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Genome-wide analysis on *Chlamydomonas reinhardtii* reveals the impact of hydrogen peroxide on protein stress responses and overlap with other stress transcriptomes

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**SUMMARY**

Reactive oxygen species (ROS) are produced by and have the potential to be damaging to all aerobic organisms. In photosynthetic organisms, they are an unavoidable byproduct of electron transfer in both the chloroplast and mitochondrion. Here, we employ the reference unicellular green alga *Chlamydomonas reinhardtii* to identify the effect of H\(_2\)O\(_2\) on gene expression by monitoring the changes in the transcriptome in a time-course experiment. Comparison of transcriptomes from cells sampled immediately prior to the addition of H\(_2\)O\(_2\) and 0.5 and 1 h subsequently revealed 1278 differentially abundant transcripts. Of those transcripts that increase in abundance, many encode proteins involved in ROS detoxification, protein degradation and stress responses, whereas among those that decrease are transcripts encoding proteins involved in photosynthesis and central carbon metabolism. In addition to these transcriptomic adjustments, we observe that addition of H\(_2\)O\(_2\) is followed by an accumulation and oxidation of the total intracellular glutathione pool, and a decrease in photosynthetic O\(_2\) output. Additionally, we analyze our transcriptomes in the context of changes in transcript abundance in response to singlet O\(_2\) (O\(_2^\bullet\)), and relate our H\(_2\)O\(_2\)-induced transcripts to a diurnal transcriptome, where we demonstrate enrichments of H\(_2\)O\(_2\)-induced transcripts early in the light phase, late in the light phase and 2 h prior to light. On this basis several genes that are highlighted in this work may be involved in previously undiscovered stress remediation pathways or acclimation responses.

**Keywords:** H\(_2\)O\(_2\), oxidative stress, stress responses, redox signaling, reactive oxygen species, RNA-seq.

**INTRODUCTION**

Reactive oxygen species (ROS) are potentially harmful but unavoidable byproducts of aerobic respiration and oxygenic photosynthesis. One of the major sources of ROS in the cell is the reduction of univalent O\(_2\) that generates superoxide (O\(_2^\bullet\)) in complexes I and III of the mitochondria. In plants, the plastid is an additional source of ROS, where energy transfer within the reaction center of photosystem II (PSII) generates singlet oxygen (O\(_2^\bullet\)), and photoreduction of O\(_2\) to O\(_2\) commonly occurs at photosystem I (PSI) (reviewed in Apel and Hirt, 2004). Although several independent detoxification systems exist, the transient accumulation of ROS is a vital signal that allows the cell to regulate the metabolic processes that generate these toxins and ensure that appropriate protective measures are taken to avoid irreversible damage to proteins, lipids and nucleic acids. The cell can orchestrate the production of enzymes to reduce ROS concentrations and repair damage; otherwise, if the damage is irrevocable, autophagy or apoptosis may ensue.

The ability of H\(_2\)O\(_2\) to elicit specific induction or repression of gene expression is well documented (Stone and Yang, 2006; Foyer and Noctor, 2009; Gough and Cotter, 2015). © 2015 The Authors

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2011). Many of these changes serve to increase the cell’s capacity to detoxify H$_2$O$_2$, usually through the activity of catalases (to produce H$_2$O and O$_2$) or peroxidases (to produce H$_2$O). Unlike catalase, peroxidase must be reduced by an external electron donor (such as glutathione for glutathione peroxidases) to reduce H$_2$O$_2$. Therefore, a common response to H$_2$O$_2$ stress is the biosynthesis of antioxidants such as glutathione and ascorbate. Other responses are not necessarily directed at managing H$_2$O$_2$ levels but at mitigating the damage produced by H$_2$O$_2$. Conserved transcriptional responses include expression of molecular chaperones and increased capacity for protein degradation (Vandenbroucke et al., 2008). Poisoning appears to originate with the Fenton reaction (Walling, 1975), whereby H$_2$O$_2$ oxidizes solvent-exposed Fe$^{2+}$, damaging Fe–S clusters (Jang and Imlay, 2010) and inactivating mononuclear Fe proteins (Sobota and Imlay, 2011; Anjem and Imlay, 2012). In addition to the injury imposed on these susceptible Fe-dependent proteins, the Fenton reaction produces the hydroxyl radical (‘OH), which cannot be enzymatically detoxified and reacts at virtually diffusion-limited rates with most biomolecules. In many ways, H$_2$O$_2$ stress can resemble an unfolded protein response, likely due to the misfolding or inactivation of oxidized proteins.

In this study, we employed the single-celled green alga *Chlamydomonas reinhardtii* as a robust system for understanding the global impact of H$_2$O$_2$ stress on the transcriptomic landscape in a photosynthesizing cell. In recent years, *Chlamydomonas* has become a favored organism for the interrogation of transcriptional responses to stress through the use of RNA-sequencing (RNA-seq) (González-Ballester et al., 2010; Blaby et al., 2013; Schmollinger et al., 2014; Wakao et al., 2014). In addition to surveying the response of global gene expression to H$_2$O$_2$, we validated several of our observations by physiological and biochemical approaches. The wealth of data from previously published transcriptomes in this organism enabled us to contextualize our results with other stresses, allowing us to identify both overlapping and unique responses.

RESULTS

Genome-wide responses to H$_2$O$_2$

To capture both intermediate and persistent transcriptional responses, RNA-seq was performed on RNA isolated immediately before (0'), 0.5 and 1 h after the addition of 1 mM H$_2$O$_2$ (Figure 1a). Based on the rapid degradation of H$_2$O$_2$ from the culture (Figure 1b), we reasoned that primary transcriptional responses would occur within the first hour. Indeed, within 0.5 h, the supernatant concentration of H$_2$O$_2$ from the cultures was reduced to about 60% and reduced to background levels after 4 h (Figure 1b). The addition of 1 mM H$_2$O$_2$ had no noticeable effect on cell number or growth rate (Figure 1c).

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The sequenced reads were aligned to the v.5 assembly of the *Chlamydomonas* genome (strain CC-503 cv92 mt*), and expression estimates were determined for 17 741 (Table S1) loci using the JGI v.5.5 gene models (Merchant
et al., 2007; Blaby et al., 2014). Roughly 7% of predicted transcripts (1278) were differentially abundant [defined as ≥2.0-fold change and ≥10 fragments per kb of exon per million fragments (FPKM)] between any two of the three time points (Table S2). The abundance of 291 transcripts changed between 0’ and 0.5 h (68 down and 223 up), 1251 transcripts between 0’ and 1 h (511 down and 740 up) and 201 transcripts between 0.5 and 1 h (68 down and 133 up).

Based on manual annotation of these transcripts using Pfam domains (Finn et al., 2014), orthology to Arabidopsis thaliana and/or Saccharomyces cerevisiae (based on reciprocal best Blastp hits) or literature-based functional curation, the highest enriched categories include protein metabolism, nucleotide/nucleic acid metabolism and transcripts that encode proteins related to organelles of the endomembrane system (Figure 2, Table S2). Stress and signaling also factored highly among upregulated functional groups, whereas the tetrapyrrole synthesis and photosynthesis groups decreased in mRNA abundance in all three permutations of comparisons (i.e. 0’–0.5, 0.5–1 and 0’–1). Roughly 13% (73 genes) of the GreenCut, a phylogenetic inventory of chloroplast-related genes (Karpowicz et al., 2011; Hein Nickel and Grossman, 2013), was differentially regulated across the time course (between 0’ and 1 h), with 28 decreasing and 45 increasing in transcript abundance (Table S3). Several of those transcripts increasing in abundance encode heat-shock proteins and other stress-related functions, and it is possible that others of currently unknown function have a role in stress remediation (see below and Table S3). Six genes belonging to the CiliaCut (conserved genes amongst ciliated organisms; Merchant et al., 2007) were differentially expressed, all of which decreased in transcript abundance between 0’ and 1 h (although only 165 of the 191 genes

Figure 2. Functional classification of transcripts involved in the H2O2 response.
(a) Functional characterization of genes differentially expressed between any two time points. The number of transcripts in each category is shown in parentheses. ROS, reactive oxygen species; ER, endoplasmic reticulum.
(b) The number of genes in each subcategory within the protein metabolism class of panel (a). Increased and decreased mRNA abundances are shown in blue and red, respectively.
(c) Relative transcript abundance of genes encoding components of the proteasome (normalized to the maximum abundance).
(d) Relative transcript abundance of genes encoding ubiquitin and proteins that mediate the ligation (or hydrolysis) of ubiquitin to target proteins. The mRNA abundances are presented normalized to maximum.
(e) Relative transcript abundance of genes encoding putative and known proteases and peptidases (normalized to peak abundance). The fragments per kb of exon per million fragments (FPKM) values can be found in Tables S1 and S2.
in the published CiliaCut, originally defined using the v.3 genome assembly, could be reliably mapped to v.5.5 gene models.

Protein stress
Using both manual annotation and automated MapMan ontologies, the largest proportion of transcripts fell into protein metabolism (Figures 2a,b and S1, Table S2), suggesting a reprioritization of cellular machinery. We found that the transcripts encoding all putative subunits of the 26S proteasome increased in abundance following the addition of H$_2$O$_2$ (Figure 2c; 1.8- to 3.3-fold between 0’ and 1 h). This coincided with an increase in 31 transcripts encoding proteins putatively involved in ubiquitination, and a reduction in transcript abundance for two putative ubiquitin-specific proteases (Figure 2d). An impact on protein degradation following the addition of H$_2$O$_2$ was not limited to the proteasome–ubiquitin pathway. We also found that the transcript abundance for 22 putative proteases and peptidases increased (while seven decreased) between 0’ and 1 h (Figure 2e). Other indications of protein stress include increased abundance of 38 transcripts encoding either known or putative molecular chaperones and chaperonins (such as HSP20, HSP90, HSP70, DnaJ proteins and FKBP-type peptidyl-prolyl cis-trans isomerases) across the time course (Table S2). While pathways were induced, presumably to assist in protein folding and initiate protein turnover, 48 transcripts encoding proteins associated with the ribosome, enzymes involved in rRNA processing and ribosome modification/assembly were downregulated, as were transcripts encoding proteins responsible for transcription and several tRNA modifications (Table S2).

Detoxification of ROS
The dual role of H$_2$O$_2$ as a harmful by-product of essential oxygenic metabolic reactions and as a signaling molecule necessitates high-fidelity modulation of its presence and concentration within each cellular compartment. Consequently, all aerobic and photosynthetic organisms encode an array of systems charged with ensuring that intracellular H$_2$O$_2$ concentrations do not become toxic. Upon exposure to H$_2$O$_2$, we observed that several ROS detoxification systems were induced. Although transcripts for some ROS remediation proteins decrease in this experiment, they were already relatively highly abundant at 0’, including transcripts encoding catalases and peroxiredoxins (Figure 3, Table S4). Of the six putative superoxide dismutases (FSD1 and MSD1–MSD5), FSD1, MSD1 and MSD2 moderately increased in transcript abundance (by about 1.5-fold) with maximal expression estimates at 839, 382 and 116 FPKM, respectively. The transcript level of MSD3 increased eight-fold between 0’ and 1 h, although transcript abundance remained relatively low (peak 24 FPKM). Neither MSD4 nor MSD5, both predicted to localize to the secretory pathway, were expressed (Table S4). The abundance of transcripts encoding three of five glutathione peroxidases did not change. However, GPX4 and GPX5 transcripts (encoding non-selenocysteine GPX, and considered to use thioredoxin rather than glutathione as electron donor; Dayer et al., 2008) increased in abundance (peak FPKM values of 20 and 193, respectively). GPX5 expression was previously shown to be induced by multiple ROS, including H$_2$O$_2$, and the corresponding protein was found to reduce H$_2$O$_2$ using cytosolic TRXh1, but not GSH, as an electron donor (Leisinger et al., 2001; Fischer et al., 2009). Transcripts encoding proteins putatively contributing to the remediation of ROS damage that were also increased included those encoding nucleoredoxin (NRX2 and NRX3), thioredoxin (TRXh1), glutaredoxin (GRX4), methionine sulfoxide reductase (Cre10.g464850) and, as noted previously (Urzica et al., 2012a), four steps in the Smirnoff–Wheeler pathway for ascorbate biosynthesis (VTc2, MDAR1, PNo1 and DHAR1). By contrast, mRNA abundances for 2-cys peroxiredoxin-, ascorbate peroxidase- or catalase-encoding genes did not increase following the addition of H$_2$O$_2$. Indeed, both CAT1 and CAT2 transcripts decreased in transcript abundance over the time course, with CAT1 transcript being highly abundant at 0’ (197 FPKM) (Table S4). In the case of 2-cys peroxiredoxins (for which the abundance of PRX1 and PRX2 transcripts was already high at 0’; FPKM > 1300), the capacity for H$_2$O$_2$ decomposition may be increased by the induction of two putative sulfiredoxins (encoded by Cre05.g232800 and Cre17.g729850), both of whose transcripts increase in abundance. These proteins can reactivate over-oxidized 2-cys peroxiredoxin, a mechanism considered to play an important role in H$_2$O$_2$ signal...
transduction in mammals (reviewed in Jeong et al., 2012). PRX1 and the sulfiredoxin Cre17.g729950 are predicted to localize to the plastid, while PRX2 and the second putative sulfiredoxin (Cre05.g232800) have no predicted signal peptide (Table S4).

We observed little impact on mRNA abundances of tocopherol (vitamin E), polyamine or carotenoid biosynthesis genes. Each of these pathways has been demonstrated to function in ROS detoxification in other organisms (Havaux et al., 2005; Schneider, 2005; Zhu et al., 2010), although tocopherol acts primarily as a scavenger of O$_2^\cdot$. Our data suggest that in Chlamydomonas these metabolites may not be important for acclimating to short-term H$_2$O$_2$ exposure.

Effect of H$_2$O$_2$ on glutathione metabolism

As in other organisms, the glutathione redox cycle is expected to play a central role in maintenance of the cellular redox balance. We observed that abundance of GSH1 mRNA, encoding γ-glutamylcysteine synthetase, the first enzyme in glutathione biosynthesis, increased more than two-fold between 0’ and 1 h (peak FPKM 122). In contrast, the abundance of the mRNA encoding the subsequent enzymatic step, glutathione synthetase (GSH2), remained low, and decreased slightly across the time course (Figure 4a, Table S4). To investigate the impact that these transcriptional responses had on glutathione levels, we determined the glutathione pool size in response to H$_2$O$_2$ and the proportion present in the oxidized state. We observed both oxidation and accumulation of glutathione after H$_2$O$_2$ treatment. The glutathione pool size increased linearly and had more than doubled within 4 h of H$_2$O$_2$ addition (Figure 4b). The redox state of glutathione was also altered concomitantly due to the accumulation of oxidized glutathione (GSSG), which resulted in a 50% decrease in the ratio of reduced to oxidized glutathione (GSH/GSSG) within 0.5 h. Within 4 h, this oxidation is partially reversed as the relative proportion of the GSH fraction begins to increase (Figure 4c).

We also observed that the abundance of GST1 and GST2 transcripts, encoding two putative glutathione-S-transferases, was significantly elevated (about 100-fold and 10-fold, respectively, between 0’ and 1 h, each peaking at >700 FPKM) (Figure 4a, Table S4). The expression of both genes was previously found to increase in response to ROS (Fischer et al., 2005). Moreover, consistent with increased sulfur requirements for cysteine production as a precursor for glutathione biosynthesis, we observed slight increases (two-fold at 1 h versus 0’) in expression of a putative cysteine transporter (AOT4) and generally for genes encoding enzymes involved in sulfur metabolism.

Photosynthesis

Previous transcriptomic studies on H$_2$O$_2$ exposure in land plants and photosynthetic microbes have noted downregulation of genes encoding proteins with roles in photosynthesis (Desikan et al., 2001; Vandenabeele et al., 2003; Zeller et al., 2005). In our dataset, although the change in abundance for most transcripts did not meet our cut-offs, we observed that in general mRNA abundance corresponding to nuclear genome-encoded subunits of photosynthetic complexes was reduced by 30–50% in response to H$_2$O$_2$ (Table S5). The abundance of transcripts encoding enzymes of chlorophyll biosynthesis are also generally

![Figure 4](image-url)
reduced to about 50% at 1 h with respect to their abundances at 0. In contrast, CHL1, which encodes the ortholog of chlorophyllase I in Arabidopsis, increased in transcript level three-fold over the same time period. The impact of H$_2$O$_2$ on these transcripts correlated with a reduction in photosynthetic efficiency evaluated by measuring the maximum efficiency of PSII and O$_2$ evolution following the addition of H$_2$O$_2$ (Figure 5a,b). Both of these parameters return to pre-H$_2$O$_2$ levels after 4 h, at which time the H$_2$O$_2$ content of the cell-free supernatant has returned to background levels (Figure 1b), consistent with observations in tobacco (Vandenabeele et al., 2003).

Exceptions to the downward trend include LHCB7, encoding a minor antenna protein associated with PSII (Peers and Niyogi, 2008), which increased in abundance by 2.5-fold, peaking at 27 FPKM at 1 h, and TBA1, encoding a putative oxidoreductase proposed to regulate translation of psbA in response to redox status (Somanchi et al., 2005), which increased in abundance by 3.3-fold, peaking at 61 FPKM. Consistent with their known/predicted role in mitigating photo-oxidative stress, several LHC-like genes rose in transcript abundance during the time course, including ELIP2 (two-fold induction, peak 13 FPKM) and ELIP8 (about three-fold induction, peak 162 FPKM) (Table S5).

**Respiration**

Having observed a reduction in photosynthetic O$_2$ evolution and in transcript abundance of many genes encoding photosynthetic subunits, we were interested to determine how the mitochondrial electron transport chain (ETC) was affected. Accordingly, we observed a decline in O$_2$ consumption, with a trough at 0.5 h before recovering to pre-H$_2$O$_2$ rates by 2-4 h. However, despite this measurable reduction, the abundance of most transcripts encoding proteins of the mitochondrial ETC does not change (although the transcript level need not necessarily correlate with changes in protein level, or activity) (Table S6). Deviating from this trend are transcripts encoding assembly factors for complex III (BCS1) and complex IV (SCO1, PET191, and CMC1), whose abundances increase between 0 and 1 h. These increases coincide with the increased abundance of transcripts encoding the alternative enzymes NDA1 (a type II NAD(P)H dehydrogenase) and AOX1 (an alternative oxidase), which increase 2.2-fold (peaking at 31 FPKM) and 4.3-fold (peaking at 96 FPKM), respectively.

**Carbon metabolism**

The abundances of many transcripts encoding enzymes of central carbon metabolism, including the glyoxylate cycle, glycolysis/glucogenesis and the Calvin-Benson (CB) cycle, generally decrease moderately over the time course (Table S7). Coincident with a reduction in O$_2$ consumption, we observed major reductions in mRNA abundance of a
few specific metabolism-related genes, specifically those involved in acetate assimilation, including a putative acetate transporter Cre17.g702990 (Goodenough et al., 2014) and ACS3 and MAST encoding acetyl-coA synthetase and malate synthase, respectively. Transcripts from these three genes exhibit some of the largest reductions in abundance in the dataset (Table S2) and may reflect a reduced flux of metabolites towards acetyl-CoA, and respiration. In contrast, we observed a slight increase in mRNA abundance of several lipid and starch synthesis genes, likely induced as part of a general stress response (Table S7). Transcript levels of GPD2, encoding glycerol-3-phosphate dehydrogenase, and linking carbon assimilation via the CB cycle and gluconeogenesis to lipid synthesis, increased five-fold between 0’ and 0.5 h (FPKM 9 and 66, respectively). Previously, it has been speculated that there may be a link between oxidative stress and the carbon-concentrating mechanism (CCM), based on observations that CAH1 and CCP1 were repressed in response to a number of ROS (Ledford et al., 2007). Our data recapitulate these observations, and we observed reduced abundance of CAH1 transcript (reduced from >500 FPKM at 0’ to 138 at 1 h), and also of CAH3 transcript (127 and 87 FPKM at 0’ and 1 h, respectively), although this appears not to be a general phenomenon of the CCM as the abundances of other known transcript-encoding components were either stable or increased slightly.

COMPARATIVE TRANSCRIPTOME ANALYSES

Overlap between H2O2 and singlet oxygen

We observed a significant overlap in the number of transcripts that increased in abundance following addition of H2O2 and rose bengal (RB), which generates O2; (Figure 6; Wakao et al., 2014). Roughly 15% of the induced H2O2 transcripts between 0' and 1 h were induced by RB (27% of the RB transcripts in H2O2; Table S2). The three most highly represented categories common to both datasets were protein metabolism, proteins related to the endoplasmic reticulum ER/Golgi/secretory pathway and lipid metabolism (Figure 6a). However, only the ER/Golgi/secretory and lipid metabolism groups were significant (P-value < 0.001). We noted that the effect on proteasome-subunit transcript abundance and transcripts encoding molecular chaperones (such as heat shock proteins, HSPs) was specific to H2O2; the only transcripts encoding HSPs affected by both stresses were HSP70E, HSP70A and CPN60C. The analysis also found that PTOX1 was increased in both datasets, as was GPX5 and two nucleoredoxins. Of the top 10 fold-changing transcripts in each pairwise comparison in our H2O2 dataset (i.e. 0’ versus 0.5 h, 0’ versus 1 and 1 h versus 0.5 h), only two transcripts were also in the RB dataset: a glutathione S-transferase and a protein bearing similarity to the ThiJ family (encoded by GST1 and Cre01.g004900, respectively). H2O2, but not RB, affected transcripts involved in tetrahydrole metabolism (Table S2).

We found a larger (and more significant) overlap between the H2O2 response and the RB response in the sak1 mutant than in the parent (4A+). SAK1 is a putative transcription factor, and disruption of the corresponding gene leads to defective transcriptional responses when the mutant is exposed to O2 stress (Wakao et al., 2014). Roughly 25% of the induced H2O2 transcripts between 0’ and 1 h were also induced by RB in sak1, versus 15% in the parent strain (Figure 6b). To qualify the over-

![Figure 6](https://example.com/figure6.png)
lap between these datasets, we also determined the overlap between the H$_2$O$_2$ response and several other transcriptomes that were noted for their enrichment of transcripts related to oxidative stress (ClpP-depletion and poor Fe nutrition; Ramundo et al., 2014; Urzica et al., 2012b) or not (rapamycin-treatment, anoxia, poor Cu nutrition and poor Zn nutrition; Castruita et al., 2011; Hemschemeier et al., 2013; Malasarn et al., 2013) (Figure 6b). We observed both ClpP-depletion and rapamycin treatment (Ramundo et al., 2014) led to a more significant enrichment of H$_2$O$_2$-induced transcripts than did addition of RB (based on the P-value calculated using the hypergeometric distribution). Conversely, both anoxia and copper depletion had relatively little overlap (Figure 6b). Among these comparisons, the Fe dataset had a moderate overlap, while this analysis revealed a surprisingly large enrichment of H$_2$O$_2$-induced transcripts in the Zn-deficient dataset (Figure 6b).

### Oxidative stress during the Chlamydomonas cell cycle

A recent diurnal transcriptome experiment identified a ‘light-stress cluster’ of 280 genes that was transiently expressed at the onset of the light phase, but remained near-undetectable across remaining time points (Zones et al., 2015). Of these 280 transcripts, 99 were significantly differentially abundant in response to H$_2$O$_2$ in our experiment (56 increased mRNA abundance, 43 decreased between 0’ and 1 h). Next, we determined the time point representing the peak transcript level for each gene in the Chlamydomonas genome (using a cutoff of at least 10 FPKM at that point) and analyzed the overlap between the transcripts at each time point and those induced by H$_2$O$_2$ addition. Again, we identified an enrichment of H$_2$O$_2$-induced transcripts at the first time point in the light. We also observed an enrichment of H$_2$O$_2$-induced transcripts with peak values towards the end of the day. We did not observe a significant enrichment during the night except, interestingly, at the 22-h time point (2 h prior to the lights coming on; Figure 7a). For transcripts to which we could assign a general function, the 1-h overlap contained transcripts involved in protein folding and nucleic acid metabolism and binding, whereas towards the end of the day there was an increasing enrichment of transcripts related to protein metabolism, in particular the proteasome. At the 22-h time point there was an enrichment of transcripts related to ROS detoxification/regulation (Figure 7b, Table S8).

For comparative purposes, we repeated this analysis using the RB treatment (parent and sak1) and anoxic transcripts in place of our H$_2$O$_2$ dataset. As with the comparisons described above, we saw some enrichment of RB-induced transcripts at the 1-h time point and toward the end of the day in both strains, but in contrast to the H$_2$O$_2$ dataset the most significant overlap was at the 22-h time point. As expected from the redox status of these cells, we saw no enrichment for anoxia transcripts until the last hour of night.

### DISCUSSION

In this study, we have leveraged our transcriptomic analysis to provide a comprehensive overview of the impact that addition of H$_2$O$_2$ has on transcript abundance in Chlamydomonas. A specific focus on photosynthetic metabolism and ROS detoxification mechanisms provides robust predictions of stress-responsive and remediating genes. In addition, by placing these data in the context of related transcriptomes (Castruita et al., 2011; Hemschemeier et al., 2013; Malasarn et al., 2013; Ramundo et al., 2014; Urzica et al., 2012b; Wakao et al., 2014, Zones et al., 2015) we have been able to identify overlapping responses, potentially highlighting the contribution of ROS stress to independent stimuli. The ability to survey the level of mRNA abundance of all genes provides a powerful tool for assessing the involvement of genes responding to a condition, and thus serves as a means towards hypothesizing roles for genes of previously unknown function.

In the presented work, we identified significant changes in the transcript abundance of 1278 genes, of which 175 have a primary gene symbol and 932 have no known biological role (Table S2). Given this, many of these genes may be of particular interest. They could encode previously undiscovered stress response/ROS detoxification systems, especially considering that the majority increased in transcript abundance; between 0’ and 0.5 h, 223 transcripts increased, as did 740 between 0’ and 1 h, whereas 68 and 511 transcripts decreased between these respective time points (Table S1). One gene that exhibits one of the greatest increases in transcript abundance (at both 0’ versus 0.5 h, and at 0’ versus 1 h) is Cre03.g152750, which was induced 80-fold. The predicted protein sequence encodes a BAG domain, associated with apoptosis and programmed cell death (Kang et al., 2006; Kabbage and Dickman, 2008), and a calmodulin-binding domain. That this gene is very lowly expressed (R/FPKM ≤ 1) in all but one other published Chlamydomonas transcriptome, including RB exposure, suggests it may be responding specifically to H$_2$O$_2$ exposure. This notion is supported by its expression profile in a recent diurnal experiment, in which mRNA is detectable at the first light time point, but at no other, and was a member of a light-stress cluster in that work (Zones et al., 2015). Cre12.g542050 and Cre04.g226138 are increased in abundance, but do not contain predicted conserved domains and bear no significant similarity to proteins encoded on genomes besides Chlamydomonas and the related green alga Volvox carteri. The absence of predicted protein domains or clear orthologs in other characterized organisms makes these genes recalcitrant to functional predictions, and future investigation employing approaches such as classical/reverse genetics will be
required to confirm their involvement in ROS detoxification. This analysis also uncovered a previously unrecognized putative ferritin-encoding gene in the Chlamydomonas genome. Cre01.g033300 encodes a protein with high similarity to bacterial DPS (a DNA-binding protein from starved cells), which are ferritin-like proteins responsible for sequestering Fe ions as a means to protect DNA from oxidative damage (Almirón et al., 1992; Pulliainen et al., 2005). Interestingly, in response to H₂O₂ Cre01.g033300 transcript was increased roughly 10-fold, but the expression estimates remained low (peak 10 FPKM; Table S2), whereas after RB treatment, this transcript increased roughly 80-fold (peak 223 FPKM; Wakao et al., 2014). Also, Cre01.g033300 is clearly regulated by SAK1 (Wakao et al., 2014), suggesting that Fe-catalyzed DNA damage may be a larger threat from singlet oxygen stress than H₂O₂. Seventy-five genes (45 of unknown function) belonging to the GreenCut inventory, whose phylogenetic profile suggests they are of importance to green photosynthetic organisms, increased in mRNA level in this study. This implicates these genes in having some role in management of ROS stress; indeed, included within this group are several genes previously identified in a light-stress cluster (Zones et al., 2015), as well as numerous characterized stress response enzymes (the largest increase in transcript level is HSP22A, whose mRNA abundance increases 176-fold to 58 FPKM at 1 h from 0.3 at 0 h). Furthermore, the Arabidopsis ortholog of one GreenCut member, LHCB7, a PSII antenna protein, has been shown to be expressed under conditions of oxidative stress (Alboresi et al., 2011) (mRNA level increase of 2.5-fold in our data across the time course).

Figure 7. H₂O₂-responsive transcripts during the diurnal cycle.
(a) Analysis of the peak transcript abundance of H₂O₂-induced transcripts during a 12-h diurnal cycle (Zones et al., 2015). P-values calculated (using the hypergeometric distribution) for enrichment of H₂O₂-induced transcripts at each time point are presented as a bar graph while the number of H₂O₂-induced transcripts as a percentage of the total number of H₂O₂-induced transcripts are shown as a circle. The 12 h in light is shaded with yellow and highlighted with a sun, while the 12 h in dark is shaded grey and highlighted with a moon. 
(b) The top five functional categories containing H₂O₂-induced transcripts at the most significant time points in the cycle. ROS, reactive oxygen species; ER, endoplasmic reticulum.
(c) Same as for panel (a) except that the percentage and associated probability of rose bengal-induced transcripts from the wild type (WT) and the sak1 mutant is shown.
(d) Same as panel (a) except that the percentage and associated probability of anoxia-induced transcripts is shown.

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In terms of redox detoxification, the transcripts for many quintessential H$_2$O$_2$-reducing enzymes did not respond to H$_2$O$_2$ treatment, such as ascorbate peroxidase and catalase. Both CAT1 and CAT2 transcript abundance were actually reduced over our time course. This is in contrast with other unicellular organisms, such as S. cerevisiae or Synechocystis PCC 6803, but similar to land plants and mammals (Vandenbroucke et al., 2008). The accumulation and oxidation of glutathione in response to H$_2$O$_2$ is commonly observed in most organisms (Vandenbroucke et al., 2008; Noctor et al., 2012). The first step in glutathione synthesis, catalyzed by γ-glutamylcysteine synthetase, is the rate-limiting step (Masip et al., 2006; Noctor et al., 2012; Lu, 2013). The increased expression of GSH1 in response to H$_2$O$_2$ in Chlamydomonas may therefore account for the observed accumulation of glutathione. A similar increased expression of this gene was reported in yeast and mammals under diverse stress conditions (Wu and Moye-Rowley, 1994; Lu, 2013). By contrast, in Arabidopsis H$_2$O$_2$ does not induce expression of either GSH1 or GSH2 but glutathione accumulation appears to involve the induction of genes encoding enzymes involved in cysteine synthesis in the chloroplast (Queval et al., 2009). Interestingly, several of these genes were also induced in Chlamydomonas after H$_2$O$_2$ addition. All of these results suggest that the mechanisms of redox signaling and detoxification of Chlamydomonas are unique since some regulations are shared with higher eukaryotes while others are related to other unicellular organisms. The specificities of Chlamydomonas redox metabolism and its responses to hydrogen peroxide may also be linked to the presence, in as mammals, of selenoproteins including glutathione peroxidases, thioredoxin reductases and methionine sulfoxide reductases. Moreover, the site of H$_2$O$_2$ accumulation also affects the transcriptional response, as recently shown in Arabidopsis (Sewelam et al., 2014), and exogenous H$_2$O$_2$ may not influence the levels of ROS within some compartments significantly. However, the addition of H$_2$O$_2$ was clearly able to elicit a generalized reduction in transcripts corresponding to nucleus-encoded photosynthetic complexes and significant remodeling of transcripts related to the endomembrane system. Alternatively, the abundance of catalase transcripts may not be directly related to catalase activity considering the hypothesized role of these proteins in gating H$_2$O$_2$ signaling, and the decrease in catalase activity upon addition of H$_2$O$_2$, most probably linked to ROS-dependent post-translational regulation (Shao et al., 2008; Michelet et al., 2013).

We noted that a large proportion of transcripts encoding proteins involved in protein folding and degradation increased coordinately with a decrease of transcripts encoding proteins involved in protein synthesis. These trends suggest that the stress caused by addition of H$_2$O$_2$ signals a general reduction in cellular activity during the period of exposure, and suggests the potential for global protein remodeling and/or clearance of oxidized proteins following H$_2$O$_2$ stress. One possible conclusion is that these systems are induced to clear oxidized proteins with the simultaneous induction of protein degradation and protein folding while protein synthesis is downregulated. Another possibility is that the induction of proteasome transcripts is a result of feedback, since the proteasome and the ubiquitin-activating/conjugating system have been shown to be inhibited by H$_2$O$_2$ (Davies, 2001) and by glutathionylation (Demasi et al., 2014). The observed induction in protease transcripts may be responsible for the degradation of oxidatively damaged proteins in compartments and/or a means to compensate for reduced activity of the proteasome.

Moderate reductions in transcript abundance of CB cycle and carbon metabolism genes are consistent with previous reports that algae experience reductions in CO$_2$ fixation in response to H$_2$O$_2$ exposure (Takeda et al., 1995). Reduced abundances of RNAs encoding subunits of the photochemistry machinery were supported physiologically by demonstrating reduced O$_2$ output subsequent to H$_2$O$_2$ exposure. The general reduction observed in transcript level of the photosynthetic machinery is correlated with reduced photosynthetic efficiency at 1 h. Reduction of transcript abundance of photosynthesis-related genes may serve to preserve redox balance and prevent production of additional ROS from photosynthetic electron transport.

Like most plant genomes, Chlamydomonas has many gene duplications with paralogs encoding multiple proteins of the same family (Merchant et al., 2007; Wu et al., 2015). As with many other biological processes, the genes encoding enzymes with roles in ROS detoxification are present in up to five or six copies. Unsurprisingly, transcript levels of known ROS-detoxification systems were generally increased, although this pattern of expression was not uniform across all associated paralogs, with transcript abundance corresponding to one gene copy far exceeding that of other copies. For example, expression estimates for GST1 and GST2 exceed 700 FPKM, compared with a GST3 peak of 70. Similar patterns were observed for many paralogous genes. While mRNA abundance does not necessarily translate to protein abundance, and therefore metabolite flux, it does suggest the possibility that some copies have more prominent roles than others.

The importance of understanding ROS-induced damage responses and protective mechanisms has resulted in gene expression studies on H$_2$O$_2$ exposure in numerous organisms (e.g. Chuang et al., 2002; Chen et al., 2003; Girardot et al., 2004; Mostertz et al., 2004), including several photosynthetic organisms (Desikan et al., 2000; Vandenabeele et al., 2003; Kobayashi et al., 2004; Li et al., 2004). In a recent meta-analysis, a number of conserved gene families was found to be induced in response to H$_2$O$_2$ across lineages.
and kingdoms (Vandenbroucke et al., 2008). Our data are largely in agreement with that study: the most highly conserved induced genes were DNAJ-like HSFs, which also increased in mRNA level in our dataset. Three other protein families were found to be induced in all eukaryotes investigated (A. thaliana, S. cerevisiae, Schizosaccharomyces pombe and Homo sapiens), including GTP-binding proteins, protein kinases and ubiquitin-conjugating enzymes. In our dataset, genes encoding each of these functions were differentially expressed. However, in our transcriptomes, mRNA abundances for genes encoding both GTP-binding proteins and several protein kinases decreased over the time course, suggesting that the direction of altered expression is not universal. Vandenbroucke et al. also observed a conserved induction of short-chain dehydrogenases/reductases and peroxiredoxins in all unicellular organisms in their study (Synechocystis, S. cerevisiae and S. pombe). Our data in Chlamydomonas, which is also unicellular, partially support this observation, where transcripts encoding proteins broadly annotated as dehydrogenases/reductases were induced several fold between 0' and 1 h.

We also compared our H2O2 transcriptomes with other relevant Chlamydomonas RNA-seq experiments. A recently published O2-response transcriptome allowed us to make a global comparison of these two oxidative stress conditions. Clearly the addition of H2O2 and RB elicit independent responses, but we did find a significant overlap highlighting a conserved cluster of stress-related genes. In particular, both ER stress and lipid stress ranked highly among the overlapping transcripts. Indeed, the Cre01.g004900 transcript level rose in response to both ROS (~50-fold between 0' and 1 h, peaking at 380 FPKM and three-fold in response to RB) and bears sequence similarity to yeast HSP31, encoding a methylglyoxylase, which is also induced in response to oxidative stress (Dubacq et al., 2006). This induction may serve to reduce build-up of methylglyoxal resulting from lipid peroxidation during both stresses.

Interestingly, the overlap between H2O2 and RB treatment of the SAK1 mutant was greater than RB treatment of the wild type. SAK1 encodes a key regulator of the RB response, and O2' exposure of the SAK1 mutant appears to generate additional oxidative stress that was not observed in the wild type. One possibility is that mis-regulation of SAK1-target transcripts in the presence of RB either exacerbates or leads to a type of oxidative cellular damage in common with H2O2. In particular we noticed a substantial increase in the number of transcripts encoding proteins putatively involved in protein turnover in the set of transcripts shared between sak1 and H2O2 (18 transcripts) versus those shared among sak1, wild type and H2O2 (seven transcripts), suggesting that protein damage could be the point at which the H2O2 and sak1 signaling intersect.

We also observed a large overlap between H2O2-responsive transcripts and both ClpP-depletion and rapamycin addition. ClpP is a stromal protease that is thought to play an essential function in the chloroplast (Nishimura and van Wijk, 2015), while rapamycin inhibits TOR signaling, a pathway that promotes cell growth and inhibits autophagy (Dennis et al., 1999). These two transcriptomes were originally performed in conjunction to identify transcriptional changes in the ClpP-depleted strain that are independent of its autophagy-like phenotype (as rapamycin treatment induces autophagy). In this study, we leveraged these datasets to reveal a potential contribution of oxidative stress to the observed transcriptional changes. Indeed, we identified roughly 197 transcripts that increase in abundance following H2O2 exposure and ClpP depletion, while nearly 144 transcripts increase in abundance following H2O2 exposure and rapamycin treatment. The increase of 94 transcripts was shared between all three datasets. At this point we can only speculate as to whether the significant overlap we observed reflects the ability of ClpP depletion and rapamycin treatment to elicit an oxidative stress response. For instance an obvious physiological intersect is induction of autophagy by the three conditions.

In the diurnal cycle, we found a significant number of H2O2-induced transcripts with peak transcript abundance after the transition from dark to light and as cells age during the day. Enrichment within the light stress cluster at the beginning of the day and towards the end of the day was shared with singlet O2'-induced transcripts, but the presence of H2O2'-induced transcripts was associated with a lower P-value. Chlamydomonas cells experience an extended G1 phase during the day, during which biomass is accumulated. Cell division is arrested until the night. Based on functional classification of the corresponding proteins, the H2O2-induced transcripts with maximum abundance at the end of day largely encode proteins involved in protein turnover. Unexpectedly, we also observed a spike in ROS-induced transcripts at 2 h prior to the transition to day, possibly highlighting clock-regulated transcripts, which are maximally abundant in anticipation of the transition to the light period. Although there was a significant enrichment of H2O2-induced transcripts with peak abundance at this time point, we found a higher significance associated with RB-induced transcripts, possibly highlighting anticipation of photo-oxidative damage associated with the shock of switching from dark to light. The abundance of the Chlamydomonas ortholog of Arabidopsis CCA1 (Cre06.g275350; circadian clock-associated 1), demonstrated to regulate the induction of oxidative stress responses in that organism (Lai et al., 2012), is slightly reduced throughout our time course. In the Chlamydomonas diurnal experiment peak abundance occurs immediately prior to the light period, suggesting that Cre06.g275350 is regulated by circadian rhythms here also, and is not influenced by the presence of ROS.
EXPERIMENTAL PROCEDURES

Strains and culture conditions

Strain CC-4532 (Mets strain 2137 mt−1) was used throughout this study. Routine growth was performed in 2-amino-2-(hydroxymethyl)-1,3-propanediol (TRIS)-acetate-phosphate (TAP) using Hutner’s trace element mix (Hutner et al., 1950). Cultures were grown in Innova incubators (New Brunswick Scientific, http://www.eyepandor.com) at 24°C, agitated at 180 rpm with continuous light (95 μmol m−2 sec−1), six cool white fluorescent bulbs at 4100K and three warm white fluorescent bulbs at 3000K per incubator. Unless indicated otherwise, freshly aliquotted H2O2 (Fisher, https://www.fishersci.com) was added to cultures to a final concentration of 1 mM at about 2 × 106 cells ml−1. Culture densities (in cells ml−1) were determined with a hemocytometer.

H2O2 RNA-seq read realignment and analysis

Transcriptomic data analyzed in this study were initially acquired as described in Urzica et al. (2012a). The sequence reads resulting from this previous study were aligned to v5.5 of the reference genome (Merchant et al., 2007) (available at ftp://ftp.jgi-psf.org/pub/compgen/phytozome/v9.0/Creinhardtii/) using STAR (Dobin et al., 2013), default parameters plus –alignIntronMax 10000 and expression estimates were determined and normalized in terms of FPKM using cuffdiff (Trapnell et al., 2010) default parameters.

Fv/Fm measurements

Fv/Fm measurements were determined using a FluorCam 800MF (Photon Systems Instruments, http://www.psi.cz/). Cells (3 × 106) were sampled from cultures and concentrated onto a Whatman 25-mm circular filter paper disk using a vacuum. Cell-containing filter disks were dark-adapted for 15 min before performing Fv/Fm measurements using FluorCam6.0 software (Photon Systems Instruments).

A2 O2 degradation (FOX) assays

The concentration of H2O2 in culture flasks was determined by FOX assays, as described by Nourooz-Zadeh (1999). Briefly 0.1-ml culture samples were taken from cultures at a cell density of 2 × 106 cells ml−1, or non-inoculated flasks for cell-free control samples. Samples containing cells were centrifuged (16 100 g for 3 min) and the supernatant transferred to fresh tubes. Then 0.9 ml of FOX assay reagent A (25 mM 93% sulfuric acid, 2.5 mM ammonium iron sulfate, 1 mM xylolol orange) was added, the mixture was incubated at room temperature 24°C for 10 min, and the absorption determined at 560 nm. H2O2 concentrations were determined by reading absorption measurements off a standard curve prepared with known concentrations. Measurements were performed on three separate cultures (biological replicates). Measurements were taken at 0 h immediately after addition of 1 mM A2 O2, and at 0.5, 1, 2, 3 and 4 h thereafter.

Oxygen consumption and evolution

Oxygen evolution and consumption rates were measured on a standard Clark-type electrode (Hansatech Oxygraph with a DW-1 chamber, http://hansatech-instruments.com/) and the data analyzed with the Hansatech OxyLab software v.1.15. Experiments were performed using 2 ml of culture at 2 × 106 cells ml−1 in the presence of 20 mM acetate and 10 mM KHCO3. Samples were taken 0.5 h prior to addition of H2O2 (added to a final concentration of 1 mM), immediately after addition (t0), and at 0.5, 1, 2 and 4 h subsequent to addition of H2O2. Measurements were performed on three separate cultures (biological repeats). The respiration rate was measured as oxygen consumption over a period of 4 min in the dark. The rate of photosynthetic O2 evolution was measured for 4 min in the light (75 μmol m−2 sec−1) after a 4-min acclimation period and was calculated as the difference between oxygen evolution in the light and oxygen consumption in the dark.

Glutathione pool measurements

Chlamydomonas cells treated with 1 mM H2O2 were collected at 0, 0.5, 1, 3 and 4 h by centrifugation (5000 g for 5 min), washed once in 50 mM sodium phosphate (pH 7.5) solution, resuspended in 0.2 N HCl and lysed by two cycles of freeze/thaw at −80°C. Crude extracts were cleared by centrifugation at 15 000 g for 20 min at 4°C. Then 50 μl of sample was neutralized by adding 50 μl of 50 mM NaH2PO4 (pH 7.5) and 0.2 N NaOH to a final pH of between 5 and 6. The neutralized sample was used directly for measuring total glutathione (GSH plus GSSG) by the recycling assay initially described by Tietze (1969) and adapted from Queval and Noctor (2007). The method relies on the glutathione reductase (GR)-dependent reduction of 5,5’-dithiobis(2-nitro-benzoic acid) (DTNB; Sigma, D8130, http://www.sigmaaldrich.com/). Oxidized glutathione was measured after treatment of a neutralized sample with 10 mM 4-vinylpyridine (VPD; Sigma V3240-4) for 30 min at 25°C. To remove excess VPD, the derivatized sample was centrifuged twice at 15 000 g for 20 min at 4°C. To measure total glutathione or GSSG, the sample was added to a mix containing 120 μM NaH2PO4 (pH 7.5), 300 μM DTNB, 500 μM NADPH, 1 mM EDTA (pH 8), 1 μM GR (Sigma, G3664), and DTNB reduction was measured at 412 nm. Different GSH (Sigma, G4251) concentrations ranging from 0 to 5 μM were used as standards. For each time point, three independent biological replicates were measured with at least three technical replicates for each sample. Data are represented as mean ± SD (n ≥ 9).

Methods for comparative transcriptome analysis

The same cutoffs as those used for the H2O2 analysis (i.e. ≥ 10 FPKM/RPKM, ≥ 2.0-fold change) were applied to the expression estimates of previously published datasets (Castruita et al., 2011; Urzica et al., 2012b; Hem schemeyer et al., 2013; Malasarn et al., 2013; Ramundo et al., 2014; Wakao et al., 2014). The list of upregulated transcripts from each dataset was then compared with the list of transcripts that increased in abundance before and 1 h after H2O2 addition (referred to as H2O2-responsive). For the ClpP-depletion H2O2 overlap, our dataset was composed of transcripts that met our cutoffs and increased in strain DCH16 after 43 h of vitamin addition. For the rapamycin analysis, our dataset was composed of transcripts that met our cutoffs and increased in A31 (the wild-type strain in that work) after 8 h of rapamycin addition. The P-value for each overlap was calculated using R, with the command sum(dhperiper(qm, k, 17 301-k, m)), where q is the number of transcripts in the overlap, m is the number of transcripts that increased in abundance following addition of H2O2 and k is the number of transcripts that increased in abundance in the dataset being compared. Because the datasets we used to perform these comparisons were all aligned to the v.4 assembly of the Chlamydomonas genome, we converted loci IDs in the H2O2 data set from v.5 to v.4 (which contains 17 301 loci) using the correspondence table available at http://genome.jgi.doe.gov/pages/dynamic/OrganismDownload.jsf?organism=Phytozoonv10. For each transcript in the diurnal cycle, we determined at which time point the maximum FPKM value is reached. Transcripts that did

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not reach 10 FPKM at any time point were excluded. We then determined the number of H$_2$O$_2$-responsive transcripts in each list of maximum FPKMs and calculated the $P$-value associated with the overlap as above.

**ACCESSION NUMBERS**

Sequenced reads are available on the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) database under accession number GSE34826.

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**SUPPORTING INFORMATION**

Additional Supporting Information may be found in the online version of this article.

**Figure S1.** Gene Ontology analysis identifies global changes in differentially expressed genes.

**Table S1.** Expression estimates for all genes.

**Table S2.** Genes whose mRNA abundance significantly changes in response to H$_2$O$_2$.

**Table S3.** GreenCut genes whose mRNA abundance significantly changes in response to H$_2$O$_2$.

**Table S4.** Messenger RNA abundances for genes encoding enzymes relating to detoxification of reactive oxygen species.

**Table S5.** Messenger RNA abundances for genes encoding enzymes relating to photosynthesis.

**Table S6.** Messenger RNA abundances for genes encoding enzymes involved in respiration.

**Table S7.** Messenger RNA abundances for genes encoding enzymes relating to carbon metabolism.

**Table S8.** Overlap of differentially expressed genes between H$_2$O$_2$ rose bengal and anoxia experiments with a *Chlamydomonas reinhardtii* diurnal cell cycle transcriptome.

**REFERENCES**


