pool of RNAs assembled from *K. fedtschenkoi* wild-type leaf pair 6 samples taken every 4 h over a light/dark cycle) was run in triplicate in order to correct any interplate variation. In addition, nontemplate controls were included in every plate to confirm the absence of contamination. Each biological replicate was assayed in triplicate, and the relative abundance of each gene was determined using Agilent MxPRO QPCR software using the manufacturer's instructions (Agilent Technologies). The results were normalized to a *K. fedtschenkoi* ortholog of a Thioesterase/thiol ester dehydrase-isomerase (TED1) superfamily protein (Phytozome Kaladp0068s0118.1; Arabidopsis ortholog AT2G30720.1). Primers for RT-qPCR analyses are listed in Supplemental Table 1.

### Immunoblotting

Total protein extracts of *K. fedtschenkoi* leaves were prepared according to Dever et al. (2015). One-dimensional SDS-PAGE and immunoblotting of leaf proteins was performed following standard methods. Blots were developed using the ECL system (GE Healthcare). Immunoblot analysis was performed using antisera to PPC (raised against PPC from *K. fedtschenkoi*, kindly supplied by H.G. Nimmo, University of Glasgow) (Nimmo et al., 1986) and the phosphorylated form of PPC (raised against a phospho-PPC peptide from barley [*Hordeum vulgare*] and kindly supplied by Cristina Echevarría, Universidad de Sevilla, Spain) (González et al., 2002; Fera et al., 2008).

### Determination of the Apparent K<sub>i</sub> of PPC for L-Malate in Rapidly Desalted Leaf Extracts

Rapidly desalted leaf extracts were prepared as described by Carter et al. (1991) using LP6 sampled from three biological replicates of the wild type, *rPPCK1-1*, and *rPPCK1-3* at 10:00 (2 h before the end of the 12-h light period) and 18:00 (middle of the 12-h dark period). The apparent K<sub>i</sub> of PPC activity for feedback inhibition by L-malate was determined for each rapidly

desalted extract according to the method described by Nimmo et al. (1984), except that the range of L-malate concentrations added to the different PPC assays was 0, 0.2, 0.5, 0.7, 1.0, 2.0, 3.0, 5.0, 8.0, and 10.0 mM. The concentration of malate needed to achieve 50% inhibition of the initial PPC activity (apparent  $K_i$ ) was determined using the plots shown in Supplemental Figure 1.

### Dry Weight Growth Measurements

Mature plants were grown from developmentally synchronized clonal leaf plantlets in greenhouse conditions for 138 d, harvested as separated aboveground and belowground tissues, and dried in an oven at 60°C until they reached a constant weight. Weights were measured to two decimal places in grams using a fine balance. Seven individual clonal plants were grown for each line (Dever et al., 2015).

### DF Measurements

The imaging system for DF was identical to the luciferase and delayed fluorescence imaging system described previously (Gould et al., 2009). DF was quantified using Imaris (Bitplane) to measure mean intensity for specific regions within an image. Background intensities were calculated for each image and subtracted, to give a final DF measurement (Gould et al., 2009).

### DF Rhythm Analysis

*K. fedtschenkoi* plants were grown in greenhouse conditions for 4 months and then entrained in 12-h light/25°C: 12-h dark/15°C cycles in a Snijders Microclima MC-1000 (Snijders Scientific) growth cabinet for 7 d as described previously (Dever et al., 2015). At dawn on the 8th day, 1.5-cm leaf discs were punched from each of leaf pair 6 for three biological replicates (i.e., two leaf discs from each biological replicate, totaling six leaf discs per line) and placed with a few drops of distilled water in one of 25 2 × 2-cm wells in a 10 × 10-cm square Petri dish. The drops of distilled water were sufficient to keep the leaf discs hydrated but were not enough for the discs to float and thus move, which would make the DF image analysis extremely challenging. The Petri dish was placed in the imaging system at 22°C in constant red-blue light (LL). DF images were collected every hour for 108 h as described previously (Gould et al., 2009). The DF images were processed as described by Gould et al. (2009). The luminescence was normalized by subtracting the Y value of the best straight line from the raw Y value. Biodare was used to carry out fast Fourier transform (nonlinear least-square) analysis and spectral resampling on each DF time-course series using the time window from 24 to 108 h in order to generate period estimates and calculate the associated RAE (Moore et al., 2014; Zielinski et al., 2014). The first 24 h of the data were excluded as this represents a bedding-in period for the leaves when transition effects, resulting from the leaves moving from light/dark cycles to constant light and temperature (LL), may be influencing the data.

### Accession Numbers

Sequence data associated with this article have been deposited in the GenBank/EMBL data libraries under the following accession numbers: *KfPPCK1*, AF162662; *KfTOC1-1*, KM078716; *KfCCA1-1*, KM078717; *KfCCA1-2*, KM078718; *KfPPCK2*, KM078720; *KfGI*, KM078724; and *KfTOC1-2*, KM078726. Other accession numbers and gene IDs are presented in Supplemental Table 1.

### Supplemental Data

**Supplemental Figure 1.** Impact of silencing *KfPPCK1* on the apparent  $K_i$  of PPC for L-malate in rapidly desalted leaf extracts.

**Supplemental Table 1.** Primers used for reverse transcription-quantitative PCR.

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### AUTHOR CONTRIBUTIONS

J.H., S.F.B., L.V.D., and P.D.G. designed the research. S.F.B. performed all experiments except the stable transformations, immunoblots, starch determinations, growth yield drought experiment, and delayed fluorescence measurements. L.V.D. performed the immunoblots, starch determinations, and the growth yield drought experiment. J.K. generated the binary constructs and carried out the stable transformation, regeneration, and initial screening of the *K. fedtschenkoi rPPCK1* transgenic lines, and also collaborated with L.V.D. on the growth yield drought experiment. P.D.G. carried out the delayed fluorescence experiments in collaboration with S.F.B. S.F.B., L.V.D., P.D.G., and J.H. analyzed the data. S.F.B. and J.H. wrote the manuscript.

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