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# Phosphoprotein SAK1 is a regulator of acclimation to singlet oxygen in Chlamydomonas reinhardtii

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**Abstract** Singlet oxygen is a highly toxic and inevitable byproduct of oxygenic photosynthesis. The unicellular green alga *Chlamydomonas reinhardtii* is capable of acclimating specifically to singlet oxygen stress, but the retrograde signaling pathway from the chloroplast to the nucleus mediating this response is unknown. Here we describe a mutant, *singlet oxygen acclimation knocked-out 1* (*sak1*), that lacks the acclimation response to singlet oxygen. Analysis of genome-wide changes in RNA abundance during acclimation to singlet oxygen revealed that SAK1 is a key regulator of the gene expression response during acclimation. The *SAK1* gene encodes an uncharacterized protein with a domain conserved among chlorophytes and present in some bZIP transcription factors. The SAK1 protein is located in the cytosol, and it is induced and phosphorylated upon exposure to singlet oxygen, suggesting that it is a critical intermediate component of the retrograde signal transduction pathway leading to singlet oxygen acclimation.

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#### Introduction

Growth of photosynthetic organisms depends on light energy, which in turn can cause oxidative damage to the cell if not managed properly (*Li et al., 2009*). Light intensity is highly dynamic in terrestrial and aquatic environments, and the cell must constantly control the dissipation of light energy to avoid photo-oxidative stress while maximizing productivity. In addition to being the site of photosynthesis, the chloroplast houses many essential biochemical reactions such as fatty acid and amino acid biosynthesis, but most of its proteins are encoded in the nucleus and must be imported after translation. Therefore the nucleus must monitor the status of the chloroplast and coordinate gene expression and synthesis of proteins to maintain healthy chloroplast functions.

It is known that signals originating from a stressed or dysfunctional chloroplast modulate nuclear gene expression, a process that is called retrograde signaling (**Nott et al., 2006**; **Chi et al., 2013**). In Arabidopsis thaliana the gun mutants have helped to define the field of chloroplast retrograde signaling, leading to the identification of GUN1, a pentatricopeptide repeat protein that is a regulator of this process (**Koussevitzky et al., 2007**), and pointing to the involvement of the tetrapyrrole biosynthetic pathway (**Vinti et al., 2000**; **Mochizuki et al., 2001**; **Larkin et al., 2003**; **Strand et al., 2003**; **Woodson and Chory, 2008**). A role for heme in retrograde signaling has been shown in **Chlamydomonas** 



**eLife digest** Plants, algae and some bacteria use photosynthesis to extract energy from sunlight and to convert carbon dioxide into the sugars needed for growth. One by-product of photosynthesis is a highly toxic molecule called singlet oxygen. Typically, organisms deal with stressful events such as the presence of toxic molecules by producing new proteins. However, protein production is generally initiated in the nucleus of the cell, and photosynthesis is carried out in structures called chloroplasts. Cells must therefore be able to alert the nucleus to the presence of toxic levels of singlet oxygen in the chloroplasts.

Like some plants that can withstand a gradual decrease in temperature, but not a sudden cold snap, the alga *Chlamydomonas reinhardtii* is capable of resisting high doses of singlet oxygen if it has previously been exposed to low doses of the molecule. Wakao et al. exploited this ability to hunt for algae that are unable to acclimate to singlet oxygen, and found that these cells are unable to produce a protein called SAK1.

Wakao et al. reveal that many factors involved in the algae's cellular response to singlet oxygen depend on the presence of SAK1. In addition, the response of the algae cells to singlet oxygen differs to the one seen in the model plant *Arabidopsis thaliana*, suggesting that the two organisms have found different ways to deal with the same problem.

The location of a protein in a cell can give clues to its function. SAK1 is present in the fluid surrounding cellular compartments—the cytosol—which is consistent with it acting as a signaling molecule between the chloroplast and the nucleus. Wakao et al. present further evidence for this hypothesis by demonstrating that the number of phosphate groups attached on SAK1 changes when exposed to singlet oxygen—a feature often seen in signaling proteins. In addition, part of SAK1 resembles proteins that can bind to DNA, which indicates that SAK1 may be directly involved in initiating protein production.

The discovery of SAK1 represents a starting point for understanding how the site of photosynthesis, the chloroplast, communicates with the nucleus. It also has implications for developing plants and algae that have a higher tolerance to environmental stress conditions for agriculture and biofuel production.

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reinhardtii as well (von Gromoff et al., 2008). Many of the gun studies were conducted in context of a dysfunctional chloroplast treated with norflurazon, an inhibitor of carotenoid biosynthesis. More recently a number of exciting advances have shed light on small molecules playing roles in retrograde stress signaling, including methylerythritol cyclodiphosphate, an intermediate of isoprenoid biosynthesis in the chloroplast (Xiao et al., 2012), 3-phosphoadenosine 5-phosphate (PAP) (Estavillo et al., 2011), as well as a chloroplast envelope transcription factor PTM (Sun et al., 2011). Plastid gene expression involving sigma factors has been implicated in affecting nuclear gene expression, although the mechanism is unknown (Coll et al., 2009; Woodson et al., 2012).

Activation of gene expression by reactive oxygen species (ROS) has been well documented (Apel and Hirt, 2004; Mittler et al., 2004; Gadjev et al., 2006; Li et al., 2009). Thus ROS have been proposed as a means for chloroplasts to signal stress to the nucleus and many examples of global gene expression changes in response to ROS have been described (Desikan et al., 2001; Vandenabeele et al., 2004; Vanderauwera et al., 2005). Singlet oxygen (102) is a highly toxic form of ROS that can be formed in all aerobic organisms through photosensitization reactions in which excitation energy is transferred from a pigment molecule to  $O_2$ . For example, porphyria in humans is caused by defects in tetrapyrrole metabolism that can lead to accumulation of photosensitizing intermediates, which generate <sup>1</sup>O<sub>2</sub> in the light (Straka et al., 1990). In oxygenic photosynthetic organisms, <sup>1</sup>O<sub>2</sub> is mainly generated at the reaction center of photosystem II, when triplet excited chlorophyll transfers energy to O<sub>2</sub> (Krieger-Liszkay, 2005). 102 is the predominant cause of lipid oxidation during photo-oxidative stress (Triantaphylidès et al., 2008) and is associated with damage to the reaction center (Trebst et al., 2002). Because of the abundance and proximity of the two elements of <sup>1</sup>O<sub>2</sub> generation, the photosensitizer chlorophyll and O2, it was hypothesized that oxygenic photosynthetic organisms must have evolved robust means to cope with this ROS (Knox and Dodge, 1985). In Arabidopsis, the EX1 and EX2 proteins in the chloroplast are required for the execution of a <sup>1</sup>O<sub>2</sub>-dependent response: growth



arrest in plants and programmed cell death in seedlings, that is distinct from cell damage (op den Camp et al., 2003; Wagner et al., 2004; Lee et al., 2007). Different players in  ${}^{1}O_{2}$  signaling have emerged recently, such as β-cyclocitral, an oxidation product of β-carotene in Arabidopsis (Ramel et al., 2012), a bZIP transcription factor (SOR1) responding to reactive electrophiles generated by  ${}^{1}O_{2}$  (Fischer et al., 2012), and a cytosolic zinc finger protein conserved in Arabidopsis and Chlamydomonas, MBS (Shao et al., 2013). In the anoxygenic photosynthetic bacterium Rhodobacter sphaeroides, a  $\sigma^{E}$  factor is responsible for the elicitation of the gene expression response to  ${}^{1}O_{2}$  (Anthony et al., 2005).

The unicellular green alga *Chlamydomonas reinhardtii* is an excellent model organism for investigation of retrograde  ${}^{1}O_{2}$  signaling. *Chlamydomonas* exhibits an acclimation response to  ${}^{1}O_{2}$ , in which exposure to a sublethal dose of  ${}^{1}O_{2}$  leads to changes in nuclear gene expression that enable cells to resist a subsequent challenge with higher levels of  ${}^{1}O_{2}$  (*Ledford et al., 2007*). We hypothesized that acclimation mutants should include regulatory mutants that are defective in sensing and responding to  ${}^{1}O_{2}$ . Here we describe the isolation of such a mutant and identification of a cytosolic phosphoprotein SAK1 that is critical for the acclimation and transcriptome response to  ${}^{1}O_{2}$ .

#### Results

### Isolation of a singlet oxygen-sensitive mutant that is defective in acclimation

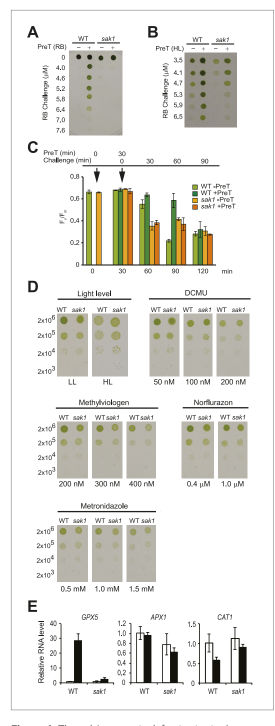
Chlamydomonas acclimates to singlet oxygen (102) generated by the exogenous photosensitizing dye rose bengal (RB) in the light (Ledford et al., 2007). As shown in Figure 1A, wild-type (WT) cells that were pretreated with RB in the light were able to survive a challenge treatment with much higher concentrations of RB, unlike cells pretreated with RB in the dark. By screening an insertional mutant population (Dent et al., 2005) for strains that were sensitive to 102, we isolated a mutant called singlet oxygen acclimation knocked-out1 (sak1) that is defective in acclimation to <sup>1</sup>O<sub>2</sub> (Figure 1A). We have previously shown that Chlamydomonas WT cells can also acclimate to RB following pretreatment with high light (Ledford et al., 2007), indicating that high light and RB induce overlapping responses to <sup>1</sup>O<sub>2</sub>. When subjected to the same conditions (high light pretreatment followed by challenge with RB), sak1 demonstrated less robust cross-acclimation (Figure 1B). We also tested conversely whether pretreatment with RB can acclimate the cells to growth in high light or in the presence of norflurazon. No increase in resistance to high light or norflurazon was induced by pretreatment with RB in either WT or sak1 (Figure 1—figure supplement 1). The viability phenotypes after RB treatment shown in Figure 1A were paralleled by changes in F<sub>v</sub>/F<sub>m</sub> values, a chlorophyll fluorescence parameter representing photosystem II efficiency (Figure 1C). In both WT and sak1, pretreatment did not cause an inhibition of photosystem II, as demonstrated by unchanged F<sub>v</sub>/F<sub>m</sub> values after 30 min. However, pretreatment increased resistance of photosystem II to the RB challenge only in WT and not in sak1 cells (Figure 1C). The pretreatment protected the cells only transiently, as by 90 min of challenge treatment both genotypes appeared to have experienced similar inhibition of photosystem II (Figure 1C), consistent with the hypothesis that sak1 is disrupted in early sensing and/or initiation of <sup>1</sup>O<sub>2</sub> response rather than its direct detoxification.

In contrast to its RB sensitivity, sak1 exhibited wild-type resistance to high light, various photosynthetic inhibitors and generators of other ROS, suggesting its defect is specific to  ${}^{1}O_{2}$  (*Figure 1D*). When tested for the gene expression response of the known  ${}^{1}O_{2}$ -specific gene GPX5 (*Leisinger et al., 2001*) during acclimation, WT cells showed a 20- to 30-fold induction, whereas a known  $H_{2}O_{2}$ -responsive ascorbate peroxidase gene (APX1) in *Chlamydomonas* (*Urzica et al., 2012*) and a catalase gene (CAT1), known to be  $H_{2}O_{2}$  responsive in *Arabidopsis* (*Davletova et al., 2005*; *Vanderauwera et al., 2005*), were unchanged. The mutant sak1 showed attenuated GPX5 induction, as expected for a mutant defective in the  ${}^{1}O_{2}$  response (*Figure 1E*).

## The global gene expression response to ${}^{1}O_{2}$ in *Chlamydomonas* is distinct from that in *Arabidopsis*

To obtain insight into the cellular processes and the genes involved in  ${}^{1}O_{2}$  acclimation, we used RNA-seq to define the transcriptome of WT cells during acclimation. The sequences were mapped to the *Chlamydomonas reinhardtii* genome version 4 (v4), and 16476 transcripts corresponding to gene models were detected (*Wakao et al., 2014*). We validated the data by quantitative reverse transcriptase PCR (qRT-PCR) for some of the differentially expressed genes during acclimation (*Figure 2*).





**Figure 1**. The *sak1* mutant is defective in singlet oxygen acclimation. **(A)** Acclimation phenotype of WT and *sak1*. The cells were pretreated in the dark (–) or under light (+) in the presence of rose bengal (RB), which requires light for generation of  ${}^{1}O_{2}$ . Pretreatment was followed by a subsequent higher concentration of RB (Challenge) as indicated under light. **(B)** Cells grown in low light were either kept in low light (–) or transferred to high light (+) for an hour before challenge in the light with increasing RB concentrations. **(C)**  $F_{v}/F_{m}$  *Figure 1. Continued on next page* 

Basal expression of some of the genes was elevated in sak1 compared to WT (Cre16.g683400 and GST1, Figure 2). Comparisons of the fold change (FC) values obtained by RNA-seq and gRT-PCR for the genes tested in Figure 2 are shown in Figure 2. The FC values are comparable between the two methods, although genes with FC greater than 20 (detected by RNA-seq) showed FC values (estimated by qRT-PCR) that were two to three times higher (Cre06.g281250. t1.1, Cre13.g566850.t1.1, Cre06.g263550.t1.1, Cre14.g623650.t1.2). Some of the genes were also induced by a transition from low light to high light, although not as strongly (Table 1), indicating that the 1O2 response elicited by addition of RB partly overlaps with that caused by increased light intensity. To examine whether the transcriptome changes were specific to <sup>1</sup>O<sub>2</sub>, we examined the expression of several previously identified H<sub>2</sub>O<sub>2</sub>-responsive genes (Urzica et al., 2012) (Table 2). Two of the seven genes, VTC2 (3.4-fold) and DHAR1 (twofold) were induced during <sup>1</sup>O<sub>2</sub> acclimation, whereas the other five genes were not differentially expressed (induced more than twofold) in our data. For these two genes, their magnitude of induction by <sup>1</sup>O<sub>2</sub> was smaller than that of H2O2-treated cells (both genes were ~ninefold induced by 1 mM H<sub>2</sub>O<sub>2</sub> treatment for 60 min) (Urzica et al., 2012). These differences suggest that our treatment with <sup>1</sup>O<sub>2</sub> did not lead to a large-scale induction of H<sub>2</sub>O<sub>2</sub>-responsive genes, and it is likely that the two abovementioned genes involved in ascorbate metabolism respond to both  $H_2O_2$  and  ${}^1O_2$ .

During acclimation of WT to <sup>1</sup>O<sub>2</sub>, 515 genes were up-regulated at least twofold with a false discovery rate (FDR) smaller than 1% (Supplementary file 1, C1), and 33% of these could be categorized into functional classes based on MapMan (Thimm et al., 2004) using the Algal Functional Annotation Tool (Lopez et al., 2011) (Figure 3A,B). The enriched classes are marked with asterisks, and the genes within those classes are listed in Table 3. Genes involved in sterol/squalene/brassinosteroid metabolism (in the hormone and lipid metabolism functional classes) were notably enriched (Table 3). A sterol methyltransferase was also detected to display differential expression in our previous microarray analysis (Ledford et al., 2007). Brassinosteroids are not known to exist in Chlamydomonas, and in plants increasing evidence indicates sterols have a signaling role independent of brassinosteroids (Lindsey et al., 2003; Boutté and Grebe, 2009). Two cyclopropane fatty acid synthases (CFAs) were among the up-regulated lipid metabolism



Figure 1. Continued

values were measured after each time point indicated. Pretreatment (PreT) with  $0.5 \mu M$  RB was applied for 30 min with (+PreT) or without (-PreT) light. After the pretreatment, RB was added to both dark and light samples to a final concentration of 3.75 µM RB (challenge), and  $F_v/F_m$  was measured for 90 min at 30 min intervals (total 120 min). First arrow: addition of pretreatment; second arrow: addition of challenge. (D) sak1 has wild-type sensitivity to other photo-oxidative stresses. Serial dilutions of WT and sak1 were spotted onto minimal (HS) plates at the indicated light intensity or on TAP plates containing the indicated inhibitor. DCMU, 3-(3,4-dichlorophenyl)-1,1-dimethylurea; low light (LL), 80 µmol photons m<sup>-2</sup> s<sup>-1</sup>; high light (HL), 450  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>. (**E**) Gene expression of a known <sup>1</sup>O<sub>2</sub>-responsive gene, GPX5, is induced during acclimation, while two genes associated with H<sub>2</sub>O<sub>2</sub> response, APX1 and CAT1, are not. WT cells were mock-pretreated without RB (white bars) or pretreated with RB in the light (black bars).

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The following figure supplements are available for figure 1:

**Figure supplement 1**. Pretreatment with RB does not increase resistance to high light or norflurazon in cells grown on plates.

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genes (Table 3). Another function that was notable among up-regulated genes, although they were not grouped to a common functional class by MapMan, were two genes coding for SOUL heme-binding domain proteins that were SAK1dependent (SOUL2 and Cre06.g299700.t1.1, formerly annotated as SOUL1) (Figure 2). Genes annotated as involved in transport comprised one of the most enriched classes (Figure 3B). These included a number of multidrug-resistant (MDR) and pleiotropic drug-resistant (PDR) type transporters as well as other various transporters for ions, peptides, and lipids (Table 3). The former types of transporters may reflect the cells' response to pump RB out. When the responses to the chemical RB and <sup>1</sup>O<sub>2</sub> were uncoupled by comparing gene expression in cultures kept in the dark with and without RB, all of the tested <sup>1</sup>O<sub>2</sub>-induced genes and ABC transporters identified from our RNA-seq remained unchanged by RB in the dark in both WT and sak1 (Table 4). This result indicates that the up-regulation of these genes when RB was added in the light was a response to <sup>1</sup>O<sub>2</sub> rather than to RB itself. Up-regulation of stress genes included those coding for chaperones and some receptor-like proteins (Figure 3B; Table 3), suggesting that the cells do mount a stress response during accli-

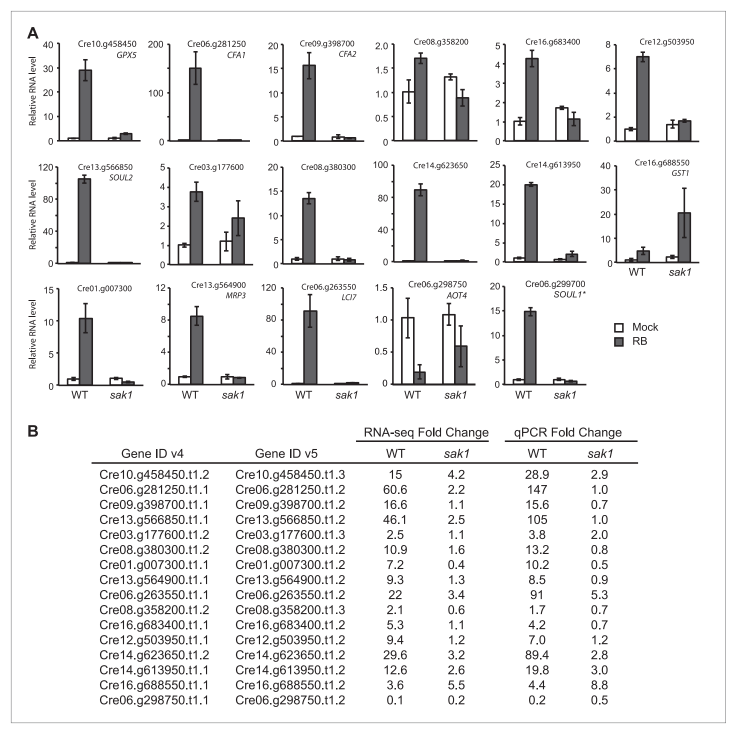
mation though not visible by gross growth phenotype (*Figure 1A*) or decrease in  $F_v/F_m$  (*Figure 1C*). A smaller number of 219 genes was down-regulated during acclimation in WT (*Supplementary file 1*, C1), only 21% of which had functional annotation. The most enriched classes of down-regulated genes were nucleotide metabolism and transport, the latter including a distinct type of transporter for small metabolites and ions, different from those found among up-regulated genes that included many MDR- and PDR-type transporters (*Figure 3B*; *Table 3*).

Although only 33% of the up-regulated genes have a functional annotation (Figure 3B), it is interesting that the <sup>1</sup>O<sub>2</sub> response in Chlamydomonas involves genes and biological processes that appear to be distinct from those that respond specifically to <sup>1</sup>O<sub>2</sub> in Arabidopsis (op den Camp et al., 2003). A total of 70 <sup>1</sup>O<sub>2</sub>-response genes have been defined using a microarray with the flu mutant in Arabidopsis (op den Camp et al., 2003). These genes include the following classes (number of genes): metabolism (11), transcription (5), protein fate (4), transport (2), cellular communication/signal transduction (17), cell rescue/defense in virulence (4), subcellular localization (2), binding function or cofactor requirement (1), transport facilitation (5) and others (19). From this list of 70 genes we found four similarly annotated genes within our 515 genes induced by 102 in Chlamydomonas: a Myb transcription factor, a mitochondrial carrier protein, an amino acid permease, and an ATPase/ aminophospholipid translocase. None of these genes in Chlamydomonas was the closest ortholog of the corresponding Arabidopsis gene. Conversely, genes similar to those strongly up-regulated in a SAK1-dependent manner such as CFAs, SOUL proteins, GPX, and sterol biosynthetic enzymes were not found among the Arabidopsis 1O2-specific genes despite having clear counterparts in Arabidopsis. Taken together, these results suggest that these two organisms may deploy distinct mechanisms in their responses to  ${}^{1}O_{2}$ .

## The sak1 mutant is defective in the global gene expression response during acclimation to <sup>1</sup>O<sub>2</sub>

In the sak1 mutant, 1020 genes were up-regulated, whereas 434 genes were down-regulated during acclimation (**Supplementary file 1**, C2). 350 of the 515 genes up-regulated in WT overlapped with the set of up-regulated genes in the mutant (**Figure 3A**). Comparing the fold changes of genes in WT and





**Figure 2.** qRT-PCR analysis of genes identified to be  ${}^{1}O_{2}$ -responsive by RNA-seq. (**A**) The error bars indicate standard deviation of biological triplicates. The locus of the transcript (v5) and gene name if annotated, are indicated. \*SOUL1 was named gene in v4 but not in v5. (**B**) Comparison of fold change values from RNA-seq data and qPCR. Fold change values were calculated for RNA-seq as described in 'Material and methods', and the values for qPCR are averages obtained from biological triplicates.

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sak1 during acclimation, we defined 104 genes as SAK1-dependent genes that displayed moderate to strong attenuation in their response (fold change ratio <0.5) (*Table 5*). Some of the genes that belong to enriched biological classes found among WT up-regulated genes are indicated in *Table 3*. Interestingly, the most strongly induced genes in WT were found among this group; 37 out of 104



**Table 1.** Moderate induction of  ${}^{1}O_{2}$  genes during high light exposure

	Fold change (SD)*	
Gene name or ID	WT	sak1
GPX5	2.86 (1.06)	1.08 (0.23)
CFA1	3.75 (0.99)	1.78 (0.52)
SOUL2	3.45 (1.25)	1.82 (0.22)
MRP3	3.10 (0.39)	2.37 (0.32)
Cre14.g613950	1.42 (0.53)	1.57 (0.46)
LHCSR1†	14.91 (4.25)	2.91 (1.35)

<sup>\*</sup>Fold change values are the average of biological triplicates and their standard deviations are indicated in parentheses.

†Known to have elevated expression in high light grown cells (**Peers et al., 2009**).

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SAK1-dependent genes were among the top 10% most strongly induced genes (*Table 5*). 33 out of these 37 most strongly induced SAK1-dependent genes displayed strong disruption in their up-regulation; reduced to 0.01-0.25 of magnitude of fold change in sak1 as compared to WT (*Table 5*). These results indicate SAK1 is required for the induction of the most strongly induced genes during acclimation reflecting its critical role in regulating the cellular acclimation response to  ${}^{1}O_{2}$ .

Classes of up-regulated genes in sak1 were distinct from those of WT and included secondary metabolism of isoprenoids (Figure 3C; Table 6), precursors to photoprotective pigments such as carotenoids and tocopherols (Li et al., 2009). Phenylpropanoids, a group of metabolites associated with defense against stresses such as ultraviolet light and herbivores (Maeda and Dudareva, 2012), also represented a larger

part of the response in sak1 as compared to WT (*Figure 3C*). Another mutant-specific class of genes was cell vesicular transport, suggesting alteration in cell organization in response to the loss of SAK1 (*Figure 3C*; *Table 6*). There were 434 genes that were down-regulated by  ${}^{1}O_{2}$  in the sak1 mutant (*Supplementary file 1*, C2), none of which overlapped with the set of down-regulated genes in WT, in contrast to the overlap of up-regulated genes in the two genotypes (*Figure 3A*). Enriched classes of genes included those involved in DNA, nucleotide metabolism, hormone metabolism (not of brassinosteroid) and tetrapyrrole metabolism (*Figure 3C*, *Table 6*).

To better understand the physiology of sak1, including the primary and secondary effects of lacking SAK1, we also focused on changes in transcript levels at the basal level, that is, without  ${}^{1}O_{2}$  treatment. At basal level 699 genes were induced, and 737 genes were repressed in the mutant compared to WT (*Supplementary file 1*, C3), displaying the genome-wide response to the loss of SAK1 function despite the mutant's wild-type appearance under normal lab growth conditions (*Figure 1D*). The enriched classes of genes that are differentially expressed are shown in *Figure 3D*. Genes induced in the mutant at basal level were enriched for those annotated to be involved in nucleotide metabolism, DNA, and RNA (*Figure 3D*; *Table 7*). Interestingly genes involved in tetrapyrrole and photosynthesis were enriched both in elevated and repressed genes at the basal level in sak1. There was no overall trend of these two pathways being up- or down-regulated, since these genes were at different steps of the pathway or encoded a select isoform of an enzyme or a subunit of a complex (*Figure 3D*; *Table 7*).

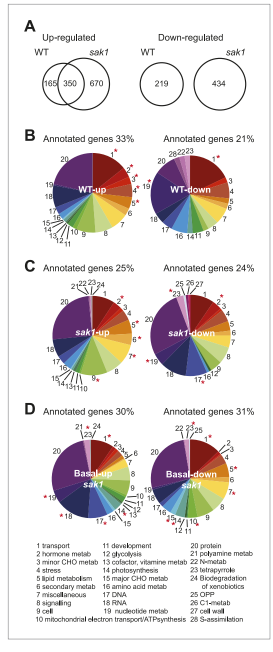
**Table 2.** Expression of H<sub>2</sub>O<sub>2</sub> response genes during <sup>1</sup>O<sub>2</sub> acclimation

	Gene ID		RPKM*				Fold changet	ŀ
Gene name	v4	v5	WT-mock	WT-RB	sak1-mock	sak1-RB	WT	sak1
APX1	Cre02.g087700.t1.1	Cre02.g087700.t1.2	49.70	36.22	79.65	58.83	0.73	0.74
MSD3	Cre16.g676150.t1.1	Cre16.g676150.t1.2	0.30	0.18	0.70	0.17	0.60	0.25
MDAR1	Cre17.g712100.t1.1	Cre17.g712100.t1.2	35.95	38.30	33.53	51.34	1.07	1.53
DHAR1	Cre10.g456750.t1.1	Cre10.g456750.t1.2	20.40	40.93	25.69	42.18	2.01	1.64
GSH1	Cre02.g077100.t1.1	Cre02.g077100.t1.2	28.27	26.91	40.42	49.95	0.95	1.24
GSHR1	Cre06.g262100.t1.2	Cre06.g262100.t1.3	19.17	19.02	19.39	22.41	0.99	1.16
VTC2	Cre13.g588150.t1.1	Cre13.g588150.t1.2	18.16	62.53	35.10	103.12	3.44	2.94

<sup>\*</sup>Average of RPKM obtained from two sequencing lanes as described in 'Material and methods'.

†Calculated as ratio of (RPKM-RB) / (RPKM-mock).

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**Figure 3**. Differentially expressed genes from pair-wise comparisons. (**A**) Venn diagram representing differentially expressed genes in WT and sak1. Mapman functional classes distribution of differentially expressed genes (passing criteria of fold change greater than  $2^1$  [up] or smaller than  $2^{-1}$  [down] with FDR <1%) during acclimation in (**B**) WT and (**C**) sak1. (**D**) Differentially expressed genes when comparing WT and sak1 in basal conditions (i.e., before exposure to  ${}^1O_2$ ). The functional classes represented by the numbers are listed; asterisks indicate classes that were enriched compared to the genome.

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We observed that some of the genes more strongly dependent on SAK1 had repressed transcript levels (e.g., CFA1 and SOUL2), indicating that SAK1 is required for their basal expression, while others had elevated basal levels (GPX5), suggesting that expression of these genes is controlled also by other pathways. As is discussed in the following section, SAK1 expression monitored by gRT-PCR followed the latter trend as the 5'UTR of the gene was elevated in the mutant (Figure 4E), which may be a result of response to other factors such as a possible oxidization product of <sup>1</sup>O<sub>2</sub>. The SAK1-dependent genes induced by <sup>1</sup>O<sub>2</sub> and repressed at basal level in the mutant (i.e., those that require SAK1 for basal expression) are indicated in Table 5.

## The sak1 mutant identifies a single nuclear gene that is itself induced during acclimation to <sup>1</sup>O<sub>2</sub>

The sak1 mutant was generated by insertional mutagenesis using a plasmid that confers resistance to zeocin (Dent et al., 2005). Progeny obtained from a backcross of sak1 with WT showed that the mutation causing the RB sensitivity phenotype was linked to zeocin resistance (Figure 4A). The site of insertion was identified by thermal asymmetric interlaced (TAIL)-PCR (Liu et al., 1995) as the second exon of the annotated gene Cre17.g741300 on chromosome 17 (Figure 4B). To test whether this gene is responsible for the mutant phenotype, a genomic fragment containing the gene with an additional ~500 bp region upstream of the predicted transcription start site was cloned and introduced into the mutant by co-transformation. Among the approximately 300 transformants screened, two clones appeared to have recovered the RB acclimation phenotype (Figure 4C). Furthermore, induction of genes we found attenuated in sak1 (Figure 2) was restored in these transformants (Figure 4D), confirming that Cre17.g741300 is the SAK1 gene required for acclimation and the gene expression response to  ${}^{1}O_{2}$ .

In WT, the SAK1 gene itself was induced by 6- to 10-fold during acclimation when probed for the 5'-and 3'-UTR of the transcript by qRT-PCR (Figure 4E). The mutant displayed elevated basal level and induction of the 5'-UTR during acclimation, whereas the 3'-UTR of the transcript was undetectable, indicating that the full-length transcript was absent in sak1 (Figure 4E). An antibody raised against an epitope of the SAK1 protein detected a single band in basal condi-

tions, whereas the SAK1 protein appeared as multiple bands with higher molecular weight in acclimated WT cells, all of which were absent in the mutant (Figure 4F). SAK1 transcript was induced



**Table 3.** Enriched functional classes among differentially expressed genes in WT during  ${}^{1}\text{O}_{2}$  acclimation

Primary MapMan	Secondary Mapman			Gene	
class	class	Gene ID (v4)	Gene ID (v5)	name	Annotation
Up-regulated	genes				
transport	ABC transporters and multidrug resistance systems	Cre03.g169300.t1.1	Cre03.g169300.t2.1		ABC transporter (ABC-2 type)
		Cre04.g220850.t1.1	Cre04.g220850.t1.2		ABC transporter (ABC-2 type)
		Cre11.g474600.t1.1§	Cre02.g095151.t1		ABC transporter (ABC-2 type)
		Cre03.g151400.t1.2	Cre03.g151400.t1.3		ABC transporter (subfamilyA member3)
		Cre14.g618400.t1.1§	Cre14.g618400.t1.2		ABC transporter
		Cre09.g395750.t1.2	Cre09.g395750.t1.3		ABC transporter (plant PDR pleitropic drug resistance)
		Cre14.g613950.t1.1§	Cre14.g613950.t2.1		ABC transporter, Lipid exporter ABCA1 and related proteins
		Cre17.g725150.t1.1	Cre17.g725150.t1.2		ABC transporter
		Cre04.g224400.t1.2§	Cre04.g224400.t1.3		ABC transporter (plant PDR pleitropic drug resistance)
		Cre13.g564900.t1.1§	Cre13.g564900.t1.2	MRP3	ABC transporter, Multidrug resistance associated protein
		Cre17.g721000.t1.1	Cre17.g721000.t1.2		ABC transporter (ABCA)
		Cre04.g224500.t1.2	Cre04.g224500.t1.3		ABC transporter (plant PDR pleitropic drug resistance)
		Cre01.g007000.t1.1§	Cre01.g007000.t1.2		ABC transporter (ABC-2 type)
	unspecified anions	Cre13.g574000.t1.2	Cre13.g574000.t1.3		Chloride channel 7
		Cre17.g729450.t1.1	Cre17.g729450.t1.2		Chloride channel 7
	amino acids	Cre04.g226150.t1.2	Cre04.g226150.t1.3	AOC1	Amino acid carrier 1; belongs to APC (amino acid polyamine organocation) family
	misc	Cre16.g683400.t1.1§	Cre16.g683400.t1.2		CRAL/TRIO domain (Retinaldehyde binding protein-related)
		Cre17.g718100.t1.1	Cre17.g718100.t1.2		Phosphatidylinositol transfer protein SEC14 and related proteins (CRAL/TRIO)
		Cre06.g311000.t1.2	Cre06.g311000.t1.3	FBT2	Folate transporte
	calcium	Cre09.g410050.t1.1§	Cre09.g410050.t1.2		Ca2+ transporting ATPase
	potassium	Cre07.g329882.t1.2	Cre07.g329882.t1.3		Ca2+-activated K+ channel proteins
	phosphate	Cre16.g686750.t1.1	Cre16.g686750.t1.2	PTA3	Proton/phosphate symporter
	metal	Cre13.g570600.t1.1	Cre13.g570600.t1.2	CTR1	CTR type copper ion transporter
	metabolite transporters at the mitochondrial membrane	Cre06.g267800.t1.2	Cre06.g267800.t2.1		Mitochondrial carrier protein
hormone metabolism*	brassinosteroid	Cre16.g663950.t1.1	Cre16.g663950.t1.2		Sterol C5-desaturase

Table 3. Continued on next page



Table 3. Continued

Primary MapMan	Secondary Mapman			Gene	
class	class	Gene ID (v4)	Gene ID (v5)	name	Annotation
		Cre02.g076800.t1.1	Cre02.g076800.t1.2		delta14-sterol reductase
		Cre12.g557900.t1.1	Cre12.g557900.t1.1	CDI1	C-8,7 sterol isomerase
		Cre02.g092350.t1.1	Cre02.g092350.t1.2		Cytochrome P450, CYP51 Sterol-demethylase
		Cre12.g500500.t1.2	Cre12.g500500.t2.1		SAM-dependent methyltransferases
	jasmonate	Cre19.g756100.t1.1	Cre03.g210513.t1		12-oxophytodienoic acid reductase
	auxin	Cre14.g609900.t1.1	Cre14.g609900.t1.1		Predicted membrane protein, contains DoH and Cytochrome b-561/ferric reductase transmembrane domains
		Cre06.g276050.t1.1	Cre06.g276050.t1.2		Aldo/keto reductase
		Cre16.g692800.t1.2	Cre16.g692800.t1.3		Aldo/keto reductase
		Cre03.g185850.t1.2	Cre03.g185850.t1.2		pfkB family, sugar kinase-related
minor CHC metabolism	) others	Cre06.g276050.t1.1	Cre06.g276050.t1.2		Aldo/keto reductase
		Cre16.g692800.t1.2	Cre16.g692800.t1.3		Aldo/keto reductase
		Cre03.g185850.t1.2	Cre03.g185850.t1.2		pfkB family, sugar kinase-related
	callose	Cre06.g302050.t1.1	Cre06.g302050.t1.2		1,3-beta-glucan synthase
	myo-inositol	Cre03.g180250.t1.1	Cre03.g180250.t1.2		Myo-inositol-1-phosphate synthase
stress	biotic	Cre01.g057050.t1.1§	Cre03.g144324.t1		Leucine Rich Repeat
		Cre01.g016200.t1.2	Cre01.g016200.t1		Mlo Family
		Cre28.g776450.t1.1§	Cre08.g358573.t1	PSMD10	26S proteasome regulatory complex
	abiotic	Cre12.g501500.t1.1	NF†		
		Cre02.g132300.t1.2	Cre09.g395732.t1		DnaJ domain
		Cre07.g339650.t1.2	Cre07.g339650.t1.3	DNJ20	DnaJ-like protein
		Cre01.g033300.t1.1§	Cre01.g033300.t2.1		No annotation‡
		Cre16.g677000.t1.1	Cre16.g677000.t1.2	HSP70E	Heat shock protein 70E
		Cre08.g372100.t1.1	Cre08.g372100.t1.2	HSP70A	Heat shock protein 70A
lipid metabolism	phospholipid synthesis	Cre13.g604700.t1.2	Cre13.g604700.t1.3	PCT1	CDP-alcohol phosphatidyltransferase/ Phosphatidylglycerol-phosphate synthase
		Cre06.g281250.t1.1§	Cre06.g281250.t1.2	CFA1	Cyclopropane fatty acid synthase
		Cre09.g398700.t1.1§	Cre09.g398700.t1.2	CFA2	Cyclopropane fatty acid synthase
	'exotics' (steroids, squalene etc)	Cre01.g061750.t1.1	Cre03.g146507.t1	SPT2	Serine palmitoyltransferase
		Cre83.g796250.t1.1	NF†	SPT1	Serine palmitoyltransferase
		Cre02.g137850.t1.1	Cre09.g400516.t1		TRAM (translocating chain- associating membrane) superfamily
	FA synthesis and FA elongation	Cre03.g182050.t1.1	Cre03.g182050.t1		Long-chain acyl-CoA synthetases (AMP-forming)

Table 3. Continued on next page



Table 3. Continued

Primary MapMan	Secondary Mapman			Gene	
class	class	Gene ID (v4)	Gene ID (v5)	name	Annotation
		Cre06.g256750.t1.1	Cre06.g256750.t1.2		Acyl-ACP thioesterase
misc	short chain dehydrogenase/ reductase (SDR)	Cre12.g556750.t1.2	Cre12.g556750.t1.3	1	Short chain dehydrogenase
		Cre27.g775000.t1.1	Cre12.g549852.t1		Short chain dehydrogenase
		Cre17.g731350.t1.2	Cre17.g731350.t1.2	!	Short chain dehydrogenase
		Cre08.g381510.t1.1§	NF†		Short chain alcohol dehydrogenase
	UDP glucosyl and glucoronyl transferases	Cre02.g144050.t1.1	Cre02.g144050.t2.1		Acetylglucosaminyltransferase EXT1/exostosin 1
		Cre16.g659450.t1.1	Cre16.g659450.t1.2		Lactosylceramide 4-alpha- Galactosyltransferase
		Cre03.g173300.t1.1	Cre03.g173300.t1.2		Lactosylceramide 4-alpha- Galactosyltransferase
	dynamin	Cre02.g079550.t1.1	Cre02.g079550.t1.2		Dynamin-related GTPase, involved in circadian rhythms
	misc2	Cre06.g258600.t1.1§	Cre06.g258600.t2.1		Predicted hydrolase related to dienelactone hydrolase
	acid and other phosphatases	Cre06.g249800.t1.1	Cre06.g249800.t1.2		Sphingomyelin synthase
Down-regula	ted genes				
nucleotide metabolism	salvage	Cre13.g573800.t1.1	Cre13.g573800.t1.2		Phosphoribulokinase / Uridine kinase family
	synthesis	Cre12.g503300.t1.1	Cre12.g503300.t1.2		Phosphoribosylamidoimidazole- succinocarboxamide synthase
		Cre06.g308500.t1.1	Cre06.g308500.t1.2	CMP2	Carbamoyl phosphate synthase, small subunit
		Cre14.g614300.t1.1	Cre14.g614300.t1.2	!	Inosine-5-monophosphate dehydrogenase
transport	ABC transporters and multidrug resistance systems	Cre06.g273750.t1.2	Cre06.g273750.t1.3	SUA1	Chloroplast sulfate transporter
		Cre02.g083354.t1.1	Cre02.g083354.t1		ATP-binding cassette, subfamily B (MDR/TAP), member 9
	calcium	Cre06.g263950.t1.2	Cre06.g263950.t1.3	}	Na+/K + ATPase, alpha subunit
	metabolite transporters at the envelope membrane	Cre08.g363600.t1.1	Cre08.g363600.t1.2	!	Glucose-6-phosphate, PEP/ phosphate antiporter
	metal	Cre17.g720400.t1.2	Cre17.g720400.t1.3	HMA1	Heavy metal transporting ATPase
	P- and V-ATPases	Cre10.g459200.t1.1	Cre10.g459200.t1.2	ACA4	Plasma membrane H + -transporting ATPase
	phosphate	Cre02.g144650.t1.1	Cre02.g144650.t1.2	PTB12	Na+/Pi symporter
	potassium	Cre06.g278700.t1.2	Cre06.g278700.t1.2		Myotrophin and similar proteins

<sup>\*</sup>Functional terms are inferred by homology to the annotation set of *Arabidopsis thaliana* (**Lopez et al., 2011**). †Corresponding gene model was not found in v5.

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<sup>‡</sup>No functional annotations found on v5 but defined by MapMan on Algal Functional Annotation Tool (**Lopez et al., 2011**). §Induction during  ${}^{1}O_{2}$  acclimation dependent on SAK1 (**Table 5**).



**Table 4.**  ${}^{1}\text{O}_{2}$  response genes are not induced when RB is added in the dark

Fold	chan	ge
+RB/	-RR	(SD)*

Gene name or ID	WT	sak1
GPX5	1.13 (0.33)	0.87 (0.31)
SAK1	1.38 (0.08)	1.29 (0.19)
CFA1	0.90 (0.04)	1.44 (0.22)
SOUL2	1.17 (0.25)	1.11 (0.19)
MRP3†,‡	1.13 (0.12)	1.07 (0.25)
Cre12.g503950†,‡	0.93 (0.06)	1.20 (0.12)
Cre14.g613950†,§	0.65 (0.06)	0.79 (0.15)
Cre04.g220850†,‡	1.00 (0.09)	1.29 (0.04)
Cre09.g395750†,‡	1.05 (0.10)	1.29 (0.12)

<sup>\*</sup>Average of fold change and standard deviation (SD) of biological triplicates.

†Annotated as transport function.

‡ABC transporter.

§Sec14-like phosphatidylinositol transfer protein.

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when probed for the 5'-UTR during high light exposure in both WT and sak1 (Figure 4G) similarly to other <sup>1</sup>O<sub>2</sub>-response genes identified by RNA-seq (Table 1), indicating that SAK1 itself is part of the endogenous response to high light.

## SAK1 contains an uncharacterized domain conserved in chlorophytes and found in some bZIP transcription factors

The predicted SAK1 protein consists of 1141 amino acid residues and has no domains with functional annotation. Only a ~150-residue region at the C-terminus, designated the SAK1 domain, has similarity to other proteins. Many predicted proteins within chlorophytes (Volvox carteri [8 proteins], Coccomyxa subellipsoidea [3 proteins], Chlamydomonas [14 proteins], Chlorella variabilis [9 proteins] and Micromonas [3 proteins]) (Table 8) contain this domain as shown in the alignment in Figure 5—figure supplement 1. Among the 37 members of the chlorophyte SAK1 domain family, 13 have possible bZIP transcription factor domains (six were significant Pfam

hits and seven were below the threshold for significance but recognizable by Pfam) (*Figure 5*). One protein contained a mitochondrial (transcription) termination factor (mTERF) domain (*Figure 5*), defined by its three leucine zipper domains required for DNA binding (*Fernandez-Silva et al., 1997*). Proteins with more distantly related SAK1 domains were found by PSI-BLAST in plants, many of which were hypothetical or unknown proteins but also included bZIP transcription factors.

Amino acid positions 900 to 1089 of SAK1, corresponding to the region aligned with other proteins in *Figure 5—figure supplement 1*, were searched for secondary structure using PHYRE, and this region was predicted to consist of mostly alpha helices with some disordered intervals. The top hit was a cobalt/nickel-binding resistance protein cnrr, and 44% of the residues were modeled with 73.6% confidence (*Figure 5—figure supplement 2*).

## SAK1 resides mainly in the cytosol and is phosphorylated during induction by <sup>1</sup>O<sub>2</sub>

To obtain insight into the function of SAK1, we isolated subcellular fractions enriched for chloroplast, ER, cytosol, and mitochondria from WT cells. The Chlamydomonas cell contains a single large chloroplast that is physically connected to other organelles such as the ER, making it particularly challenging to fractionate. The patterns of markers specific for chloroplast, ER, cytosol, and mitochondria showed that each target fraction was enriched as expected, although with some cross contamination (Figure 6A,B). The distribution of SAK1 in these fractions resembled most closely that of the cytosolic marker NAB1 (Mussgnug et al., 2005), although the SAK1 signal was not as enriched as NAB1 in the cytosolic fraction, possibly due to partial degradation of SAK1 during the fractionation. The localization was the same in cells with and without RB treatment (Figure 6A). Because SAK1 was required for the induction of many genes during acclimation to 102 and the list of proteins with similarity to SAK1 included those predicted to be bZIP transcription factors, we tested whether SAK1 protein was dually targeted to the nucleus and cytosol, which would account for the lack of enrichment of SAK1 in the cytosolic fraction (Figure 6A). As shown in Figure 6C although a faint SAK1 signal was detected in nuclear fraction, there was no enrichment as seen for the nuclear marker histone H3 (H3). The distribution of the cytosolic marker NAB1 indicated the contamination of the nuclear fraction by cytosolic proteins (Figure 6C). Therefore we conclude that the low signal of SAK1 in the nuclear fraction is likely to be due to cytosolic contamination. Attempts to detect the protein by immunofluorescence using anti-SAK1 antibodies as well as anti-FLAG and anti-HA antibodies against tagged proteins in transgenic lines were unsuccessful due to a very low signal-to-noise ratio even in bleached cells.

**Table 5.** Genes that require SAK1 for induction by  $^1\mathrm{O}_2$ 

Gene ID (v4)	Gene ID (v5)	Gene name	Annotation	FC WT* (log <sub>2</sub> )	FC sak1 (log <sub>2</sub> )	Attenuation (FC-sak1/FC-WT)†	Basal repression in sak1 (log <sub>2</sub> )
Cre02.g137700.t1.1‡	Cre09.g400404			6.49	1.80	0.04	-3.35
Cre06.g281250.t1.1‡	Cre06.g281250	CFA1	Cyclopropane fatty acid synthase	5.92	1.16	0.04	-2.10
Cre27.g775950.t1.2	Cre12.g557928			5.83	0.81	0.03	
Cre01.g033300.t1.1	Cre01.g033300			5.72	-0.39	0.01	
Cre13.g566850.t1.1#	Cre13.g566850	SOUL2	SOUL heme-binding protein	5.53	1.33	0.05	-2.60
Cre14.g623650.t1.1	Cre14.g623650		Alcohol dehydrogenase	4.89	1.67	0.11	
Cre13.g600650.t1.1	Cre06.g278245		Rieske 2Fe-2S domain	4.76	1.64	0.12	
Cre06.g263550.t1.1	Cre06.g263550	TCI7	R53.5-related protein	4.46	1.77	0.15	
Cre07.g342100.t1.1	Cre07.g342100			4.43	1.40	0.12	
Cre06.g299700.t1.1#	Cre06.g299700	SOUL1	SOUL heme-binding protein	4.32	0.43	0.07	-1.13
Cre09.g398700.t1.1#	Cre09.g398700	CFA2	Cyclopropane fatty acid synthase	4.05	0.18	0.07	-1.00
Cre12.g492650.t1.1‡	Cre12.9492650	FAS2	Fasciclin-like protein	4.01	0.07	0.07	-1.24
Cre08.g381510.t1.1	<b>L</b> Z			3.94	0.73	0.11	
Cre10.g458450.t1.2	Cre10.9458450	GPX5	Glutathione peroxidase	3.91	2.06	0.28	
Cre11.g474600.t1.1	Cre02.g095151		ABC transporter (ABC-2 type)	3.90	0.44	0.09	
Cre13.g600700.t1.1	Cre06.9278246			3.78	1.48	0.20	
Cre14.g613950.t1.1	Cre14.g613950			3.65	1.38	0.21	
Cre06.g269300.t1.1	Cre06.g269300		DUF1365	3.50	0.40	0.12	
Cre08.g380300.t1.2	Cre08.g380300	MSRA3	Peptide methionine sulfoxide reductase	3.45	99.0	0.14	
Cre28.g776450.t1.1	Cre08.g358573	TRP7	Transient receptor potential ion channel	3.31	-0.79	90.0	
Cre01.g031650.t1.2	Cre01.g031650	CGLD12	Potential galactosyl transferase activity	3.30	0.67	0.16	
Cre14.g629061.t1.1	ШZ		DUF2177	3.25	0.08	0.11	
Cre12.g503950.t1.1	Cre12.g503950		CRAL/TRIO domain	3.24	0.31	0.13	
	) ) ) )						

Table 5. Continued on next page

Gene ID (v4)	Gene ID (v5)	Gene name	Annotation	FC WT* (log <sub>2</sub> )	FC sak1 (log <sub>2</sub> )	Attenuation (FC-sak1/FC-WT)†	Basal repression in sak1 (log <sub>2</sub> )
Cre13.g564900.t1.1	Cre13.g564900		ABC transporter transmembrane region	3.22	0.34	0.14	
Cre02.g139500.t1.1	Cre09.9401701		DUF1295	3.04	-0.16	0.11	
Cre14.g618400.t1.1	Cre14.g618400			2.97	1.15	0.28	
Cre17.g715150.t1.1	Cre17.g715150			2.89	0.13	0.15	
Cre17.g741300.t1.2#	Cre17.g741300	SAK1		2.88	99.0	0.21	-2.77
Cre01.g007300.t1.1	Cre01.g007300			2.85	-1.15	0.06	
Cre16.g648700.t1.2#	Cre16.g648700		ABC transporter (ABC-2 type)	2.79	0.26	0.17	-1.26
Cre13.g566900.t1.2	Cre13.g566900			2.76	-0.38	0.11	
Cre02.g137750.t1.2	Cre09.9400441		JmjC domain	2.72	-0.31	0.12	
Cre06.g263500.t1.1	Cre06.g263500		Archease protein family (DUF101)	2.67	1.02	0.32	
Cre01.g016150.t1.1#	Cre01.g016150		ADP-ribosylglycohydrolase	2.65	0.17	0.18	-1.26
Cre08.g380000.t1.1	Cre08.g380000		Formylglycine-generating sulfatase enzyme	2.59	1.53	0.48	
Cre14.g615600.t1.1	Cre14.g615600		Putative serine esterase (DUF676)	2.53	-0.54	0.12	
Cre11.9472900.t1.2	Cre02.g095113		CAP-Gly domain	2.45	-0.05	0.18	
Cre06.g269250.t1.1	Cre06.g269250			2.44	0.55	0.27	
Cre02.g120600.t1.1	Cre09.9403071			2.44	0.94	0.35	
Cre06.9261200.t1.1	Cre06.g261200	ERG25	Sterol desaturase	2.42	0.64	0.29	
Cre16.g683400.t1.1	Cre16.g683400		CRAL/TRIO domain	2.40	0.08	0.20	
Cre22.g765150.t1.1	Cre11.g467725		hypothetical protein	2.30	0.46	0.28	
Cre13.g571800.t1.2	Cre13.9571800		DUF1336	2.27	0.72	0.34	
Cre13.g579450.t1.2	Cre13.g579450	CST1	Membrane transporter	2.27	1.23	0.49	
Cre08.9380350.t1.1	Cre08.9380350			2.21	-0.01	0.21	
Cre16.g649250.t1.2	Cre16.g649250			2.08	0.58	0.35	

Table 5. Continued on next page

Table 5. Continued

Gene ID (v4)	Gene ID (v5)	Gene name	Annotation	FC WT* (log <sub>2</sub> )	FC sak1 (log <sub>2</sub> )	Attenuation (FC-sak1/FC-WT)†	Basal repression in sak1 (log <sub>2</sub> )
Cre11.g476250.t1.1	Cre11.g476250			2.08	0.49	0.33	
Cre02.g108000.t1.2	Cre02.g108000			2.08	1.03	0.49	
Cre13.g583300.t1.1	Cre13.g583300			1.98	-0.48	0.18	
Cre04.g215300.t1.2	NF			1.97	0.57	0.38	
Cre02.g139450.t1.1	Cre09.g401663		DUF947	1.95	-0.62	0.17	
Cre03.g194750.t1.2	Cre03.g194750			1.95	0.73	0.43	
Cre06.g258600.t1.1	Cre06.g258600		Dienelactone hydrolase family	1.91	-0.95	0.14	
Cre10.g418700.t1.1	Cre10.g418700		Probable N6-adenine methyltransferase	1.87	-0.03	0.27	
Cre10.g444550.t1.1	Cre10.9444550	SPP1A	Signal peptide peptidase	1.81	0.51	0.41	
Cre01.g060050.t1.2	Cre03.g145807			1.78	-0.11	0.27	
Cre09.g410050.t1.1	Cre09.g410050		Calcium transporting ATPase	1.76	0.51	0.42	
Cre03.g163400.t1.2	Cre03.g163400			1.76	-0.17	0.26	
Cre01.g008450.t1.1	Cre01.g008450		Nuf2 family	1.73	-0.54	0.21	
Cre12.g536650.t1.1	Cre12.g536650			1.72	0.35	0.39	
Cre02.g114900.t1.2	Cre02.g114900	ANK23	predicted protein	1.71	0.08	0.32	
Cre16.g661850.t1.2	Cre16.g661850		Calcium/calmoduline dependent protein kinase association	1.69	0.03	0.32	
Cre14.g615500.t1.2	Cre14.g615500		Glycoprotease family	1.68	-0.76	0.18	
Cre11.g483100.t1.2	Cre11.9483100		Protein kinase	1.66	-0.49	0.22	
Cre28.g776650.t1.1	Cre08.g358569			1.64	0.33	0.40	
Cre07.g340250.t1.2	Cre07.9340250		Protein kinase	1.63	-0.41	0.24	
Cre06.9296250.t1.2	Cre06.g296250	SYK1	tRNA synthetase, class II	1.60	0.54	0.48	
Cre06.g310500.t1.1	Cre06.9310500			1.57	0.18	0.38	
Cre07.g342800.t1.2	Cre07.9342800	CGL16	Predicted protein	1.49	0.32	0.44	

Table 5. Continued on next page

Table 5. Continued

Basal repression in sak1 (log<sub>2</sub>) 9 (FC-sak1/FC-WT)† **Attenuation** 0.46 0.37 0.39 0.43 0.35 0.35 0.36 0.45 0.47 0.46 0.47 0.38 0.37 0.32 0.40 0.28 0.46 0.26 0.27 0.30 FC sak1 (log<sub>2</sub>) 0.35 0.03 -0.12-0.12-0.130.16 -0.49 -0.70 90.0 0.19 0.22 0.16 0.17 -0.14 -0.24 0.00 -0.84 -0.03 -0.50 -0.58 -0.48-0.17 0.21 FC WT\* (log<sub>2</sub>) 1.38 1.38 1.19 1.17 1.17 1.14 1.14 1.13 1.1 1.47 1.35 1.34 1.33 1.32 1.32 1.28 1.26 1.47 1.42 1.29 1.27 1.43 DNA polymerase delta, subunit 4 Calcium/calmoduline dependent Microtubule associated protein ABC transporter (ABC-2 type) ABC transporter (ABC-2 type) Diaminopimelate epimerase Ubiquitin carboxyl-terminal hydrolase Inosine-uridine preferring protein kinase association Truncated hemoglobin nucleoside hydrolase No annotation Protein kinase Annotation **DUF1619** Gene name MAP65 THB3 DAE1 Gene ID (v5) Cre03.g181450 Cre14.g614050 Cre16.g661750 Cre01.g007000 Cre04.g224400 Cre16.g680790 Cre05.g237400 Cre14.g609600 Cre03.g179200 Cre10.9417730 Cre03.g159700 Cre12.g540150 Cre01.9006550 Cre03.g159950 Cre12.g557503 Cre09.g387208 Cre04.g218800 Cre35.q759497 Cre06.g292950 Cre05.9234850 Cre07.9315050 Cre04.g217500 ¥ Cre01.g006550.t1.2# Cre03.g181450.t1.2 Cre14.g614050.t1.2 Cre04.g224400.t1.2 Cre01.g068400.t1.2 Cre14.g609600.t1.2 Cre05.g234850.t1.2 Cre03.g159700.t1.2 Cre12.g540150.t1.2 Cre03.g159950.t1.2 Cre27.g775900.t1.2 Cre07.g315050.t1.2 Cre04.g218800.t1.2 Cre04.g217500.t1.1 Cre16.g661750.t1.1 Cre10.g417730.t1.1 Cre02.g121600.t1.1 Cre14.g609550.t1.1 Cre66.q793601.t1.1 Cre06.g292950.t1.1 Cre01.g007000.t1.1 Cre05.g237400.t1.1 Cre03.g179200.t1.1 Gene ID (v4)

Table 5. Continued on next page

Table 5. Continued

Table 5. Continued

Gene ID (v4)	Gene ID (v5)	Gene name	Annotation	FC WT* (log <sub>2</sub> )	FC sak1 (log <sub>2</sub> )	Attenuation (FC-sak1/FC-WT)†	Basal repression in sak1 (log <sub>2</sub> )
Cre02.g133300.t1.1	Cre09.g396624			1.11	-0.43	0.34	
Cre01.g060650.t1.2	Cre03.g146067			1.10	-0.42	0.35	
Cre01.g057050.t1.1	Cre03.g144324			1.10	0.04	0.48	
Cre06.g304950.t1.1	Cre06.9304950			1.07	-0.65	0.30	
Cre08.g358200.t1.2	Cre08.9358200	A4	Protein kinase	1.07	-0.82	0.27	
Cre16.g689550.t1.2	Cre16.g689550	PTK8	Putative tyrosine kinase	1.06	-0.17	0.43	
Cre17.g720950.t1.1	Cre17.g720950		3-oxo-5-alpha-steroid 4-dehydrogenase	1.05	-0.26	0.40	
Cre02.g090950.t1.2	Cre02.g090950			1.05	-0.27	0.40	
Cre16.g683350.t1.1	Cre16.g683350			1.03	-0.67	0.31	
Cre02.g109450.t1.1	Cre02.g109450			1.01	-0.03	0.48	
Cre16.g652750.t1.1	Cre16.g652750			1.01	-0.29	0.41	
Cre03.g190000.t1.1	Cre03.9190000			1.00	-0.99	0.25	

\*Data were ordered by FC in WT.

10f the 52 most highly induced genes in WT (the top 10%), 37 were SAK1-dependent, and the induction of 33 of these genes was strongly attenuated to only 0.01-0.25 of magnitude of FC found in the WT. Dashed line indicates cutoff of FC for the top 10% most strongly induced genes.

‡Genes that are repressed at basal level in sak1.

NF, not found in v5.

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**Table 6.** Enriched functional classes among differentially expressed genes in sak1 during  $^{1}O_{2}$  acclimation

Primary Mapman class	Secondary Mapman class	Gene ID (v4)	Gene name	Annotation
Up-regulated genes				
Secondary metabolism	isoprenoids	Cre13.g565650.t1.1		Geranylgeranyl pyrophosphate synthase/Polyprenyl synthetase
		Cre06.g267600.t1.1		Lycopene epsilon cyclase
		Cre09.g407200.t1.1		Phytoene desaturase
		Cre06.g267600.t1.1		Lycopene epsilon cyclase
		Cre01.g011100.t1.1		Prenyltransferase and squalene oxidase repeat, Oxidosqualene- lanosterol cyclase and related proteins
	N misc	Cre08.g381707.t1.1		NF*
	phenylpropanoids	Cre03.g207800.t1.1		Alcohol dehydrogenase, class V
		Cre14.g623650.t1.1		Alcohol dehydrogenase, class V (Zinc-binding)
		Cre01.g039350.t1.1		Cytochrome P450 reductase, possibly CYP505B family
	sulfur-containing	Cre06.g299400.t1.1		NF*
	wax	Cre17.g722150.t1.1	PKS3	Type III polyketide synthase
		Cre07.g318500.t1.2		FAE1/Type III polyketide synthase- like protein, Chalcone and stilbene synthases
Lipid metabolism	'exotics' (steroids, squalene etc)	Cre01.g061750.t1.1		serine palmitoyltransferase
		Cre02.g137850.t1.1		NF*
		Cre83.g796250.t1.1		NF*
		Cre01.g011100.t1.1		Prenyltransferase and squalene oxidase repeat, Oxidosqualene- lanosterol cyclase and related proteins
	FA synthesis and FA elongation	Cre06.g256750.t1.1		Acyl carrier protein thioesterase
		Cre03.g182050.t1.1		Long-chain acyl-CoA synthetases (AMP-forming)
		Cre02.g074650.t1.1		Kelch repeat-containing proteins, Acyl-CoA binding protei
	glycerol metabolism	Cre01.g053000.t1.1	GPD2	Glycerol-3-phosphate dehydrogenase/dihydroxyacetone- 3-phosphate reductase
	glycolipid synthesis	Cre13.g583600.t1.1	DGD1	Digalactosyldiacylglycerol synthase
	lipid degradation	Cre01.g057450.t1.2		NF*
		Cre02.g126050.t1.1		NF*
	phospholipid synthesis	Cre06.g281250.t1.1	CFA1	Cyclopropane fatty acid synthase
		Cre01.g038250.t1.1	SDC1	Serine decarboxylase
		Cre11.g472700.t1.1		NF*
		Cre13.g604700.t1.2		CDP-alcohol phosphatidyltransferase/ Phosphatidylglycerol-phosphate synthase

Table 6. Continued on next page



Table 6. Continued

Primary Mapman	Secondary		Gene	
class	Mapman class	Gene ID (v4)	name	Annotation
Cell	vesicle transport	Cre18.g744100.t1.1		NF*
		Cre17.g721900.t1.1	COG5	Component of oligomeric golgi complex
		Cre01.g003050.t1.1	SEC8	Component of the Exocyst Complex
		Cre04.g224800.t1.1		Endosomal R-SNARE protein, Vamp7/Nyv1-family
		Cre17.g728150.t1.1		Endosomal R-SNARE protein, Yky6-family
		Cre12.g507450.t1.1		Trans-Golgi network Qa-SNARE protein, Syntaxin16/Syx16/Tlg2/ Syp4-family
		Cre03.g210600.t1.1		NF*
		Cre04.g225900.t1.1		Endosomal R-SNARE protein, Vamp7/Nyv1-family
		Cre02.g101400.t1.1	CHC1	Clathrin Heavy Chain
		Cre17.g709350.t1.1		Late endosomal Qc-SNARE protein, Syx8/Syntaxin8-family
		Cre07.g342050.t1.1		Endosomal Qb-SNARE, Npsn- family
		Cre16.g692050.t1.1		ER-Golgi Qa-SNARE protein, Syntaxin5/Syx5/Sed5/Syp3-family
		Cre16.g676650.t1.1	AP1G1	Gamma1-Adaptin
		Cre02.g099000.t1.1		Late endosomal Qc-SNARE protein, Syx6/Tlg1/Syp5/6-family
		Cre12.g554200.t1.2		ER-Golgi Qb-SNARE, Memb/GS35/ Bos1-family
		Cre06.g310000.t1.1	AP4E1	Epsilon4-Adaptin
		Cre10.g421250.t1.1	EXO70	Hypothetical Conserved Protein. Similar to Exo70, a subunit of the exocyst complex
		Cre07.g330950.t1.1	AP4S4	Sigma4-Adaptin
		Cre12.g488850.t1.2		Adaptin, alpha/gamma/epsilon
	division	Cre06.g269950.t1.1	CDC48	Protein involved in ubiquitin- dependent degradation of ER-bound substrates
		Cre08.g359200.t1.2		Regulator of chromosome condensation (RCC1)
	organisation	Cre13.g588600.t1.2		Kinesin (SMY1 subfamily)
		Cre12.g513450.t1.1	TUH1	Eta-Tubulin
		Cre01.g010950.t1.2		26S proteasome regulatory complex, subunit PSMD10 (Ankyrin repeat)
		Cre16.g679650.t1.2		Fimbrin/Plastin
		Cre06.g261950.t1.1		Myotrophin and similar proteins (Ankyrin repeat)
		Cre06.g291700.t1.1	RSP3	Radial spoke protein 3
		Cre10.g446700.t1.1	ANK28	Ankyrin repeat and DHHC-type Zn-finger domain containing proteins

Table 6. Continued on next page



Table 6. Continued

Primary Mapman	Secondary	C 15 / 4)	Gene	
class	Mapman class	Gene ID (v4)	name	Annotation
Hormone metabolism†	abscisic acid	Cre16.g657800.t1.2	CCD3	Carotenoid cleavage dioxygenase
	auxin	Cre14.g609900.t1.1		Predicted membrane protein, contains DoH and Cytochrome b-561/ferric reductase transmembrane domains
	brassinosteroid	Cre16.g663950.t1.1		Sterol C5 desaturase
		Cre02.g092350.t1.1		Cytochrome P450, CYP51 superfamily; sterol 14 desaturase
		Cre12.g557900.t1.1	CDI1	C-8,7 sterol isomerase
		Cre02.g076800.t1.1		Delta14-sterol reductase, mitochondrial
		Cre12.g500500.t1.2		24-methylenesterol C-methyltransferase
	ethylene	Cre02.g108450.t1.1	FAP280	Flagellar Associated Protein, transcriptional coactivator-like, putative transcription factor
	jasmonate	Cre19.g756100.t1.1		NF*
Misc	acid and other phosphatases	Cre09.g396900.t1.1		NADH pyrophosphatase I of the Nudix family of hydrolases
		Cre06.g259650.t1.1		Calcineurin-like phosphoesterase, Acid-phosphatase-related
		Cre06.g249800.t1.1		Sphingomyelin synthetase -related
	cytochrome P450	Cre05.g234100.t1.1		Cytochrome P450, CYP197 superfamily
	dynamin	Cre02.g079550.t1.1	DRP2	Dynamin-related GTPase, involved in circadian rhythms
		Cre05.g245950.t1.1	DRP1	Dynamin-related GTPase
	glutathione S transferases	Cre03.g154950.t1.1		Glutathione S-transferase
	misc2	Cre12.g538450.t1.1	EPT1	CDP-Etn:DAG Ethanolamine phosphotransferase
	short chain dehydrogenase/ reductase (SDR)	Cre12.g556750.t1.2		Short-chain dehydrogenase/ reductase
		Cre08.g384864.t1.1		SH3 domain, protein binding
		Cre27.g775000.t1.1		NF*
		Cre17.g731350.t1.2		Short chain dehydrogenase
	UDP glucosyl and glucoronyl transferases	Cre02.g111150.t1.2	ELG26	Exostosin-like glycosyltransferase
		Cre02.g144050.t1.1		Acetylglucosaminyltransferase EXT1/exostosin 1
		Cre03.g204050.t1.2	ELG6	Exostosin-like glycosyltransferases
		Cre11.g474450.t1.1		NF*
		Cre03.g173300.t1.1		Lactosylceramide 4-alpha- galactosyltransferase (alpha- 1,4-galactosyltransferase)
		Cre02.g116600.t1.1	ELG23	Exostosin-like glycosyltransferase
Down-regulated gene	es			
Hormone metabolism†	cytokinin	Cre18.g744950.t1.2		NF*

Table 6. Continued on next page



Table 6. Continued

Primary Mapman class	Secondary Mapman class	Gene ID (v4)	Gene name	Annotation
		Cre16.g678900.t1.1		Response regulator receiver domain
		Cre01.g040450.t1.1	HDT1	Histidine-aspartic acid phosphotransferase 1 (phosphorylation cascade)
	ethylene	Cre09.g403550.t1.1		Iron/ascorbate family oxidoreductases
Nucleotide metabolism	deoxynucleotide metabolism	Cre12.g491050.t1.1	RIR2	Ribonucleotide reductase (RNR), small subunit
		Cre12.g492950.t1.1	RIR1	Ribonucleotide reductase (RNR), large subunit, class I
		Cre16.g667850.t1.1		dUTP pyrophosphatase
	synthesis	Cre14.g614300.t1.1		Inosine-5-monophosphate dehydrogenase/GMP reductase
		Cre07.g318750.t1.1		Phosphoribosylformylglycinamidine cyclo-ligase
Tetrapyrrole synthesis	porphobilinogen deaminase	Cre16.g663900.t1.1		Porphobilinogen deaminase
	protochlorophyllide reductase	Cre01.g015350.t1.1		Light-dependent protochlorophyllide reductase
	urogen III methylase	Cre02.g133050.t1.2		NF*
DNA	repair	Cre16.g670550.t1.2		XP-G/RAD2 DNA repair endonuclease
	synthesis/chromatin structure	Cre07.g338000.t1.1	МСМ2	Minichromosome maintenance protein
		Cre07.g314900.t1.2		ATP-dependent RNA helicase, DEAD/DEAH helicase
		Cre03.g172950.t1.1	CBF5	Centromere/microtubule binding protein
		Cre01.g015250.t1.1		Eukaryotic DNA polymerase delta
		Cre27.g774200.t1.2		NF*
		Cre07.g316850.t1.1	МСМ4	Minichromosome maintenance protein
	unspecified	Cre10.g451250.t1.2		Adenylate and guanylate cyclase catalytic domain, 3-5 exonuclease
		Cre01.g059950.t1.2		NF*

<sup>\*</sup>Corresponding gene model was not found in v5.

†Functional terms are inferred by homology to the annotation set of *Arabidopsis thaliana* (**Lopez et al., 2011**). DOI: 10.7554/eLife.02286.012

By SDS-PAGE and immunoblot analysis, SAK1 appeared in multiple forms with higher molecular weight during acclimation compared to that observed in control cells (*Figures 4F and 6A,C*). When the extracted protein samples were treated with phosphatase, the diffuse pattern of multiple forms collapsed into a single band detected by immunoblot analysis that had an even higher mobility that that of untreated cells (*Figure 6D*). This result indicates that SAK1 is a phosphorylated protein during basal conditions, and it is further phosphorylated upon exposure of cells to  ${}^{1}O_{2}$ .

#### **Discussion**

#### SAK1 is necessary for acclimation of Chlamydomonas cells to <sup>1</sup>O<sub>2</sub>

To understand the retrograde signal transduction pathway involved in the cellular response to  ${}^{1}O_{2}$ , we focused on the unique ability of *Chlamydomonas* to acclimate to  ${}^{1}O_{2}$  stress (*Ledford et al., 2007*), and



 Table 7. Enriched functional classes among differentially expressed genes in sak1 at basal level

Primary Mapman class	Secondary Mapman class	Gene ID (v4)	Gene name	Annotation
Elevated in sak1				
nucleotide metabolism	deoxynucleotide metabolism	Cre12.g491050.t1.1	RIR2	Ribonucleotide reductase (RNR), small subunit
		Cre12.g492950.t1.1	RIR1	Ribonucleotide reductase (RNR), large subunit, class I
		Cre16.g667850.t1.1		dUTP pyrophosphatase
	phosphotransfer and pyrophosphatases	Cre02.g122450.t1.1		NF*
		Cre02.g093950.t1.1	PYR5	Uridine 5'- monophosphate synthase/orotate phosphoribosyltransferase
		Cre12.g519950.t1.1		Flagellar Associated Protein similar to adenylate/guanylate kinases
		Cre26.g772450.t1.1		NF*
	synthesis	Cre65.g793400.t1.1		NF*
		Cre02.g079700.t1.1	PYR2	Aspartate carbamoyltransferase
		Cre01.g048950.t1.1		dUTP pyrophosphatase
		Cre07.g318750.t1.1		Phosphoribosylformylglycinamidine cyclo-ligase.
DNA	repair	Cre07.g314650.t1.1		Chloroplast RecA recombination protein
	synthesis/chromatin structure	Cre04.g214350.t1.2		Eukaryotic DNA polymerase alpha, catalytic subunit
		Cre07.g314900.t1.2		ATP-dependent RNA helicase (DEAD/DEAH)
		Cre04.g223850.t1.1		Cytoplasmic DExD/H-box RNA helicase
		Cre01.g015250.t1.1		Eukaryotic DNA polymerase delta, catalytic subunit.
		Cre07.g342506.t1.1		Ubiquitin-protein ligase
		Cre07.g338000.t1.1	МСМ2	Minichromosome maintenance protein
		Cre03.g178650.t1.1	МСМ6	MCM6 DNA replication protein
		Cre07.g312350.t1.2		DNA polymerase alpha, primase subunit
		Cre01.g009250.t1.2	TOP2	DNA topoisomerase II
		Cre26.g772150.t1.1		NF*
		Cre07.g316850.t1.1	МСМ4	Minichromosome maintenance protein 4
		Cre06.g263800.t1.2		tRNA-splicing endonuclease positive effector (SEN1)
		Cre06.g295700.t1.2	МСМ3	Minichromosome maintenance protein
		Cre06.g251800.t1.1	RFC4	DNA replication factor C complex subunit 4
	unspecified	Cre07.g322300.t1.2		DNA repair helicase of the DEAD superfamily
		Cre17.g718100.t1.1		Phosphatidylinositol transfer protein SEC14 and related proteins (CRAL/TRIO)

Table 7. Continued on next page



Table 7. Continued

Primary	Secondary		Gene	
Mapman class	Mapman class	Gene ID (v4)	name	Annotation
Tetrapyrrole synthesis	Glu-tRNA reductase	Cre07.g342150.t1.1	HEM1	Glutamyl-tRNA reductase
	Glu-tRNA synthetase	Cre44.g788000.t1.1		Glutamyl-tRNA reductase
		Cre06.g306300.t1.1	CHLI1	Magnesium chelatase subunit I
	magnesium chelatase	Cre07.g325500.t1.1		Magnesium chelatase subunit H
	protochlorophyllide reductase	Cre01.g015350.t1.1	POR1	Light-dependent protochlorophyllide reductase
Photosynthesis	Calvin-Benson cycle	Cre05.g234550.t1.1		Fructose-biphosphate aldolase
	light reaction	Cre07.g330250.t1.1	PSAH	Subunit H of photosystem I
		Cre07.g334550.t1.1		Photosystem I subunit PsaO
		Cre06.g261000.t1.1	PSBR	10 kDa photosystem II polypeptide
	photorespiration	Cre12.g542300.t1.1	GYK1	Glycerate kinase
		Cre06.g253350.t1.1	GCSH	Glycine cleavage system, H-protein
		Cre06.g293950.t1.1	SHMT2	Serine hydroxymethyltransferase 2
Transport	ABC transporters and multidrug resistance systems	Cre04.g222700.t1.1		ATPase component of ABC transporters with duplicated ATPase domains/Translation elongation factor EF-3b
		Cre17.g728400.t1.2		ABCtransporter (ABC-2 type)
		Cre05.g241350.t1.2		ABCtransporter (ABC-2 type)
		Cre03.g169300.t1.1		ABCtransporter (ABC-2 type)
		Cre11.g474600.t1.1		NF*
	amino acids	Cre04.g226150.t1.2	AOC1	Amino acid carrier 1; belongs to APC (Amino acid Polyamine organo Cation) family
	calcium	Cre09.g388850.t1.1	ACA1	P-type ATPase/cation transporter, plasma membrane
	metabolite transporters at the envelope membrane	Cre06.g263850.t1.2	TPT2	Triose phosphate/phosphate translocator
	metabolite transporters at the mitochondrial membrane	Cre10.g449100.t1.1		Mitochondrial oxodicarboxylate carrier protein
		Cre01.g069350.t1.1		NF*
		Cre15.g641200.t1.1		Mitochondrial fatty acid anion carrier protein/Uncoupling protein
		Cre09.g396350.t1.1		Mitochondrial carrier protein PET8
	misc	Cre06.g311000.t1.2	FBT2	Folate transporte
		Cre17.g718100.t1.1		Phosphatidylinositol transfer protein SEC14 and related proteins (CRAL/TRIO)
	phosphate	Cre16.g686750.t1.1	PTA3	Proton/phosphate symporter
		Cre16.g675300.t1.2		Sodium-dependent phosphate transporter, major facilitator superfamily
	potassium	Cre12.g553450.t1.2		NF*
	sulphate	Cre17.g723350.t1.1	SUL2	Sulfate anion transporter
	unspecified cations	Cre13.g573900.t1.1		Na+:iodide/myo-inositol/ multivitamin symporters

Table 7. Continued on next page



Table 7. Continued

Primary Mapman class	Secondary Mapman class	Gene ID (v4)	Gene name	Annotation
	sugars	Cre16.g675300.t1.2		Sodium-dependent phosphate transporter, major facilitator superfamily
RNA	processing	Cre10.g427700.t1.1		ATP-dependent RNA helicase, DEAD/DEAH box helicase
		Cre12.g538750.t1.1	LSM1	U6 snRNA-associated Sm-like protein LSm1, RNA cap binding; (SMP6d)
		Cre10.g433750.t1.2	PAP1	Nuclear poly(A) polymerase
		Cre03.g182950.t1.1		NF*
		Cre08.g375128.t1.1		NF*
	regulation of transcription	Cre17.g728200.t1.2		YL-1 protein (transcription factor-like 1)
		Cre06.g275500.t1.1		AP2 Transcription factor
		Cre28.g777500.t1.2		NF*
		Cre13.g572450.t1.1		Response regulator receiver domain (sensor histidine kinase-related, regulation of transcription)
		Cre14.g620500.t1.1		AP2 Transcription factor
		Cre16.g673150.t1.1		Histone deacetylase complex, catalytic component RPD3
		Cre02.g078700.t1.2		DNA damage-responsive repressor GIS1/RPH1, jumonji superfamily
		Cre03.g198800.t1.1		Myb-like DNA-binding domain
		Cre04.g218050.t1.2		RWP-RK domain
		Cre07.g324400.t1.1	VPS24	Subunit of the ESCRT-III complex, vaculoar sortin protein
		Cre11.g481050.t1.1		SWI/SNF-related chromatin binding protein
		Cre02.g101950.t1.1	TMU2	tRNA (uracil-5)-methyltransferase
		Cre10.g459600.t1.2		CAATT-binding transcription factor/60S ribosomal subunit biogenesis protein
		Cre01.g018650.t1.2		NF*
		Cre01.g012200.t1.2		NF*
		Cre02.g129750.t1.1		NF*
		Cre10.g461750.t1.2		DNA (cytosine-5-)- methyltransferase
		Cre01.g004600.t1.2	RWP12	Putative RWP-RK domain transcription factor
		Cre09.g400100.t1.1		Predicted Zn-finger protein, zinc and DNA binding domains
		Cre07.g335150.t1.2		SBP domain
	RNA binding	Cre16.g662700.t1.1		NF*
		Cre07.g330300.t1.1		RNA-binding protein musashi/ mRNA cleavage and polyadenylation factor I complex, subunit HRP1

Table 7. Continued on next page



Table 7. Continued

Primary	Secondary		Gene	
Mapman class	Mapman class	Gene ID (v4)	name	Annotation
		Cre06.g275100.t1.1		RNA-binding protein musashi/ mRNA cleavage and polyadenylation factor I complex, subunit HRP1
	transcription	Cre07.g322200.t1.1		NF*
Repressed in sak1				
Transport	ABC transporters and multidrug resistance systems	Cre02.g097800.t1.2		ABC transporter (MDR)
		Cre17.g725200.t1.1		ABC transporter, peptide exporter
		Cre13.g580300.t1.1		ABC transporter family protein
		Cre10.g439000.t1.2		Long-chain acyl-CoA transporter, ABC superfamily (involved in peroxisome organization and biogenesis)
	amino acids	Cre06.g292350.t1.1	AOC4	Amino acid carrier
	calcium	Cre06.g263950.t1.2		Sodium/potassium-transporting ATPase subunit alpha
		Cre16.g681750.t1.2		Calcium transporting ATPase
	metabolite transporters at the mitochondrial membrane	Cre03.g172300.t1.1		Mitochondrial phosphate carrier protein
		Cre09.g394800.t1.2		Mitochondrial substrate carrier protein
	metal	Cre03.g189550.t1.2	ZIP3	Zinc transporter, ZIP family
		Cre11.g479600.t1.2		Sodium/calcium exchanger NCX1 and related proteins
		Cre06.g281900.t1.1	ZIP7	Zinc transporter and related ZIP domain-containing proteins
	misc	Cre02.g089900.t1.1		Secretory carrier membrane protein
		Cre10.g448050.t1.1		Retinaldehyde binding protein- related (CRAL/TRIO domain)
		Cre03.g177750.t1.2		Multidrug resistance pump
	NDP-sugars at the ER	Cre02.g112900.t1.1		GDP-fucose transporter (Triose- phosphate transporter family)
	P- and V-ATPases	Cre01.g027800.t1.1	ATPvH	Vacuolar ATP synthase subunit H
		Cre10.g446550.t1.1	ATPvF	Vacuolar ATP synthase subunit F
		Cre03.g176250.t1.1	ATPvD1	Vacuolar ATP synthase subunit D
		Cre06.g250250.t1.1	ATPvC	Vacuolar ATP synthase subunit C
		Cre10.g459200.t1.1	ACA4	P-type ATPase/cation transporter, plasma membrane (Low CO2 inducible gene)
	phosphate	Cre12.g515750.t1.2		Sodium-dependent phosphate transporter-related
		Cre08.g379550.t1.2		Sodium-dependent phosphate transporter, major facilitator superfamily
		Cre12.g489400.t1.1	PTB7	Putative phosphate transporter, sodium/phosphate transporter

Table 7. Continued on next page



Table 7. Continued

Primary	Secondary		Gene	
Mapman class	Mapman class	Gene ID (v4)	name	Annotation
		Cre02.g144650.t1.1	PTB12	Sodium/phosphate symporter
	unspecified anions	Cre09.g404100.t1.1		CI- channel CLC-7 and related proteins (CLC superfamily)
		Cre17.g729450.t1.1		CI- channel CLC-7 and related proteins (CLC superfamily)
		Cre01.g037150.t1.2		Voltage-gated chloride channel activity
	sugars	Cre03.g206800.t1.2	HXT1	Hexose transporter
	P- and V-ATPases	Cre03.g176250.t1.1	ATPvD1	Vacuolar ATP synthase subunit D
		Cre10.g446550.t1.1	ATPvF	Vacuolar ATP synthase subunit F
		Cre01.g027800.t1.1	ATPvH	Vacuolar ATP synthase subunit H
Mitochondrial electron transport / ATP synthesis	cytochrome c reductase	Cre01.g051900.t1.1	RIP1	Rieske iron-sulfur protein of mitochondrial ubiquinol- cytochrome c reductase (complex III)
		Cre06.g262700.t1.2		Ubiquinol cytochrome c reductase, subunit 7
	F <sub>1</sub> -ATPase	Cre02.g116750.t1.2		F0F1-type ATP synthase, alpha subunit
		Cre01.g018800.t1.1	ATP6	Mitochondrial F1F0 ATP synthase subunit 6
		Cre10.g420700.t1.1		Mitochondrial F1F0-ATP synthase, subunit epsilon/ATP15
		Cre16.g680000.t1.1	ATP5	Mitochondrial ATP synthase subunit 5, OSCP subunit
	NADH-DH	Cre10.g434450.t1.1	NUOA9	Putative NADH:ubiquinone oxidoreductase (Complex I) 39 kDa subunit
		Cre08.g378900.t1.1	NUO3	NADH:ubiquinone oxidoreductase ND3 subunit
		Cre10.g450400.t1.1	NUO5	NADH:ubiquinone oxidoreductase (Complex I) 24 kD subunit
Lipid metabolism	'exotics' (steroids, squalene etc)	Cre14.g615050.t1.1		3-oxo-5-alpha-steroid 4-dehydrogenase, Steroid reductase required for elongation of the VLCFAs (enoyl reductase)
		Cre12.g530550.t1.2	KDG2	Diacylglycerol kinase, sphingosine kinase
		Cre02.g137850.t1.1		NF*
	FA desaturation	Cre17.g711150.t1.1		Omega-6 fatty acid desaturase (delta-12 desaturase)
	glyceral metabolism	Cre13.g577450.t1.2		Glycerol-3-phosphate dehydrogenase
	glycolipid synthesis	Cre13.g583600.t1.1	DGD1	Digalactosyldiacylglycerol synthase
		Cre16.g656400.t1.1	SQD1	UDP-sulfoquinovose synthase
	lipid degradation	Cre06.g252801.t1.2		CGI-141-related/lipase containing protein (TAG lipase)
		Cre03.g164350.t1.2		Lysophospholipase, putative drug exporter of the RND superfamily

Table 7. Continued on next page



Table 7. Continued

Primary Mapman class	Secondary Mapman class	Gene ID (v4)	Gene name	Annotation
	phospholipid synthesis	Cre06.g281250.t1.1	CFA1	Cyclopropane fatty acid synthase
		Cre09.g398700.t1.1	CFA2	Cyclopropane fatty acid synthase
		Cre11.g472700.t1.1		NF*
		Cre06.g262550.t1.1		Zinc finger MYND domain containing protein 10
Photosynthesis	Calvin-Benson cycle	Cre12.g511900.t1.1	RPE1	Ribulose phosphate-3-epimerase
		Cre02.g120100.t1.1	RBCS1	Ribulose-1,5-bisphosphate carboxylase/oxygenase small subunit 1
	light reaction	Cre05.g243800.t1.1	CPLD45	Photosystem II Psb27 protein
		Cre10.g420350.t1.1	PSAE	Photosystem I reaction center subunit IV
		Cre01.g071450.t1.2		NF*
		Cre06.g291650.t1.1		Ferredoxin
		Cre05.g242400.t1.1		No functional annotation
	photorespiration	Cre09.g411900.t1.2	SHMT3	Serine hydroxymethyltransferase 3
		Cre06.g295450.t1.1	HPR1	Hydroxypyruvate reductase
Major CHO metabolism	degradation	Cre09.g415600.t1.2		Starch binding domain
		Cre11.g473500.t1.2		NF*
		Cre09.g415600.t1.2		Starch binding domain
	synthesis	Cre06.g289850.t1.2	SBE1	Starch Branching Enzyme
		Cre17.g721500.t1.1		Granule-bound starch synthase I
misc	acid and other phosphatases	Cre13.g568600.t1.2		Multiple inositol polyphosphate phosphatase-related, Acid phosphatase activity
	alcohol dehydrogenases	Cre13.g569350.t1.1		Sterol dehydrogenase-related, Flavonol reductase/cinnamoyl-CoA reductase
	cytochrome P450	Cre07.g356250.t1.2		Cytochrome P450 CYP4/CYP19/ CYP26 subfamilies, beta-carotene 15,15'-monooxygenase
		Cre07.g356250.t1.2		Cytochrome P450 CYP4/CYP19/ CYP26 subfamilies, beta-carotene 15,15'-monooxygenase
	dynamin	Cre17.g724150.t1.1	DRP3	Dynamin-related GTPase
	GCN5-related N-acetyltransferase	Cre16.g657150.t1.2		N-acetyltransferase activity (GNAT) family
	gluco-, galacto- and mannosidases	Cre03.g171050.t1.2	GHL1	Glycosyl hydrolase
	misc2	Cre14.g614100.t1.1	GTR26	Dolichyl- diphosphooligosaccharide-protein glycosyltransferase
	rhodanese	Cre07.g352550.t1.1	RDP3	Putative rhodanese domain phosphatase
	short chain dehydrogenase/ reductase (SDR)	Cre07.g352450.t1.1		Corticosteroid 11-beta- dehydrogenase and related short chain-type dehydrogenases, 3-hydroxybutyrate dehydrogenase

Table 7. Continued on next page



Table 7. Continued

Primary Mapman class	Secondary Mapman class	Gene ID (v4)	Gene name	Annotation
		Cre12.g559350.t1.1		1-Acyl dihydroxyacetone phosphate reductase and related dehydrogenases
		Cre03.g191850.t1.1		Short chain dehydrogenase
	UDP glucosyl and glucoronyl transferases	Cre11.g474450.t1.1		NF*
		Cre03.g205250.t1.2	ELG4	Exostosin-like glycosyltransferase
		Cre16.g659500.t1.1		Lactosylceramide 4-alpha- galactosyltransferase
		Cre11.g483400.t1.2	ELG10	Exostosin-like glycosyltransferase
Tetrapyrrole synthesis	Glu-tRNA synthetase	Cre12.g510800.t1.1	CHLI2	Magnesium-chelatase subunit chll
	magnesium protoporphyrin IX methyltransferase	Cre12.g498550.t1.2		Magnesium protoporphyrin IX S-adenosyl methionine O-methyl transferase (Magnesium- protoporphyrin IX methyltransferase) (PPMT)
	unspecified	Cre12.g516350.t1.1	COX10	Cytochrome c oxidase assembly protein Cox10
	urogen III methylase	Cre02.g133050.t1.2		NF*

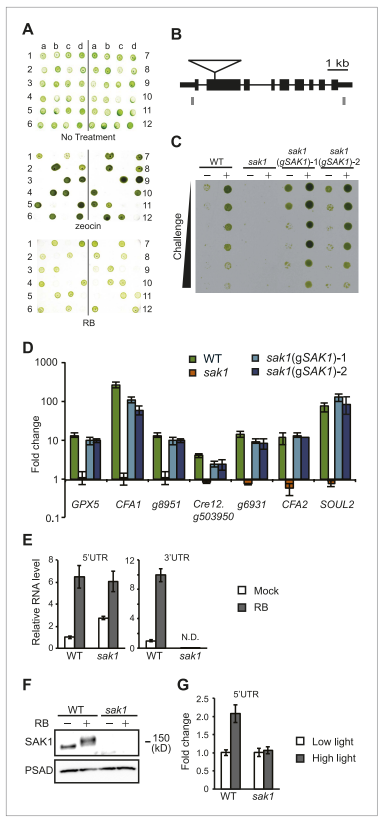
\*Corresponding gene model was not found in v5.

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we isolated a regulatory mutant that is unable to acclimate. Several previous genetic screens aimed at dissecting the mechanisms of  ${}^{1}O_{2}$  signaling have concentrated on the nuclear gene expression response to  ${}^{1}O_{2}$ , often relying on the response of a single marker gene (*Baruah et al., 2009a; Brzezowski et al., 2012; Fischer et al., 2012; Shao et al., 2013*). In contrast, our screen exploited a physiological response to sublethal levels of  ${}^{1}O_{2}$ , which induces the wild type to survive a subsequent, otherwise lethal treatment with the  ${}^{1}O_{2}$  generator RB (*Ledford et al., 2007*). The *sak1* mutant completely lacks this ability to acclimate to  ${}^{1}O_{2}$  (*Figure 1A*). An analogous phenotype is exhibited by the *yap1* $\Delta$  mutant of *Saccharomyces cerevisiae*, which is unable to acclimate to hydrogen peroxide stress (*Stephen et al., 1995*).

In contrast to the complete loss of acclimation to RB, sak1 acclimates (but less effectively than WT) when pretreated with high light and challenged with RB (Figure 1B). This result suggests that the high light pretreatment induces a broader response than that elicited by RB and that sak1 is still able to respond to other signals besides  ${}^{1}O_{2}$  (e.g., plastoquinone redox state,  $H_{2}O_{2}$ , and/or superoxide) that are involved in the response to high light. When tested on TAP agar plates for photoheterotrophic growth in the presence of various photosynthetic inhibitors, the sak1 mutant displayed sensitivity to RB but not to other inhibitors (Figure 1D). In particular, sak1 is not more sensitive than WT to high light or norflurazon (an inhibitor of the biosynthesis of carotenoids, which function as quenchers of  ${}^{1}O_{2}$ ). We speculate that the lack of 102-sensitive phenotype in these plate experiments is attributable to the time-scale of the treatments involved. <sup>1</sup>O<sub>2</sub> generated by RB or during a transfer to higher light intensity is transient, whereas NF requires longer time to exert its effect because it needs to enter the cell, inhibit biosynthesis, and deplete cells of existing carotenoids. During this time, the cell is likely able to acclimate by detoxifying and reducing the generation of 102 by various means such as changing the composition of the photosynthetic apparatus. We have previously shown that acclimation to <sup>1</sup>O<sub>2</sub> is transient and is dissipated by 24 hr post-treatment (Ledford et al., 2007). Consistent with this, pretreatment with RB does not acclimate the cells to stresses such as growth in high light or norflurazon that require a period of days to assess an effect on viability (Figure 1—figure supplement 1). We have also observed that under our experimental conditions, the induction of target gene expression upon exposure to <sup>1</sup>O<sub>2</sub> lasts up to 90 min and then declines. We conclude that SAK1 functions mainly during transient perturbations that generate 102. However, during steady-state growth under high light or norflurazon, the cell is able to cope by other means that do not involve SAK1.





**Figure 4**. Genetic and molecular analysis of *sak1*. (**A**) The insertion of a zeocin resistance gene and the RB sensitivity phenotype are linked. Twelve complete tetrads from a backcross of *sak1* to wild type are shown. Numbers indicate independent tetrads, and letters (a-d) indicate the individual progeny from tetrads. (**B**) Gene Figure 4. Continued on next page



#### Figure 4. Continued

structure of SAK1 and the insertion site. Gray boxes indicate positions of primers used for qPCR. **(C)** Transformation of sak1 with a genomic fragment containing SAK1 rescues the acclimation phenotype. sak1(gSAK1)-1 and sak1(gSAK1)-2 are two independent transformants. **(D)** sak1(gSAK1)-1 and sak1(gSAK1)-2 show recovery of  $^{1}O_{2}$  target gene expression. Y-axis indicates fold change during acclimation to  $^{1}O_{2}$ . **(E)** qRT-PCR of SAK1 in WT and sak1 mutant using primers for 5'- and 3'-UTR shown in panel **B**. **(F)** SAK1 protein is induced in WT and detected as higher molecular weight bands during acclimation to  $^{1}O_{2}$  generated by RB. **(G)** SAK1 transcript probed for 5'-UTR in cells transferred from low light to high light for 1 hr. Error bars indicate standard deviation of biological triplicates. DOI: 10.7554/eLife.02286.014

## SAK1 is necessary for a subset of the genome-wide response to ${}^{1}\text{O}_{2}$ in Chlamydomonas

A physiological acclimation response that results in such an evident growth phenotype (**Figure 1A**) likely involves large-scale changes in gene expression, and transcriptome analysis of wild-type cells showed that hundreds of nuclear genes are up- or down-regulated during acclimation to  ${}^{1}O_{2}$  (**Figure 3A,B**; **Supplementary file 1**, C1). The *sak1* mutant is specifically impaired in regulation of a notable subset of these genes, that is, those that are most strongly induced in the wild type (**Table 5**), suggesting that these genes play a key role in the acclimation response to  ${}^{1}O_{2}$ .

In particular, many genes involved in sterol and lipid metabolism were induced by  ${}^{1}O_{2}$  in Chlamydomonas (Figure 3B; Table 3). For example, two genes encoding putative cyclopropane fatty acid synthase (CFA1 and CFA2) exhibited SAK1-dependent induction (Figure 2). Cyclopropane fatty acids have been found in large amounts in the seeds of Sterculia foetida (Bao et al., 2002), although its biological function is unknown. In bacteria, it has been implicated in oxidative stress responses (Guerzoni et al., 2001; Kim et al., 2005) and particularly in the anoxygenic photosynthetic bacterium Rhodobacter sphaeroides, CFA gene expression is induced during  ${}^{1}O_{2}$  stress by a  ${}^{6}$  factor (Ziegelhoffer and Donohue, 2009). Interestingly CFA mutants of R. sphaeroides are compromised in the induction of genes in response to  ${}^{1}O_{2}$ , suggesting a regulatory role of the gene, protein, or the product of its enzymatic function (cyclopropane fatty acids, Bao et al., 2002) in gene expression rather than solely a biochemical stress response (Nam et al., 2013).

Another intriguing class of up-regulated genes enriched during  ${}^{1}O_{2}$  acclimation in WT and not in sak1 was a group of genes encoding transporters, especially ABC transporters related to the MDR and PDR types. This was not surprising considering that  ${}^{1}O_{2}$  exists in aquatic and terrestrial environments, where it is generated by photosensitizing humic substances (*Frimmel et al., 1987*; *Steinberg et al., 2008*), which are known to affect microbial populations including phytoplankton (*Glaeser et al., 2010, 2014*). Assuming that some of these transporters function to export photosensitizing molecules from the cell, our results suggest that removal of photosensitizers is an integral part of the  ${}^{1}O_{2}$  response in *Chlamydomonas*, rather than simply a response to the presence of a xenobiotic compound such as RB (*Table 4*). It is likely that *Chlamydomonas*, a soil-dwelling microalga, needs to respond to  ${}^{1}O_{2}$  that is generated not only in the chloroplast, but also in other compartments. In this context, it is noteworthy that a recent study has demonstrated light-independent  ${}^{1}O_{2}$  generation in multiple organelles other than the chloroplast under various biotic and abiotic stresses in plants (*Mor et al., 2014*).

Two proteins with SOUL heme-binding domains were among SAK1-dependent up-regulated genes (SOUL2 and Cre06.g299700.t1.1, formerly annotated as SOUL1 in v4). Aside from their ability to bind various porphyrins (*Blackmon et al., 2002*; *Sato et al., 2004*), SOUL heme-binding proteins have been described in diverse biological functions in mice, such as in apoptosis by interacting with a mitochondrial anti-apoptotic factor Bcl-xL (*Ambrosi et al., 2011*) or an isoform-specific role in retina and pineal gland (*Zylka and Reppert, 1999*). The latter form is suggested to play a role in transporting heme or by binding free heme to prevent oxidative stress (*Sato et al., 2004*). In *Arabidopsis* a chloroplast-localized SOUL5 protein has been shown to interact with a heme oxygenase, HY1, and mutation of the gene encoding SOUL5 causes oxidative stress (*Lee et al., 2012*). *Chlamydomonas* contains five putative SOUL heme-binding proteins, only one of which contains an amino-terminal chloroplast transit peptide. The two SOUL protein genes induced by <sup>1</sup>O<sub>2</sub> in our study do not seem to be targeted to the chloroplast, and they may function in the cytosol where SAK1 resides. It would be interesting to test whether these proteins bind porphyrins and are required for <sup>1</sup>O<sub>2</sub> acclimation.



**Table 8.** SAK1 domain containing proteins in chlorophytes

Transcript/

Number in	Organism	Transcript/ Protein IDaTranscript/ Protein IDaTranscript/ Protein ID*
alignment	Organism Volvox carteri	Vocar20009235
2	Volvox carteri	Vocar20007233
3	Volvox carteri	Vocar20002437
4	Volvox carteri	Vocar20004923
5	Volvox carteri	Vocar20012349
6	Volvox carteri	Vocar20005988
7	Volvox carteri	Vocar20007158
8	Volvox carteri	Vocar200071883
9		57405
9	Coccomyxa subellipsoidea	5/405
10	Coccomyxa subellipsoidea	59655
11	Coccomyxa subellipsoidea	57694
12	Chlamydomonas reinhardtii	Cre16.g652650.t1.3
13	Chlamydomonas reinhardtii	Cre06.g271000.t1.2
14	Chlamydomonas reinhardtii	Cre06.g285800.t1.2
15	Chlamydomonas reinhardtii	Cre06.g275600.t1.2
16	Chlamydomonas reinhardtii	Cre06.g285750.t1.3
17	Chlamydomonas reinhardtii	Cre06.g270950.t1.2
18	Chlamydomonas reinhardtii	g9774.t1
SAK1	Chlamydomonas reinhardtii	KF985242
20	Chlamydomonas reinhardtii	Cre03.g179150.t1.2
21	Chlamydomonas reinhardtii	g3701.t1
22	Chlamydomonas reinhardtii	Cre03.g179250.t1.2
23	Chlamydomonas reinhardtii	Cre03.g179200.t1.2
24	Chlamydomonas reinhardtii	Cre01.g004800.t1.2
25	Chlamydomonas reinhardtii	Cre01.g048550.t1.3
26	Chlorella variabilis	EFN51260
27	Chlorella variabilis	EFN53496
28	Chlorella variabilis	EFN55618
Table 8. Conti	inued on next page	

A recent study reported the role of bilins in retrograde signaling in Chlamydomonas through characterization of heme oxygenase mutants disrupted in bilin biosynthesis and transcriptome analyses during dark to light transitions (Duanmu et al., 2013). The transcriptome changes indicated that much of the cell's response during a dark-to-light transition (DL) involves photooxidative stress. Interestingly, among the 515 genes up-regulated in WT during <sup>1</sup>O<sub>2</sub> acclimation, 144 genes overlapped with those that are induced during DL (Table 9). Focusing on the 104 genes that we defined as SAK1-dependent (Table 5), 31 genes overlapped (Table 9). CFA1, CFA2, and SOUL2 were among these genes, suggesting that a part of the gene expression response to DL in Chlamydomonas is a response to <sup>1</sup>O<sub>2</sub>. SAK1 itself was also up-regulated during DL as was SOR1, which encodes a more broadly oxidative stress-responsive bZIP transcription factor (Fischer et al., 2012). We found that 64 of the genes induced during acclimation to <sup>1</sup>O<sub>2</sub> were also up-regulated in the gain-of-function sor1 mutant (Fischer et al., 2012). However, the most strongly induced SAK1-dependent genes were not among these genes, except for GPX5, consistent with the idea that SAK1 and SOR1 function in different pathways.

## SAK1 is a key intermediate component in the retrograde signaling pathway for <sup>1</sup>O<sub>2</sub> acclimation

Cloning of the SAK1 gene revealed that it encodes a large previously uncharacterized phosphoprotein located primarily in the cytosol (Figure 6A,D), suggesting that it functions as an intermediate in the retrograde signaling pathway from the chloroplast to the nucleus that leads to <sup>1</sup>O<sub>2</sub> acclimation. Previous genetic screens in Arabidopsis have identified proteins in the chloroplast, such as EX1 and EX2 (Wagner et al., 2004; Lee et al., 2007), and in the nucleus, such as PLEIOTROPIC RESPONSE LOCUS 1 (Baruah et al., 2009b) and topoisomerase VI (Simková et al., 2012), that are involved in <sup>1</sup>O<sub>2</sub> signaling. By screening for mutants that are unable to induce a <sup>1</sup>O<sub>2</sub>-responsive reporter gene (HPS70A) in Chlamydomonas, a small zinc finger protein (Cre09.q416500.t1.2) called MBS was recently identified as having a role in ROS signaling in both Chlamydomonas and Arabidopsis (Shao et al., 2013). Like SAK1, MBS in Chlamydomonas is located in the cytosol, raising a question about the relationship of these two proteins in <sup>1</sup>O<sub>2</sub> signaling. As expected, we found HSP70A among the genes induced by RB

Table 8. Continued

Number in alignment	Organism	Transcript/ Protein IDaTranscript/ Protein IDaTranscript/ Protein ID*
29	Chlorella variabilis	EFN57652
30	Chlorella variabilis	EFN55658
31	Chlorella variabilis	EFN54262
32	Chlorella variabilis	EFN54510
33	Chlorella variabilis	EFN55806
34	Chlorella variabilis	EFN53492
35	Micromonas sp. RCC299	ACO61347
36	Micromonas pusilla CCMP1545	EEH57791
37	Micromonas sp. RCC299	ACO65814

<sup>\*1–25,</sup> as defined on phytozome.net; 26–37, CrSAK1, genbank accession numbers.

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treatment of *Chlamydomonas* (**Table 3**) however in *sak1* it was not significantly induced above the twofold threshold, suggesting that SAK1 might function in the same signaling pathway as MBS. The *MBS* gene itself is not induced by  ${}^{1}O_{2}$  (*Shao et al., 2013*), and we will investigate the genetic and biochemical relationship of SAK1 and MBS in future research.

SAK1 contains a novel domain of ~150 amino acid residues that is found in several chlorophyte species (Table 8). The sequence of this domain is not highly conserved (Figure 5-figure supplement 1), and is even less conserved among land plant proteins, although it is detectable by PSI-BLAST, indicating that it has diverged in sequence in plants and algae. We identified 37 proteins that have the SAK1 domain, 13 of which also contained a bZIP transcription factor domain, consistent with a function in regulating gene expression. Under our standard laboratory growth conditions, SAK1 appears to have a relatively low level of phosphorylation, but it becomes hyperphosphorylated during <sup>1</sup>O<sub>2</sub> acclimation (Figure 6D). Phosphorylation prediction software NetPhos 2.0 (http://www.cbs.dtu.dk/

services/NetPhos/) predicted 24 serine, 9 threonine, and one tyrosine residue as possible sites throughout the protein (*Figure 5—figure supplement 3*). One of these serine residues is within the conserved SAK1 domain and is relatively conserved for polar amino acids. At this position, 18 SAK1 family members had threonine, and three had serine residues including SAK1 (*Figure 5—figure supplement 1*). We speculate that phosphorylation of SAK1 in the cytosol is a necessary intermediate step in <sup>1</sup>O<sub>2</sub> acclimation. Through further analysis of the transcriptome data, isolation of proteins that physically interact with SAK1, and characterization of additional, non-allelic *sak* mutants, we hope to identify the kinase that is responsible for the direct modification of SAK1 as well as other upstream and downstream components of this retrograde signaling pathway in *Chlamydomonas*.

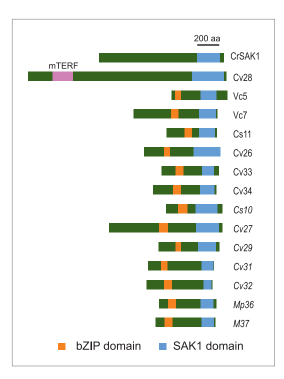
#### Material and methods

#### Chlamydomonas strains and culture conditions

The sak1 mutant was generated by insertional mutagenesis as described previously (**Dent et al., 2005**) from WT strain 4A+. Cells were grown at 22°C photoheterotrophically in Tris-acetate phosphate media (TAP) unless otherwise stated (**Harris, 2009**).

#### RB sensitivity screen and acclimation assays

For systematic screening of large number of strains for increased or decreased resistance to RB, individual strains were inoculated into 180-200  $\mu$ l TAP medium in 96-well plates, grown for a at least 3 days to saturation under light intensity of 60–80  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>, spotted onto TAP plates with 2.7, 3.0, or 3.3  $\mu$ M RB, and scored for their growth compared to WT and *sak1*. For more quantitative evaluation of RB sensitivity, the cells were grown to saturation in 1 ml of TAP medium because we have observed rapidly growing cells to have more variable sensitivity to RB (data not shown). The cells were counted and adjusted to equal cell density then dispensed into aliquots in duplicate 96-well plates. One of the duplicates was pretreated in dark while the other was placed in light for 40 min with 1  $\mu$ M RB. For challenge treatments, 4.5, 5.1, 5.7, 6.3, 6.9, and 7.5  $\mu$ M RB was added to both plates, which were placed under light for 1 hr and then spotted onto TAP agar media with no RB. All treatments were applied under light intensity of 60–80  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>, which is the light intensity described as low light unless stated otherwise.



**Figure 5.** SAK1 contains an uncharacterized domain present in some bZIP transcription factors. Schematic of relative positions of SAK1 and bZIP domains. One protein (Cv28) contains a mitochondrial termination factor (mTERF) domain. The letters and numbers in the abbreviated names represent initials of the species and numbers listed in **Table 8.** Proteins with italicized names contain bZIP domains that were recognized by Pfam but scored below significance.

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The following figure supplements are available for figure 5:

**Figure supplement 1**. Multiple sequence alignment of SAK1 domains.

DOI: 10.7554/eLife.02286.017

**Figure supplement 2**. Secondary structure prediction of SAK1 domain.

DOI: 10.7554/eLife.02286.018

**Figure supplement 3**. Prediction of phosphorylation sites in SAK1.

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## Pretreatment and challenge with RB and $F_v/F_m$ measurement

Cells were grown under 100 µmol photons  $m^{-2}$  s<sup>-1</sup>, adjusted to 2 × 10<sup>6</sup> cells  $ml^{-1}$ , and treated with RB at a final concentration of 0.5 µM for 30 min (pretreatment) in light (+) or dark (–). After the pretreatment all the cultures were exposed to an additional 3.75 µM RB (challenge) in low light and collected for measurement of  $F_v/F_m$  at 30, 60, and 90 min. The cells were dark-acclimated for at least 30 min before applying a saturating light pulse of 2000 µmol photons  $m^{-2}$  s<sup>-1</sup> and measuring the chlorophyll fluorescence yield using an FMS2 fluorometer (Hansatech Instruments, Norfolk, UK).

## Culture conditions for gene expression analyses by qRT-PCR and RNA-seq

Cultures were grown for at least two light-dark cycles (12 hr light-12 hr dark), and then cell density was adjusted to  $2-2.5 \times 10^6$  cells ml<sup>-1</sup> and split into two flasks (one control and the other for RB treatment) at least an hour prior to adding RB to a final concentration of 1 µM. An equal volume of H<sub>2</sub>O was added to the control. RB was added ~6 hr after the start of the light cycle under light intensity of ~100 µmol photons m<sup>-2</sup> s<sup>-1</sup> and the treatment lasted for an hour before harvest. The cells were cooled and harvested by centrifugation at 1200×g for 3 min at 4°C, frozen with liquid nitrogen and stored at -80°C until extraction of RNA. For low light to high light transfer experiment, cultures were grown in continuous light in minimal (HS) medium for 3 days to cell density of  $3 \times 10^6$  cells ml<sup>-1</sup> at 45 µmol photons m<sup>-2</sup> s<sup>-1</sup>. The light intensity was increased to 500 µmol photons  $m^{-2}$  s<sup>-1</sup> for 1 hr before harvest.

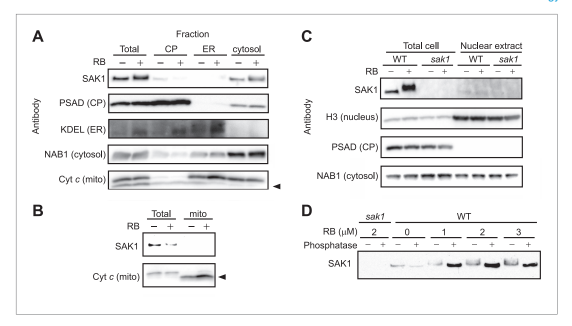
### Gene expression analysis by qRT-PCR

RNA was extracted with TRIzol (Life Technologies, Carlsbad, CA) following manufacturer's instructions and treated with DNasel (Promega, Madison, WI), then cleaned up using Qiagen RNeasy col-

umns (Qiagen, Germantown, MD). cDNA was synthesized using Omniscript (Qiagen, Germantown, MD) starting with 2–3  $\mu$ g DNA-free RNA per 20  $\mu$ l reaction. qPCR was performed using a 7300 FAST qPCR machine (Life Technologies, Carlsbad, CA). The primers were designed with a  $T_m$  of 60°C using Primer3 or PrimerExpress (Life Technologies, Carlsbad, CA) (*Table 10*). All primer pairs described in this study were confirmed as having 90–105% amplification efficiency and linear amplification within their dynamic range in experimental samples using serial dilutions of cDNA prior to the experiments. Relative transcript levels were calculated by  $\Delta\Delta$ Ct method (*Livak and Schmittgen, 2001*) using  $C\beta LP$  as internal reference.

#### RNA-seq library preparation and analysis

RNA was extracted (**Schmollinger et al., 2014**) and the quality was determined using a 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA). The triplicate RNA was pooled and 10 µg total



**Figure 6.** SAK1 is a phosphorylated protein that is in the cytosol. (**A** and **B**) SAK1 is detected in the cytosol and not in other subcellular fractions. (**C**) SAK1 is not enriched in nuclear extracts. Approximately 30  $\mu$ g of protein was loaded into each well except for mitochondrial fractions that were loaded approximately 7.5  $\mu$ g protein due to low protein yield in isolated fractions. Subcellular markers: Chloroplast (CP), PSAD; Endoplasmic reticulum (ER), KDEL; Cytosol, NAB1; Mitochondria (mito), cytochrome c (Cyt c); Nuclear, histone 3 (H3). The arrowhead indicates the band corresponding to Cyt c. (**D**) Protein extracts from cells treated with increasing concentrations of RB were then treated with phosphatase (+) or only with buffer (–) before detection of SAK1 by immunoblot analysis. DOI: 10.7554/eLife.02286.020

RNA was used to prepare RNA-seq library according to the manufacturer's protocol (Illumina, San Diego, CA). The quality of the library was assessed using a 2100 Bioanalyzer before sequencing with Genome Analyzer (Illumina, San Diego, CA). Each sample was run in replicates on two lanes. RNA-Seq data was analyzed as before (*Duanmu et al., 2013*). On average, 75% of the sequences could be assigned unambiguously to Augustus v10.2 gene models to generate the matrix of counts per gene. This matrix was used for differential expression analysis using *DESeq* (*Anders and Huber, 2010*) using *per-condition* dispersion estimates and variance stabilization to compute moderate fold changes. Genes were classified as differentially expressed based on a (moderate) twofold regulation and a false discovery rate (FDR) <1%.

### Amplification of cDNA and genomic region of SAK1 and transformation of sak1

Near full-length cDNA was isolated by RT-PCR (described in above section; Gene expression analysis by qRT-PCR) and rapid amplification of cDNA ends (RACE) using GeneRACER (Life Technologies, Carlsbad, CA) as previously described (*Molnar et al., 2009*). Despite multiple attempts the 5' end of the transcript could not be amplified by 5'-RACE. Because the experimentally obtained CDS differed from the most current v5, it has been deposited to genbank (accession KF985242). Though some differences exist at the nucleotide level, the protein sequence of the resulting CDS was identical to that of au5.g7871\_t1. Genomic DNA containing *SAK1* was amplified using primers 5'-CAGGACCGGGCACTGAGTGAAGGTTA-3' (+) and 5'-ATGATGCACTGTGGGACACGCTGAGT-3' (-) using PrimeStar HS with GC buffer (Takara/Clontech, Palo Alto, CA) and cloned into pGEMTeasy after adding an adenine. The resulting plasmid was co-transformed with pBC1 and selected with 1 µM paromomycin. Transformation of *sak1* was performed as described previously (*Kindle et al., 1989*).

#### SAK1 antibody generation and protein detection by immunoblotting

To raise antibodies against SAK1, an epitope at the N-terminus of the translated coding sequence of SAK1 (DTLLTPLREDATAESGGDA) was designed, synthesized and injected into rabbits, and the



**Table 9.** Genes up-regulated during both  ${}^{1}O_{2}$  acclimation and dark to light transition

Gene ID (v4)	Gene name	Annotation	RB (log <sub>2</sub> )	DL (log <sub>2</sub> ) (Duanmu et al., 2013)
Cre02.g137700.t1.1*	Gene name	Amotation	6.49	2.34
Cre06.g281250.t1.1*	CFA1	cyclopropane fatty acid synthase	5.92	4.49
Cre01.g033300.t1.1*	CIAI	cyclopropane latty acid synthase	5.72	3.62
Cre13.q566850.t1.1*	SOUL2	SOUL heme-binding protein	5.53	2.25
Cre13.g600650.t1.1*	30012	500E herne binding protein	4.76	3.26
Cre06.g263550.t1.1*	LCI7	R53.5-related protein	4.46	5.27
Cre07.g342100.t1.1*	LCII	N33.3-related protein	4.43	1.84
Cre09.g398700.t1.1*	CPLD27	coclaurine N-methyltransferase	4.45	1.36
Cre12.g492650.t1.1*	FAS2	fasciclin-like protein	4.03	9.24
Cre08.g381510.t1.1*	1 A32	тазсісіті-тке ргосеті	3.94	3.27
	GPX5	alutathiana naravidasa	3.94	3.08
Cre10.g458450.t1.2*	GPAS	glutathione peroxidase		
Cre11.g474600.t1.1*			3.90	1.99
Cre13.g600700.t1.1*			3.78	5.79
Cre14.g613950.t1.1*			3.65	2.68
Cre06.g269300.t1.1*			3.50	1.99
Cre08.g380300.t1.2*	MSRA3	peptide methionine sulfoxide reductase	3.45	1.79
Cre01.g031650.t1.2*	CGLD12	protein with potential galactosyl transferase activity	3.30	4.90
Cre14.g629061.t1.1*			3.25	1.88
Cre13.g564900.t1.1*			3.22	3.38
Cre13.g586450.t1.1			3.21	3.50
Cre02.g139500.t1.1*			3.04	2.12
Cre19.g756100.t1.1			3.04	6.53
Cre01.g036000.t1.2			3.02	1.16
Cre14.g618400.t1.1*			2.97	2.16
Cre17.g741300.t1.2*			2.88	1.92
Cre16.g648700.t1.2*			2.79	2.35
Cre17.g729950.t1.1			2.77	2.61
Cre17.g721000.t1.1			2.70	2.12
Cre06.g263500.t1.1*			2.67	3.37
Cre01.g016150.t1.1*			2.65	2.92
Cre08.g380000.t1.1*			2.59	3.74
Cre04.g224800.t1.1	VAMP74	R-SNARE protein, VAMP72-family	2.58	3.34
Cre03.g210150.t1.1		, ,	2.57	3.44
Cre14.g615600.t1.1*			2.53	2.40
Cre06.g293100.t1.1		Qc-SNARE SYP6-like protein	2.48	4.90
Cre08.g368950.t1.1	DHQS	3-dehydroquinate synthase	2.39	2.49
Cre10.g424350.t1.2	51140	metalloprotease	2.37	3.18
Cre12.g537225.t1.1		stanoprotease	2.34	3.39
Cre07.g336900.t1.2			2.34	2.31
Cre16.g664050.t1.1			2.31	1.88
Cre16.g677750.t1.1			2.04	2.22
Cre12.g537227.t1.1			2.00	3.46

Table 9. Continued on next page

Table 9. Continued

Gana ID (v4)	Gene name	Annatation	DD /log \	DL (log <sub>2</sub> ) (Duanmu et al.,
Gene ID (v4)	Gene name	Annotation	RB (log <sub>2</sub> )	2013)
Cre17.g737050.t1.1		RabGAP/TBC protein	1.99	2.32
Cre06.g297450.t1.1			1.93	1.46
Cre06.g258600.t1.1*		COED OF A LLL	1.91	3.63
Cre16.g663950.t1.1		SC5D, C-5 sterol desaturase	1.89	2.03
Cre13.g588150.t1.1	DIVCO		1.86	6.21
Cre17.g722150.t1.1	PKS3	type III polyketide synthase	1.85	1.61
Cre16.g688550.t1.1	GSTS1	glutathione-S-transferase	1.84	1.20
Cre03.g207800.t1.1			1.84	7.09
Cre10.g444550.t1.1*	SPP1A	signal peptide peptidase	1.81	5.33
Cre13.g602500.t1.2			1.76	1.59
Cre03.g163400.t1.2*			1.76	2.15
Cre10.g450000.t1.1			1.74	2.18
Cre01.g015500.t1.1			1.72	1.55
Cre02.g105750.t1.2			1.71	3.23
Cre01.g061750.t1.1	SPT2	serine palmitoyltransferase	1.71	2.29
Cre83.g796250.t1.1			1.68	1.59
Cre16.g656150.t1.1			1.67	3.55
Cre01.g002050.t1.2			1.66	3.15
Cre12.g556750.t1.2	Tic32-like 1	Short-chain dehydrogenase, classical family, similar to PsTic32	1.66	3.15
Cre12.g559100.t1.1			1.66	3.11
Cre09.g411750.t1.2			1.61	1.96
Cre11.g482650.t1.2			1.57	3.40
Cre06.g310500.t1.1*			1.57	6.23
Cre09.g397900.t1.1		transmembrane protein	1.56	2.02
Cre04.g215600.t1.1			1.53	2.64
Cre02.g093800.t1.1			1.51	4.99
Cre02.g093750.t1.1	NRX2	Nucleoredoxin 2	1.50	6.26
Cre01.g004350.t1.1			1.50	2.29
Cre01.g034600.t1.1			1.50	2.22
Cre11.g472600.t1.2			1.48	2.00
Cre12.g500500.t1.2	SMT1	sterol-C24-methyltransferase	1.46	3.05
Cre13.g577950.t1.1	VPS6	subunit of the ESCRT-III complex	1.45	2.36
Cre02.g118200.t1.1			1.44	2.79
Cre01.g012500.t1.1	PRA1	prenylated rab acceptor family protein	1.43	2.46
Cre12.g521600.t1.2			1.42	2.89
Cre03.g179100.t1.1		ubiquitin fusion degradation protein	1.41	3.38
Cre09.g413150.t1.2			1.39	4.31
Cre13.g572200.t1.1		tyrosine/tryptophan transporter protein	1.39	2.57
Cre03.g185850.t1.2		PfkB-type carbohydrate kinase	1.37	3.05
Cre18.g743600.t1.1		7)F - 12 2y 2atoaoo	1.37	1.65
Cre02.g076800.t1.1		sterol reductase	1.36	2.41
Cre06.g256750.t1.1	FAT1	acyl carrier protein thioesterase	1.35	1.67
	1711	asyr carrier protein timoesterase	1.00	1.07

Table 9. Continued on next page

Table 9. Continued

Gene ID (v4)	Gene name	Annotation	RB (log <sub>2</sub> )	DL (log₂) (Duanmu et al., 2013)
Cre17.g729450.t1.1	Gene name	Amotation	1.34	1.90
Cre11.g471550.t1.1			1.34	3.29
Cre09.g395750.t1.2			1.33	2.87
Cre14.g617100.t1.1			1.33	3.33
Cre16.g691500.t1.1		Sec14p-like lipid-binding protein	1.33	2.28
Cre02.g079550.t1.1	DRP2	Dynamin-related GTPase	1.32	2.34
Cre02.g079300.t1.1	VPS4	AAA-ATPase of VPS4/SKD1 family	1.32	1.96
Cre05.g231700.t1.2	V1 34	AAA-ATT ase of VI 34/3KDT failing	1.31	2.40
Cre02.g132300.t1.2	DNJ12	DnaJ-like protein	1.30	2.24
Cre69.g794101.t1.1	DINJIZ	Dilag-like protein	1.30	2.65
Cre13.g565600.t1.2			1.29	3.42
Cre13.g593700.t1.1		monooxygenase, DBH-like	1.29	1.81
		monooxygenase, DBH-like	1.28	3.88
Cre12.g498000.t1.2				
Cre06.g292900.t1.2	LICDZOA	11+-hl	1.28	2.16
Cre08.g372100.t1.1	HSP70A	Heat shock protein 7A	1.27	2.28
Cre01.g039350.t1.1	NCR2	NADPH-cytochrome P45 reductase	1.26	2.19
Cre03.g211100.t1.1			1.26	2.11
Cre17.g731800.t1.1			1.25	1.78
Cre17.g730650.t1.1			1.25	2.28
Cre02.g123000.t1.2			1.24	1.42
Cre05.g247700.t1.2			1.24	2.71
Cre08.g360800.t1.2		haloacid dehalogenase-like hydrolase	1.23	4.39
Cre07.g350750.t1.1	PTOX1	alternative oxidase	1.22	3.32
Cre17.g703750.t1.1			1.20	2.21
Cre06.g306041.t1.1			1.20	2.90
Cre02.g116650.t1.1			1.20	2.83
Cre08.g379400.t1.2			1.18	3.04
Cre16.g677000.t1.1	HSP70E	Heat shock protein 7E	1.18	2.50
Cre06.g283900.t1.1			1.18	5.24
Cre14.g626750.t1.1			1.17	4.12
Cre01.g010700.t1.1			1.16	2.10
Cre01.g002000.t1.2		predicted proteim	1.15	1.68
Cre04.g213150.t1.1			1.15	2.78
Cre16.g694250.t1.1			1.15	2.92
Cre05.g246400.t1.1			1.15	2.74
Cre02.g128450.t1.1			1.13	2.82
Cre03.g180250.t1.1		Myo-inositol-1-phosphate synthase	1.13	2.05
Cre03.g186150.t1.1			1.13	1.78
Cre02.g137800.t1.1			1.13	2.00
Cre11.g471500.t1.1	MFT10	predicted protein	1.11	1.40
Cre10.g435200.t1.1			1.10	2.13
Cre13.g593850.t1.2			1.10	3.91
-				

Table 9. Continued on next page

Table 9 Continued

				DL (log₂) (Duanmu et al.,
Gene ID (v4)	Gene name	Annotation	RB (log	
Cre19.g754000.t1.2			1.10	2.33
Cre13.g593869.t1.1			1.10	3.90
Cre08.g377300.t1.2			1.09	3.27
Cre04.g225050.t1.2		predicted protein	1.09	3.55
Cre07.g330300.t1.1			1.08	2.22
Cre12.g500450.t1.2			1.08	3.00
Cre06.g262000.t1.1			1.08	1.87
Cre10.g441550.t1.2	<i>МАМ3В</i>	predicted protein	1.07	1.54
Cre06.g249800.t1.1		unknown conserved protein	1.07	2.08
Cre01.g038250.t1.1	SDC1	serine decarboxylase	1.06	1.92
Cre44.g788200.t1.1			1.06	2.13
Cre08.g359200.t1.2			1.03	2.69
Cre05.g245950.t1.1	DRP1	Dynamin-related GTPase	1.03	2.15
Cre05.g234100.t1.1	CYP745A1	cytochrome P45	1.01	2.61
Cre07.g328700.t1.2			1.01	1.56
Cre10.g440250.t1.2			1.01	2.14
Cre17.g725200.t1.1		MDR-like ABC transporter	1.01	3.30
Cre82.g796100.t1.1			1.01	2.49

\*Genes defined as SAK1-dependent in Table 4.

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resulting crude serum was affinity purified (Open Biosystems/Thermo Scientific, Waltham, MA). For immunoblot detection of SAK1, proteins were separated with NuPAGE 3–8% Tris Acetate gels (Life Technologies, Carlsbad, CA) and transferred to nitrocellulose membranes. All other blots were prepared from running the protein on 10–20% Tris-glycine gels and transferring to a PVDF membrane. The membranes were blocked for several hours in 5% milk in TBS-T, incubated with the primary antibody overnight, then with secondary antibody for several hours in 1% milk TBS-T before washing and developing with a chemiluminescence detection kit. Commercial antibodies were anti-histone H3 (ab1791; Abcam, Cambridge, UK) and anti-KDEL (ab12223; Abcam, Cambridge, UK). Other antibodies were generous gifts from Jean-David Rochaix (anti-PSAD), Olaf Kruse (anti-NAB1), and Patrice Hamel (anti-cytochrome c).

#### Subcellular fractionation and protein quantification

Nuclear fractions were prepared from 450 ml of synchronized cultures with ~2 × 10 $^6$  cells ml $^{-1}$  that had been incubated with or without 2  $\mu$ M RB under light for 40 min. The cells were collected and treated with autolysin for 40 min and examined for the removal of cell walls by addition of 1 volume of 0.1% Triton-X. Nuclear extract was prepared as described previously (*Winck et al., 2011*) using CelLytic PN kit (Sigma-Aldrich, St. Louis, MO). Because there were bands detected in the nuclear extract close to the size of SAK1, nuclear extract was prepared from WT (4A+) and *sak1* rather than a cell wall-deficient strain (*cw15*). Chloroplasts were isolated from cell wall-less strain *cw15* as described previously (*Klein et al., 1983*). Mitochondria were isolated as described (*Eriksson et al., 1995*). After unbroken cells, chloroplasts, and mitochondria were collected, the ER fraction was collected by centrifugation at 100,000×g for 90 min at 4°C. The remaining supernatant was enriched for cytosol. Protein was extracted and prepared for SDS-PAGE as described (*Calderon et al., 2013*) with minor modifications. Protein was quantified by using BCA1 kit (Sigma-Aldrich, St. Louis, MO) after extraction with the methanol-chloroform method (*Wessel and Flügge, 1984*).

Table 10. Primers used for qRT-PCR analyses

v4 ID	v5 ID	Gene name	Forward	Reverse
Cre01.g007300.t1.1	Cre01.g007300.t1.2		AGCATGTGCGTGTGGAGTAG	CCTTACCATAGGCCTGACCA
au5.g10700_t1a	Cre03.g177600.t1.3		CTGGACATGTCGGCTATGAA	GCTCATGTCGTACTCCAGCA
au5.g13389_t1*	Cre06.g299700.t1	SOUL1†	TGCGTATGGGTGTCCACTAA	TGGGGATCTTCTTCATGTCC
Cre06.g263550.t1.1	Cre06.g263550.t1.2	CCI7	TTTGGTTGCGTTGCATGTAT	TCAACGCGGTGTCAAACTTA
Cre06.g281250.t1.1	Cre06.g281250.t1.2	CFA1	CCTACAACGACGACGTG	GGAAGTTCCAGGATGACCAG
Cre06.g298750.t1.1	Cre06.g298750.t1.2	AOT4	CCGTGTGCACAGATTCAAAG	CACACAGCGCCTCCTACATA
Cre08.g358200.t1.2	Cre08.g358200.t2.1		TGTGGCATCAAGGTGTGTTGT	AACCCCACACCCCTCTTT
Cre09.g398700.t1.1	Cre09.g398700.t1.2	CFA2	CGACCTGCTCTACTTCC	GTGTAGGCGGTGGTCAAGAT
Cre10.g458450.t1.2	Cre10.g458450.t1.3	GPX5	AACCAATCGCCTAACACCTG	CACTTGCTAGCCACGTTCAC
Cre12.g503950.t1.1	Cre12.g503950.t1.2		GGAGGGAGTACCACGAGACA	GATTGCTGTAAGGCCGGATA
Cre13.g564900.t1.1	Cre13.g564900.t1.2	MRP3	TCATGACGTACATCTCGATTCTCA	AGGGAATGTAGTAGCGCTGAATG
au5.g4402_t1*	Cre13.g566800.t1.2		TGCTTGGAAGACCCACTTTT	GAGCTGGAGTTGCAGTTGTG
Cre13.g566850.t1.1	Cre13.g566850.t1.2	SOUL2	CCCTCCCTTCAGACTA	CGTACCTGAGGCGCATATTT
Cre14.g613950.t1.1	Cre14.g613950.t2.1		CGCCCAACCCCATGATC	CCGCAACGTACCGTGATG
Cre16.g683400.t1.1	Cre16.g683400.t1.2		CCTGAACAACACGGATGG	GAACGCCGTCAAATCATCTT
Cre16.g688550.t1.1	Cre16.g688550.t1.2	GST1	AGTGCGGAGGAAGTCGTAAA	GTAAAAGACGTGCGTGCAAA
	g6364.t1	CBLP(RCK1)	GAGTCCAACTACGGCTACGC	GGTGTTCAGGTCCCACAGAC
Cre14.g623650.t1.1	Cre14.g623650.t1		GACAACGCGGCCTACAAGA	CCGAGCTGGCGGTGTTAA
au5.g2281_t1*	g16723.t1	MKS1	GCTTGAGCGCGAGACGAA	CGCTGAAAGCATTGCAGAAG
Cre08.g380300.t1.2	Cre08.g380300.t1.2		ACCACCAGCAGTACCTGTCC	CGCTCCAATAAAGCCTTCAG
au5.g7871_t1‡	(Cre17.g741300.t1.2)‡	SAK1(5'UTR)	CAAGTGCTCATGAGAGGCCTTA	TACGTCATCCAGTTCCACATCC
au5.g7871_t1‡	(Cre17.g741300.t1.2)‡	SAK1(3'UTR)	TCAAGCGTGTGGGTAAGAGCTA	ACGCTATCTCCGTCCTAATCCA
Cre08.g365900.t1.1	Cre08.g365900.t1.2	LHCSR1	CACACAATTCTGCCAACAGC	ATCTGCTTCACGGTTTGGTC
Cre04.g220850.t1.1	Cre04.g220850.t1.2		TAATGGTATGGATGCGGTCA	ACTGCCAGTTATGGGTCCTG
Cre09.q395750.t1.2	Cre09.q395750.t1.3		ACCGTCCGTGAACCTTACTG	CGCAAACACGTCTCAAAGAA

\*Was originally mapped and identified as augustus version 5 models within Chlamydomonas genome v4.

†50UL1 was given the name in v4 but not v5.

‡Primers were designed against experimentally obtained cDNA (Genbank accession KF985242) and differs from v5. Closest gene model is au5.g7871\_t1. DOI: 10.7554/eLife.02286.022



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#### **Author contributions**

SW, Conception and design, Acquisition of data, Analysis and interpretation of data, Drafting or revising the article; BLC, RMD, Acquisition of data, Drafting or revising the article; HKL, Acquisition of data; DC, Acquisition of data, Analysis and interpretation of data, Drafting or revising the article; MP, Acquisition of data, Analysis and interpretation of data; SSM, Analysis and interpretation of data, Drafting or revising the article; KKN, Conception and design, Analysis and interpretation of data, Drafting or revising the article

#### **Additional files**

#### Supplementary file

• Supplementary file 1. Genes that display significant differential expression by pair-wise comparisons. DOI: 10.7554/eLife.02286.023

#### Major dataset

The following dataset was generated:

Author(s)	Year	Dataset title	Dataset ID and/or URL	Database, license, and accessibility information
Wakao S, Chin BL, Ledford HK, Dent RM, Casero D, Pellegrini M, Merchant SS, Niyogi KK	2014	Data from: Phosphoprotein SAK1 is a regulator of acclimation to singlet oxygen in Chlamydomonas reinhardtii	http://dx.doi.org/ doi:10.5061/dryad. h7pm2	Available at Dryad Digital Repository under a CC0 Public Domain Dedication.

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