# Phototropin connects blue light perception to starch metabolism in green algae

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Abstract: In photosynthetic organisms light acts as an environmental signal to control their development and physiology, and as energy source to drive the conversion of CO<sub>2</sub> into carbohydrates used for growth or storage. The main storage carbohydrate in green algae is starch, which accumulates during the day and is broken down at night to meet cellular energy demands. The signalling role of light quality in the regulation of starch accumulation remains unexplored. Here, we report that in the model green alga *Chlamydomonas reinhardtii* blue light perceived by the photoreceptor PHOTOTROPIN causes dephosphorylation of the PHOTOTROPIN-MEDIATED SIGNALLING KINASE 1 that then suppresses starch accumulation by inhibiting the expression of GLYCERALDEHYDE-3-PHOSPHATE DEHYDROGENASE. Our results provide an in-depth view of how photoreceptor-mediated signalling controls microalgal carbon metabolism.

**One-Sentence Summary:** Blue light perception by PHOTOTROPIN triggers kinase-mediated signaling to inhibit starch accumulation in the green alga *Chlamydomonas*.

#### Introduction

Photosynthetic microalgae convert light into chemical energy in the form of ATP and NADPH to fuel the CO<sub>2</sub> fixation in the Calvin–Benson cycle during photosynthesis (*I*). Light is also a spatiotemporal signal that regulates important cellular functions, including: gene expression, sexual life cycle, phototaxis, and photoprotection (2–9). This is achieved through a network of specialized photoreceptors (*I*0). Fixed CO<sub>2</sub> in combination with nitrogen is used to synthesize amino acids—the building blocks of proteins that drive biochemical reactions. It is also employed in the synthesis of cellular reserves that ensures carbon and energy supply during waning period. The most abundant carbon reserve in the model photosynthetic green microalga *Chlamydomonas reinhardtii* is starch; its synthesis occurs during the day and its degradation is triggered at night to sustain energy-demanding cellular functions (*II*). Little is known about the molecular mechanisms underlying the control exerted by light on carbon storage in microalgae, and current knowledge is limited to factors impacting starch accumulation under adverse environmental conditions, such as: nitrogen (*I2*) and phosphorus (*I3*) limitation.

A link between light perception and starch accumulation has been suggested in vascular plants. For instance, Arabidopsis mutants devoid of the red/far-red photoreceptor phytochrome B have impaired carbon partitioning and although they have reduced CO<sub>2</sub> uptake, they over-accumulate daytime sucrose and starch at the expense of growth (14). Further, the blue light receptor PHOTOTROPIN (PHOT) has been found to mediate starch degradation in guard cells in the light, thus energizing stomatal opening in Arabidopsis (15). Despite early findings reporting that light quality impacts carbohydrate accumulation and metabolism in green algae (16-18), the underlying molecular mechanism connecting light perception to starch accumulation remains unexplored. Here we address this gap by combining genetics, proteomics and phosphomimetics to unveil a signalling cascade linking blue light perception by PHOT with starch accumulation in Chlamvdomonas. PHOT-dependent phosphorylation at a single serine PHOTOTROPIN-MEDIATED SIGNALLING KINASE 1 (PMSK1) represses GAP1 (also known as GAPDH; glyceraldehyde-3-phosphate dehydrogenase), which we found to act as an enhancer of starch metabolism.

## The *phot* mutant is a starch hyperaccumulator

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We compared starch levels in wild type CC125 cells (WT) exposed to white and red light, and found that red light favored starch accumulation, in agreement with prior work (16). Interestingly, when low fluence blue light was superimposed on red light, the beneficial effect of red light was lost and cells accumulated starch levels similar to those of cells exposed to white light (Fig. S1). Therefore, we reasoned that blue light acts as a repressor of starch accumulation likely via a blue light receptor. To test this hypothesis, we measured starch content in WT and mutant cells lacking different blue-light receptors, including: acry, lacking the ANIMAL-TYPE CRYPTOCHROME ((2), generated in this study), pcry lacking the PLANT-TYPE CRYPTOCHROME ((19), generated in this study), the double acrypcry mutant (generated in this study) and the phot mutant, devoid of PHOTOTROPIN (generated in (20)). When grown asynchronously under continuous light, the starch content of the cryptochrome mutants was indistinguishable from that of the WT (Fig. 1A) and the same was true when cultures were synchronized under 12h light/12h dark cycles (Fig. S2), favoring cryptochrome accumulation in Chlamydomonas (2, 19). In contrast to the cryptochrome mutants, the *phot* mutant accumulated approximately three times more starch than the WT under continuous illumination (Fig. 1A). This phenotype was rescued by ectopic expression of the WT PHOT gene; it was fully rescued in strain phot-C1 (21) (Fig. 1B), which accumulates PHOT protein at WT levels (Fig. S3A), and partially rescued in strain phot-C2 (Fig. 1B), which accumulates PHOT protein but to a lesser extent than the WT (Fig. S3A). Complete rescue of the phenotype was also achieved by complementation of a permanently active PHOT mutant lacking the LOV sensory domains (phot-kin strain; Fig. 1B, S3A and B); a phot-kin strain in a different genetic background has previously been shown to exhibit PHOT activity in a lightquality-independent manner (22). The accumulation of high levels of starch in the phot mutant was also confirmed by transmission electron microscopy (TEM), revealing that the chloroplast of phot is filled with starch while the pyrenoid is surrounded by very thick starch sheaths, in contrast to the chloroplast of the WT and phot-C1 strains grown under 12h light/12h dark synchronized conditions (Fig. 1C).

We further explored the link between PHOT and starch accumulation in synchronized cultures under white, blue or red light. Starch accumulated during the light phase and degraded during the dark phase, in accord with (23, 24), under all light qualities and in all different strains tested (Fig. 1D). We found that starch levels in WT are higher under red light, where PHOT is inactive, than under white or blue, where PHOT is active (Fig. 1D). In the *phot* mutant, starch levels are high in all three light qualities, while the fully complemented phot-C1 behaves like WT and the partially complemented phot-C2 is intermediate between WT and phot. In contrast to the phot mutant, starch levels in strain phot-kin, in which PHOT is active, are low and unaffected by the light quality (Fig. 1D). These data (Fig. 1) indicate that PHOT acts as suppressor of starch accumulation in Chlamydomonas. We provided further evidence for this claim by using another set of phot mutants in a different genetic background, the cw15-302 (22, 25). In accordance with the data in Fig. 1, we found that the cw15-302 phot mutant accumulated more starch than cw15-302 (Fig. S3C). This phenotype was only partially rescued in the complemented *pphot* strain, which accumulates lower levels of PHOT protein than the cw15-302 WT strain (22), and was fully rescued in the cw15-302 phot mutant expressing the PHOT kinase domain (pkin strain; Fig. S3C). Conversely, complementation of cw15-302 phot cells with truncated gene carrying only the photosensory domains LOV1 and 2 (ploy strain) did not rescue starch accumulation. Importantly, we observed a similar phenotype when the cwl5-302 phot mutant was complemented with the dead kinase PHOT (pkin-D strain; Fig. S3C), demonstrating that the suppression of starch accumulation by

PHOT requires its kinase activity. Growth (**Fig. S4A**) and photosynthesis (**Fig. S4B**) were not affected by the overaccumulation of starch in *phot* (**Fig. 1C**). Finally, *phot* showed no difference to WT in terms of total protein (**Fig. S5**), lipid content (**Fig. S6**) and composition (**Fig. S7**).

# GAP1 is a key regulator of starch metabolism and is controlled by PHOT

To gain more insight into the molecular mechanism underlying the PHOT-mediated light perception and starch accumulation, we applied mass spectrometry (MS)-based quantitative proteomics to compare WT and *phot* cells grown in asynchronous photoautotrophic conditions under continuous white light. Gene ontology enrichment analyses revealed that carbohydrate metabolic processes are upregulated in the phot mutant (Fig. 2A). We notably found that GLYCERALDEHYDE-3-PHOSPHATE DEHYDROGENASE (GAP1) was expressed 27.5-fold higher in phot compared to WT (Fig. 2A, Data S2). GAP1 is a chloroplast isoform of GAPDH in Chlamydomonas, a Calvin-Benson cycle enzyme predominantly localized in the stromal region surrounding the pyrenoid (26, 27). The significant accumulation of GAP1 in phot was in accord with the high accumulation of GAP1 mRNA (Fig. 2B). While mRNA accumulation of all tested starch-related genes was high in the phot mutant in the middle of the light phase (Fig. S8), GAP1 was the most highly expressed, prompting further investigation into the link between PHOT and GAP1. We found that GAP1 mRNA overaccumulation phenotype of phot was completely rescued in phot-C1 and phot-kin and partially rescued in phot-C2 (Fig. 2B), in accord with the PHOT expression levels in the different complemented lines (Fig. S3A and B), suggesting that PHOT acts as suppressor of GAP1 upon illumination. Similar to the PHOT-mediated suppression of starch (Fig. S3C), the PHOT-dependent suppression of GAP1 requires the kinase activity of PHOT (Fig. S9). The pkin-D strain, a phot mutant expressing a dead kinase PHOT with point mutations in the ATP binding site (25) and the plov strain expressing only the sensory domains LOV, both accumulate GAP1 mRNA at the level of phot. Complementation of the phot mutant with the fulllength PHOT (pphot strain) or its kinase domain (pkin strain) rescues GAP1 mRNA to WT levels (Fig. S9).

We also measured *GAP1* mRNA in cells synchronized to a 12/12 light-dark cycle. *GAP1* mRNA in WT was found to be strongly influenced by the diurnal cycle, starting at low levels after the initial onset of white light, rising to maximal levels at the end of the light phase, and then progressively declining during the night phase (**Fig. 2C**); this is in accordance with data from previous studies (28, 29) on the impact of diel cycle on the genome-wide gene expression (replotted in **Fig. S10**). *GAP1* mRNA accumulation profile was identical in white and blue light, but reached maximal levels at the middle of the light phase under red light conditions (**Fig. 2C**), suggesting that the low levels of *GAP1* mRNA after the initial onset of light are a result of suppression by blue light. Indeed, in all light qualities tested, *GAP1* mRNA levels in the synchronized *phot* mutant (**Fig. 2D**) were identical to those of the WT in red light (**Fig. 2C**). The fully complemented *phot-C1* behaved like WT (**Fig. 2E**) while the partially complemented *phot-C2* only partially rescued the observed phenotype (**Fig. 2F**). Importantly, the *phot-kin* strain in which PHOT is always active regardless of light quality, expressed very low levels of *GAP1* (**Fig. 2G**; note the log scale).

Our results so far show a strong association between starch and *GAP1* mRNA accumulation in the different *phot* mutants (**Fig. 1B** and **Fig. 2B**). WT, *phot-C1* and *phot-kin* accumulate low amounts of starch (**Fig. 1B**) and low mRNA *GAP1* (**Fig. 2B**), *phot* accumulates high levels of starch (**Fig. 1B**) and high *GAP1* mRNA levels (**Fig. 1B**) and finally the partially complemented *phot-C2* (**Fig. 1B**)

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S3A) accumulates intermediate levels of starch and *GAP1* mRNA (Fig. 1B and Fig. 2B). To further confirm the positive connection between *GAP1* expression level and starch amount, we generated WT strains overexpressing *GAP1* (Fig. S11). Strains *gap1-oe1* and *gap1-oe2* accumulated approximately 15-fold more *GAP1* mRNA than WT (Fig. 2H), resulting in 1.5-fold higher starch content (Fig. 2I). The strain *gap1-oe3*, with a more modest overexpression of *GAP1* mRNA (2.5-fold compared to WT; Fig. 2H), had starch content at WT levels (Fig. 2I). For comparison, we included the *phot* mutant in these analyses, which expressed 80-fold more *GAP1* and accumulated 3-fold more starch than the WT. We also downregulated *GAP1* in the *phot* mutant and generated strains *phot-gap1-i1*, 2 and 3 that accumulated 1.6-, 3.1- and 5.5-fold less *GAP1* mRNA (Fig. 2H) and 1.3-, 1.5- and 2-fold less starch, respectively, compared to *phot* (Fig. 2I). In conclusion, overexpression of *GAP1* in WT led to overaccumulation of starch whereas downregulation of *GAP1* in *phot* decreased starch accumulation (Fig. 2H and I). These results (Fig. 2) strongly suggest that GAP1 plays a key role in starch metabolism in *Chlamydomonas*, acting under the control of PHOT.

# PHOT alters phosphorylation state of PHOTOTROPIN-MEDIATED SIGNALLING KINASE 1

To identify missing components in the phototropin-mediated signalling pathway that suppress starch accumulation in *Chlamydomonas*, we compared the phosphoproteome of WT and *phot* cells after an overnight dark acclimation and a five-minute exposure to blue light. We identified 1119 phosphopeptides, belonging to 747 phosphoproteins, and applied unsupervised hierarchical clustering of z-transformed relative abundance of phosphopeptides to obtain an overview of the and phosphorylation (Fig. condition-specific phototropin-dependent changes Phosphopeptides falling within cluster "E" were highly phosphorylated in the dark and became de-phosphorylated after blue light illumination, but only in the case of WT; in the phot mutant they remained highly phosphorylated after blue light illumination. One such peptide was the ADGVSpSPHELTR, phosphorylated at serine 120 (S120), which belongs to the gene product of Cre16.g659400 encoding a Ser/Thr protein kinase (Fig. 3B), localized in the cytosol, plasma membrane and flagella (Fig. S13). We named this kinase Phototropin-mediated signalling kinase 1 (PMSK1).

To validate the phosphoproteomic findings, we generated WT and phot lines expressing PMSK1 fused to a FLAG epitope. These lines (WT/PMSK1-FLAG and phot/PMSK1-FLAG) were dark acclimated prior exposure eto red or blue light and samples were taken at 5 min intervals for a period of 20 min to assess the PMSK1-FLAG phosphorylation status using phos-tag SDS-PAGE, followed by immunodetection against FLAG (Fig. S14). In accordance with the role of PHOT in the light-dependent dephosphorylation of PMSK1 detected in our phosphoproteomic analyses (Fig. 3A), the phos-tag data revealed a rapid dephosphorylation of PMSK1-FLAG (Fig. 3C and **D**) upon exposure to light. In the WT background, this dephosphorylation only occurred under blue light (Fig. 3C and D), whereas in the phot background, no dephosphorylation was observed under either blue or red light (Fig. 3C), supporting our hypothesis that dephosphorylation of PMSK1 requires blue light-activated PHOT. Interestingly, the observed mobility shift of PMSK1-FLAG (Fig. 3C) can be exclusively attributed to the phosphorylation of S120 of PMSK1-FLAG (Fig. S15). Indeed, in blue-light exposure experiments with WT cells expressing PMSK1<sup>S120A</sup>-FLAG, where S120 was substituted by alanine, and with PMSK1S120D-FLAG, where S120 was substituted by aspartic acid, the modified PMSK1-FLAG had the same apparent molecular weight as the phosphatase-treated unmodified PMSK1 (labeled "-P control"; Fig. S15).

## Phosphorylation state of serine 120 of PMSK1 controls starch accumulation

We next set out to investigate the functional significance of the phosphorylation status of S120 in vivo taking advantage of the phosphomimetic mutation S120D and the non-phosphorylatable mutation S120A in PMSK1-FLAG, expressed in WT and in phot (Fig. S14). We measured GAP1 mRNA and starch accumulation in WT, phot and all above-mentioned generated mutants, synchronized under white, blue or red light in a 12h light/12h dark regime. Overexpression of PMSK1<sup>S120D</sup>-FLAG in WT, resulted in an enhanced *GAP1* expression level and increased starch content under all three light qualities, exceeding those recorded in the phot mutant (Fig. 4A and **B, Fig. S16 and S17**). Overexpression in *phot* of PMSK1<sup>S120D</sup>-FLAG resulted in an even further higher GAP1 expression level and increased starch content as compared to phot (Fig. 4A and B, Fig. S17). Conversely, overexpression in phot or WT of PMSK1<sup>S120A</sup>-FLAG, resulted in low, WTlevel or even lower, GAP1 mRNA and starch levels, across all three light quality tests (Fig. 4A and B, Fig. S16 and S17). We also analyzed the impact of transgene expression of the unmodified PMSK1-FLAG in both WT and phot mutant (Fig. 4A and B, Fig. S14 and S17); WT and WT/PMSK1-FLAG behaved very similarly in all light colors with respect to GAP1 mRNA and starch accumulation but phot/PMSK1-FLAG accumulated higher GAP1 mRNA and starch (Fig. 4A and B, Fig. S17), likely because in the absence of PHOT all PMSK1 expressed in the cells, endogenous PMSK1 and transgenic PMSK1-FLAG would remain phosphorylated (Fig. 3C and **D**).

Taken together, our data (**Fig. 4**) reveal that the phosphorylation state of S120 of PMSK1 controls starch metabolism through transcriptional regulation of GAP1. To investigate whether this role is mediated by the kinase activity of PMSK1 we combined an inactive-kinase version of PMSK1, in which Asp-442 was replaced with Asn (D442N) to inactivate the ATP-binding site, with the phosphomimetic mutation S120D or the non-phosphorylatable mutation S120A. Neither case resulted in an effect on starch content (**Fig. S18**). Thus, PMSK1 fulfills the observed regulatory role on starch metabolism through its kinase activity.

## PMSK1 acts downstream of PHOT to regulate starch metabolism in response to blue light

We applied CRISPR-CAS9 to disrupt *PMSK1* in the WT and in *phot*, thus generating the single *pmsk1* and the double *phot pmsk1* mutants. In comparison to WT, the *pmsk1* mutant accumulated 3-, 4- and 10-fold lower *GAP1* mRNA levels in white, blue and red light, respectively. Nevertheless, *GAP1* expression in *pmsk1* remained dependent on light-quality used, reaching higher expression levels under illumination with red light (**Fig. 5A and Fig. S19**). However, this overaccumulation of *GAP1* mRNA in red light was not sufficient for overaccumulation of starch (**Fig. 5B and Fig. S20**), as in the case of the *gap1 oe-3* line, which slightly overaccumulated *GAP1* mRNA but starch levels remained comparable to WT levels (**Fig. 2H and I**). As a result, we concluded that starch accumulation in *pmsk1* was light-quality-independent. The double mutant *phot pmsk1* accumulated 3-fold less *GAP1* mRNA and 2-fold less starch as compared to *phot* (**Fig. 5A and B, Fig. S19, Fig. S20**). Taken together, our results show that PMSK1 plays a critical role in transducing the PHOT-mediated blue light signal to regulate starch metabolism in *Chlamydomonas*.

## **Discussion**

In this work we report the discovery of a novel light-signalling pathway linking blue-light perception by PHOT with starch accumulation in *Chlamydomonas*. Our data solve a four-decadeslong question about the reasons for starch accumulation in green algae predominantly under red light (16). We showed that PHOT controls the phosphorylation state of a single serine residue S120 of a newly identified kinase, termed PMSK1, which in turn transduces the signal to activate transcription of *GAP1* (**Fig. 5C**). Since GAP1 is involved in the generation of phosphorylated sugars, as precursors of starch synthesis, our findings demonstrate how this light-signalling pathway leads to improvement of starch metabolism.

Our work establishes PMSK1 as the first kinase described in to mediate PHOT-dependent signalling. Yet, PHOT may not be the only blue-light responsive protein suppressing starch accumulation in *Chlamydomonas*; as our data show, application of red light in the *phot* mutant results in higher accumulation of starch compared to white or blue illuminated cells (**Fig. 1D**). How PHOT, a kinase, is involved in the dephosphorylation of PMSK1 is an intriguing question that needs further investigation. In *Arabidopsis*, blue light induces the dephosphorylation of NPH3 (NON-PHOTOTROPIC HYPOCOTYL3) in a PHOT1-dependent manner via an unidentified PHOT1-controlled protein phosphatase. The dephosphorylated NPH3 then heterodimerizes with CUL3 (Cullin homolog 3) and serves as an anchor for the PHOT1 substrate, subsequently triggering the ubiquitination of PHOT1 (*30*). We hypothesize that a protein phosphatase (indicated PPase in **Fig. 5C**) is a missing component in our proposed model; the activity of this PPase could be mediated by PHOT to dephosphorylate S120 of PMSK1.

PMSK1 is found to belong to a family of Serine/threonine-protein kinases (named PMSK-like family, see material and methods) which is conserved in green algae and vascular plants. When searching for PMSK1-like sequences in the Arabidopsis genome in the NCBI database (31), HIGH LEAF TEMPERATURE 1 (HT1) and CONVERGENCE OF BLUE LIGHT AND CO2 1/2 (CBC1/2) are found in addition to the PMSK-like Arabidopsis members (**Fig. S21-24**, **Supplementary Text**), proteins that have been shown to mediate responses to CO<sub>2</sub> and blue light in *Arabidopsis* (32). Yet, while *Arabidopsis* CBC1/2/HT1 act to stimulate stomatal opening by inhibiting S-type anion channels, our findings demonstrate that the regulatory function of its Chlamydomonas counterpart PMSK1 controls starch metabolism through the transcriptional regulation of *GAP1*.

Importantly, the higher starch accumulation in the *phot* mutant under the low light intensity conditions used in our experiments, does not affect growth and photosynthesis (**Fig. S4**). This finding indicates that this PHOT-mediated light-signaling pathway overcomes a key resource allocation trade-off present in *Chlamydomonas*, whereby carbon, fixed by photosynthesis, is allocated to energy reserves (e.g. starch) at the cost of growth (11). Our findings pave the way for the application of precise kinase engineering of PMSK1 as a sustainable way to produce starch from green microalgae in biotechnological applications.

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- 29 Writing – review & editing: Y.Y., D.P. with contributions of all coauthors.

## **Competing interests:**

Authors declare that they have no competing interests.

## Data and materials availability:

- 35 All biological material described in this study is available upon request. All data are available in
- 36 the main text or the supplementary materials. Global MS-based proteomic data have been deposited
- 37 to the ProteomeXchange Consortium via the PRIDE partner repository (33) with the dataset identifier
- 38 PXD046943. Phosphoproteomics data have been deposited to the ProteomeXchange Consortium via the
- 39 PRIDE partner repository (33) with the dataset identifier PXD045599.

## **Materials and Methods**

## Algal Material

The strains used in this study included *Chlamydomonas phot* (defective in *PHOT*; gene ID: Cre03.g199000) and phot-C1 (phot strain complemented with WT PHOT gene), as well as their background strain CC-125, which have been previously described (21). Additionally, Chlamydomonas acry (34) (defective in animal-type cryptochrome, aka aCRY; gene ID: Cre06.g278251), pcrv (defective in plant-type cryptochrome, aka pCRY; gene ID: Cre06.g295200), and acrypcry (defective in both animal-type and plant-type cryptochrome) were generated through CRISPR-CAS9 provided by following the protocol described in (34). The pmsk1 (defective in phototropin-mediated signaling kinase 1, aka PMSK1; gene ID: Cre16.g659400) and photpmsk1 (defective in both PHOT and PMSK1) mutants were generated using insertional CRISPR-Cas9 RNP method described by Kim et al. (35) with a few modifications. The target sgRNA sequence of PMSK1 was designed by Cas-Designer (http://www.rgenome.net/cas-designer) and selected considering the recommendation guideline. To induce early termination of translation, the sgRNA targets were selected in exon 2 (Data S1). To form an RNP complex in vitro, 100 µg of purified Cas9 protein (Cas9 expression plasmid: Plasmid #62934, addgene, US) and 70 µg of sgRNA synthesized by using GeneArt<sup>TM</sup> Precision gRNA Synthesis Kit (ThermoFisher, US), were mixed gently. For efficient and fast screening, 0.5 ug of paromomycin-resistance gene cassette was co-transformed with RNP complex. The Chlamydomonas cell wall was permeabilized by treatment of Max Efficiency buffer (ThermoFisher, US) following the manufacturer's protocol. The *Chlamydomonas* transformation was performed in the 4 mm gap electroporating cuvette by electroporation with the specific parameter (600 V, 50  $\mu$ F, 200  $\Omega$ ). One day after transformation, cells were plated on TAP medium containing 1.5% agar and paromomycin (25 µg/ml). Once colonies appear after transformation, genomic DNA PCR and Sanger sequencing were performed to validate knockout events.

To prepare transgenic lines with a knockdown of GAP1, pChlamiRNA3int-GAP1 was transformed into the *phot* strain. Generation of amiRNA plasmids was performed according to (36). The oligonucleotides designed for targeting GAP1 (Data S1) using the WEB MicroRNA Designer platform (WMD3: <a href="http://wmd3.weigelworld.org/cgi-bin/webapp.cgi">http://wmd3.weigelworld.org/cgi-bin/webapp.cgi</a>. Ossowski Stephan, Fitz Joffrey, Schwab Rebecca, Riester Markus and Weigel Detlef, personal communication) were annealed and ligated into pChlamiRNA3int (SpeI digested) to create pChlamiRNA3int-gap1. Transformed cells were selected and further checked by RT-qPCR.

For the preparation of transgenic lines overexpressing GAP1(Glyceraldehyde 3-phosphate dehydrogenase, aka *GAP1*; gene ID: Cre12.g485150) or different versions of PMSK1, the genomic sequences of *GAP1* and *PMSK1* were PCR amplified from genomic DNA of *Chlamydomonas* CC-125 and cloned into pLM005 in-frame with a C-terminal Venus-3Xflag using Gibson Assembly (37), and then transformed into WT or *phot* strains. The primers used for different gene amplification and point mutations are described in Data S1.

## Chlamydomonas reinhardtii cultivation

All *Chlamydomonas* strains were maintained on solid Tris-acetate-phosphate (TAP) (38) agar plates with or without appropriate antibiotic at 22°C and 5  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>. Prior to the start of the experiments, cells were cultured in 50 mL TAP medium in 250 ml Erlenmeyer flasks at 23 °C, 120 rpm/min and 15  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>. The experiments were conducted in Sueoka's high salt medium (HSM) (39) at an initial cell density of 1 million cells/ml at 50  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> unless otherwise stated.

For continuous light experiments, the cells were transferred to HSM medium and grew at 23 °C, 120 rpm/min and 50  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>.

For synchronized experiments, cells were grown in HSM for at least 5 days under a 12h light/12h dark cycle under white light or different light qualities (light intensity was set at 50  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>; temperature was 18°C in the dark and 23°C in the light). The light spectrum of the LED lighting system used in this study is the same as previously described (40).

# <u>Transformation of Chlamydomonas reinhardtii</u>

 The transformation was performed by electroporation, which follows the protocol of Zhang et al (41) with minor modification. Cells for transformation were collected at 1-2 h before the end of the light phase in a synchronized (12 h light/ 12h dark) culture. For three reactions 11 ng/kb linearized plasmid was mixed with 400  $\mu$ l of 1.0 x10<sup>7</sup> *Chlamydomonas reinhardtii* cells/ml and electroporated at a volume of 125 ml in a 2-mm-gap electro cuvette using a NEPA21 square-pulse electroporator, using two poring pulses of 250 and 150 V for 8 msec each, and five transfer pulses of 50 msec each starting at 20 V with a "decay rate" of 40% (i.e., successive pulses of 20, 12, 7.2, 4.3, and 2.6 V). Electroporated cells were immediately transferred to a 15ml centrifugation tube containing 9 ml TAP plus 40mM sucrose. After overnight dark incubation, cells were collected by centrifugation and spread on TAP agar plates which contain the appropriate antibiotic (20  $\mu$ g/ml paromomycin or 7.5  $\mu$ g/ml zeocin or 20  $\mu$ g/ml hygromycin B). Transformants typically appear after 5-7 days.

The putative antibiotic-resistant transformants were transferred into individual wells of a 96-well, flat-bottom transparent microplate, with each well containing 250 µl of TAP medium. Cultures were grown for 3 days under 15 µmol photons m<sup>-2</sup> s<sup>-1</sup> light without shaking, refreshed by replacing half of the culture with fresh medium, and allowed to grow for an additional day. Transformants were screened for Venus expression using a fluorescent microplate reader (Tecan Group Ltd., Switzerland), with parameters including Venus (excitation 515/12 nm and emission 550/12 nm) and chlorophyll (excitation 440/9 nm and emission 680/20 nm). The fluorescence signal was normalized to the chlorophyll fluorescence signal, and colonies with a high Venus/chlorophyll value were selected as putative complemented strains. These putative positive transformants were further validated by western blotting and RT-qPCR.

## RNA extractions and RT-qPCR analysis

Total RNA for RNA-seq and RT-qPCR was extracted using RNeasy Mini Kit (Qiagen, Germany) and treated with the RNase-Free DNase Set (Qiagen, Germany). 1  $\mu$ g total RNA was reverse transcribed with oligo dT using Sensifast cDNA Synthesis kit (Meridian Bioscience, US). qPCR reactions were performed and quantitated in a Bio-Rad CFX96 system using SsoAdvanced Universal SYBR Green Supermix (BioRad, US). The *CBLP* gene (42) served as the housekeeping control and relative fold differences were calculated on the basis of the  $\Delta C_t$  method ( $2^{-(Ct \text{ target gene }-Ct CBPL)}$ ) (43–45). All primers used for the RT-qPCR analyses were synthesized by ThermoFisher (US) or IDT (Integrated DNA Technologies, Inc. Coralville, Iowa, US) and were presented in Data S1.

## Analyses of Total Starch content

The total starch content of samples collected daily was determined using Total Starch Assay Kit (K-TSTA-100A, Megazyme, Ireland) as described in its instruction with modifications. The results were calculated according to the standard curve made with glucose solution after starch digestion. To prepare the samples for glucose determination, 10 mL of the liquid culture was

pelleted by centrifugation and resuspended in 40  $\mu$ L of 80% (v/v) ethanol. Next, 400  $\mu$ L of cold 1.7 M sodium hydroxide solution was added, and the samples were incubated on ice for 15 min. Following this, 1.6 mL of sodium acetate buffer containing calcium chloride (5 mM) was added and mixed well. Subsequently, 20  $\mu$ L of  $\alpha$ -amylase and 20  $\mu$ L of amyloglucosidase were added, and the samples were incubated at 50 °C for 30 min. The supernatant was collected by centrifugation at 13,000 rpm for 5 min and will be ready for glucose determination.

# Analyses of Total Protein and Lipid content

The total protein content of samples collected daily was determined using BCA Protein Assay Kit (ThermoFisher, US) with the standardized protocol. The total lipid content of samples collected daily was determined using sulfo-phospho-vanillin (SPV) method (46)

# <u>Transmission electron microscopy</u>

 Cells were harvested by centrifugation at 700g for 5 min, washed two times in 0.1 M PB (phosphate buffer, pH 7.4) and then were fixed in 0.1 M PB containing 2.5% (v/v) glutaraldehyde for 2h at room temperature and stored overnight at 4 °C. The cells were then washed five times in 0.1 M PB before being fixed by a 1h incubation on ice in 0.1 M PB containing 2% osmium and 1.5% ferricyanide potassium. After being washed five times with 0.1 M PB , the samples were resuspended in 0.1 M PB containing 0.1% (v/v) tannic acid and incubated for 30 min in the dark at room temperature. The cells were washed five times with 0.1 M PB , dehydrated in ascending sequences of ethanol, infiltrated with an ethanol/Epon resin mixture, and finally embedded in Epon. Ultrathin sections (50–70 nm) were prepared with a diamond knife on a PowerTome ultramicrotome (RMC Boeckeler, US) and collected on 200  $\mu$ m nickel grids. The ultrathin sections were examined on a Philips CM120 transmission electron microscope operating at 80 kV.

# Confocal Microscopy

The preparation of samples for confocal microscopy followed the protocol reported by Mackinder et al (47). The confocal microscope used in the study was from the cell-imaging platform at IBS, Grenoble, France. All confocal microscopy images were analyzed using Fiji.(48)

## **Immunoblotting**

Protein samples of whole cell extracts (5 μg protein) were loaded on 4-20% SDS-PAGE gels (Mini-PROTEAN TGX Precast Protein Gels, Bio-Rad, US) and blotted onto nitrocellulose membranes. Antisera against ATPB (AS05085, 1:15000) was from Agrisera (Sweden); antiserum against FLAG (F3165, 1:2000) was from Sigma-Aldrich (US); antiserum PHOT (LOV1 domain, 1:5000) was previously described<sup>10</sup>. ATPB was used as a loading control. The anti-rabbit or mouse horseradish peroxidase–conjugated antiserum (Jackson Immuno Research, US) was used for detection at 1:10000 dilution. The blots were developed with ECL detection reagent, and images of the blots were obtained using ImageQuant 800 (Cytiva, UK). For the densitometric quantification, data were normalized with ATPB.

## Phos-Tag Gel Electrophoresis

Double-layer Phos-tag gels with a concentration of 12% (w/v) acrylamide/bisacrylamide 37.5:1 and 65 mM of Phos-Tag (Wako Pure, US) were prepared as in (49), with the exception that Zn(NO<sub>3</sub>)<sup>2</sup> was added equimolarly to the samples to compensate for the absence of EDTA in the lysis buffer. The gels were denatured for 30 min at 37°C prior to loading. In vitro dephosphorylation involved resuspending a cell pellet in 5 mM of HEPES at pH 7.5, 10 mM of

EDTA, and 1% (v/v) TritonX 100. An aliquot containing 10 mg of protein was then subjected to lambda protein phosphatase reaction mix following the manufacturer's instructions (New England Biolabs, US) for 1 hour at 30°C, in accordance to (22).

## Fluorescence-based measurements

Fluorescence-based photosynthetic parameters were measured with a pulse modulated amplitude fluorimeter (MAXI-IMAGING-PAM, HeinzWaltz GmbH, Germany). Prior to the onset of the measurements, cells were acclimated to darkness for 15 min. Chlorophyll fluorescence was recorded under different intensities of actinic light; starting with measurements in the dark (indicated as D below the x-axis of the graphs), followed by measurements at 21  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> (indicated as L1 below the x-axis of the graphs) and 336  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> (indicated as L2 below the x-axis of the graphs) and finishing with measurements of fluorescence relaxation in the dark. The effective photochemical quantum yield of photosystem II was calculated as Y(II) = (Fm'-F)/Fm'; F and Fm' are the fluorescence yield in steady state light and after a saturating pulse in the actinic light, respectively;

# Phosphoproteomics analysis

Protein extraction. C. reinhardtii pellets were resuspended in 2000 μL of lysis buffer (100 mM Tris-HCl, PhosphoSTOP inhibitors, protease inhibitors) and ultrasonicated in the Covaris for 4 min each. Samples were diluted by adding 2000 μL of dilution buffer (100 mM Tris-HCl, 5 mM TCEP, 30 mM chloroacetamide, 1 mM sodium orthovanadate, phosphoSTOP inhibitors, 1 mM magnesium chloride) and 1 μL Benzonase. Lysates were shaken at 25 °C for 1 h. 8 mL of methanol was added to each sample, followed by 3 mL chloroform and 3 mL water, with vortexing after each subsequent addition. Samples were centrifuged for 10 min at 3,220 x g and the top layer was removed, leaving the interphase intact. An additional 10 mL of methanol was added and the samples were centrifuged for 20 min at 3,220 x g and the supernatant was removed. Protein pellets were allowed to dry at RT and resuspended in 1 mL of digestion buffer (100 mM Tris-HCl, 2 M urea). Proteins were digested with 12 μg of trypsin overnight and cleaned up via C18 SPE cartridges. Samples were resuspended in 250 μL of water and a BCA assay was performed to determine peptide concentration. 10 μg of digested protein was taken for global analysis, and 500 μg was used for phosphopeptide enrichment.

Phosphopeptide enrichment. Phosphopeptides were enriched using a ProPac Fe-IMAC column (ThermoFisher, US) on a Shimadzu Prominence HPLC system. Before enrichment, the column was charged with 25 mM FeCl<sub>3</sub> in 100mM acetic acid. Mobile phase A consisted of 30% acetonitrile in water (v/v) with 0.07% trifluoroacetic acid (v/v). Mobile phase B consisted of 0.3% ammonium hydroxide in water (v/v). Tryptic peptides were diluted to 30% acetonitrile and injected on the column at a flow rate of 0.2 ml/min. After 3 min of loading, flow rate was increased to 2 mL/min. Peptides were eluted by rapidly ramping the gradient to 50% B. Fractions containing phosphopeptides were cleaned up with C18 SPE cartridges (Waters, US) and resuspended in 20  $\mu$ L of LC-MS water prior to mass spectrometry analysis.

*LC-MS analysis.* Phosphopeptide samples were analyzed using a nanoACQUITY UPLC (Waters, US) coupled to a TripleTOF 5600 mass spectrometer (Sciex, Canada). Mobile phase A consisted of water with 0.1% formic acid and mobile phase B was acetonitrile with 0.1% formic acid. Injections were made to a Symmetry C18 trap column (100 Å, 5  $\mu$ m, 180  $\mu$ m x 20 mm; Waters, US) with a flow rate of 5  $\mu$ L/min for 3 min using 99% A and 1% B. Peptides were then separated on an HSS T3 C18 column (100 Å, 1.8  $\mu$ m, 75  $\mu$ m x 250 mm; Waters, US) using a linear gradient of increasing mobile phase B at a flow rate of 300 nL/min. Mobile phase B increased

from 5% to 40% in 90 min before ramping to 85% in 5 min, where it was held for 5 min before returning to 5% in 2 min and re-equilibrating for 13 min.

The mass spectrometer was operated in positive polarity mode. MS survey scans were accumulated across an m/z range of 350-1600 in 250 ms optimized at  $\geq$  30,000 resolution. For data-dependent acquisition, the mass spectrometer was set to automatically switch between MS and MS/MS experiments for the first 20 features above 150 counts having +2 to +5 charge state. Precursor ions were fragmented using rolling collision energy and accumulated in high sensitivity mode for 85 ms across an m/z range of 100-1800 optimized at  $\geq$  30,000 resolution. Dynamic exclusion for precursor m/z was set to 8 s.

Bioinformatic analysis. Raw data files were imported into Progenesis for peak alignment and quantification. Spectra were searched in Mascot against the *C. reinhardtii* phytozome database (v6.1) using a precursor / fragment tolerance of 15 ppm / 0.1 Da, trypsin specificity, two possible missed cleavages, fixed modification cysteine carbamidomethylation, and variable modifications of methionine oxidation, protein N-term acetylation, and phosphorylation (STY). Identifications were imported back into Progenesis for peak assignment, and statistical analysis was performed using the QuantifyR workflow, which can be found on the Hicks Lab Github (github.com/hickslab/QuantifyR). Phosphoproteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository (33) with the dataset identifier PXD045599.

# **Lipidomics**

 Glycerolipids were extracted from freeze-dried cell pellets frozen immediately in liquid nitrogen after harvesting. Once freeze-dried, cell pellets were resuspended in 4 mL of boiling ethanol for 5 minutes to prevent lipid degradation and lipids were extracted according to (50) by addition of 2 mL methanol and 8 mL chloroform at room temperature. The mixture was then saturated with argon and stirred for 1 hour at room temperature. After filtration through glass wool, cell remains were rinsed with 3 mL chloroform/methanol 2:1, v/v and 5 mL of NaCl 1% were then added to the filtrate to initiate biphase formation. The chloroform phase was dried under argon before solubilizing the lipid extract in pure chloroform. Total glycerolipids were quantified from their fatty acids: in an aliquot fraction, a known quantity of 15:0 was added and the fatty acids present were transformed as methyl esters (FAME) by a 1-hour incubation in 3 mL 2.5% H<sub>2</sub>SO<sub>4</sub> in pure methanol at 100°C (51). The reaction was stopped by addition of 3 mL water and 3 mL hexane. The hexane phase was analyzed by gas chromatography-flame ionization detector (GC-FID) (Perkin Elmer, US) on a BPX70 (SGE; Trajan Scientific and Medical location, Australia) column. FAME were identified by comparison of their retention times with those of standards (Sigma, US) and quantified by the surface peak method using 15:0 for calibration.

The lipid extracts corresponding to 25 nmol of total fatty acids were dissolved in 100  $\mu$ L of chloroform/methanol [2/1, (v/v)] containing 125 pmol of each internal standard. Internal standards used were PE 18:0-18:0 and DAG 18:0-22:6 from Avanti Polar Lipid and SQDG 16:0-18:0 extracted from spinach thylakoid (52) and hydrogenated as described in (53). Lipids were then separated by HPLC and quantified by MS/MS.

The HPLC separation method was adapted from (54). Lipid classes were separated using an Agilent 1200 HPLC system using a 150 mm×3 mm (length × internal diameter) 5  $\mu$ m diol column (Macherey-Nagel, Germany), at 40°C. The mobile phases consisted of hexane/isopropanol/water/ammonium acetate 1M, pH5.3 [625/350/24/1, (v/v/v/y)] (A) and isopropanol/water/ammonium acetate 1M, pH5.3 [850/149/1, (v/v/v)] (B). The injection volume was 20  $\mu$ L. After 5 min, the percentage of B was increased linearly from 0% to 100% in 30 min

and stayed at 100% for 15 min. This elution sequence was followed by a return to 100% A in 5 min and an equilibration for 20 min with 100% A before the next injection, leading to a total runtime of 70 min. The flow rate of the mobile phase was 200  $\mu$ L/min. The distinct glycerophospholipid classes were eluted successively as a function of the polar head group.

Mass spectrometric analysis was done on a 6470 triple quadrupole mass spectrometer (Agilent, US) equipped with a Jet stream electrospray ion source under following settings: Drying gas heater: 230°C, Drying gas flow 10 L/min, Sheath gas heater: 200°C, Sheath gas flow: 10L/min, Nebulizer pressure: 25 psi, Capillary voltage:  $\pm 4000$  V, Nozzle voltage  $\pm 2000$ . Nitrogen was used as collision gas. The quadrupoles O1 and O3 were operated at widest and unit resolution respectively. DGTS analysis was carried out in positive ion mode by scanning for precursors of m/z 236 at a collision energy (CE) of 55 eV. SQDG analysis was carried out in negative ion mode by scanning for precursors of m/z -225 at a CE of -55eV. PE, PI, PG, MGDG and DGDG measurements were performed in positive ion mode by scanning for neutral losses of 141 Da, 277 Da, 189 Da, 179 Da and 341 Da at CEs of 29 eV, 21 eV, 25 eV, 8 eV and 11 eV, respectively. Quantification was done by multiple reaction monitoring (MRM) with 30 ms dwell time. DAG and TAG species were identified and quantified by MRM as singly charged ions [M+NH<sub>4</sub>]<sup>+</sup> at a CE of 19 and 26 eV respectively with 30 ms dwell time. Mass spectra were processed by MassHunter Workstation software (Agilent, US) for identification and quantification of lipids. Lipid amounts (pmol) were corrected for response differences between internal standards and endogenous lipids and by comparison with a quality control (QC). QC extract correspond to a known lipid extract from Chlamydomonas cell culture qualified and quantified by TLC and GC-FID as described in (55).

## Proteomics analysis

 Proteins from total extracts of three biological replicates of WT and *phot Chlamydomonas* reinhardtii were solubilized in Laemmli buffer and heated for 10 min at 95°C. They were then stacked in the top of a 4-12% NuPAGE gel (ThermoFisher, US), stained with Coomassie blue R-250 (Bio-Rad, US) before in-gel digestion using modified trypsin (Promega, US) as previously described (*56*). The resulting peptides were analyzed by online nanoliquid chromatography coupled to MS/MS (Ultimate 3000 RSLCnano and Q-Exactive HF, ThermoFisher, US) using a 180-min gradient. For this purpose, the peptides were sampled on a precolumn (300 μm x 5 mm PepMap C18, ThermoFisher, US) and separated in a 75 μm x 250 mm C18 column (Reprosil-Pur 120 C18-AQ, 1.9 μm, Dr. Maisch, Germany). The MS and MS/MS data were acquired using Xcalibur (V2.8, ThermoFisher, US).

Peptides and proteins were identified by Mascot (V2.8.0, Matrix Science) through concomitant searches against the C. reinhardtii phytozome database (V5.6) (19526 sequences), the mitochondrion and chloroplast protein sequences (downloaded from NCBI, respectively 69 and 8 proteins), and a homemade database containing the sequences of classical contaminant proteins found in proteomic analyses (human keratins, trypsin..., 126 sequences). Trypsin/P was chosen as the enzyme and two missed cleavages were allowed. Precursor and fragment mass error tolerances were set at respectively at 10 and 20 ppm. Peptide modifications allowed during the search were: Carbamidomethyl (C, fixed), Acetyl (Protein N-term, variable) and Oxidation (M, variable). The Proline software (57) (V2.2.0) was used for the compilation, grouping, and filtering of the results (conservation of rank 1 peptides, peptide length  $\geq$  6 amino acids, false discovery rate of peptide-spectrum-match identifications < 1% (58), and minimum of one specific peptide per identified protein group). MS data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository (59)with the dataset identifier PXD046943. Proline was then used

to perform a MS1 label-free quantification of the identified protein groups based on razor and specific peptides.

Statistical analysis was performed using the ProStaR software (60) based on the quantitative data obtained with the three biological replicates analyzed per condition. Proteins identified in the contaminant database, proteins identified by MS/MS in less than two replicates of one condition, and proteins quantified in less than three replicates of one condition were discarded. After log2 transformation, abundance values were normalized using the variance stabilizing normalization (vsn) method, before missing value imputation (SLSA algorithm for partially observed values in the condition and DetQuantile algorithm for totally absent values in the condition). Statistical testing was conducted with limma, whereby differentially expressed proteins were selected using a log2(Fold Change) cut-off of 1 and a p-value cut-off of 0.00912, allowing to reach a false discovery rate inferior to 1% according to the Benjamini-Hochberg estimator. Proteins found differentially abundant but identified by MS/MS in less than two replicates, and detected in less than four replicates, in the condition in which they were found to be more abundant were invalidated (p-value = 1).

## GO enrichment analysis

GO term enrichment was tested for all proteins significantly differential abundant at a false discovery rate below 5% using the Benjamini-Hochberg estimator (61). GO annotation of proteins was obtained from phytozome database (v.5.6) and all ancestral GO terms were added to a protein using the R package GO.db. P-values were obtained according to the null hypothesis, that the number of differential abundance proteins bearing a GO term is a random variable whose probability distribution is described by the hypergeometric distribution. The false discovery rate was controlled below 5% using the Benjamini-Hochberg-estimator (61). Only GO terms linked to at least 6 measured proteins were tested.

## Statistical analysis

Prism (GraphPad Software) was used for statistical analysis and all error bars represent standard deviation. ANOVA tests and t-tests were performed, with the p-values or degree of significance provided in the figures and the legends.

## **Supplementary Text**

## Phylogenetic analysis

PMSK1 is found to belong to a family of Serine/threonine-protein kinases (named the PMSK-like family, see material and methods) which is conserved in green algae and vascular plants (Fig. S21 and 22.). In particular, the PMSK-like family is consistent with the known Chlorophyta evolution (62). Interestingly, when searching for PMSK1 similar sequences into the Arabidopsis thaliana genome in the NCBI database (31), next to the PMSK-like Arabidopsis members come the Arabidopsis HT1 (At1g62400, NP 176430.2), CBC1 (At3g01490, NP 186798.1) and CBC2 (At5g50000, NP 199811.1). It has been found that (i) the best reciprocal best hit for these three proteins when searching for Chlamydomonas reinhardtii was PMSK-like family members, especially PMSK1 (ii) this was true when searching for any Chlorophyta species and that (iii) regarding into the three family CBC, HT and PMSK-like, Chlorophyta was only present in the latest (Fig. S23). A simple molecular clock analysis phylogenetic analysis revealed that the root is likely to be placed between the CBC and the HT/PMSK1 family (Fig. S24) suggesting that a representative for these three families could have been present in the common ancestor of the Chlorophyta. One explanation for the observed distribution of the sequences is that both HT and CBC representative have been lost before the radiation of the Chlorophyta. It must be noticed that most of the time, duplication in CBC, HT and PMSK-like family seems to have happened after recent speciation events, suggesting that a few members of this family could have been present in the last common ancestor of the Chlorophyta/Charophyta/Land Plants Phylum.

## Identification of the PMSK-like family

The gene product of Cre16.g659400, identified as encoding a Ser/Thr protein kinase (**Fig. 3B**) and named kinase Phototropin-mediated signalling kinase 1 (PMSK1), was translated and the amino acid sequence used as a query for a series of BLAST searches in the publicly available National Center for Biotechnology Information (NCBI) <u>database (31)</u>. Both BLASTp and PSI-BLAST (63, 64) were performed together with a human-curated process in order to obtain the widest dataset possible and the most robust one. The final dataset (#1) contained 111 amino acid sequences from at least 10 orders of Chlorophyta, Charophyta, Bryopsida, Pteridophyta, Gymnospermae and Angiosperm.

## Identification of a relation between the PMSK-like family and both the HT1 and the CBC family

Because both HT1, CBC1 and CBC2 were identified in the BLAST research as first remotely related *Arabidopsis thaliana* sequences in the previous homologous sequences search, a reciprocal best hit research was done using these three sequences together with a human-curated process and results was append to the previous dataset. This new dataset (#2) contained 278 amino acid sequences from at least 10 orders of Chlorophyta, Charophyta, Bryopsida, Pteridophyta, Gymnospermae and Angiosperm. Interestingly, homologous sequences investigations revealed that while the four family Charophyta, Bryopsida, Pteridophyta, Gymnospermae are present in the both the HT and the CDC family, the most closely related Chlorophyta sequences of the CBC/HT *Arabidopsis* thaliana representative were actually the PMSK-like family sequences. A subset of dataset #2 containing 11 sequences was used to compute both a clock-model and a relaxed-clock model to infer the possible root of the three PMSK-like, HT and CBC family.

## Multiple Sequence Alignments

 Amino acid multiple sequence alignments for each protein segment were created using MUSCLE (65) and then adjusted using Gblocks (version 0.91.1) (66) or the MEGA ML analysis (see below) or manually using Jalview (version 2.11.1.4) (67) for both the PhyML and the Bayesian analysis (see below).

## Dataset #1 and #3. Bayesian Inferences

Two independent Metropolis-coupled Markov Chain Monte Carlo (MCMC) analyses were performed using the final curated datasets in the MrBayes (version 3.2.7.a) software (68). The two datasets were used for Bayesian inference. The first was include in a nexus file resulting from the MUSCLE alignment. The dataset #3 was include in the 278 sequences multiple alignment nexus file from the dataset #2 from which, following a command line in the MrBayes software, it was extracted before setting the algorithm parameters. The Bayesian posterior probabilities (BPP) were estimated by two independent runs of four Metropolis Coupled chains (MCMC): 2 independent runs, together with one cold chain and three "heated" (temperature parameter 0.1, Dataset 3) chains drove the analyses. The analyses were allowed to switch among all the substitution models implemented in the software to identify the best model without any a priori (aamodelpr = mixed). The prior on the branch length was set to unconstrained for the dataset #1, leading to an unrooted tree whereas it was set on both a uniform and a relaxed clock model to infer the root position of the HT, CBC and PMSK-like families.

80,000 and 160,000 generations with sampling every 100 generations were set for Dataset #1 and Dataset #3, respectively. The standard deviation of the split frequency between the two parallel analyses at the end of each analysis were 0.021973 and 0.005348 for Dataset #1 and Dataset #3, respectively. The first 25% of the trees produced were discarded in order to let the analyses stabilize (burnin = 0.25). The convergence of the runs was estimated and the potential scale reduction factor (PSRF +) were all around 1 for all the parameters indicating that the associated chains converged to one target distribution. Large PSRF (PSRF +>1) indicate convergence failure. Nodes with a posterior probability  $\geq$ 0.90 were considered as well supported. Alignments in fasta and trees in newick format are available upon request.

## Dataset #2. Maximum Likelihood inferences

Unrooted maximum likelihood (ML) phylogenetic trees were then inferred using both the MEGA (version 11) software package Whelan And Goldman ML model (69) and PhyML (version 3.3.1) (70) based on the best-fit models of amino acid substitution determined by Find Best Protein Model (67). Non-uniformity of evolutionary rates among sites was modeled using a discrete Gamma distribution (+G) and a certain frac-tion of sites were considered to be evolutionarily invariable (+I). Initial trees for the heuristic search were obtained automatically by applying Neighbor Joining (NJ) and BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and then selecting the topology with superior log likelihood value. All positions with less than 95% site coverage were eliminated. Statistical tests for branch support for the MEGA ML algorithem was estimated with a Bootstrap procedure (71), using 100 replicates. Statistical tests for branch support concerning the PhyML algorithm was estimated with the aLRT SH-like branch support method (72). Trees were drawn to scale with the branch length measured in the number of substitutions per site.

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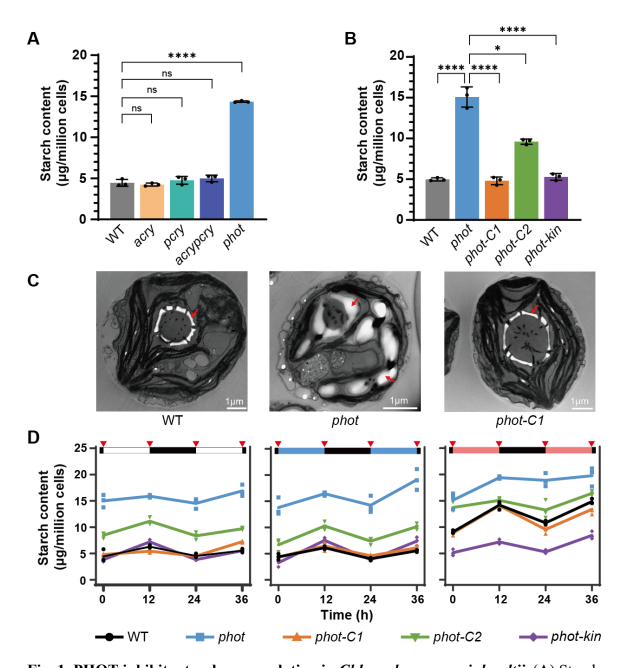
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**Fig. 1. PHOT inhibits starch accumulation in** *Chlamydomonas reinhardtii.* (A) Starch content of WT, *acry, pcry, acrypcry* lines under continuous light. (B) Starch content of WT and variously *phot*-complemented lines under continuous white light. (C) Transmission electron microscopy pictures of WT, *phot* and *phot-C* synchronised to a12/12 dark/light cycle. Samples were collected in the end of the light phase. Red arrows indicate starch granules. (D) Starch content of WT and *phot*-complemented lines synchronised to 12/12 dark/light cycle. Red triangles indicate sample collection time. The dark phase is indicated by black bars above the graphs; the light phase by white, blue or red bars, depending on the light quality used. In some cases, the error bars are smaller than the data point symbols. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Asterisks indicated the p-values (\*, p<0.05; \*\*\*\*, p < 0.0001; ns, not significant). In some cases, the error bars are smaller than the data point symbols. Detailed statistical analyses are presented in Data S4.

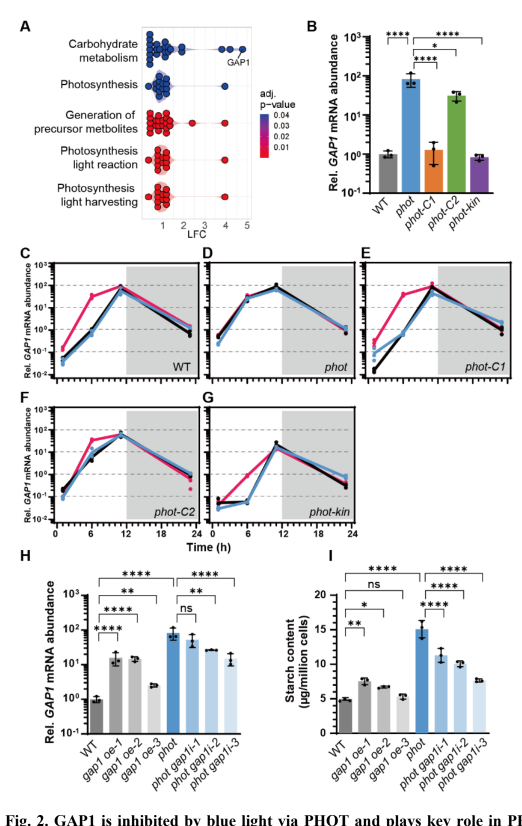
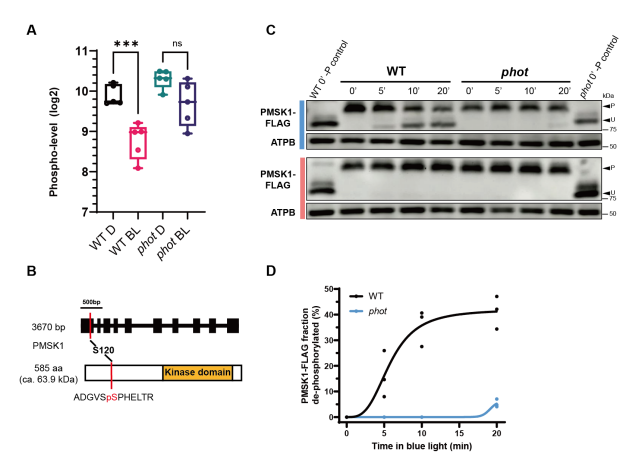


Fig. 2. GAP1 is inhibited by blue light via PHOT and plays key role in PHOT-dependent starch metabolism in *Chlamydomonas reinhardtii*. (A) Gene ontology enrichment analyses for differentially abundant proteins in *phot* compared to WT based on whole cell proteomics data.

LFC, log2 fold change (enriched GO terms were only found for proteins with increased abundance in *phot* in comparison to WT); color code indicates adjusted p-value of GO set enrichment (null hypothesis: number of differentially abundant proteins in GO set is hypergeometric random distributed. P-value were adjusted according to Benjamini-Hochberg procedure (**B**) GAPI relative mRNA abundance of WT and *phot*-complemented lines under continuous white light. (**C-G**) relative mRNA abundance of GAPI in WT and *phot*-complemented lines synchronized to a 12/12 dark/light cycle. Phases are indicated by white and gray shading. Line colors indicated light qualities. Black, white light; Red, red light; Blue, blue light. The GAPI transcription level (**H**) and starch content (**I**) of WT, GAPI overexpression and *photgap1* knockdown lines under continuous light. Data are presented as mean  $\pm$  SD (n = 3 biologically independent samples). Asterisks indicated the p-values (\*, p <0.05; \*\*, p < 0.01; \*\*\*\*, p < 0.0001; ns, not significant). In some cases, the error bars are smaller than the data point symbols. Detailed statistical analyses are presented in Data S4.



**Fig. 3. Identification of PMSK1 as a key protein on PHOT-dependent starch metabolism in** *Chlamydomonas reinhardtii.* (A) Changes in the phosphorylation level of S120 of PMSK1 in response to BL in WT and *phot*, quantified by phosphoproteomics. Samples were collected after 24h of acclimation to darkness and 5 min after the start of blue light. D, dark; BL, blue light. (B) Upper: Genomic structures of PMSK1. Black boxes and lines indicate exons and introns, respectively. Lower: Schematic structures of PMSK1. Red lines indicate the phosphorylated residue in PMSK1. The orange box indicates the kinase domain. (C) Phosphorylation level of PMSK1-FLAG as a function of time in WT/PMSK1-FLAG and *phot/PMSK1-FLAG*, exposed to blue or red light after 24h acclimation to darkness (indicated as t=0 in the graph). Detection was performed by phos-tag SDS-PAGE; ATPB was used as a loading control. Phosphatase-treated WT/PMSK1-FLAG and *phot/PMSK1-FLAG* samples were also loaded on the gels. "U" and "P" indicate the unphosphorylated and phosphorylated PMSK1-FLAG respectively. The blue or red bar to the left of the immunoblots indicates the light quality used. (D) PMSK1 phosphorylation levels, quantified by densitometric analyses of the phos-tag SDS-PAGE in (C), plotted as a function of time under blue light.

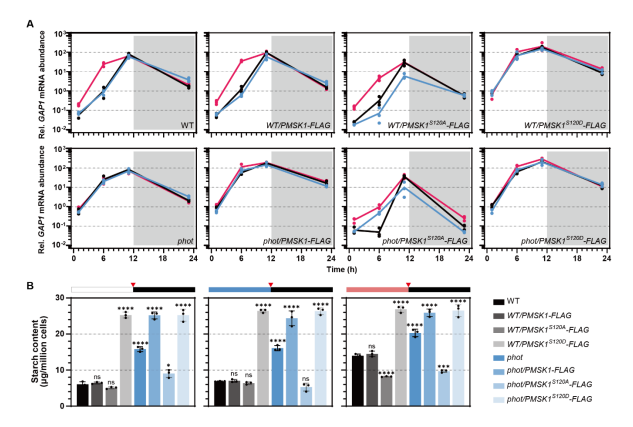


Fig. 4. PMSK1 phosphorylation status controls starch metabolism in *Chlamydomonas* reinhardtii. *GAP1* transcription level (**A**) and starch content (**B**) of various PMSK1 overexpression lines synchronized to a 12/12 light dark cycle under different light qualities. Data are presented as mean  $\pm$  SD (n = 3 biologically independent samples). (**A**) Phases are indicated by white and gray shading. Line colors indicated light qualities. Black, white light; Red, red light; Blue, blue light. (**B**) Red triangles indicate sample collection time. The dark phase is indicated by black bars above the graphs; the light phase by white, blue, or red bars, depending on the light quality used. Asterisks indicated the p-values compare to WT (\*, p<0.05; \*\*\*, p < 0.001; \*\*\*\*, p < 0.0001; ns, not significant). In some cases, the error bars are smaller than the data point symbols. Detailed statistical analyses are presented in Data S4.

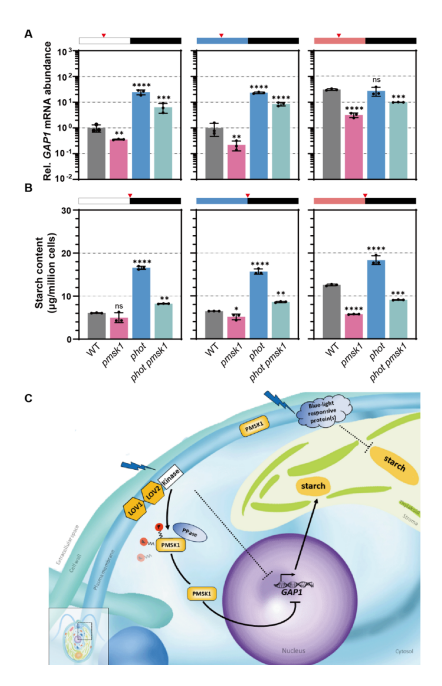


Fig. 5. PHOT regulates starch metabolism via PMSK1 and GAP1 in *Chlamydomonas reinhardtii*. *GAP1* transcription level (A) and starch content (B) in single and double *phot* and *pmsk1* mutants synchronized to a 12/12 light dark cycle under different light qualities. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Red triangles indicate sample collection time. Dark phase is indicated by black bars above the graphs; the light phase by white, blue or red bars, depending on the light quality used. Red triangles indicated samples collection time. Asterisks indicated the p-values compare to WT (\*, p <0.05; \*\*, p < 0.01; \*\*\*\*, p < 0.001; \*\*\*\*, p < 0.0001; ns, not significant). In some cases, the error bars are smaller than the data point symbols. Detailed statistical analyses are presented in Data S4. (C) A hypothetic model depicting the mechanisms of blue-light dependent regulation of starch metabolism in *C. reinhardtii*.

Supplementary Materials for

Phototropin connects blue light perception to starch metabolism in green algae

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The PDF file includes:

Figs. S1 to S24

Other Supplementary Materials for this manuscript include the following:

Data S1 to S5 are in a combined Excel file to be downloaded separately

Data S1. Oligo primers used in this study.

Data S2. Proteomics data of WT and phot.

Data S3. Phosphoproteomics data of WT and phot.

Data S4. Raw values and statistical analysis of figures

Data S5. All Chlamydomonas reinhardtii strains used in this study

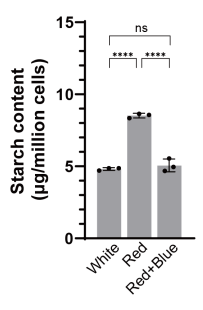


Fig. S1. Effect of superimposing low intensity blue light on the red light on starch content. The starch content of WT under different continuous light conditions. White,  $50 \mu E$  white light; Red,  $50\mu E$  red light; Red+Blue,  $45\mu E$  red light plus  $5\mu E$  blue light. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Asterisks indicated the p-values (\*\*\*\*, p < 0.0001; ns, not significant)

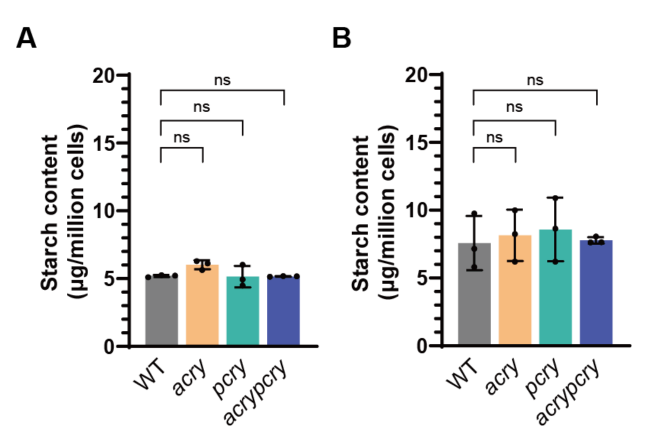


Fig. S2. Mutants devoid of animal- or plant-like cryptochromes accumulate WT-levels starch. The starch content of acry, pcry and acrypcry synchronized to a 12 h/12 h light dark cycle (white light). Samples were collected at the start (A) and at the end of the light phase (B). Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). ns, not significant.

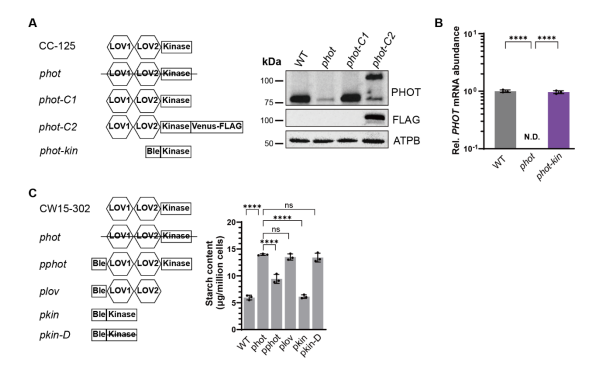


Fig. S3. PHOT inhibits starch accumulation in *Chlamydomonas reinhardtii*. (A) Left: Domains of the *PHOT* gene in WT and complemented *phot* lines. LOV1 and LOV2, photosensory domains; Kinase, kinase domain; Venus-FLAG, yellow fluorescent protein with FLAG tag; Ble, gene conferring resistance to bleomycin. Right: Immunoblot analyses of PHOT level in WT (CC125), *phot* and complements strains. (B), RT-qPCR analysis of *PHOT-Kinase* transcription level in WT and *phot-kin*. N.D., not detected. (C), Left: Domains of the *PHOT* gene in WT (*cw15-302*) and complemented *phot* lines. Right: Starch content of WT, *phot* and various *phot*-complemented lines grown under continuous white light. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Asterisks indicated the p-values (\*, p<0.05; \*\*, p<0.01; \*\*\*\*, p<0.001; \*\*\*\*\*, p<0.001; \*\*\*\*, p<0.001; \*\*\*\*, p<0.001; \*\*\*\*, p<0.001; \*\*\*\*\*, p<0.001; \*\*\*\*\*,

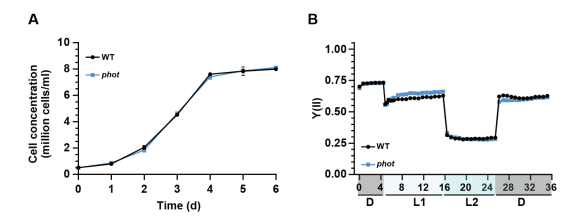


Fig. S4. The *phot* mutant shows WT-like growth and photosynthetic efficiency. Growth curves (A) and photosystem II efficiency (YII) (B) of WT and *phot* grown under continuous white light. *In vivo* chlorophyll fluorescence was recorded in the dark (labelled as "D"), at 21 (labelled as "L1") and 336 (labelled as "L2")  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup> as indicated in the graphs. Y(II) values calculated as (Fm'-F)/Fm' (n = 3 biological samples, mean  $\pm$  s.d.). Please note that in some cases the error bars are smaller than the data point symbols.

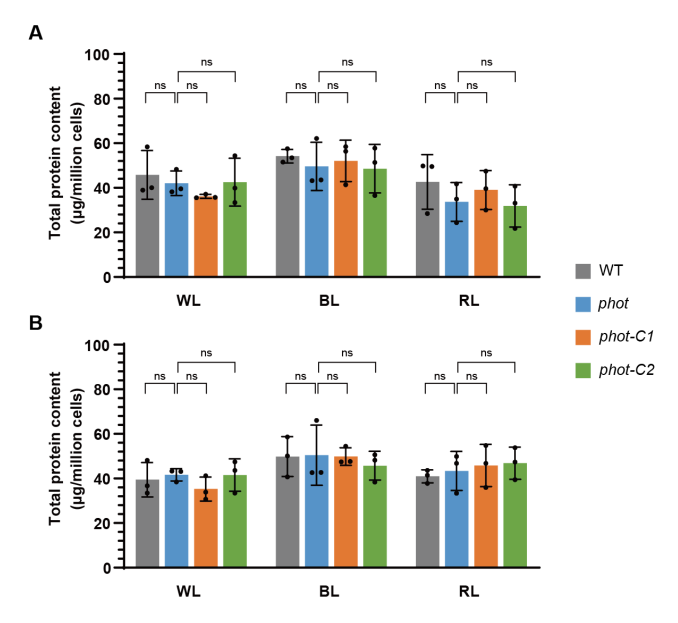


Fig. S5. The *phot* mutant has WT-like total protein content. The total protein content of WT, *phot* and complemented *phot* lines synchronized to a 12 h/12 h light dark cycle under different light qualities (WL, white light; BL, blue light; RL, red light). Samples were collected at the start of the start (A) and at the end (B) of the light phase. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). ns, not significant.

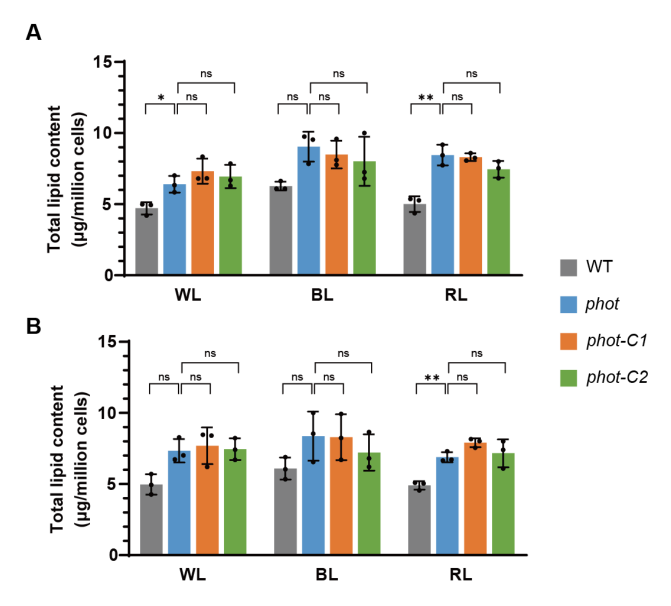
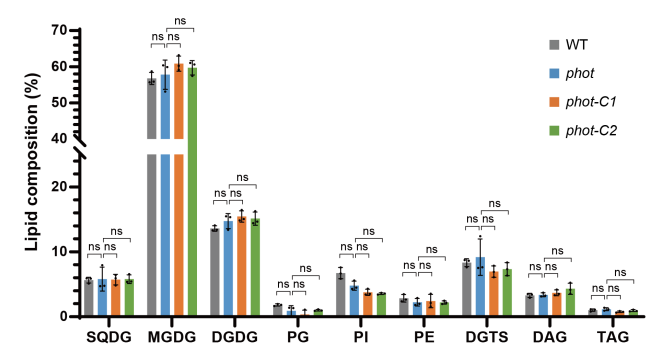
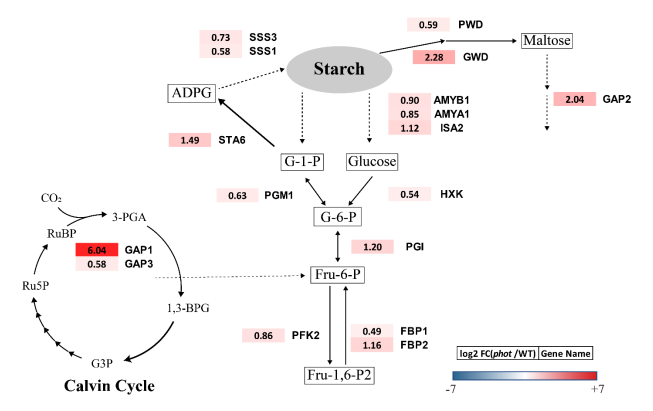


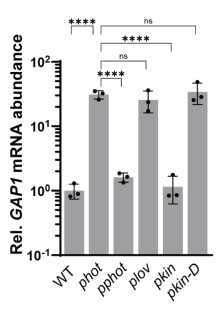
Fig. S6. The *phot* mutant has WT-like total lipid content. The total lipid content of WT, *phot* and complemented *phot* lines synchronized to a 12 h/12 h light dark cycle under different light qualities (WL, white light; BL, blue light; RL, red light). Samples were collected at the start of the start (A) and at the end (B) of the light phase. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Asterisks indicated the p-values (\*\*, p < 0.01; ns, not significant).



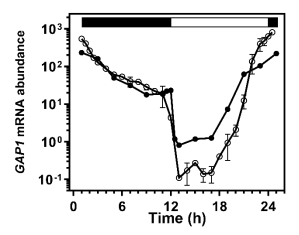
**Fig. S7. The** *phot* **mutant has WT-like lipid composition.** The lipid composition of WT, *phot and* complemented *phot lines* under 12h/12h light dark conditions. Samples are collected at the end of the light phase. Data are represented as mean ± SD (n = 3 biologically independent samples). SQDG, sulfoquinovosyl diacylglycerol; MGDG, monogalactosyl diacylglycerol; DGDG, digalactosyl diacylglycerol; PG, Phosphatidylgylcerols; PI, Phosphatidylinositols; PE, Phosphatidylethanolamines; DGTS, diacylglycerol-trimethyl homoserine; DAG, diacylglycerols; TAG, triacylglycerols. In some cases, the error bars are smaller than the data point symbols. ns, not significant.



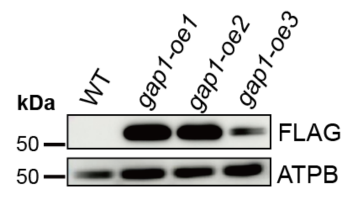
**Fig. S8.** The mRNA abundance of genes involved in starch metabolism is increased in the *phot* mutant. WT and *phot* were synchronized to a 12h/12h light dark cycle under white light. Samples were taken in the middle of the light phase (6h) and the extracted RNA was analyzed by RT-qPCR to quantify mRNA levels of the following genes: GAP, glyceraldehyde-3-phosphate dehydrogenase; PFK, phosphofructokinase; FBP, Fructose-1,6-bisphosphatase; PGI, phosphoglucose isomerase; PGM, phosphoglucomutase; HXK, Hexokinase; STA6, ADP-glucose pyrophosphorylase small subunit; SSS, starch synthase; AMY, α-amylase; ISA, isoamylase; PWD, phosphoglucan water dikinase; GWD, α-glucan water dikinase. Fold change between *phot* and WT was calculated as log2 FC (*phot*/WT).



**Fig. S9. PHOT suppresses** *GAP1* **mRNA accumulation.** *GAP1* relative mRNA abundance in WT (cw15-302) and in different phot-complemented lines, grown under continuous white light. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Asterisks indicated the p-values (\*, p<0.05; \*\*, p<0.01; \*\*\*, p<0.001; \*\*\*\*, p<0.001; \*\*\*\*, p<0.0001; ns, not significant).



**Fig. S10**. *GAP1* transcription level reaches its peak at dusk. *GAP1* expression profile in cells synchronized to a 12h/12h light dark cycle under white illumination, from *Strenkert et al.* (ref. 28, closed circles) and *Zones et al.* (ref. 27, open circles). The dark phase is indicated by black bars above the graphs; the light phase by white.



**Fig. S11**. **Immunoblot analyses of** *gap1-oe* **lines.** Samples were collected from cultures grown in mixotrophic medium containing acetate (TAP medium) under continuous white light. Total protein extracts were probed with anti-FLAG antibody to quantify GAP1-FLAG levels. ATPB served as a loading control.

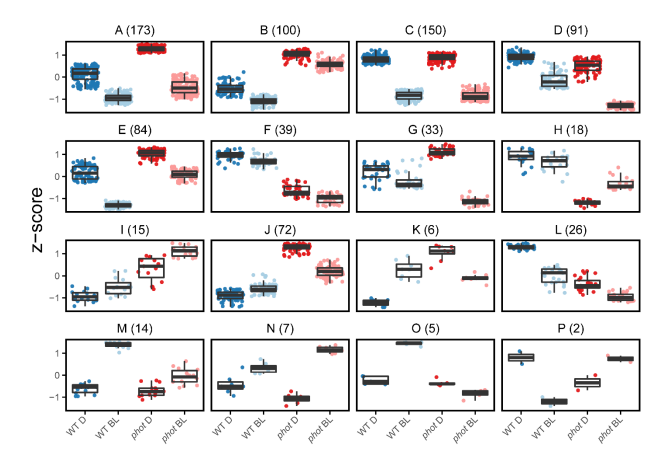


Fig. S12. Unsupervised hierarchical clustering analyses of phosphoproteome. Significantly changing phosphopeptides across treatments based on a one-way ANOVA (FDR-adjusted p-value < 0.05). Each cluster is labelled A-P, and the number of phosphopeptides contained in each cluster are in parenthesis. Z-scores represent the relative abundance of each peptide across condition.

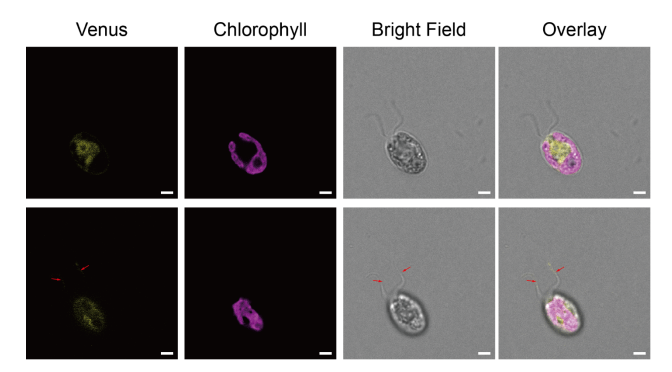


Fig. S13. PMSK1 subcellular localization. Representative confocal fluorescent microscopy images PMSK1-Venus (Yellow) constitutively expressed in WT. Scale bar: 2  $\mu$ m, red arrows indicate flagella. Two focal planes representative of the volume at 2.1  $\mu$ m in distance were shown.



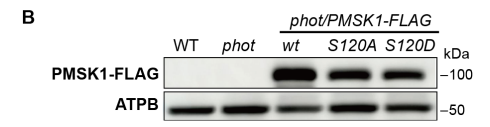
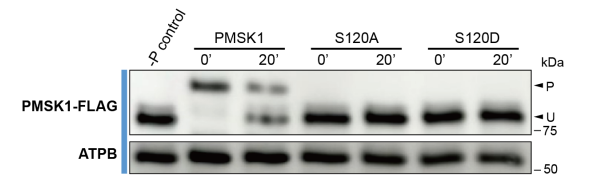


Fig. S14. PMSK1-FLAG protein levels in WT/PMSK1-FLAG and phot/PMSK1-FLAG lines. Samples were collected from cultures grown in mixotrophic medium containing acetate (TAP medium) under continuous white light. Total protein extracts were probed with anti-FLAG antibody to quantify PMSK1-FLAG levels in the WT (A) or phot (B) background. ATPB served as a loading control.



**Fig. S15.** The blue-light triggered mobility shift of the PMSK1-FLAG protein is mediated by serine 120. Changes in the phosphorylation level of PMSK1-FLAG in WT/PMSK1-FLAG, WT/PMSK1<sup>S120A</sup>-FLAG and WT/PMSK1<sup>S120D</sup>-FLAG lines. Samples were collected after 24h of acclimation to darkness (t=0') and 20' after exposure to blue light (100 μmol photons m<sup>-2</sup> s<sup>-1</sup>). Detection was performed by phos-tag SDS-PAGE; ATPB was used as a loading control. Phosphatase-treated WT/PMSK1-FLAG sample was also loaded on the gels. "U" and "P" indicate the unphosphorylated and phosphorylated PMSK1-FLAG respectively.

## WT/PMSK1 S120D-FLAG

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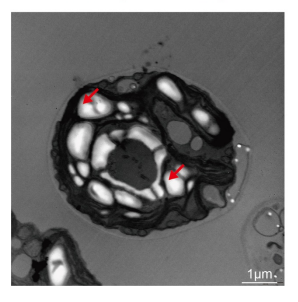
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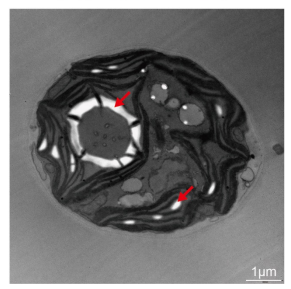
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## phot/PMSK1 S120A-FLAG





**Fig. S16.** Transmission electron microscopy pictures of WT/pmsk1<sup>S120D</sup> and phot/pmsk1<sup>S120A</sup>. The two strains were synchronized to a 12/12 light dark cycle under white light and samples were collected at the end of the light phase. Bar scale: 1 μm. Red arrows indicate starch granules.

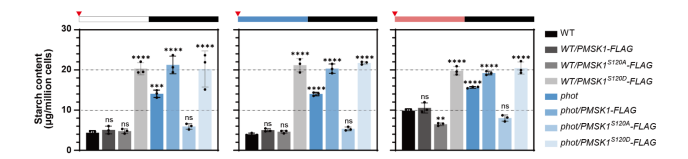


Fig. S17. Impact of PMSK1-FLAG overexpression in WT and phot on starch metabolism. Starch content in WT or phot overexpressing PMSK1-FLAG. Cells were synchronized to a 12h/12h light dark cycle under different light qualities. The dark phase is indicated by black bars above the graphs; the light phase by white, blue or red bars, depending on the light quality used. Red triangles indicate sample collection time. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Asterisks indicate the *P* values compared to WT. (\*, p<0.05; \*\*, p<0.01; \*\*\*, p<0.001; \*\*\*\*, p<0.001; \*\*\*\*, p<0.0001; ns, not significant). Please note that in some cases the error bars are smaller than the data point symbols.

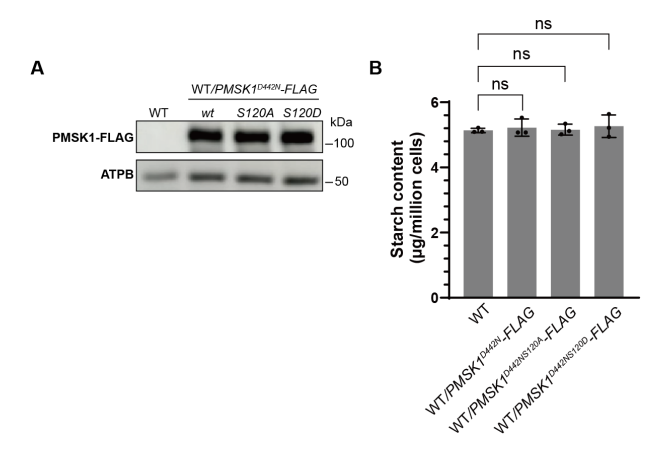


Fig. S18. PMSK1 kinase activity is necessary to mediate light quality dependent starch metabolism in *Chlamydomonas reinhardtii* (A) Immunoblot analyses of PMSK1-FLAG in various PMSK1-FLAG overexpressing lines. ATPB served as a loading control. (B) Starch content of various dead kinase PMSK1-FLAG overexpression lines grown under continuous white light. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). ns, not significant.

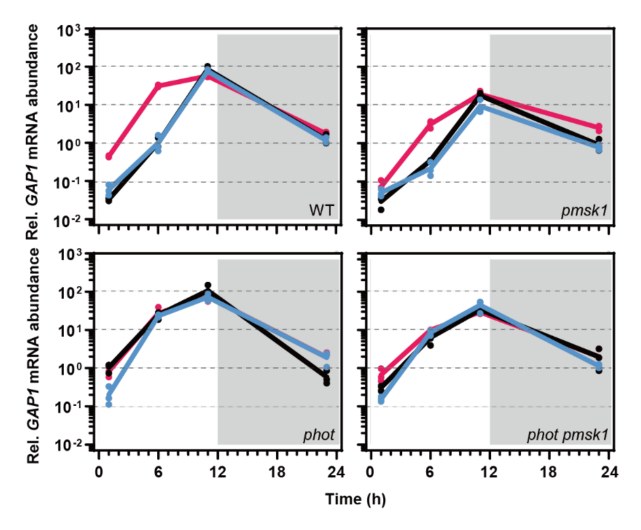


Fig. S19. GAP1 relative mRNA abundance in WT and in single and double phot pmsk1 mutants synchronized to a 12h/12h light dark cycle under different light qualities. Phases are indicated by white and gray shading. Line colors indicated light qualities. Black, white light; Red, red light; Blue, blue light. Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Please note that in some cases the error bars are smaller than the data point symbols.

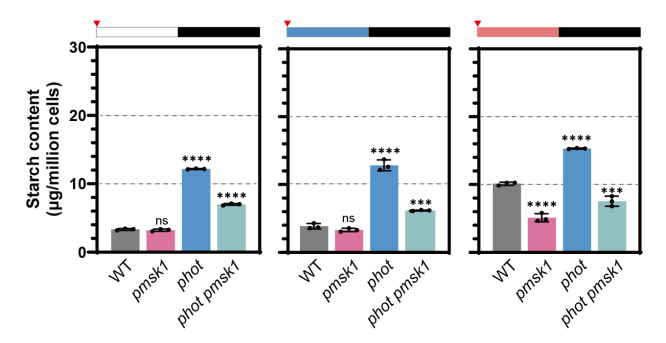
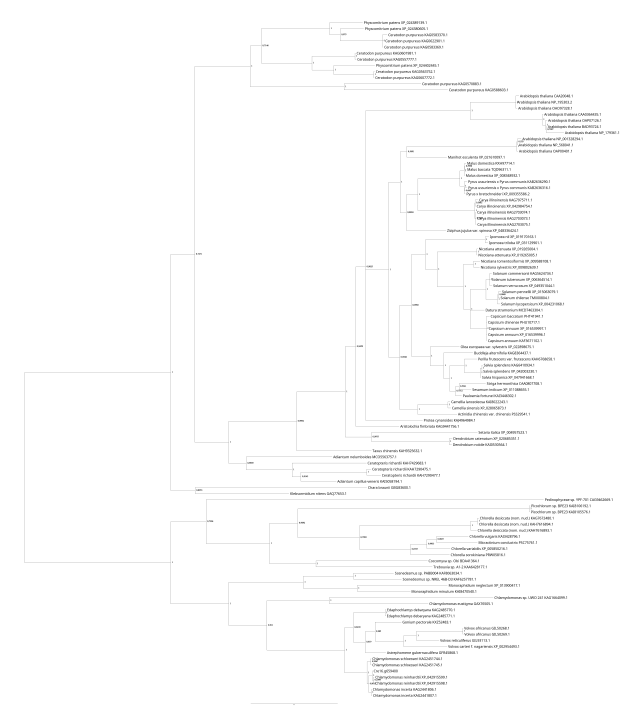
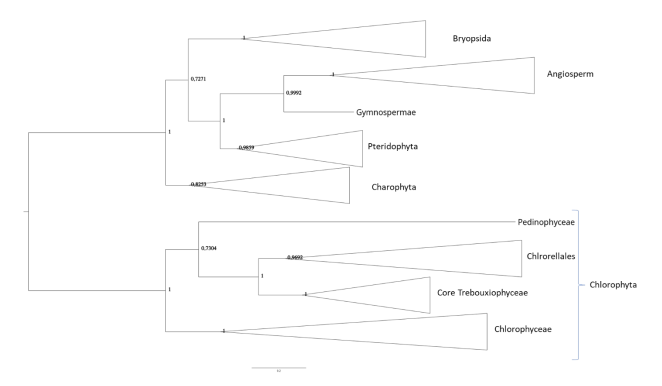


Fig. S20. Starch content in WT and in single and double *phot pmsk1* mutants sampled at dawn. Cells were synchronized to a 12h/12h light dark cycle under different light qualities. The dark phase is indicated by black bars above the graphs; the light phase by white, blue or red bars, depending on the light quality used. Red triangles indicate sample collection time. Asterisks indicated the p-values compared to WT. (\*, p<0.05; \*\*, p<0.01; \*\*\*\*, p<0.001; \*\*\*\*, p<0.001; ns, not significant). Data are represented as mean  $\pm$  SD (n = 3 biologically independent samples). Please note that in some cases the error bars are smaller than the data point symbols.



**Fig. S21. Phylogenetic tree of the PMSK-like family.** Complete Unrooted phylogenetic tree of the PMSK-like family (dataset #1) identified as described in the "Identification of the PMSK-like familly" from Chlorophyta representatives (Chlorophyceae, core Trebouxiophyceae, Chlorellales, Pedinophyceae), Charophyta, Briopsida, Pteridophyta, Gymnosperma and Angiosperma. The tree presented was inferred by Bayesian analysis as described in the "Methods" section. Bayesian Posterior Probability (BPP) values are reported at each node.



**Fig. S22.** Evolutionary relationship between major green lineage clades inside the PMSK-like family. Simplified version of the S21phylogenetic tree of the PMSK-like family (dataset #1). The tree presented was inferred by Bayesian analysis as described in the "Methods" section. Bayesian Posterior Probability (BPP) values are reported at each node. This tree highlights the monophyletic origin of the PMSK family. The root represented on this tree is here to help the identification of major clades with the S21 figures is not indicative of an ancestral state.

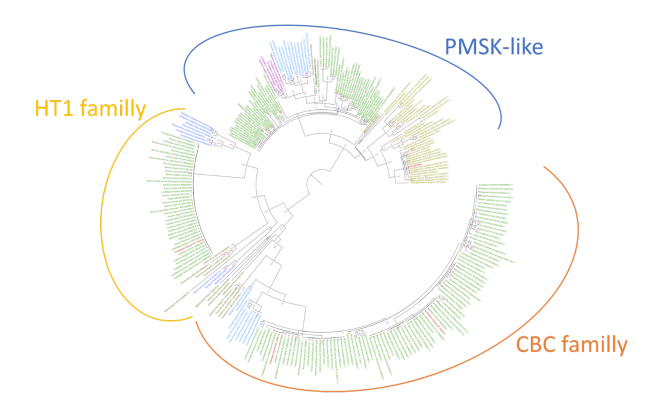


Fig. S23. Evolutionary analysis by Maximum Likelihood method of the PMSK-like family with the HT1 and the CBC family

The evolutionary history was inferred from the dataset #2 by using the PhyML maximum likehood method from Chlorophyta representatives (Chlorophyceae, core Trebouxiophyceae, Chlorellales, Pedinophyceae), Charophyta, Briopsida, Pteridophyta, Gymnosperma and Angiosperma. The tree with the highest log likelihood (-4797.84912) is shown. The percentage of trees in which the associated taxa clustered together is shown next to the branches. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the WAG model, and then selecting the topology with superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites (4 categories (+G, parameter = 1.283)). The rate variation model allowed for some sites to be evolutionarily invariable ([+I], 15.3% sites). The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. This analysis involved 278 amino acid sequences. Evolutionary analyses were conducted in MEGA11.

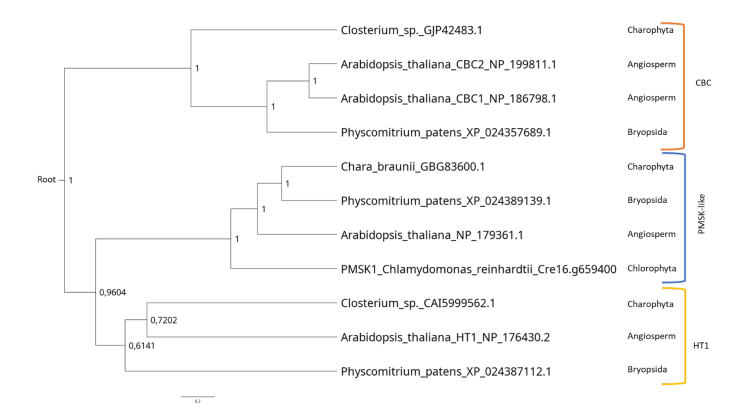


Fig. S24. Phylogenetic Clock tree of representatives of the PMSK-like, the HT1 and the CBC families. Rooted phylogenetic tree of representatives of the PMSK-like, the HT1 and the CBC families. The tree presented was inferred by Bayesian analysis with a clock-uniform method as described in the "Methods" section. Bayesian Posterior Probability (BPP) values are reported at each node. The root position was evaluated during the phylogenetic computation and is indicative of the evolutionary relationship between the three represented proteins families.