

Insights into the Survival of *Chlamydomonas reinhardtii* during Sulfur Starvation Based on Microarray Analysis of Gene Expression†‡

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Responses of photosynthetic organisms to sulfur starvation include (i) increasing the capacity of the cell for transporting and/or assimilating exogenous sulfate, (ii) restructuring cellular features to conserve sulfur resources, and (iii) modulating metabolic processes and rates of cell growth and division. We used microarray analyses to obtain a genome-level view of changes in mRNA abundances in the green alga *Chlamydomonas reinhardtii* during sulfur starvation. The work confirms and extends upon previous findings showing that sulfur deprivation elicits changes in levels of transcripts for proteins that help scavenge sulfate and economize on the use of sulfur resources. Changes in levels of transcripts encoding members of the light-harvesting polypeptide family, such as LhcSR2, suggest restructuring of the photosynthetic apparatus during sulfur deprivation. There are also significant changes in levels of transcripts encoding enzymes involved in metabolic processes (e.g., carbon metabolism), intracellular proteolysis, and the amelioration of oxidative damage; a marked and sustained increase in mRNAs for a putative vanadium chloroperoxidase and a peroxiredoxin may help prolong survival of *C. reinhardtii* during sulfur deprivation. Furthermore, many of the sulfur stress-regulated transcripts (encoding polypeptides associated with sulfate uptake and assimilation, oxidative stress, and photosynthetic function) are not properly regulated in the *sac1* mutant of *C. reinhardtii*, a strain that dies much more rapidly than parental cells during sulfur deprivation. Interestingly, sulfur stress elicits dramatic changes in levels of transcripts encoding putative chloroplast-localized chaperones in the *sac1* mutant but not in the parental strain. These results suggest various strategies used by photosynthetic organisms during acclimation to nutrient-limited growth.

Sulfur (S) is an essential element present in proteins, lipids, and various metabolites. It is critical for the association of metal ions to proteins (electron carriers and redox controllers) and is a component of metabolites that function in photoprotection (14, 29) and signal transduction (such as in symbiosis) (45). Because most organisms have limited S storage, their growth and development is dependent upon a continuous supply of this nutrient from the environment. The majority of accessible S in soil solutions is in the form of the SO_4^{2-} anion. However, in some cases the majority of soil SO_4^{2-} may not be readily available to the biota, since the SO_4^{2-} anion is often complexed with cations as insoluble salts that are tightly adsorbed onto the surface of soil particles or exists as a soluble anion that rapidly leaches through the soil matrix. Furthermore, a large proportion of soil SO_4^{2-} may be covalently bonded to organic molecules in the form of sulfate esters and sulfonates.

The acquisition of SO_4^{2-} by plants and microorganisms is facilitated by specific transport systems. Following uptake, the anion is either used for the direct sulfation of compounds or is reduced and converted to cysteine and methionine, which is

incorporated into both proteins and specific cellular metabolites (1, 25, 41, 49). Cysteine is also an important building block for the synthesis of both glutathione and phytochelatin. The former serves as a source of reductant for many physiological processes and as an antioxidant that enables cells to withstand oxidative stress (34, 35), while the latter helps combat heavy metal toxicity (16, 63).

S can be limiting in the environment and strongly influence ecosystem composition. It may also limit plant productivity in certain agricultural settings (27, 28, 57), which can result in reduced quality and yields of seeds, and upon severe S limitation, the growth of the plant may be stunted. We have used the unicellular alga *Chlamydomonas reinhardtii* to identify and elucidate S limitation responses. *C. reinhardtii* synthesizes a prominent, extracellular arylsulfatase (ARS) in response to S limitation (26, 44). An ARS associated with the proteinaceous cell wall of the alga has been purified to homogeneity and characterized (12). This ARS polypeptide has at least three O-linked oligosaccharides, is very stable in the extracellular space (stability may be conferred on the polypeptide by oligosaccharide decorations), and is synthesized as a preprotein with a signal sequence that is cleaved as it is exported from the cell (11). The extracellular location of ARS allows it to hydrolyze soluble SO_4^{2-} esters in the medium, releasing free SO_4^{2-} for assimilation by the cells. Two ARS genes arranged in tandem, but in the opposite orientation on the *C. reinhardtii* genome, have been described (11, 40), but the gene family may contain a number of additional members.

There are several transcripts, in addition to those encoding

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ARS, that are synthesized by *C. reinhardtii* in response to S deprivation. Some of these transcripts encode enzymes involved in the uptake and assimilation of SO_4^{2-} (62). Levels of mRNAs for enzymes of the SO_4^{2-} assimilation pathway in plants also increase during S starvation (15, 42, 47, 52). S starvation of *C. reinhardtii* causes a dramatic increase in the V_{\max} and a decrease in the K_m for SO_4^{2-} transport (61). Eukaryotic organisms have multiple systems for SO_4^{2-} uptake, and generally high-affinity SO_4^{2-} uptake increases when they are deprived of S (3, 4, 17, 24, 61). Recently, a *C. reinhardtii* gene encoding a putative SO_4^{2-} transporter was isolated and appears to be activated during S-limited growth (J. Davies, personal communication), but genes encoding potential SO_4^{2-} transporters (designated *SULTR1*, *SULTR2*, and *SULTR3*) have also been identified in cDNA and genomic databases (<http://genome.jgi-psf.org/chlre2/chlre2.home.html>). The subcellular locations of these putative protein products have not been determined. Furthermore, in *C. reinhardtii* there are prominent extracellular polypeptides of apparent molecular masses of 76 kDa (designated extracellular polypeptide 76, or ECP76) and 84 kDa (ECP84) that have been shown to be synthesized in response to S deprivation (51). The deduced ECP76 and ECP84 polypeptide sequences have significant similarity to those of various cell wall proteins but contain either no or few S-containing amino acids (between mature ECP76 and ECP84 there is only a single S-containing amino acid). These results suggest that a highly regulated process tailors the protein-rich cell wall of *C. reinhardtii* for S deprivation.

There are a number of factors that control the assimilation of SO_4^{2-} and the regulation of genes involved in that assimilation. *O*-acetylserine, the carbon backbone used for the synthesis of cysteine, functions as a positive effector for the transcription of genes encoding enzymes that participate in the uptake and assimilation of SO_4^{2-} (22) in both bacteria and vascular plants (20, 50). Increases in *O*-acetylserine levels appear to occur when *Arabidopsis thaliana* plants are starved for S or provided with excess N (19). In *C. reinhardtii*, specific regulatory elements, including SAC1 and SAC3, control S deprivation responses (8–10). A number of physiological responses of *C. reinhardtii* to S limitation require the SAC1 protein (8). The *sac1* mutant of *C. reinhardtii* exhibits abnormal SO_4^{2-} uptake and is unable to synthesize extracellular ARS, as well as other extracellular proteins, in response to S deprivation. Furthermore, there was little or no induction of *ECP76*, *ECP84*, and *ATSI* (encoding ATP sulfurylase, the enzyme that activates free SO_4^{2-} , allowing reduction to occur) genes in a *sac1* null mutant background (51, 62). The responses of the *sac1* mutant to P and N limitation appear normal. The inability of the *sac1* strain to properly respond to S limitation is also reflected in a rapid decline in the viability of mutant cells following exposure to S deprivation (8); the decreased viability is sensitive to environmental light levels and the activity of photosynthetic electron transport. Since the *sac1* mutant cannot properly control photosynthetic activity during S deprivation, it may accumulate reactive oxygen species, which could cause extensive cellular damage, and the electron transport chain would become hyperreduced (e.g., a highly reduced plastoquinone pool), which could have adverse effects on the control of metabolic processes.

Interestingly, the *SAC1* gene product is predicted to be a

polypeptide with similarity to anion transporters (reference 9 and our unpublished work). The deduced SAC1 polypeptide sequence and phenotype of the *sac1* mutant have some similarities with the Snf3 polypeptide of yeast and the phenotype of the *snf3* mutant, respectively. Snf3 is a yeast transporter-like regulatory protein that governs expression of genes involved in hexose utilization (37, 38). The putative regulatory functions of SAC1 and Snf3 raise the possibility that polypeptides that originally functioned in binding and transporting various substrates into cells may have evolved into regulatory elements.

In this study we used microarray analyses to obtain a genome-based picture of *C. reinhardtii* responses to S limitation and to determine how the *sac1* lesion alters S deprivation-triggered changes in gene expression. The results presented are discussed in the context of physiological changes that may be elicited by S limitation and the regulatory circuits that may operate during S starvation.

MATERIALS AND METHODS

Cell culture. The *C. reinhardtii* arginine auxotroph CC425 and the *sac1* mutant (carrying *ars5-5*) (8) were cultured in S-replete, Tris-acetate-phosphate (TAP) medium to mid-logarithmic phase on a rotating platform (120 rpm) and continuously illuminated ($80 \mu\text{mol of photon m}^{-2} \text{s}^{-1}$) at 25°C. To assay global changes in transcript levels in response to S deprivation, the cells were harvested at room temperature ($5,000 \times g$, 5 min), washed twice with TAP medium without S, and resuspended in TAP medium without S to 5×10^6 to 10×10^6 cells per ml. The cultures were maintained on a rotating platform, illuminated with $80 \mu\text{mol of photon m}^{-2} \text{s}^{-1}$ for 2, 4, 8, 12, and 24 h, and then rapidly cooled in liquid N_2 and harvested by centrifugation ($5,000 \times g$, 5 min, 4°C).

RNA preparation. Cell pellets were resuspended in 3 ml of extraction buffer (50 mM Tris, 300 mM NaCl, 0.5 mM EDTA, 2% sodium dodecyl sulfate [SDS], pH 8.0, that had been treated with 0.1% diethyl pyrocarbonate), followed by the addition of 5 μl of proteinase K (40 $\mu\text{g/ml}$). The suspension was gently agitated for 30 min at room temperature and then extracted with an equal volume of phenol-chloroform (1:1) until the aqueous phase became clear. Nucleic acid in the aqueous phase was precipitated with 2 volumes of ethanol, dried, and dissolved in diethyl pyrocarbonate-treated double-distilled H_2O (ddH_2O). RNA was purified by adding an equal volume of 4 M LiCl to the nucleic acid solution, incubating the solution on ice for 8 h, and collecting the precipitated RNA by centrifugation at $10,000 \times g$ for 15 min. Two RNA preparations for each time point were generated from independent experiments. RNA samples were further purified with the RNeasy MiniKit (QIAGEN, Valencia, Calif.).

Fluorescent labeling of probe. Total RNA served as the template for labeling cDNA by direct incorporation of Cy3-dUTP or Cy5-dUTP (Amersham Pharmacia Biotech, Piscataway, N.J.) by using reverse transcription. RNA (10 μg total) was denatured at 70°C for 10 min in the presence of 1 μg of Oligo(dT)12-18 (Invitrogen, Carlsbad, Calif.) in a reaction volume of 23 μl . The reaction was cooled on ice prior to the addition of 8 μl of $5 \times$ Superscript II reverse transcriptase buffer (Invitrogen), 1 μl of 0.1 M dithiothreitol, 4 μl of deoxynucleoside triphosphates (10 mM dTTP, 25 mM dATP, 25 mM dGTP, and 25 mM dCTP), 2 μl of Superscript II reverse transcriptase (200 U/ μl), and 2 μl of 1 mM Cy3-dUTP or Cy5-dUTP. The reaction was incubated at 42°C in a thermocycler (MJ Research, Waltham, Mass.) for 2 h and then stopped by the addition of 2 μl of 500 mM EDTA and 2 μl of 500 mM NaOH. The alkalinized solution was incubated at 70°C for 10 min to degrade RNA prior to neutralization by the addition of 2 μl of 500 mM HCl. The Cy5-dUTP-labeled probe from each time point following the initiation of starvation was combined with the Cy3-dUTP-labeled control RNA (zero time; unstarved cells), purified with a QiaQuick PCR purification kit (QIAGEN), and recovered in 17 μl of ddH_2O . Incorporation of the fluorescent dyes into cDNA was visualized with a Typhoon 8600 (following electrophoresis of the cDNA on a 0.8% agarose gel). It should be emphasized that the reference RNA was isolated from our parental strain (CC425) prior to S deprivation (0 h). Samples prepared from S-deprived (2, 4, 8, 12, and 24 h) CC425 and the unstarved (0 h) and starved (2, 4, 8, 12, and 24 h) *sac1* mutant were compared to this sample. The dye labeling was reversed in a separate experiment, and the experiments were duplicated with independently isolated RNA samples.

Microarray preparation. *C. reinhardtii* cDNA clones from the Core (TAP light, TAP dark, HS+CO₂, and HS) and Stress I Libraries (NO₃⁻ to NH₄⁺ [30 min, 1 h, 4 h], NH₄⁺ to NO₃⁻ [30 min, 1 h, 4 h], TAP without N [30 min, 1 h, 4 h], TAP without S [30 min, 1 h, 4 h], and TAP without P [4 h, 12 h, 24 h]) were sequenced (48), and the sequences generated from the 3' ends were assembled into 2,761 unique contigs that were used to construct a microarray. Further details of the "2.7-k array" are given at the website <http://nostoc.stanford.edu/jeff/lab/chlamyarray/index.html> and in both Table 1 and Table S1 in the supplemental material. The 3' sequences were amplified with a universal primer (forward primers: 631u24, 5'-CGACTCACTATAGGGCGAATTGGG-3', 1 bp longer than the T7 primer; or M13-21, 5'-TGTAACGACGGCCAGT-3') and a specific reverse primer designed to anneal to the first strand of each clone in a region 200 to 500 bp away from 3' end of the transcript. PCRs were performed in a 96-well format according to the protocol 94°C for 2.5 min, 94°C for 30 s, 62°C or 50°C for 30 s, and 72°C for 1 min for 40 cycles. The quality and specificity of the amplification products were determined by electrophoresis on a 1% agarose gel followed by visualization of nucleic acid by ethidium bromide staining; 96.6% of the clones gave a single product of the expected size. The PCR products were cleaned by ethanol precipitation and dissolved in 40 µl of 3× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate), and 5 µl was transferred from the 96-well microtiter plates to printing plates, which were in the 384-well microtiter format (MJ Research). Printing plates were wrapped with moistened paper towels, sealed with Saran Wrap, and stored at -20°C until they were used. PCR products were arrayed from the 384-well microtiter plates onto CMT-gamma amino propyl silane slides coated with γ amino propyl silane (Corning Microarray Technology, Corning, N.Y.) by using an Omnigridd microarrayer (GeneMachines, San Carlos, Calif.) with ChipMaker 2 pins (TeleChem International, Sunnyvale, Calif.). Array spots were ~150 µm in diameter, and the center-to-center distance was 212.5 µm. Four complete cDNA sets were printed onto each slide, with each set consisting of eight subsets of 349 spots each (18 rows by 20 columns). Printed slides were baked at 80°C for 2 h and cross-linked by UV irradiation in the Stratalinker 1800 (Stratagene, La Jolla, Calif.) at a total power of 300 mJ. The arrays were blocked by incubating them for 15 min in a succinic anhydride-NaBO₄⁻ solution. This solution was freshly prepared by dissolving 5 g of succinic anhydride (Sigma, St. Louis, Mo.) in 315 ml of *N*-methylpyrrolidinone and adding 35 ml of 0.2 M NaBO₄⁻ (boric acid dissolved in ddH₂O and made pH 8.0 with NaOH). The arrays were then soaked for 2 min with gentle agitation in 95°C ddH₂O, immersed in 95% ethanol (high-performance liquid chromatography grade) for 1 min, and dried by centrifugation for 5 min in a SpeedVac Plus model SC210A (Savant, Holbrook, N.Y.). The arrays were stored in a desiccator at room temperature until they were used.

Hybridization, washing, and scanning. Arrays were prehybridized in 3× SSC-0.1% SDS-10-mg/ml bovine serum albumin for 20 min at 50°C, followed by immersion of the slides into ddH₂O and then isopropanol for 2 min each. Slides were dried by centrifugation for 5 min in a SpeedVac. The hybridization solution was prepared by adding 15 µl of 2× hybridization buffer [6× SSC, 0.2% SDS, 0.4-µg/µl poly(dA), 0.4-µg/µl tRNA] to 15 µl of the labeled probes, followed by passage of the probe through a QiaQuick PCR purification column. The probe solution was incubated at 95°C for 3 min, cooled to room temperature, applied to a prehybridized microarray, covered with a LifterSlip (Erie Scientific Company, Portsmouth, N.H.), placed in a hybridization chamber, and hybridized at 50°C for 16 h. Following hybridization, the slides were washed for 5 min each in successive solutions A (2× SSC, 0.1% SDS), B (1× SSC), and C (0.05× SSC) at room temperature and dried by centrifugation in a SpeedVac Plus model SC210A (Savant) for 5 min. The arrays were scanned at 532 and 635 nm in an Axon GenePix 4000A scanner (Foster City, Calif.) at 10-µm resolution. Photomultiplier tube voltages were adjusted to minimize background and saturation of the hybridization signals. Images of the fluorescence at 532 nm for Cy3 and 635 nm for Cy5 were recorded and analyzed for eight complete sets of all cDNAs (four sets on each of two slides). RNA samples used to synthesize the probes that were hybridized to the slides were from independent experiments.

Data selection and analysis. Microarray images representing spot intensities from scanned slides were imported into the GenePix Pro 3.0 program (Axon Instruments) and quantified. Spot positions were defined according to a pre-defined microarray layout that was subsequently adjusted by eye to help optimize spot recognition. Spot signals that were distorted by dust or locally high backgrounds were not included in the analyses. The data were imported into the Stanford Microarray Database (13, 46) and normalized by using that database's standard computed algorithm (for details, see <http://genome-www.stanford.edu/microarray/>). The experiment identification numbers within the database are 31781 to 31790, 37207, 37266 to 37269, 37271 to 37276, and 37356.

Only those spots with an intensity/background ratio of >2 in either channel, and with a minimum normalized net intensity for the median (channel 1 net

median intensity or normalized channel 2 net median intensities) of >350 in at least one of the two channels, were included in the analysis. Log₂ of the 635 nm/532 nm normalized ratio for the median was retrieved. Genes for which the transcripts appeared to increase or decrease by ≥3-fold (averaged ratio per array over the duplicate samples) during S deprivation, for ≥4 of the 8 array sets analyzed, and for at least two of the time points following the imposition of S deprivation were chosen for further analysis. Gene annotations were retrieved from the Unigene Set stored in the Chlamydomonas EST database (http://www.biology.duke.edu/chlamy_genome/unigene.html) and further confirmed manually by BLAST analyses by using the Chlamydomonas Genome Sequence Database (<http://genome.jgi-psf.org/chlr2/chlr2.home.html>), the predicted gene models within the database, and NCBI protein databases. All alignments were visually inspected.

At each time point, for each gene, a one-way *t* test was conducted to assess the significance of the relative transcript abundance levels. Each *t* test considered the unaveraged (but normalized) log₂ data for each gene. The following *t*-score threshold levels were set: for α = 0.05, df(1), *t* = 12.6; and for α = 0.10, df(7), *t* = 2.8. The 2.8 *t* value represents the least stringent significance measure, where each sample is considered to be independent of all others. The 12.6 *t* value represents the most stringent significance measure, where the test is considered to have only two independent samples. The + and - patterns displayed in Table 1 depict the least stringent threshold (*t* = 2.8), although both are reported in the larger Table S1 in the supplemental material.

A significance analysis of microarrays (SAM) (39, 55) was also used to correct for false discovery rates. All eight replicates (two biological replicates by four on-array replicates) at each time point were analyzed separately, and the median false discovery rate was set as close to 1 as possible in order to compare results between time points. The SAM results, which are reported in Tables S3 and S4 in the supplemental material, were largely congruent with the results reported in Table 1, although SAM selected fewer genes.

RESULTS

Modulation of transcript abundance in response to S deprivation. Among the 2,761 cDNA fragments included on the microarray, 2,565 were amplified as 3' fragments of cDNAs and had the expected sizes, while the characteristics of the remaining 196 PCR products were not acceptable (no band, multiple bands, or smearing along the lane) and were excluded from analysis. The number of genes represented on the array is probably about one-sixth of those present on the *C. reinhardtii* genome and may be biased toward highly expressed genes, as the microarray was designed from expressed sequence tags (ESTs) selected at an early stage of the EST sequencing project (48). Since the cDNA library from S-starved cells was among the first to be sequenced, there is also presumably some bias toward genes whose transcripts increase during S starvation. The RNA used for synthesizing labeled cDNA to probe the array was isolated from cells at five different times (2, 4, 8, 12, and 24 h) following the imposition of S deprivation. We analyzed both the parental strain, CC425 (Chlamydomonas Genetics Center), and the *sacI* mutant (carrying *ars5-5*; equivalent to CC3794; this strain was generated by mutagenesis of CC425). The reference for all time points for the parental cells and the *sacI* mutant (including the *sacI* zero time) was RNA from the parental strain isolated at time zero (after pelleting the cells by centrifugation and just as SO₄²⁻ was eliminated from the medium).

There were 171 cDNAs for CC425 and 132 for the *sacI* mutant for which transcripts exhibited a ≥3-fold alteration (ratios were either ≥3 or ≤0.33) for at least 4 of the 8 array sets analyzed and two of the five time points sampled following the imposition of S deprivation (Fig. 1, left panel). Forty-seven of the selected clones were common to CC425 and the *sacI* mutant. If the cutoff was set to select genes for which transcript

TABLE 1. Transcripts showing a ≥ 3 -fold change in abundance during sulfur deprivation^a

Clone (Fig. 3 panel) ^b	ACE ^c	Name	Gene annotation	Direction of change in transcript level		Transcript level (fold increase) for											
				CC425 ^d		CC425						<i>sac1</i> mutant					
				CC425 ^d	<i>sac1</i> ^d	2 h	4 h	8 h	24 h	0 h	2 h	4 h	8 h	12 h	24 h		
Sulfur metabolism																	
'963038E01	1936	<i>ARS1</i>	Arylsulfatase	+++++	8.5	17.52	42.81	41.59	3.39	1.32	1.1	0.8	0.77	0.99	NA	
'963046D01	9009	<i>SIR1</i>	Sulfite reductase	+++++	-.+.+.+	1.48	1.63	2.05	2.62	3.16	0.39	3.30	1.42	1.24	1.47	1.20	
'963038A06	4494		Sulfite reductase	+++++	+++++	2.69	2.48	3.18	3.59	2.39	0.75	2.37	1.85	1.76	1.91	1.89	
'963042H10	8444	<i>SIR3</i>	Sulfite reductase, bacterial type	+++++	0.92	1.03	1.11	0.96	0.99	0.93	2.16	NA ^e	0.49	1.72	NA	
'894033F10	[7588]	<i>ATS1</i>	ATP sulfurylase	+++++	+.+.+.+	4.94	5.29	8.62	10.58	7.69	1.05	2.65	2.43	2.04	1.7	0.9	
'963046C09	1668	<i>ATS2</i>	ATP sulfurylase	+++++	+++++	3.63	4.69	5.91	8.64	7.74	0.9	1.61	1.23	0.85	1.35	0.42	
'894044E12	9443	<i>APR</i>	5'-Adenylylsulfate reductase	+++++	+++++	2.35	1.57	1.31	1.40	1.34	1.01	0.90	0.85	0.99	1.21	0.97	
'894077C01	1669	<i>AKN2</i>	APS kinase	+++++	0.77	1.17	0.70	1.42	1.07	0.57	NA	NA	NA	NA	2.18	
'963045D11	3138	<i>SATI</i>	Serine O-acetyltransferase	+++++	-.+.+.+	6.69	6.25	10.59	14.77	9.06	0.88	1.77	1.44	1.24	1.44	2.18	
'963024B05	8367	<i>OASTL4</i>	O-Acetylserine(thiol)lyase	+++++	+++++	6.01	7.00	10.03	9.16	8.3	0.77	2.34	2.29	2.59	2.23	1.17	
'963076G10	6009	<i>CDO1</i>	Cysteine dioxygenase	+++++	+++++	5.94	5.68	7.75	8.9	7.54	0.71	2.29	1.92	2.01	2.01	2.01	
'963035D06	8168	<i>SOD1</i>	UDP-SQ synthase	+++++	+++++	5.63	6.64	7.73	9.5	9.57	0.97	4.43	2.4	2.45	2.36	4.2	
'894101C06	None ^f	<i>SOD2a</i>	UDP-SQ; diacylglycerol SO ₂ transferase	+++++	+++++	5.16	6.13	8.44	8.31	6.89	0.83	1.69	2.35	1.9	1.75	1.12	
'963017H04	[5161]	<i>ECP76</i>	Extracellular polypeptide 76	+++++	-.+.+.+	4.83	9.49	34.07	22.14	21.45	0.85	1.12	0.78	1.28	1.41	1.28	
'894020C12	5894	<i>SBDP</i>	Selenium-binding protein	+++++	+++++	7.89	5.52	9.66	8.23	5.19	1.29	1.7	1.53	1.39	1.52	1.6	
'963027A09	5894	<i>SBDP</i>	Selenium-binding protein	+++++	+.+.+.+	12.72	14.16	16.13	20.4	15.71	0.74	1.75	1.53	1.57	1.86	0.89	
'963046G02	7825		SAC1-like proteins (2 overlapping genes)	+++++	+++++	27.52	18.06	40.38	45.9	26.56	0.77	3.66	2.13	1.86	2.41	1.63	
'894081F11	[7825]		SAC1-like protein	+++++	-.+.+.+	60.83	48.89	37.09	84.46	53.56	0.64	1.62	2.12	2.34	2.95	1.23	
Photosynthesis																	
'963024B11 (C)	5736	<i>LHC41</i>	Light-harvesting complex, PS I	+.+.+-	-----	1.21	1.38	1.2	0.7	0.39	0.38	0.41	0.25	0.25	0.26	0.06	
'963047H05 (C)	2886	<i>LHC42</i>	Light-harvesting complex, PS I--	-----	1.27	1.16	1.1	0.58	0.3	0.39	0.2	0.16	0.15	0.09	0.03	
'894033H06 (C)	4209	<i>LHC43</i>	Light-harvesting complex, PS I--	-----	1.04	1.13	1.09	0.55	0.22	0.37	0.31	0.22	0.17	0.12	0.01	
'963042A01 (C)	9189	<i>LHC45</i>	Light-harvesting complex, PS I	+.+.+-	-----	1.55	1.67	1.28	0.85	0.39	0.34	0.38	0.28	0.26	0.16	0.02	
'894041D11 (C)	7340	<i>LHC45</i>	Light-harvesting complex, PS I--	-----	1.22	1.67	1.19	0.69	0.45	0.66	0.37	0.29	0.29	0.32	0.22	
'894087C09 (C)	4579	<i>LHC46</i>	Light-harvesting complex, PS I	+++++	-----	1.46	1.42	1.36	0.84	0.36	0.66	0.56	0.33	0.37	0.42	0.41	
'894044B07 (C)	5739	<i>LHC46</i>	Light-harvesting complex, PS I	+++++	-----	1.2	1.15	1.21	0.68	0.41	0.69	0.57	0.39	0.32	0.26	0.1	
'894078C01 (C)	[6907]	<i>LHC49</i>	Light-harvesting complex, PS I	+++++	-----	1.28	1.27	1.22	0.79	0.26	0.78	0.34	0.39	0.33	0.18	0.01	
'894076B06 (C)	7340	<i>LHC49</i>	Light-harvesting complex, PS I	+.+.+-	-----	1.2	1.13	1.18	0.67	0.37	0.47	0.19	0.13	0.11	0.1	NA	
'963069C06 (C)	[8250]	<i>LHCBM1</i>	Light-harvesting complex, PS II	+++++	-----	1.26	1.46	1.33	1.11	0.41	0.54	0.19	0.23	0.17	0.07	0.00	
'894080G01 (C)	1951	<i>LHCBM3</i>	Light-harvesting complex, PS II	+.+.+-	-----	NA	1.18	0.79	0.84	0.52	0.57	0.25	0.19	0.14	0.09	0.01	
'894062E07 (C)	7231	<i>LHCB4</i>	Light-harvesting complex, PS II	+++++	-----	1.38	1.34	1.49	0.95	0.73	0.82	0.52	0.36	0.47	0.39	0.07	
'894052A01 (C)	1436	<i>LHCB5</i>	CP26, minor light-harvesting complex, PS II	+++++	-----	1.21	1.3	1.36	0.69	0.63	0.55	0.28	0.22	0.22	0.18	0.07	
'894097E05 (E)	5770	<i>LhcSR2</i>	Light-harvesting family polypeptide	+++++	+.+.+.+	10.22	13.94	19.99	20.54	22.38	0.82	1.4	0.67	NA	1.62	NA	
'894005B12	[8181]	<i>LIL</i>	Light-harvesting family polypeptide	+.+.+-	1.55	0.97	0.85	0.58	0.94	1.00	0.7	0.74	0.87	0.72	0.38	
'963047E03 (A)	[2510]	<i>PSAD</i>	PS I subunit IV--	-----	1.16	1.13	1.18	1.01	0.65	0.84	0.45	0.43	0.44	0.29	0.03	
'894083B07 (A)	[3606]	<i>PSAE</i>	PS I subunit V--	-----	1.03	0.9	1.00	0.69	0.24	0.75	0.36	0.36	0.44	0.28	0.03	
'894041H01 (A)	1029	<i>PSAF</i>	PS I subunit III	+.+.+-	-----	1.07	1.21	1.31	0.91	0.42	0.69	0.57	0.49	0.44	0.37	0.08	
'894065A07 (A)	[4807]	<i>PSAG</i>	PS I subunit G--	-----	1.05	0.88	0.96	0.57	0.24	0.63	0.24	0.23	0.17	0.11	0.03	
'894100A05 (A)	[4807]	<i>PSAG</i>	PS I subunit G--	-----	1.03	1.15	1.01	0.63	0.27	0.63	0.27	0.24	0.17	0.11	0.05	
'894014A05 (A)	6118	<i>PSAH</i>	PS I subunit H	+++++	-----	1.17	1.26	1.21	0.86	0.36	0.74	0.49	0.63	0.6	0.4	0.16	
'894086C09 (A)	4991	<i>PSAK</i>	PS I subunit K--	-----	0.99	0.7	0.98	0.57	0.41	1.03	0.51	0.54	0.51	0.42	0.39	
'894004A09 (A)	[7107]	<i>PSAL</i>	PS I subunit L--	-----	1.09	1.1	1.13	0.8	0.4	0.55	0.39	0.39	0.33	0.32	0.16	
'894019E07 (A)	683	<i>PSAO</i>	PS I subunit O--	-----	1.19	1.13	1.19	0.65	0.27	0.81	0.31	0.3	0.24	0.16	0.03	
'894068A11 (B)	3356	<i>PSBO</i>	OEE1, oxygen evolution enhancer 1	+.+.+-	-----	1.08	1.22	1.18	0.9	0.58	0.68	0.36	0.35	0.42	0.32	0.09	
'894006E05 (B)	7935	<i>PSBP1</i>	OEE2, oxygen evolution enhancer 2--	-----	1.02	1.00	0.94	0.64	0.66	0.78	0.27	0.26	0.32	0.23	0.07	
'963041E04 (B)	[5978]	<i>PSBQ</i>	OEE3, oxygen evolution enhancer 3	+.+.+-	-----	0.93	1.09	1.22	0.86	0.93	0.75	0.15	0.4	0.48	0.31	0.07	
'894035D06	2796	<i>PSBP3</i>	Similar to OEE2--	-----	0.73	0.84	0.81	0.99	0.82	1.04	1.14	1.00	1.04	1.08	1.02	
'894084F07	[3599]	<i>PSBR</i>	10-kDa PS II polypeptide--	-----	1.24	1.19	1.31	1.67	1.37	1.13	1.31	1.26	1.21	1.25	1.25	
'894069B11	6084	<i>PSBW</i>	PS II reaction center W protein	+.+.+-	-----	0.97	1.21	1.27	1.01	0.68	1.06	0.53	0.52	0.61	0.46	0.14	

'894072A03	'[2051]	PSB28	Ycf79, also called PsbW-like	0.78	0.76	0.56	0.64	0.77	1.02	0.90	0.84	0.78	0.81	0.79
'894100F04 (D)	'[5600]	PETC	Cytochrome <i>b_f</i> Rieske subunit	1.22	1.01	0.85	0.49	0.47	0.83	0.65	0.54	0.47	0.43	0.33
'963053C08 (D)	9358	PETM	Cytochrome <i>b_f</i> M subunit	1.27	1.45	1.25	0.6	0.42	1.49	0.57	0.68	0.51	0.43	0.15
'894089E08 (D)	5858	PETN	Cytochrome <i>b_f</i> N subunit	1.03	1.10	0.84	0.48	0.37	1.11	0.83	0.63	0.57	0.46	0.36
'963092G08 (D)	5888	PETO	Cytochrome <i>b_f</i> subunit V	0.86	0.78	0.55	0.34	0.5	0.74	0.38	0.23	0.23	0.1	NA
'894002C07 (D)	5888	PETO	Cytochrome <i>b_f</i> subunit V	0.94	0.75	0.64	0.38	0.46	0.61	0.36	0.23	0.24	0.12	0.02
'894069E01 (D)	5927	PETE	Plastocyanin (PCY1)	0.8	0.81	0.64	0.58	0.18	0.76	0.38	0.27	0.16	0.1	0.02
'894017C09 (D)	1713	PETF1	Ferredoxin	0.77	0.62	0.39	0.23	0.2	1.34	0.28	0.33	0.26	0.18	0.04
'963046B11 (D)	4881	PETF5	Ferredoxin	0.04	0.05	0.04	0.04	0.05	0.13	0.09	0.23	0.18	0.08	0.06
'963025F07 (D)	6292	FNR	Ferredoxin-NADP reductase	1.08	0.87	0.63	0.26	0.21	0.6	0.32	0.17	0.2	0.12	0.15
'963048F01	7947	FTRV	Ferredoxin-thioredoxin reductase variable chain	0.83	0.67	0.67	0.62	0.54	1.04	0.21	0.54	0.6	0.52	0.25
'894006G05	8211	ATPC	ATP synthase gamma chain, short transcript	1.18	1.27	1.05	0.67	0.59	0.87	0.85	0.78	0.71	0.74	0.64
'894041H06	6174	ATPC	ATP synthase gamma chain, longer transcript	1.23	1.05	0.97	0.36	0.41	0.71	0.54	NA	NA	NA	0.38
'894073D05	8211	ATPC	ATP synthase gamma chain, longer transcript	1.21	1.02	0.91	0.35	0.49	0.88	0.68	0.78	0.64	0.60	0.50
'963039D06	6174	ATPC	ATP synthase gamma chain, longest transcript	1.52	1.11	1.13	0.69	0.60	0.89	1.69	0.53	0.44	0.44	NA
'894021B01	'[6167]	ATPD	ATP synthase delta chain	1.39	1.55	1.40	0.49	0.66	1.39	0.96	1.26	1.01	0.81	0.90
'963089H03	302	ATPG	ATP synthase CFO subunit II	1.74	1.63	1.35	0.59	0.45	1.49	0.90	1.11	0.84	0.59	0.42
Carbon metabolism														
'894022H12	2613	PRK	Phosphoribulokinase	1.11	1.00	0.76	0.37	0.37	1.04	0.43	0.35	0.44	0.34	0.05
'963079E12	'[8511]	RBCS1/2	Ribulose biphosphate carboxylase small chain	NA	1.25	1.28	1.03	0.63	1.99	0.76	0.76	0.75	0.45	0.19
'894098E09	1399	TPI	Chloroplast triosephosphate isomerase	0.57	0.55	0.51	0.69	0.61	0.57	0.6	0.47	0.27	0.47	0.41
'963014G05	6790		SERA, putative D-3-phosphoglycerate dehydrogenase	2.06	2.77	4.03	5.15	3.3	0.65	1.71	1.09	1.47	1.84	0.61
'963028B11	6790		SERA, putative D-3-phosphoglycerate dehydrogenase	2.06	2.66	4.46	4.38	3.1	0.35	1.93	0.96	1.5	1.88	0.47
'894006B02	4808		ALD3, fructose-1,6-bisphosphate aldolase	2.05	2.69	3.6	3.08	1.22	1.35	1.8	2.14	2.07	1.25	0.9
'894040F03	9404	PGK	Phosphoglycerate kinase	0.82	0.81	0.73	0.59	0.6	0.91	0.32	0.27	0.27	0.18	0.05
'894082E07	6137		Putative transketolase	0.96	0.91	0.61	0.47	0.53	0.67	0.45	0.37	0.41	0.45	0.22
'894022G03	5947	PCK	Phosphoenolpyruvate carboxykinase	1.31	1.12	1.07	0.5	0.35	1.03	0.81	0.91	0.74	0.47	0.16
'963017B07	'[6111]	GND	6-Phosphogluconate dehydrogenase	2.25	3.07	4.47	6.76	6.55	1.41	4.95	3.93	4.85	5.63	5.33
'894037H05	4431	TAL1	Transaldolase	2.98	3.16	2.38	1.14	0.59	0.81	0.86	0.92	0.5	0.28	NA
'894029C08	6695		Similar to PGLS (6-phosphogluconolactonase)	2.87	3.23	3.78	4.44	2.61	0.58	0.85	0.76	0.86	0.96	0.67
'894080B03	1974	GAP1	Cytosolic glyceraldehyde 3-P dehydrogenase	0.42	0.69	1.79	2.44	1.84	1.29	0.51	1.69	2.31	2.13	0.37
'963029F05	6610	SHMT1	Serine hydroxymethyltransferase	1.39	1.4	1.33	1.32	1.5	0.8	1.71	2.29	1.98	1.88	2.49
'963036C03	6493		GLX1, putative glyoxal oxidase	0.03	0.09	0.04	0.04	0.05	0.11	0.04	0.03	0.02	0.02	NA
'963070H06	'[9345]		SDC1, serine decarboxylase	1.24	0.95	0.78	0.81	0.97	0.44	0.4	0.38	0.34	0.37	0.31
'963045H04	5544		ACS3, putative acetyl-coenzyme A (CoA) synthetase	3.15	2.58	1.47	0.78	0.3	1.68	1.55	1.43	1.14	0.68	0.08
'894038D07	7098		PEL, pyruvate formate lyase	0.21	0.21	0.31	0.34	0.32	0.25	0.06	0.3	0.42	0.31	0.06
'963048D10	1813	ST42	Granule-bound starch synthase I	3.25	4.04	6.16	4.96	5.07	0.85	1.41	1.5	1.4	1.58	0.19
'894008B02	7333	GLPV	Glycogen/starch phosphorylase	0.99	1.25	2.34	3.11	4.17	0.84	1.07	1.82	2.07	2.08	2.08
Respiration alternative electron transfer pathways, ATPases, transporters														
'894024D07	8491	COX2a	Cytochrome oxidase subunit IIa, N terminus	0.77	1.03	1.41	1.38	0.93	0.77	0.4	0.81	0.92	0.73	0.38
'894102D01	'[7853]	COX2a	Cytochrome oxidase subunit IIb, C terminus	0.76	1.02	1.10	1.51	1.18	1.17	1.14	1.37	1.40	1.19	0.88
'894076G10	6296	COX90	Cytochrome oxidase, <i>Chlamydomonas</i> -specific subunit	0.72	1.03	1.09	1.28	0.76	0.93	0.43	0.72	0.67	0.66	0.27

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TABLE 1—Continued

Clone (Fig. 3 panel) ^b	ACE ^c	Name	Gene annotation	Direction of change in transcript level		Transcript level (fold increase) for											
				<i>sac1^d</i>		CC425						<i>sac1</i> mutant					
				CC425 ^d	<i>sac1^d</i>	2 h	4 h	8 h	12 h	24 h	0 h	2 h	4 h	8 h	12 h	24 h	
'963044H03	1715	<i>ATP6</i>	Subunit 6 mitochondrial ATP synthase	-. . . -	-----	0.86	0.94	0.99	0.82	0.53	0.51	0.48	0.64	0.68	0.68	0.37	
'894008C03	1860	<i>AOX4</i>	Chloroplast plastoquinol-oxygen oxidoreductase	+++++	+++++	2.1	2.24	3.28	3.33	9.52	1.58	5.68	6.39	7.52	7.35	4.16	
'894076B10	4448	<i>HYD1</i>	Chloroplast Fe-hydrogenase	-----	-----	0.27	0.25	0.19	0.23	0.22	0.56	0.16	0.17	0.24	0.14	0.22	
'894044F09	8476	<i>AOX1</i>	Mitochondrial alternative oxidase	-. . . .	-. . . .	0.57	0.59	0.77	0.89	1.14	0.76	0.25	0.5	0.66	0.58	0.2	
'894072B06	'[4402]		Mitochondrial fatty acid carrier/uncoupling protein	+. . . -	+++++	2.83	3.48	1.41	0.67	0.29	5.35	10.56	5.29	1.89	0.67	NA	
'894059B10	'[3328]		MPC1, mitochondrial phosphate transporter	+++++	+++++	1.66	1.9	2.52	2.57	1.42	1.12	2.61	3.05	1.8	1.47	1.15	
'894026D10	4026		Plastid ATP/ADP transporter	-. . . -	-. . . -	0.73	0.79	1.05	0.98	0.81	0.35	0.38	0.53	0.49	0.6	0.33	
'894092F04	2863		Plastid ATP/ADP transporter	0.73	0.85	1.04	1.08	1.15	0.42	0.8	0.56	0.54	0.7	0.42	
'894063B07	3927		Plasma membrane proton ATPase	-. . . .	-. . . .	0.32	0.3	0.38	0.48	0.53	0.79	0.41	0.55	0.68	0.73	0.7	
'894040E11	592		Vacuolar ATP synthase, subunit A	1.2	1.77	1.08	1.12	1.19	0.76	1.35	1.14	1.25	1.5	1.78	
'894018B07	4771	<i>ATPvE</i>	Vacuolar ATPase chain E	1.05	1.15	1.05	1.18	1.44	1.03	1.72	1.42	1.24	1.31	1.44	
'894022G07	'[4453]	<i>ATPvL1</i>	Vacuolar ATPase proteolipid subunit	1.26	1.40	1.42	1.70	2.01	1.15	1.38	1.55	1.51	1.68	1.60	
'963017F05	'[4552]		Proton-translocating, vacuolar pyrophosphatase	-. . . -	-. . . -	0.84	0.73	0.99	0.91	0.45	0.61	0.52	0.43	0.56	0.65	0.34	
'894054D08	'[7558]		Putative transporter/GTP binding	+. . . +	1.28	1.37	1.15	1.46	1.32	1.18	1.31	1.28	1.12	1.23	1.24	
'894071E08	'[4402]		Similar to mitochondrial substrate carrier	+++++	+++++	3.63	2.92	1.66	0.43	0.17	3.74	2.36	4.48	1.4	0.46	NA	
'894099G11	'3494		PTB4, phosphate transporter	-. . . .	1.07	1.04	1.04	1.07	0.96	0.74	0.55	0.47	0.43	0.62	0.35	
Oxidative stress, chaperones, proteolysis																	
'894081G12	3424	<i>FSD1</i>	Iron chloroplast superoxide dismutase	+++++	+++++	1.58	1.6	1.78	1.98	2.83	0.87	1.4	1.89	1.92	2.19	3.11	
'963046E04	5826	<i>GPXI</i>	Glutathione peroxidase	+++++	+++++	3.14	3.32	3.17	2.6	2.7	1.2	2.53	3.72	4.15	3.14	2.2	
'894082E09	6986		Thioredoxin peroxidase/hydroperoxide reductase	+. . . +	+. . . +	1.95	1.49	2.43	2.07	1.96	1.18	1.92	2.14	1.78	1.95	1.16	
'963032D03	5314		Putative vanadium chloroperoxidase	+++++	+++++	15.15	12.76	46.99	40.92	5.64	0.92	1.56	0.57	0.83	1.16	0.58	
'963025C05	4324		Putative peroxidoxin O	+++++	+. . . +	2.33	3.71	4.75	2.24	4.35	0.96	NA	0.97	0.23	0.25	3.9	
'894062A07	8645		PDx1, pyridoxin biosynthesis protein	+++++	+++++	1.71	1.71	1.7	1.63	1.00	0.86	2.64	2.35	1.9	1.98	2.49	
'894024F09	5909	<i>HSP90A</i>	Heat shock protein 90, cytosolic	-. . . .	-. . . .	0.2	0.32	0.36	0.46	0.73	0.45	0.21	0.52	0.59	0.57	0.58	
'963036F10	4194	<i>HSP70C</i>	Mitochondrial HSP70 chaperone	0.88	0.90	0.69	0.79	0.91	1.06	1.16	1.13	1.07	1.12	1.20	
'894078G05	4517	<i>HSP70B</i>	Chloroplast HSP70 chaperone	+. . . +	+. . . +	1.23	1.62	1.34	1.49	1.38	0.84	0.96	1.53	1.64	1.49	1.77	
'963026B05	7505	<i>CGE1</i>	Cochaperone of chloroplast HSP70	1.35	0.79	0.79	0.91	1.81	1.02	0.87	1.67	1.13	1.27	2.97	
'963041D04	1067	<i>VIPP1</i>	Membrane-associated 30-kDa chloroplast protein	+++++	+++++	1.96	2.06	2.13	1.87	2.80	1.10	4.46	5.10	3.94	8.75	3.37	
'894038F08	8733	<i>HCF136</i>	PS II stability/assembly factor	+++++	+. . . +	4.46	3.89	4.11	4.76	3.68	0.76	5.58	1.7	3.3	2.78	1.4	
'894044H01	2747	<i>ALB3.1</i>	Required for LHC integration	+. . . -	1.81	1.44	0.86	0.82	0.81	0.84	1.33	0.74	0.81	0.82	0.62	
'963035G03	'[4315]	<i>CPN60A</i>	Alpha subunit Rubisco-binding protein, GroEL	-. . . .	-. . . .	0.29	0.26	0.17	0.27	0.42	0.7	0.31	0.53	0.68	0.7	0.72	
'894062E01	5669	<i>CPN60B</i>	Beta subunit Rubisco-binding protein, GroEL	-.	0.35	0.23	0.22	0.3	0.44	0.64	0.21	0.73	0.76	0.9	1.15	
'894039G09	1845		GROES-like protein	0.9	0.96	0.63	0.86	0.85	1.19	1.39	1.36	1.32	1.42	1.46	
'894076H02	'[4562]		Rubisco activase	+++++	+. . . +	1.90	1.77	1.42	1.07	1.39	1.77	0.49	0.68	1.36	1.06	0.44	
'963050F02	3862	<i>HSP22C</i>	Chloroplast 22-kDa heat shock protein	0.77	0.73	1.13	0.88	1.17	1.10	2.21	1.25	1.52	2.09	2.96	
'963068F01	5588	<i>HSP22E</i>	Chloroplast 22-kDa heat shock protein	+. . . +	+. . . +	1.74	2.68	2.39	2.14	1.1	0.37	1.33	3.55	4.06	5.39	18.74	
'894096A07	'[7754]	<i>HSP22F</i>	Chloroplast 22-kDa heat shock protein	+++++	+++++	1.61	3.23	1.83	2.42	0.89	0.49	1.8	5.11	5.69	8.22	42.22	
'894008D03	2231		Cytosolic peptidyl-prolyl <i>cis-trans</i> isomerase	-. . . .	-. . . .	0.33	0.39	0.43	0.42	0.34	1.03	0.53	0.68	0.67	0.7	0.37	
'894086F07	344		Chloroplast peptidyl-prolyl <i>cis-trans</i> isomerase	-.	0.31	0.43	0.34	0.41	0.32	1.08	0.72	0.63	0.72	0.8	0.46	
'963042H09	344		Chloroplast peptidyl-prolyl <i>cis-trans</i> isomerase	-.	0.42	0.51	0.37	0.44	0.30	1.04	0.54	0.57	0.77	0.71	0.50	
'894005A06	'[2563]		Chloroplast peptidyl-prolyl <i>cis-trans</i> isomerase	-. . . .	-. . . .	0.59	0.46	0.36	0.36	0.26	0.72	0.27	0.33	0.31	0.33	0.18	

TABLE 1—Continued

Clone (Fig. 3 panel) ^b	ACE ^c	Name	Gene annotation	Direction of change in transcript level		Transcript level (fold increase) for											
				CC425 ^d		CC425						sacI mutant					
				CC425 ^d	sacI ^d	2 h	4 h	8 h	12 h	24 h	0 h	2 h	4 h	8 h	12 h	24 h	
'894097B09	2841		IGPD (imidazoleglycerol-phosphate dehydratase)	. + . . .	-. + . . . +	1.04	1.17	0.91	1.15	1.03	0.7	1.76	1.31	1.54	1.96	1.84	
'894103C12	'[2773]		SAHI, S-adenosyl homocysteine hydrolase	-----	-----	0.46	0.3	0.28	0.12	0.06	0.25	0.69	0.39	0.64	0.69	0.27	
'894056F03	'[6078]		THIH-like protein, thiazole biosynthesis	-----	0.17	0.25	0.16	0.42	0.38	1.03	1.03	0.94	0.79	0.83	1.07	
'894096H02	1312		HISC2, putative histidinol-phosphate transaminase	-. - . . .	-. + + + + + + +	0.82	0.96	0.89	1.22	0.88	0.73	2.26	2.25	2.28	2.53	2.88	
'963030A07	2652	<i>FER1</i>	FER1, ferritin	-----	-----	0.77	0.61	0.68	0.58	0.29	0.56	0.54	0.42	0.4	0.43	0.17	
'963042H06	6157		ACH, aconitate hydratase	+. +. - - -	. . . - - - -	1.65	1.2	0.9	0.74	0.63	0.94	0.94	0.93	0.63	0.63	0.27	
'963044D08	2010		APOC, apoprotein-associated protein	. . . - - - -	-----	1.01	1.07	1.13	0.73	0.54	0.8	0.57	0.56	0.34	0.33	0.25	
'894037G08	8636		ALS, acetolactate synthase	+. + + + + +	1.39	1.29	1.47	1.55	1.33	1.14	1.23	1.23	1.25	1.35	1.38	
'894064A10	468		ILV3, dihydroxy-acid dehydratase	-. - - - -	. . + + + + +	0.49	0.55	0.43	0.8	0.57	0.83	1.41	1.98	2.00	2.07	3.88	
'894080E01	773		Putative acetyl-CoA carboxylase subunit	. - - - - +	-----	0.96	0.78	0.75	0.69	1.33	0.83	0.49	0.33	0.5	0.54	0.38	
'894062A07	8645		PDX1, pyridoxine biosynthesis protein	+. + + + +	. + + + + + +	1.71	1.71	1.7	1.63	1.00	0.86	2.64	2.35	1.9	1.98	2.49	
'894102B02	'[2302]		Putative Igr3p cargo protein	-----	+ -	0.28	0.29	0.32	0.33	0.32	3.95	NA	NA	NA	NA	0.72	
'894019G08	4767		Adenylosuccinate synthetase (AMP biosynthesis)	-----	-	0.3	0.33	0.32	0.45	0.4	0.76	0.62	0.84	0.87	0.92	1.01	
'894068F05	415		dTDP-glucose 4-6-dehydratase	-----	- - - - -	0.5	0.46	0.6	0.6	0.43	0.9	0.39	0.5	0.62	0.66	0.28	
Signal transduction, transcription, translation																	
'894096C06	467		Sensory opsin A	-----	-----	0.63	0.56	0.74	0.79	0.67	0.59	0.38	0.59	0.69	0.63	0.07	
'963038D05	7033		RABF1, RabH/Rab5 type of small GTPase	+. + + + + +	. + + + + + +	1.49	1.73	1.87	2.2	2.07	0.99	2.2	2.23	1.9	1.74	1.64	
'963048H07	'[6103]		GBLP, guanine nucleotide binding protein +	. . . + + .	0.93	1.07	1.09	1.01	1.44	1.42	1.3	1.53	1.63	1.47	1.46	
'894006F01	'[9232]		SKS1-like protein kinase	-----	. - - - - -	0.54	0.53	0.58	0.43	0.37	0.77	0.2	0.4	0.42	0.36	0.18	
'963045F12	'[2130]		PP1, serine/threonine protein phosphatase	-----	-----	0.45	0.42	0.44	0.54	0.55	0.53	0.31	0.4	0.45	0.45	0.31	
'894006C11	7297		XPO1, exportin-	. - . . .	-----	0.72	0.9	0.83	0.81	0.84	0.45	0.49	0.12	0.36	0.25	0.36	
'894006C08	7660		GBP1p, G-strand-binding protein	-----	-----	0.24	0.17	0.13	0.2	0.15	0.59	0.22	0.25	0.26	0.29	0.3	
'963046C02	8379		Putative nucleosome assembly protein 1-like protein	+. + + + + + + +	2.24	2.47	2.86	3.96	4.34	0.98	1.08	1.19	1.26	1.32	1.37	
'894027A08	None		ATP dependent helicase	-----	. -	0.28	0.19	0.27	0.41	0.37	0.74	0.63	0.73	0.73	0.75	0.84	
'963047C11	5558		Putative transcription factor IIB	-----	+ +	0.22	0.28	0.3	0.37	0.3	2.69	1.22	NA	1.15	3.12	NA	
'963087F05	7852		Probably DNA-binding protein	-----	-----	0.4	0.31	0.26	0.32	0.35	0.75	0.5	0.83	1.05	1.01	1.14	
'894081G03	8592		Potential RNA-binding protein	+. +. - - -	. - . - - - -	1.56	1.29	1.08	0.8	0.68	0.99	0.73	0.76	0.55	0.53	0.26	
'894010G03	8151		Similar to <i>Arabidopsis</i> SC35-like splicing factor SCL35	-----	-----	0.39	0.47	0.55	0.54	0.39	0.48	0.27	0.3	0.32	0.41	0.31	
'963079B08	6042		EF1A1, putative eukaryotic translation initiation factor	. . - - - -	. . . + + + +	0.94	0.88	0.55	0.61	0.63	0.92	1.25	1.43	1.65	1.53	1.72	
'894081B08	7498		EF1A2, putative elongation factor 1 a 2	+. + + + + +	+. + + + + + +	1.24	1.23	1.6	1.44	1.13	1.86	2.32	2.67	2.4	2.45	2.38	
'963077H03	1273		Protein translation factor, similar to SUH1	- + . + + + +	1.18	1.25	1.12	0.97	0.9	1.19	1.72	1.44	1.93	2.29	5.69	
'894080F07	5613		NOP56-like, nucleolar rRNA processing protein	-----	0.39	0.49	0.28	0.48	0.4	1.23	0.74	1.36	1.06	1.27	1.24	
'963030C08	8384		Fibrillarin-related nucleolar RNA-binding protein	-----	. - . . . +	0.22	0.25	0.21	0.52	0.34	0.89	0.27	0.75	0.84	1.07	1.33	
'894069A01	6666		HMGBl, high-mobility-group DNA-binding protein	-----	-----	0.48	0.57	0.67	0.64	0.25	0.45	0.44	0.56	0.52	0.47	0.37	
'963047E09	115		NHP2, high-mobility-group nucleolar protein	-----	. - . . . +	0.38	0.39	0.36	0.57	0.46	1.14	0.41	1.23	1.09	1.25	1.61	
Cytoplasmic and chloroplast ribosomal proteins ^g																	
'894081C01	6021		RPS3a-like	. + + + + +	+. + + + + + +	1.02	1.21	1.42	1.63	1.7	2.69	1.79	2.66	2.49	2.63	4.00	
'963045C05	448		RPS5	. +	+ -	0.98	1.15	1.02	1.31	1.31	1.28	0.51	0.84	0.77	0.72	0.73	

'894070B04	2212	RPL22	0.97	1.02	0.91	1.14	1.37	1.96	1.54	2.17	1.74	1.42	1.94
'963038E05	2202	RPS20	1.11	0.96	1.06	0.97	1.29	2.03	0.97	1.23	1.42	1.22	1.19
'894011F03	[8867]	RPS16	0.72	1.06	0.4	1.98	2.77	1.26	3.91	4.88	1.26	2.81	2.85
'894006G10	[9571]	RPS17	1.00	1.09	1.1	1.24	1.35	2.42	0.96	1.57	1.81	1.66	2.24
'894064E10	5820	RPS19	0.87	1.07	1.01	1.21	1.54	2.08	1.62	1.97	1.86	1.83	1.79
'894037B07	8122	RPS25	1.17	0.95	1.19	1.05	1.13	2.00	0.7	1.25	1.23	1.09	1.00
'894014F12	4106	RPS30	1.01	1.27	1.22	1.65	1.7	4.07	1.68	2.4	2.48	2.04	2.22
'894017F08	5393	RPS27	0.92	1.33	1.28	1.22	1.25	1.89	1.31	1.76	2.1	1.97	2.36
'963036H09	6746	RPL6	1.24	1.09	1.36	1.04	1.07	1.46	1.68	1.8	2.23	2.1	1.93
'894027B04	2452	RPL9	1.00	1.02	1.02	1.24	1.07	2.72	1.16	1.49	1.69	1.49	1.6
'963038C05	[3133]	RPL10	0.92	0.94	0.99	0.93	1.15	1.65	1.28	1.81	1.75	1.58	2.19
'894065B09	9108	RPL11	1.11	0.79	1.14	1.13	1.29	2.56	0.93	1.25	1.73	1.59	1.57
'894064D04	3624	RPL12	0.95	1.03	0.97	1.23	1.32	2.35	1.16	1.57	1.67	1.54	1.99
'894019C08	7238	RPL15	1.2	1.2	1.45	1.29	1.46	3.48	1.26	1.76	1.96	1.85	1.5
'963063C08	3	RPL17	1.07	0.86	0.99	0.96	1.17	2.05	0.7	0.93	1.29	1.11	1.22
'894039B07	2305	RPL23	1.07	0.94	1.11	1.16	1.25	2.09	0.66	1.07	1.63	1.22	1.1
'963048E03	2072	RPL24	1.11	0.86	0.98	1.13	1.33	2.27	0.84	1.39	1.63	1.3	1.31
'963017B08	[8598]	RPL27	1.06	0.96	1.00	1.12	1.11	1.85	0.9	1.27	1.28	1.2	1.21
'894006A06	111	RPL30	1.1	0.98	1.06	1.12	1.67	3.35	0.97	1.39	1.9	1.52	1.5
'894066H09	6057	RPL35	0.48	0.41	0.31	0.3	0.55	0.9	0.49	0.7	0.77	0.8	1.22
'894026G08	2805	RPSCL1	0.25	0.2	0.21	0.19	0.21	0.32	0.1	0.11	0.13	0.12	0.05
'894044G05	5258	RPSCL3	0.38	0.41	0.19	0.29	0.52	0.95	0.76	0.74	0.83	0.95	1.27
'894037C03	9418	RPSCL13	0.46	0.33	0.32	0.34	0.62	1.02	0.61	0.71	0.87	1.01	1.4
'894002G05	825	RPSCL20	0.37	0.27	0.19	0.28	0.49	1.14	0.38	0.51	0.77	0.78	0.88
'963039A05	8056	RPLCL1	0.35	0.29	0.2	0.27	0.59	0.96	0.59	0.82	0.82	0.99	1.94
'894037B11	2083	RPLCL6	0.47	0.38	0.25	0.27	0.54	1.3	0.47	0.64	0.85	0.99	1.26
'894089F04	4407	RPLCL9	0.48	0.48	0.26	0.3	0.54	1.25	1.01	0.88	1.00	0.88	1.2
'963032C10	3366	RPLCL12	0.34	0.23	0.14	0.18	0.4	0.93	0.49	0.54	0.69	0.86	1.15
'894055G04	[2967]	RPLCL18	0.4	0.36	0.2	0.24	0.5	1.3	0.53	0.74	0.8	0.93	1.48
'894050B05	[6564]	RPLCL28	0.34	0.31	0.2	0.3	0.68	1.14	0.57	0.74	1.02	1.06	2.01
'894073D04	'9202	RPLCL31	0.46	0.45	0.24	0.29	0.54	1.14	0.74	0.82	0.94	1.02	1.2

Putative structural, surface, and matrix proteins

'894017B12	7239	VFL2, centrin (caltractin)	0.65	0.54	0.66	0.61	0.74	1.51	0.42	0.6	0.59	0.54	0.34
'894032G12	7345	TUA2, a tubulin 2	0.38	0.34	0.28	0.36	0.33	1.29	0.48	0.5	0.58	0.68	0.57
'894006H07	82	CHLRE_650068, microtubule-associated protein	2.73	3.08	2.93	3.08	3.3	1.31	5.63	5.27	5.43	5.59	8.45
'963046E08	8405	TUA1, alpha tubulin 1	0.46	0.32	0.3	0.29	0.26	0.77	0.06	0.08	0.12	0.07	NA
'894056C11	[7442]	TUB1, beta tubulin 1	0.27	0.19	0.21	0.27	0.16	1.18	0.1	0.22	0.23	0.24	0.02
'894093H02	[3445]	TUB2, beta tubulin G1	0.54	0.36	0.29	0.23	0.12	4.16	0.81	0.56	0.35	0.16	NA
'894056E01	[1531]	FAP15, flagellar proteome	0.24	0.24	0.37	0.39	0.39	0.49	0.03	0.07	0.2	0.06	NA
'894083H05	[285]	SWP71, syntaxin	1.37	1.27	1.3	1.65	1.82	1.19	1.9	2.57	2.42	2.44	1.32
'963026C08	568	FLA14, dynein light chain	0.36	0.33	0.35	0.36	0.31	1.85	0.67	0.74	0.86	0.59	0.32
'963077B03	2780	FMG1b, flagellar membrane glycoprotein 1b	0.19	0.17	0.17	0.18	0.12	0.16	0.07	0.02	0.02	0.04	0.01
'963046G04	835	Putative cell adhesion protein	3.34	2.1	3.96	4.01	5.51	1.13	1.25	1.27	1.18	1.28	1.36
'963024H07	8951	KATA, kinesin-like motor protein	1.95	1.14	3.24	2.11	1.41	0.84	0.68	0.88	1.14	1.05	0.95
'963046H09	7877	Putative sulfated surface glycoprotein	1.19	1.32	1.12	0.89	0.83	0.34	0.34	0.32	0.36	0.44	0.09
'894078A07	1239	Similar to pterophorin of <i>Volvox</i>	0.91	1.04	1.23	0.86	0.2	0.37	0.17	0.19	0.24	0.3	0.17
'963029B12	1971	GAS31, hydroxyproline-rich glycoprotein	0.4	0.36	0.37	0.31	0.2	0.62	0.35	NA	0.26	0.25	NA
'894068G03	[6102]	Putative prolyl 4-hydroxylase, a subunit	0.15	0.18	0.19	0.23	0.31	1.27	0.37	0.99	0.78	0.31	0.14
'894068C01	4654	Putative prolyl 4-hydroxylase, a subunit	0.04	0.05	0.04	0.05	0.08	0.21	0.04	0.08	0.11	0.05	0.01

Continued on following page

TABLE 1—Continued

Clone (Fig. 3 panel) ^b	ACE ^c	Name	Gene annotation	Direction of change in transcript level		Transcript level (fold increase) for												
				<i>sac1</i> ^d		CC425						<i>sac1</i> mutant						
				CC425 ^d	<i>sac1</i> ^d	2 h	4 h	8 h	12 h	24 h	0 h	2 h	4 h	8 h	12 h	24 h		
'894042G04	7452		Putative prolyl 4-hydroxylase, a subunit	-----	0.07	0.19	0.07	0.1	0.15	1.02	0.48	0.15	1.02	0.56	0.84	0.79	0.88
'894101C02	[428]		Putative prolyl 4-hydroxylase, a subunit	-----	-.	0.17	0.24	0.19	0.16	0.19	0.48	0.13	0.48	0.13	0.28	0.25	0.2	0.05
Unknown ^r																		
'963016B04	1549		Unknown	+++++	8.95	10.94	18.57	22.72	12.48	1.04	NA	4.17	3.29	3.72	1.4		
'963033D03	None		Unknown	+++++	-	13.46	15.01	22.74	23.51	22.3	0.39	4.5	5.34	4.58	3.85	2.16		
'963032A08	6781		Unknown	+++++	35.69	30.83	59.23	50.06	34.64	0.68	22.69	16.05	17.99	21.9	19.76		
'963047F05	2729		Unknown	+++++	2.07	4.42	16.00	18.91	15.01	1.05	1.74	5.6	4.76	3.56	1.21		
'963096E10	1549		Unknown	+++++	16.98	21.88	32.67	35.42	23.93	0.95	3.61	3.89	3.32	4.62	1.47		
'894063D10	3295		Unknown, starch-binding domain of glycoside hydrolase	+++++	5.18	4.33	16.54	14.59	13.15	0.92	1.54	1.16	1.53	1.71	1.63		
'963027G01	6569		Unknown	+++++	1.56	4.35	16.00	23.01	18.73	1.01	1.4	0.35	0.31	1.36	5.06		
'963063B08	5533		Unknown	+++++	-	10.05	17.49	23.73	21.99	18.27	0.41	2.96	4.27	3.85	3.25	2.39		
'963068A11	None		Unknown	+++++	6.97	8.97	17.49	18.08	12.56	1.24	4.32	2.15	1.7	2.8	6.36		
'963032A01	'9229		Unknown proline-rich protein	+++++	+++++	5.4	6.85	18.01	23.03	24.96	1.26	4.61	7.67	6.57	10.67	8.17		

^a The ≥ 3 -fold change is for at least two points in either CC425 and/or the *sac1* mutant. Some genes that did not satisfy these criteria but were discussed in the text are also included. This table also has a finer partitioning of the genes into categories than Table S1 in the supplemental material, but the genes can be tracked by the clone ID numbers.

^b The identifier of the array element, based on the cDNA clone used as a template for PCR amplification. Clone numbers followed by a letter in parentheses belong to a category used to generate Fig. 3A through E, as indicated.

^c The identification number in the 20021010 assembly of ESTs for a predicted unique gene (ACE) in which this clone is a member. The bracketed numbers are for clones not used in the ACE assembly but that match a specific ACE as determined by BLAST analysis.

^d Difference in transcript abundance (+ or -; a dot indicates no statistically significant change) relative to RNA from CC425 at time zero. The five symbols represent 2, 4, 8, 12, and 24 h after initiation of sulfur starvation of CC425; the data for the *sac1* mutant include a comparison of the *sac1* zero time point (initial symbol) with the wild-type zero time point.

^e NA, not enough high-quality data for statistical tests.

^f The read did not match any of the ACEs and thus was not included in the assembly of reads.

^g RP, ribosomal protein; L, large subunit; S, small subunit; CL, chloroplast; the number represents the molecular mass of the subunit.

^h A selection of 10 of the 187 "unknowns" that satisfied the ≥ 3 -fold selection criterion. The abundance of the transcripts from these genes within the unknown category exhibited the greatest increase following S deprivation.

ⁱ SO, sulfoquinovose.

^j SO, sulfoquinovosyl.

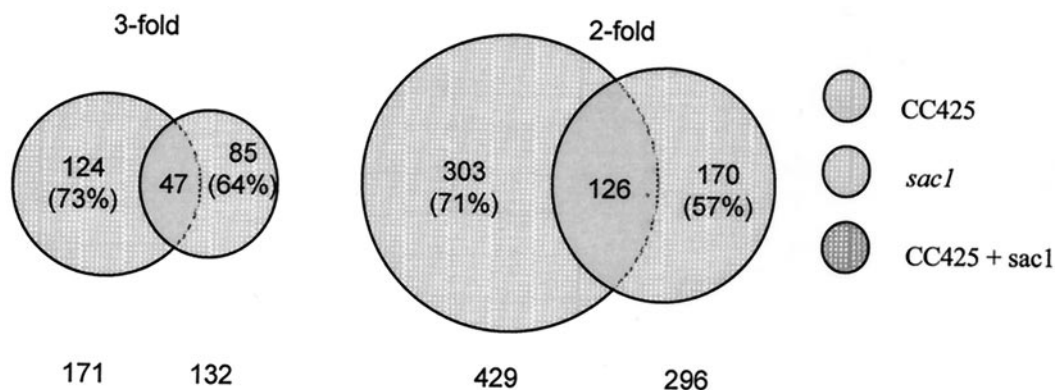


FIG. 1. Interlocking diagrams showing the number of genes from CC425 (left circles) and the *sac1* mutant (right circles) that respond to S-deprivation conditions. Left panel, transcript levels altered ≥ 3 -fold; right panel, transcript levels altered ≥ 2 -fold.

abundance changed ≥ 2 -fold (Fig. 1, right panel), more genes were selected, but the proportion of regulated genes for both strains remained similar. All of the CC425 and *sac1* mutant common transcripts were regulated in a similar direction (e.g., those that increased in CC425 also increased in the *sac1* mutant), although the kinetics and the extent of the changes could be somewhat different.

These results demonstrate that S deprivation leads to extensive changes in transcript abundance (transcripts from $>20\%$ of the CC425 genes represented on the array exhibit a change of twofold or more), reflecting a change in the physiological state of the cell. Furthermore, while the *sac1* mutant is defective for the control of a number of genes, there are still 30% of the sulfur-responsive genes in CC425 (126 out of 429) that appear to be properly regulated in this strain. We performed further analyses of those cDNAs for which the transcripts changed ≥ 3 -fold; the levels of all of these transcripts at the different times following S deprivation are shown in Table 1. This table also contains the data for a number of other genes, providing information relevant to the discussion of S deprivation responses. The data presented are an average of eight replicates in two separate experiments. The direction of change is indicated in the table for each time point (by + or -) when judged significant based on a *t* test. The ratio values for all of the genes represented on the array is available in the supplemental material (Table S1).

Genes involved in sulfur assimilation. Most transcripts for genes encoding polypeptides involved in the acquisition or assimilation of SO_4^{2-} increased following the imposition of S deprivation conditions. Relative transcript abundance for all of the genes in this category is presented in Table 1 (under the heading Sulfur metabolism); the data are also visually depicted in a bar graph presented as Fig. 2. Genes from this group that have been previously studied encode ARS1, ATP sulfurylase (ATS1), and ECP76; transcripts from these genes were previously shown to increase in response to S deprivation (7, 51, 62). Elevated levels of these transcripts were observed by 2 h following the imposition of S deprivation. Temporal changes in the levels of these transcripts during S deprivation, as determined from analyses of the microarrays, were consistent with the evaluations of transcript levels by RNA blot hybridizations (7, 51, 62). This category also contains genes for all steps of the

sulfate assimilation pathway, including two distinct sulfite reductases (*SIR1* and a distinct unannotated *SIR* gene; note that the transcript from the *SIR3* gene, which encodes another type of sulfite reductase, did not increase), the two ATP sulfurylases, and adenylyl sulfate reductase. Surprisingly, the single adenosine-*S*-phosphosulfate kinase gene present in the genome was not induced, suggesting that 3'-phosphoadenosine-5'-phosphosulfate synthesis does not become limiting under sulfur stress conditions. Transcripts for UDP-sulfoquinovosynthase (*SQD1*) and for a sulfolipid synthase (*SQD2a*), which probably utilizes sulfite for the synthesis of sulfolipids in the thylakoid membranes, increase during S starvation. In addition, the transcripts for serine acetyltransferase (*SAT*) and *O*-acetylserine(thiol)lyase (*OASTL*), which are involved in the biosynthesis of cysteine, are upregulated, as is the transcript for cysteine dioxygenase (963076G10), which converts cysteine to cysteine sulfinic acid (58) and may be involved in reclaiming S from amino acids. Furthermore, there is a rise in the levels of transcripts encoding a selenobinding protein (*SBDP*), which may be critical for sequestering selenate (by binding it and depositing it in the vacuole and/or facilitating its excretions), a potentially toxic SO_4^{2-} analog. Selenate would be more likely to accumulate in S-starved cells since the transport machinery of such cells would have a high affinity for SO_4^{2-} (and potentially selenate) and an elevated capacity for its uptake. It is also possible that the *SBDP* is involved in acclimation of cells to selenate deprivation (most selenate in the culture medium would originate as contamination from the sulfate stock solution), allowing for efficient acquisition when the levels of selenate are low in the environment. Recent work strongly suggests the presence of a number of selenoproteins in *C. reinhardtii* (36). The transcripts corresponding to the two array elements for *SBDP* (clone identification numbers 894020C12 and 963027A09) behave similarly and appear to represent the same gene. Finally, two genes coding for *SAC1*-like polypeptides are also induced (*SAC1* itself is not on the array).

These results demonstrate that transcripts from most genes involved in the acquisition, assimilation, and utilization of SO_4^{2-} (almost all of those known to be on the array; note that none of the sulfate transporter genes are on this array) increase during S deprivation and that SO_4^{2-} is likely scavenged from external and internal stores. Furthermore, as observed in

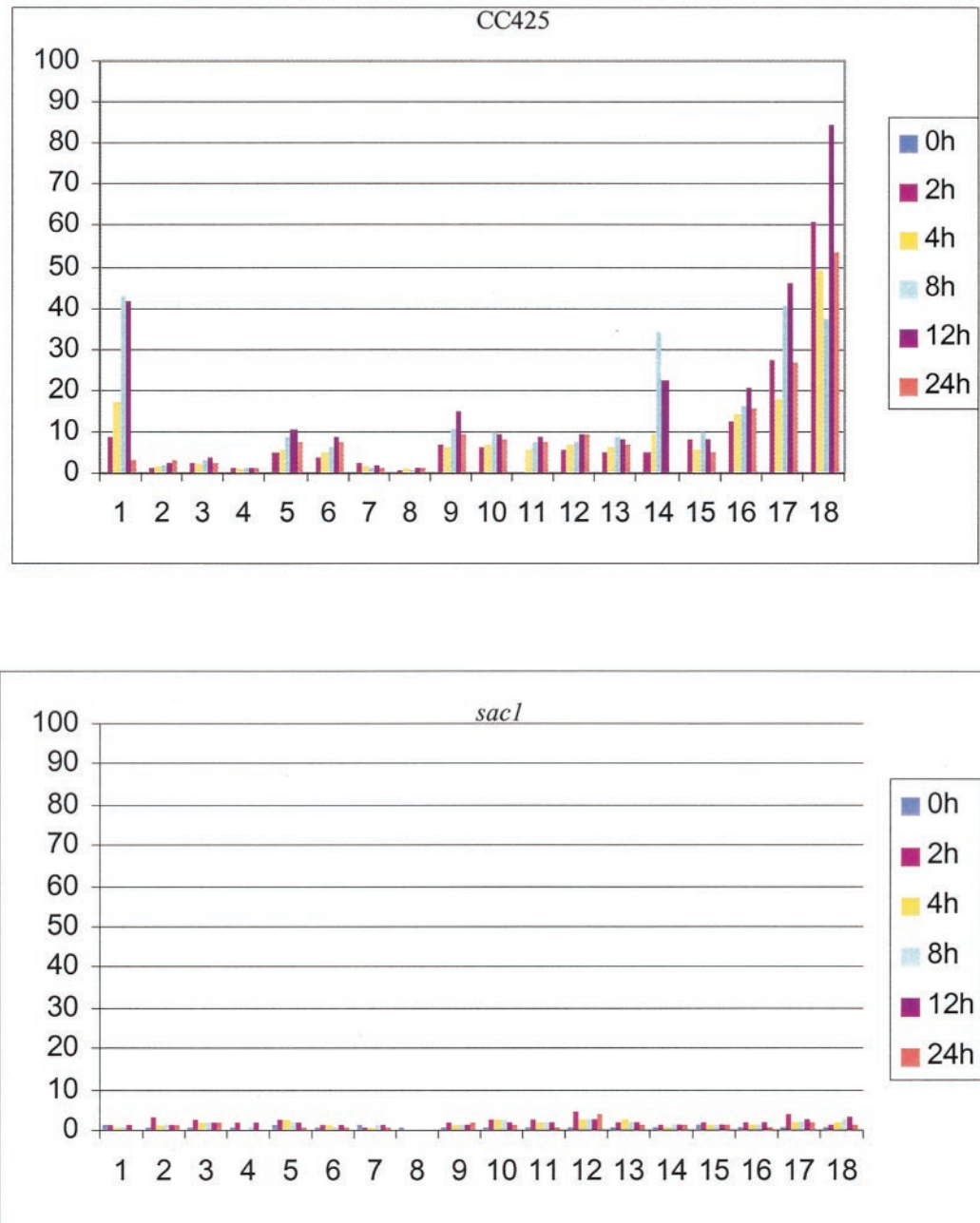


FIG. 2. Bar graph showing changes in transcript levels for genes involved in sulfur metabolism in CC425 (top graph) and the *sac1* mutant (bottom graph) during S deprivation. The responses, shown as the change (n -fold) relative to RNA from CC425 at time zero (which is set at 0), at the different times following S deprivation are given in different colors, as indicated on the graph, and the different genes represented encode the following proteins: 1, arylsulfatase (ARS1); 2, sulfite reductase (SIR1); 3, sulfite reductase (no gene or protein designation given); 4, sulfite reductase (SIR3); 5, ATP sulfurylase (ATS1); 6, ATP sulfurylase (ATS2); 7, 5'-adenylyl sulfate reductase (APR); 8, adenosine 5'-phosphosulfate kinase (AKN2); 9, serine *O*-acetyltransferase (SAT1); 10, *O*-acetylserine(thio)lyase (OASTL4); 11, cysteine dioxygenase (CDO1); 12, UDP-sulfoquinovose synthase (SQD1); 13, UDP-sulfoquinovose:diacylglycerol sulfoquinovosyltransferase (SQD2a); 14, extracellular polypeptide 76 (ECP76); 15, selenium-binding protein (SBDP); 16, selenium-binding protein (SBDP) (15 and 16 represent the same gene); 17, SAC1-like protein; and 18, SAC1-like protein.

the lower panel of Fig. 2, there was very little increase in the levels of many of these transcripts in the *sac1* mutant, suggesting that these genes may be under the control of SAC1 (either directly or indirectly).

Photosynthesis genes. Many transcripts encoding proteins involved in photosynthesis decline by 12 to 24 h of S deprivation in both wild-type cells and the *sac1* mutant. This decline is often more severe in the *sac1* mutant strain, although the levels

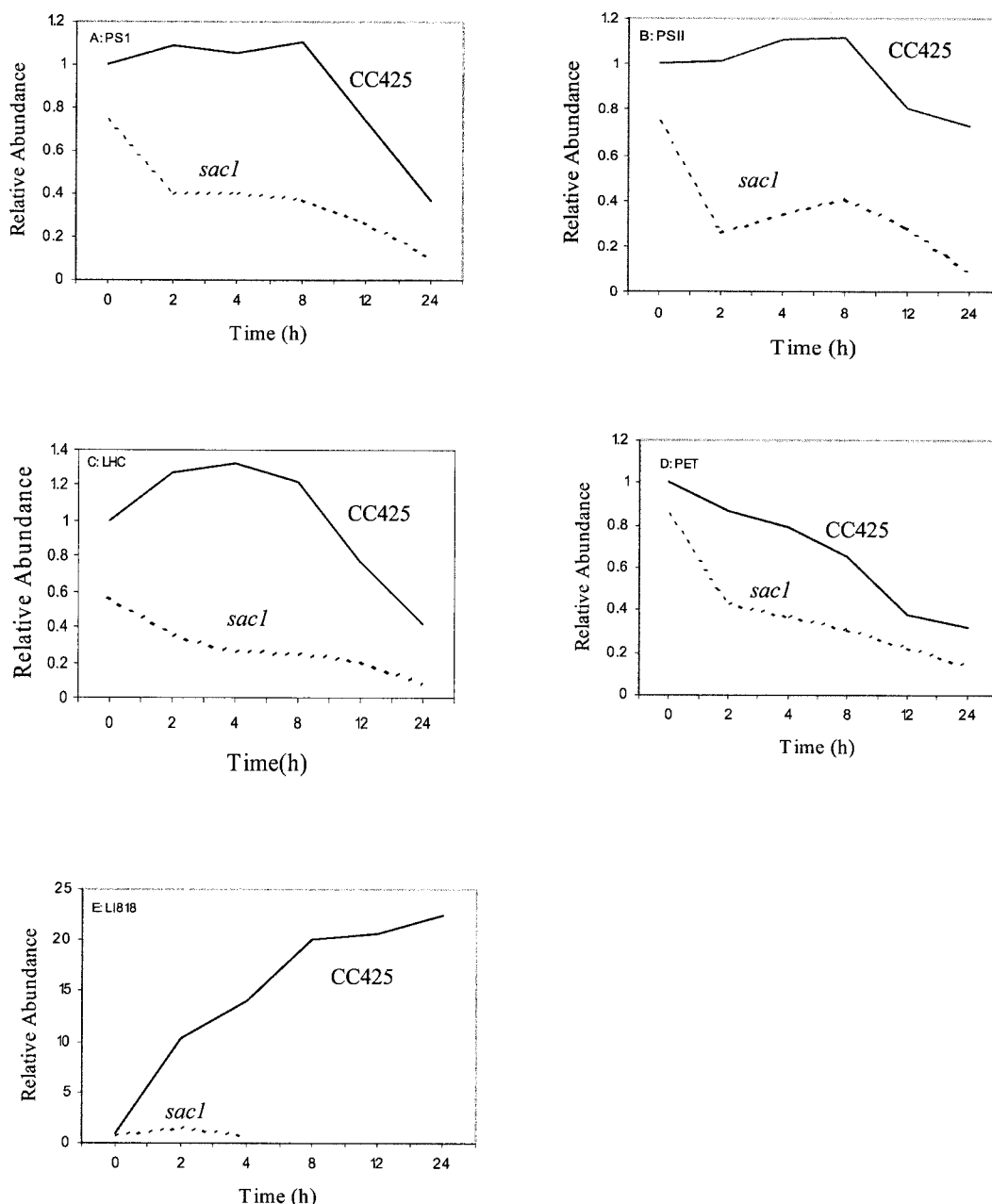


FIG. 3. Influence of S deprivation on the abundance of transcripts encoding proteins that function in photosynthesis in both CC425 and the *sacI* mutant strains. The average values represented on the graphs are for each time point for all of the transcripts listed in the different categories, given below. All of the values are relative to the time zero value of RNA from CC425, which is set at 1. Transcripts encode the following proteins. (A) PS I: PSAD (963047E03), PSAE (894083B07), PSAF (894041H01), PSAG (894065A07 and 894100A05), PSAH (894014A05), PSAK (894086C09), PSAL (894004A09), and PSAO (894019E07). (B) PS II (oxygen-evolving complex): PSBO (894068A11), PSBP (894006E05), and PSBQ (963041E04). (C) LHC: LHCA1 (963024B11), LHCA2 (963047H05), LHCA3 (894033H06), LHCA5 (963042A01 and 894041D11), LHCA6 (894044B07), LHCA7 (894076B06), LHCA8 (894087C09), LHCA9 (894078C01), LHCBM1 (963069C06), LHCBM3 (894080G01), LHCB (894052A01), and LHCB4 (894062E07). (D) Photosynthetic electron transport: PETC (894100F04), PETE (894069E01), PETN (894089E08), PETF1 (894017C09), PETF5 (963046B11), PETM (963053C08), PETO (963092G08 and 894002C07), and FNR (963025F07). (E) LI1818-1 (894097E05).

of these transcripts are somewhat lower in the mutant strain even under nutrient-replete conditions. The graphs presented in Fig. 3 represent an average of the change in transcript abundances of genes encoding subunits of the different complexes of the photosynthetic apparatus; the values for the tran-

script levels used in these analyses are presented in Table 1 (under Photosynthesis) and Table S1 in the supplemental material. On average, the levels of transcripts for genes encoding subunits of photosystem I (PS I) (Fig. 3A; Table 1), PS II (Fig. 3B; Table 1), cytochrome b_{6f} (Fig. 3C; Table 1), light-harvest-

ing complex (LHC) polypeptides (Fig. 3D; Table 1), and chloroplast ATP synthase (Table 1) either increase slightly or stay approximately the same as the level observed for nutrient-replete CC425 cells during the first 8 h of S deprivation. However, after 12 and 24 h, the transcript levels usually decline to 25 to 50% of the prestress level (Fig. 3). The only exceptions on the whole microarray were the *PSBR* and *PSBP3* genes (the changes in transcript levels for these genes did not make the threefold cutoff but were added to Table 1), coding for PS II polypeptides that remained nearly constant in both strains over the time course. For PS II and cytochrome *b₆*, some of the genes (*PSB28* and *PETO*) declined before the others, suggesting that they may play a role in the inactivation or disassembly of these complexes under stress conditions.

Interestingly, nearly all of the transcripts encoding components of the photosynthetic complexes exhibited an earlier and more pronounced decline in the *sac1* mutant than in the parental strain, and this decline could be highly significant even after 2 h of S starvation. These results suggest that SAC1 may influence the stoichiometry of complexes involved in the photosynthetic electron transport chain, possibly in an indirect manner. The overall rapid downregulation of photosynthesis genes during S deprivation of the *sac1* mutant may reflect the extreme stress that this strain experiences as it is unable to acclimate to S deprivation.

In contrast to the results described above, the level of transcripts for LhcSR2 (LI818r-1), an unusual member of the LHC gene family with unknown functions, increased dramatically in CC425 but not in the *sac1* mutant during S deprivation (Fig. 3E; Table 1). Some of the LhcSR2 transcript data for the *sac1* mutant is not shown in the figure because the normalized net intensity for the median is below 350 pixels for both the red and green channels, the criterion used to filter dim spots from further analysis. One aspect of the SAC1-dependent responses may be to restructure the light-harvesting apparatus to accommodate a reduction in electron flow caused by S deprivation; LhcSR2 may be involved in this process.

Reorganization of C metabolism. A decline in photosynthetic electron transfer is an early event in the suite of responses exhibited by *C. reinhardtii* during S starvation, occurring even before the major complexes cease to be synthesized. This result is exemplified by early declines in the levels of mRNAs encoding plastocyanin, two ferredoxins, and a ferredoxin-thioredoxin reductase subunit. In addition, transcripts for polypeptides associated with the Calvin-Benson cycle also decline during S deprivation (*PRK*, *RBCS1/2*, *TPI*, and *PGK*), while the level of serine hydroxymethyltransferase mRNA becomes elevated, suggesting a stimulation of photorespiration, which can participate in photoprotection (59).

There is an increase in the level of transcripts encoding polypeptides of the oxidative pentose phosphate cycle (*GND*, *TAL1*, and 6-phosphogluconolactonase) and cytosolic glyceraldehyde 3-P dehydrogenase after a transient decline. In addition, two types of complementary changes are observed that tend to reduce the redox pressure on the photosynthetic electron transfer chain, a predictable effect of cessation of growth in an environment in which the cells are absorbing considerable excitation energy. First, transcripts for polypeptides that function in starch synthesis accumulate; these include starch synthase and starch phosphorylase. Second, the *AOX4* (or

PTOX1) transcript, which encodes a chloroplast alternative oxidase, rapidly increases. This enzyme diverts electrons away from the photosynthetic electron transfer chain when carbon fixation is limited (5, 6). Together, these responses may serve as electron valves that help control redox poising in the thylakoid membrane and allow for the maintenance of an appropriate cellular redox environment. Not all alternative electron pathways are stimulated during S deprivation, as demonstrated by the findings that levels of transcripts encoding the mitochondrial *AOX1* and the chloroplast Fe-hydrogenase declined.

Stress genes. Several transcripts involved in limiting the potential damage caused by oxidative stress increase in both CC425 and the *sac1* mutant following exposure of the cells to S deprivation. Within this group are transcripts encoding glutathione peroxidase (*GPX1*; 963046E06), the chloroplast iron superoxide dismutase (*FSD1*; 894081G12), a putative thioredoxin/periredoxin (894082E09), and serine hydroxymethyltransferase (*SHMT1*; 963029F05) (Table 1). *GPX* helps limit oxidative damage by catalyzing the reduction of H_2O_2 , lipid peroxides, and organic hydroperoxides, while the chloroplast-localized *FSD1* catalyzes the dismutation of superoxides which can form when the photosynthetic apparatus is absorbing excess excitation energy.

Metabolic stress usually triggers activation of the chaperone systems, which function in protein repair and remodeling. However, each chaperone system will react differently, depending on its functions, substrates, and subcellular localization. Compared to the relatively stable transcript levels associated with mitochondrial *HSP70C*, the modest increase in mRNA associated with the chloroplast *HSP70B* isoform may be significant with respect to the response of the organelle to stress conditions. The observed increase in transcript accumulation for *VIPP1* (vesicle-inducing protein in plastids), a coiled-coiled protein involved in the biogenesis of thylakoid membranes (23), may be indicative of increased turnover of thylakoid lipids and proteins during S deprivation. The transcript for *HCF136*, a chaperone of unknown function present in the thylakoid lumen and necessary for PS II biogenesis (30), significantly increases, as do transcripts encoding two of the three chloroplast 22-kDa heat shock proteins. These *HSP22* polypeptides are similar in sequence and are encoded by genes oriented head-to-head on the genome. In contrast, the transcript encoding the *CPN60* subunits of the chloroplast chaperonin exhibits a marked decline, suggesting a reduction in chloroplast translation and the synthesis of organellar complexes. The levels of transcripts for several peptidyl-prolyl *cis-trans* isomerases of the cyclophilin family, involved in protein folding and other functions, are reduced. Note also the dramatic decrease in the level of mRNA for the cytosolic *HSP90A*. The transcripts encoding several proteases of the aspartic, cysteine, and serine (a chloroplast *DEGP*) types increase in both CC425 and the *sac1* mutant during S deprivation (Table 1 and Table S1 in the supplemental material). These findings could reflect initiation of specific protease-dependent regulatory processes and/or the need to redistribute the amino acid resources of the cell. Furthermore, the finding that the transcript levels for a number of these genes appear to be properly regulated in the *sac1* mutant, at least over a short term following the initiation of S deprivation, again suggests

that some regulatory aspects of the acclimation response still operate in the *sac1* mutant.

There are also several transcripts encoding stress-related polypeptides that attain high levels in CC425 but not the *sac1* mutant. One of these transcripts encodes a putative vanadium chloroperoxidase (963032D03), an enzyme that reduces alkyl peroxides, potentially repairing damaged molecules in the cell (56). Other transcripts that exhibit a SAC1-dependent increase encode an aspartyl aminopeptidase (963065A08), LCI5 (894082D09), which is also elevated during CO₂ limitation, and the nucleosome assembly protein I-like protein (963046C02). The levels of these transcripts may be directly controlled by SAC1, like those of most of the sulfur assimilation genes.

There are also stress-associated transcripts that increase in the *sac1* mutant but not in CC425. An increase in some of these might reflect the fact that the mutant strain cannot acclimate to S deprivation, which could result in an extreme stress response. Interestingly, some of the transcripts in this category encode components of the ubiquitin and proteasome protein degradation machinery. There is an increase in levels of transcripts encoding two E2 ubiquitin-conjugating enzymes, a SKP1-like E3 ubiquitin ligase and the proteasome α -6 subunit (PAF1) (Table 1 and Table S1 in the supplemental material). In the case of the chloroplast 22-kDa heat shock proteins, the increase in their mRNA levels is much stronger in the *sac1* mutant than in CC425; the increase is as high as 19- and 42-fold after 24 h of S deprivation in the mutant strain. Previous studies have suggested that the absorption of excitation energy by the photosynthetic apparatus is lethal to the *sac1* mutant strain during S deprivation, and death of the cells is observed between 1 and 2 days after they are transferred to medium devoid of S. In contrast, wild-type cells survive under these conditions for prolonged periods of time (8). The sharp increases in HSP22E and HSP22F transcript abundance occur when the cells are beginning to lose viability.

Other significant changes. Examination of Table 1 shows that there are significant declines in the levels of many transcripts encoding chloroplast ribosomal polypeptides and that this decline is often more pronounced in CC425 than in the *sac1* mutant. Another intriguing case relates to the CHL27A and CHL27B polypeptides (also known as CTH1 and CRD1 [31, 33]), two isoforms of the oxidative Mg-ProtoIX monomethyl ester cyclase involved in chlorophyll synthesis (53). While transcripts for the latter severely decline in both strains, transcripts for the former markedly increase, even after 2 h of starvation. This increase is reduced to some extent in the *sac1* mutant, suggesting that SAC1 participates in regulation of the abundance of this mRNA. The expression patterns of these genes appear to be orthogonal, with CHL27B being expressed in Cu-limited, hypoxic conditions and CHL27A being expressed in Cu-replete, oxygenic conditions (33). Although the CHL27B and CHL27A polypeptides are very similar and appear to have similar functions, they are not able to completely substitute for each other with respect to the formation of pigment-protein complexes in the photosynthetic apparatus (32). CHL27B and CHL27A may have distinct functions in the delivery of chlorophyll to different complexes of the photosynthetic apparatus and help tailor pigment binding and photosynthetic function to different environmental conditions.

Another intriguing link with Cu metabolism is provided by

the observation that the transcript for CUTA1, a chloroplast copper-binding protein, is reduced in CC425 but significantly elevated in the *sac1* mutant. Also, while levels of transcripts for many regulatory factors decrease during S deprivation, some, including those encoding an AAA-type ATPase of unknown function (894037G11), an elongation factor (894081B08), and a RAB-type GTPase (963038D05), appear to increase. It should be emphasized that the function of these potential gene products need to be more clearly established before they can be ascribed a role in the acclimation process.

Finally, it is interesting that numerous transcripts for genes encoding polypeptides of unknown function increase dramatically during S deprivation. The full list of those genes whose transcripts satisfied the ≥ 3 -fold filter is given in Table S2 in the supplemental material. A short list of 10 of the unknowns that exhibit very high transcript levels are given in Table 1 and include 963016B04 (23-fold at 12 h), 963033D03 (24-fold at 12 h), 963032A08 (50-fold at 12 h), 963047F05 (18-fold at 12 h), 963096E10 (35-fold at 12 h), 894063D10 (15-fold at 12 h), 963027G01 (23-fold at 12 h), 963063B08 (22-fold at 12 h), 963068A11 (18-fold at 12 h), and 963032A01 (23-fold at 12 h). There are many other transcripts that exhibit statistically significant increases in abundance during S deprivation, and some of these appear to be under the control of SAC1 (e.g., 894063D10). This result emphasizes how little we know about acclimation processes and also presents us with the challenge of identifying these unknown genes and defining their function in the acclimation process.

DISCUSSION

Examination of gene expression using high-density DNA microarrays is a potent means of examining how organisms modulate gene activity in response to environmental conditions. We have used this technology to explore the ways in which *C. reinhardtii* acclimates to S deprivation. While there are approximately 2,565 genes represented on the arrays used for these experiments (slightly less because of some duplications on the array), the array elements were derived from cDNAs defined in recombinant libraries from cells grown both under nutrient-replete conditions (Core Library) and under S, phosphorus, and nitrogen starvation conditions (Stress I Library) (48). Therefore, the microarray was likely to contain a number of stress-responsive genes. This probability was substantiated by the findings that both the *ARS* and *ECP76* genes were represented on the array (present only in the Stress I and not the Core Library); these genes only become active when *C. reinhardtii* cells are starved for S (8, 11, 51). The presence of numerous stress-related array elements was also substantiated by the results of the experiments in which 171 of the represented genes exhibited a ≥ 3 -fold change in gene expression in CC425 after the cells were placed into medium devoid of S; the levels of transcripts encoded by many of these genes increased.

The patterns of gene expression in *C. reinhardtii* cells following S deprivation reveal both the general and specific responses that enable a cell to survive extended periods of time under specific deprivation conditions. Furthermore, the use of mutant strains with aberrant acclimation responses that die much more rapidly than wild-type cells following the imposition of the stress are beginning to reveal extreme responses of

cells when their normal acclimation responses become compromised. This is the case for the *sac1* mutant, which exhibits rapid death (relative to wild-type cells) and an inability to accumulate the ARS transcript and transcripts encoding many other polypeptides during S deprivation (8).

There are a number of different generalizations that can be made from the data presented in this study. First, the levels of transcripts for many genes change substantially during S deprivation. A number of the transcripts that are extremely sensitive to S deprivation encode proteins involved in the conservation, acquisition, and assimilation of SO_4^{2-} , including ARS, ECP76, ATS, SAT, OASTL, polypeptides involved in sulfolipid biosynthesis, a selenium-binding protein, and SAC1-like polypeptides (which may be involved in regulation or in the transport of SO_4^{2-} into the cell). Transcripts encoding a number of these polypeptides increase to well over 10-fold the level observed in nutrient-sufficient cells. However, a number of these transcripts are not detected in nutrient-replete cells (e.g., this is the case for the ARS and ECP76 transcripts), which causes imprecision in the quantification, making the ratio values a semiquantitative estimate. Based on our experience, the increase measured in the microarray analyses is often less than that determined by using real time reverse transcription-PCR (54). Furthermore, while in many instances changes in transcript levels reflect changes in corresponding protein and activity levels, as has been shown for ARS and ECP76 (8, 51), this correlation may not always hold. For example, even though transcripts encoding the sulfolipid biosynthesis enzymes SQD1 and SQD2a accumulate during sulfur deprivation (Table 1), the sulfolipid content of *C. reinhardtii* membranes appears to decline (K. Sugimoto and M. Tsuzuki [Tokyo University of Pharmacy and Life Science], personal communication).

Many of the genes, including ARS1, ATS2, APR, SQD2a, and ECP76, that are sensitive to S-limited growth conditions appear to be under SAC1 control, since there is no significant increase in their transcript levels in the *sac1* mutant at any time point analyzed, even the early 2-h and 4-h time points. In other cases (SAT1, OASTL4, ATS1, SIR1, CDO1, SQD1, SBDP, and genes encoding SAC1-like proteins) an increase in transcript levels is observed in the *sac1* mutant, but the increase is generally much more modest and less sustained than that in wild-type cells. These results suggest that SAC1 may play a direct role in the control of a number of genes. While other regulatory mechanisms may result in some changes in transcript levels during the early phase of starvation, sustained expression requires a functional SAC1 gene and may reflect the fact that SAC1 is required to maintain cell vitality. There are a number of other transcripts whose levels are potentially controlled by SAC1, some of which encode proteins that might function in the chloroplast, including a putative vanadium chloroperoxidase (although the similarity to other vanadium chloroperoxidases is not that high), aspartyl amino peptidase, and a specific light-harvesting polypeptide (LhcSR2; see below).

While the levels of transcripts from a number of genes dramatically change in CC425 and are no longer properly regulated in the *sac1* mutant, it is clear that other genes are properly controlled in both the parental and the mutant strains. For example, transcripts for genes encoding the chloroplast quinol oxidase, the glutathione peroxidase, and the putative periredoxin Q increase to a similar extent in CC425 and the *sac1*

mutant strain during the 24 h of S deprivation. Hence, regulatory elements other than SAC1 are involved in modulating S deprivation-elicited changes in gene expression in *C. reinhardtii*. This result is not surprising, since there are mutations in at least four other loci that result in a *sac1* mutant-like phenotype (J. Davies, S. Pollock, W. Pootakham, and A. R. Grossman, unpublished data). Isolating the altered genes and examining how lesions in these genes modulate global transcript levels during S deprivation will help define the nature of the gene products and their potential relationships with each other.

There are also many polypeptides for which the transcript levels dramatically decline in CC425 during S deprivation. Perhaps most striking is the finding that a number of transcripts encoding proteins involved in photosynthesis and chloroplast ribosomal polypeptides decline markedly in CC425 during S deprivation. In the case of the chloroplast ribosomal protein transcripts, the decrease is generally not nearly as great in the *sac1* mutant. These results suggest that it might be critical to control the activity of the chloroplast ribosomes during S deprivation and that such control may be conferred to ribosomes by changing the complement of ribosomal proteins. Changes in ribosomal proteins have also been associated with nitrogen starvation (21). Similarly, many transcripts for proteins involved in photosynthesis decline in both CC425 and the *sac1* mutant, but the decline is generally more precipitous in *sac1* mutant cells. Such transcripts include those encoding polypeptides of PS I, PS II, the photosynthetic electron transport chain, and the LHC. While the reason for the earlier decline in the *sac1* mutant is not clear, it may reflect the inability of the cells to scavenge S from the environment and tune cellular metabolism to decreased nutrient availability, which could elicit a highly stressed condition and nonspecific degradation of many of the less stable transcripts.

A highly stressed state of the S-depleted *sac1* mutant is also suggested by elevated levels of specific transcripts encoding proteins associated with stress conditions. For example, the *sac1* mutant exhibits a dramatic increase in transcripts encoding two putative chloroplast 22-kDa heat shock proteins (18); the increase in transcripts encoding these polypeptides in CC425 is much less pronounced. This type of chaperone is involved in preventing protein denaturation and aggregation in plastids. Furthermore, *sac1* mutant cells exhibit an increase in transcripts for proteins involved in proteolysis relative to CC425 cells, although the increases are not that marked. Some of these transcripts encode proteins of the ubiquitin and proteasome systems, including two E2 ubiquitin-conjugating enzymes, an E3 ubiquitin ligase, and a proteasome α -6 subunit (Table 1). There is also a significant increase in a cathepsin Z protease.

While, as mentioned previously, transcripts for many PS and LHC polypeptides decrease to some extent in CC425 and more dramatically in the *sac1* mutant during S deprivation, there is one exception to this trend. The transcript encoding the LhcSR2 polypeptide increases in CC425 nearly 20-fold relative to the level observed in unstarved cells. This increase is not observed in the *sac1* mutant (indeed, the level of the transcript is so low in the *sac1* mutant at some of the time points following S deprivation that we could not reliably quantify it). The transcript for this member of the LHC gene family has been pre-

viously shown to be transiently expressed following the transfer of cells from darkness to light, which would cause high light stress (2, 43). Hence, the organism is retailoring the photosynthetic apparatus to accommodate S deprivation conditions. The inability of the *sac1* mutant to scavenge S and restructure its metabolic machinery (e.g., the photosynthetic apparatus) during S deprivation may cause severe functional and structural damage, especially within chloroplasts (which may be reflected in the increase in the HSP22-related polypeptide). This conjecture is supported by the facts that nutrient stress causes dramatic changes in the activity of the electron transport chain (60) and that the rapid death of the *sac1* mutant during S deprivation can be ameliorated by placing the strain in the dark or blocking photosynthetic electron flow with specific inhibitors (8).

In sum, experiments presented in this report augment our understanding of how photosynthetic organisms respond to S deprivation and provide clues concerning processes that are important for acclimation. This work also highlights the fact that there are many genes of unknown function whose transcripts markedly increase in response to S deprivation. Furthermore, the study has revealed processes triggered when cells are unable to acclimate to the limitation conditions and as cell viability declines. It is now important to try to understand some of the other changes in transcript levels that have been observed (for which there are no obvious explanations) and whether all of these changes are observed at the level of transcript abundance that reflects changes in protein accumulation and activity.

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