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Long-term acclimation to cadmium exposure reveals extensive phenotypic plasticity in Chlamydomonas

Stanislas Thiriet-Rupert, ^{1,†} Gwenaëlle Gain, ^{1,2} Alice Jadoul, ¹ Amandine Vigneron, ¹ Bernard Bosman , ³ Monique Carnol, ³ Patrick Motte, ¹ Pierre Cardol , ² Cécile Nouet , ¹ and Marc Hanikenne , ¹ **, **

- 1 InBioS-PhytoSystems, Functional Genomics and Plant Molecular Imaging, University of Liège, 4000 Liège, Belgium
- 2 InBioS-PhytoSystems, Genetics and Physiology of Microalgae, University of Liège, 4000 Liège, Belgium
- 3 InBioS-PhytoSystems, Laboratory of Plant and Microbial Ecology, University of Liège, 4000 Liège, Belgium

Research Article

M.H. conceived and directed the research. M.H., S.T., C.N., A.J., G.G., and P.C. designed experiments. A.J., G.G., S.T., C.N., P.C., B.B., and M.C. performed experiments. S.T., G.G., M.H., P.C., A.J., P.M., and A.V. analyzed and interpreted the data. S.T., G.G., and M.H. generated all Figures and Supporting Information material. S.T., M.H., and P.C. wrote the manuscript, with comments of all authors.

The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (https://academic.oup.com/plphys/pages/general-instructions) is Marc Hanikenne.

Abstract

Increasing industrial and anthropogenic activities are producing and releasing more and more pollutants in the environment. Among them, toxic metals are one of the major threats for human health and natural ecosystems. Because photosynthetic organisms play a critical role in primary productivity and pollution management, investigating their response to metal toxicity is of major interest. Here, the green microalga Chlamydomonas (*Chlamydomonas reinhardtii*) was subjected to short (3 d) or chronic (6 months) exposure to 50 µM cadmium (Cd), and the recovery from chronic exposure was also examined. An extensive phenotypic characterization and transcriptomic analysis showed that the impact of Cd on biomass production of short-term (ST) exposed cells was almost entirely abolished by long-term (LT) acclimation. The underlying mechanisms were initiated at ST and further amplified after LT exposure resulting in a reversible equilibrium allowing biomass production similar to control condition. This included modification of cell wall-related gene expression and biofilm-like structure formation, dynamics of metal ion uptake and homeostasis, photosynthesis efficiency recovery and Cd acclimation through metal homeostasis adjustment. The contribution of the identified coordination of phosphorus and iron homeostasis (partly) mediated by the main phosphorus homeostasis regulator, Phosphate Starvation Response 1, and a basic Helix-Loop-Helix transcription factor (Cre05.g241636) was further investigated. The study reveals the highly dynamic physiological plasticity enabling algal cell growth in an extreme environment.

Introduction

The release of trace metal elements by anthropogenic activities into the environment is one of the most significant threats to human health, as well as water and soil ecosystems (Nagajyoti et al., 2010). Essential metals [iron (Fe),

copper (Cu), zinc (Zn), manganese (Mn), etc.] are micronutrients of critical importance for the correct functioning of living cells through their roles as cofactors for many enzymes (Palmer and Guerinot, 2009; Nouet et al., 2011;

^{*}Author for communication: marc.hanikenne@uliege.be

[†]Present address: Unité de Génétique des Biofilms, Département Microbiologie, Institut Pasteur, Paris, France.

Senior author

Thomine and Vert, 2013). However, these essential micronutrients are toxic at supraoptimal concentrations. In contrast, nonessential metals, such as lead (Pb), mercury (Hg), or cadmium (Cd), have toxic effects at low concentrations (Warren, 1989). With the exception of the diatom *Thalassiosira weissflogii* grown under Zn deficiency, where Cd can act as a cofactor of a carbonic anhydrase (Lane and Morel, 2000; Alterio et al., 2015), these nonessential metals have no known physiological function.

Cd is highly toxic, especially upon long-term (LT) exposure to low concentrations (Satarug et al., 2003, 2010), with carcinogenic, embryotoxic, teratogenic, and mutagenic properties (Hartwig, 2010) and has been associated to severe health disorders in human (Pan et al., 2010; Bernard, 2008). Widespread low-level contamination of soil and water is stemming mostly from anthropogenic sources, e.g. mining, nonferrous metal manufacturing, or the application of phosphate fertilizers and sewage sludge on soils. Although Cd emissions into the environment have declined in many industrialized countries, it is still prevalent in emerging countries and low-level contamination of soils still results in important contamination of the food chain (Clemens et al., 2013). Although Cd has multiple harmful effects, affecting the cell from its ultrastructure to its metabolism, the mechanisms underlying Cd toxicity are not clear. Cd exposure seems to induce oxidative stress (Stohs and Bagchi, 1995; Nagajyoti et al., 2010; Yadav, 2010), but since Cd is a nonredox-active metal (Stoiber et al., 2013), reactive oxygen species (ROS) are produced indirectly. Lipid, protein, and nucleic acid damages were also reported under Cd exposure (Watanabe et al., 2003) and, in photosynthetic organisms, growth inhibition and a decrease of chlorophyll (Chl) content were observed (Nagajyoti et al., 2010; Yadav, 2010).

Investigating the response of photosynthetic organisms to metal toxicity is cardinal because of their critical role in primary productivity and pollution management (Pinto et al., 2003; Monteiro et al., 2011; Rawat et al., 2011). In this context, Chlamydomonas (Chlamydomonas reinhardtii), as a model unicellular photosynthetic organism, is a highly relevant experimental model (Hanikenne, 2003; Merchant et al., 2007). Cd toxicity in Chlamydomonas is characterized by ultrastructural modifications including starch accumulation, cytoplasmic vacuolization, formation of cytoplasmic electron-dense granules (Nishikawa et al., 2003; Aguilera and Amils, 2005), cell volume increase, development of membranous organelles, or cell-wall volume increase (Visviki and Rachlin, 1994), as well as growth inhibition (Prasad et al., 1998; Hanikenne et al., 2001, 2005b; Jamers and Coen, 2009). Since a fraction of Cd accumulates in the unique chloroplast of Chlamydomonas (Nagel et al., 1996), functions related to this organelle are strongly affected. Indeed, about one-third of the soluble proteins involved in photosynthesis decrease upon Cd exposure, including Rubisco, ferredoxin, ferredoxin-NADP⁺ reductase, and enzymes involved in the Calvin cycle and Chl biosynthesis (Gillet et al., 2006). Moreover, Cd²⁺ ions were shown to compete for binding to Photosystem II (PSII) calcium (Ca²⁺) site during the assembly of the water-splitting complex (Faller et al., 2005). These disturbances of plastid-related functions result in impaired photosynthesis (Faller et al., 2005; Vega et al., 2006; Perreault et al., 2011). An impact on proteins involved in carbohydrate and lipid metabolism, amino acid, and protein biosynthesis was also identified (Gillet et al., 2006).

So far, no transporter specific to Cd^{2+} ions has been identified. However, as a divalent cation, Cd2+enters the cell via essential metal transporters (Shahid et al., 2017). For instance, in Arabidopsis (Arabidopsis thaliana), Cd can be transported by Iron-Regulated Transporter 1 (IRT1), a ferrous ion transporter of the Zrt-, Irt-like proteins (ZIPs) family, and that an increase of Fe availability induces a decrease of Cd uptake (Rogers et al., 2000; He et al., 2017). In Chlamydomonas, the homolog ZIP transporters IRT1 and IRT2 are induced by Fe deficiency, suggesting an involvement in Fe uptake (Hanikenne et al., 2009; Urzica et al., 2012). Similar to plants, ATP-binding cassette (ABC) transporters were also involved in Cd tolerance in Chlamydomonas (Hanikenne et al., 2001; Faller et al., 2005; Hanikenne et al., 2005b; Wang and Wu, 2006). A higher concentration of Ca, cobalt, or Zn in the culture medium provides protection to Chlamydomonas by decreasing Cd uptake (Mosulén et al., 2003; Lavoie et al., 2012). This competition of Cd for divalent cation transporters perturbs essential metal homeostasis of the cell, which in turn impacts the whole cell functioning.

When exposed to pollutants such as Cd, organisms are able to induce a variety of stress mechanisms allowing acclimation. In Chlamydomonas, the first level of defence is the hydroxyproline and glycoprotein-rich cell-wall, which has high affinity for metal cations (Collard and Matagne, 1990). However, this passive barrier seems to be saturated at environmental Cd concentration (Macfie et al., 1994) and cell-wall-deficient strains accumulate less Cd than the wild-type (WT) (Macfie et al., 1994; Penen et al., 2017). Once inside the cell, different types of mechanisms are used to cope with Cd. Antioxidant defence is enhanced, with the induction of the expression and activity of antioxidant enzymes (Gaur and Rai, 2001; Pinto et al., 2003; Aksmann et al., 2014). Strategies including active efflux, bioconversion into less harmful forms, such as the intracellular chelation by thiolated peptides like phytochelatins, were also widely studied (Howe and Merchant, 1992; Gaur and Rai, 2001; Hanikenne et al., 2005b; Wang and Wu, 2006; Lavoie et al., 2009; Bräutigam et al., 2011). Other effective intracellular chelators used by cells to sequester metal ions are polyphosphates (Rachlin et al., 1984; Seufferheld et 2008; de Lima et al., 2013). In the microalgae Chlamydomonas acidophila and Chlamydomonas, polyphosphates are thought to bind Cd in the cytoplasm allowing its sequestration into acidic vacuoles (Nishikawa et al., 2003; Blaby-Haas and Merchant, 2014; Penen et al., 2017). More generally, a crosstalk seems to connect metal and phosphorus (P) homeostasis since intracellular concentrations of essential metal under Cd exposure differ depending on P availability in Chlamydomonas (Wang and Dei, 2006; Webster et al., 2011).

Little data are available on the effect of LT, chronic, exposure to Cd in Chlamydomonas (Yu et al., 2018, 2020). In this study, we used growth assays, ionomic and transcriptomic profiling, photosynthesis analysis, as well as mutant characterization to examine in detail the response to 6-month Cd exposure compared to a 3-d Cd exposure. We examined whether a chronic exposure triggers acclimation mechanisms allowing algae to better face Cd-induced toxicity. We also evaluated the reversibility of these responses. We confirmed that the cell wall and the photosynthetic apparatus are key targets of Cd and showed that coordinated regulation of the P and Fe homeostatic networks is required for proper acclimation to Cd.

Results

Cultures of Chlamydomonas were grown for 6 months in liquid TAgP medium in the absence or presence of 50 μ M Cd. TAgP is a modification of the classical Tris-Acetate-Phosphate (TAP) medium (Harris et al., 2009) where inorganic phosphate was replaced by β-glycerophosphate (TAgP) to avoid Cd-P complex formation (Collard and Matagne, 1990). Following this acclimation time, the response of cells to four growth conditions were compared: (1) control condition (Ctrl, six months growth without Cd); (2) control cultures exposed for 3 d to 50 µM Cd (shortterm [ST] exposure); (3) 6 months acclimated cultures (LT exposure); and (4) the exact same acclimated cultures subsequently grown for three days in the absence of Cd (Recovery) to investigate the recovery phase upon Cd removal (Supplemental Figure S1). Note that "ST" is used here in opposition to "LT" and already represents an acclimation response in a unicellular organism dividing every 8-12 h. Detailed phenotyping was then conducted to examine the impact of these treatments on algae physiology, including cell growth, total Chls, cell morphology, ionomic profile, and photosynthesis efficiency, supported by a transcriptomic analysis by RNA sequencing (RNA-Seq).

Pre-acclimation decreases the phenotypic impact of Cd exposure

ST exposure (50 μ M Cd) during 3 d resulted in 30% lower biomass relative to the Ctrl. In contrast, this reduction was only \sim 12% upon LT exposure, similar to the recovery condition (Figure 1A; Supplemental Figure S2).

In agreement with previous reports showing that Cd induced chlorosis (Nagajyoti et al., 2010; Yadav, 2010), a 30% decrease of total Chl content was observed after ST exposure, whereas the LT acclimated cultures had similar Chl levels than the Ctrl cells (Figure 1B). Recovery induced a 15% increase of the Chl content (Figure 1B).

Cd induced the formation of cell aggregates, upon both ST and LT exposures, which persisted upon recovery (Supplemental Figure S3A). In addition, groups of two or four cells surrounded by a common matrix were observed in these three conditions. Such formations were previously observed in Chlamydomonas cultures exposed to stress

conditions and are called palmella (Figure 1, C and D; Supplemental Figure S3B; Iwasa and Murakami, 1969; Lurling and Beekman, 2006; Jamers and Coen, 2009; Boyd et al., 2018; Samadani and Dewez, 2018; Cheloni and Slaveykova, 2021; Li et al., 2021). Palmella are originated from dividing cells that remain grouped within the matrix after mitosis (Nakamura et al., 1975). Very few of such structures were observed in the Ctrl, but they appeared in ST cultures, became largely dominant upon LT exposure to Cd and their frequency decreased during recovery (Figure 1, C and D). In addition, loss of motility and increased cell volume were observed in presence of Cd, as described (Visviki and Rachlin, 1994; Hessen et al., 1995; Khona et al., 2016).

Next, the effects of Cd on the ionome of the cells (Salt et al., 2008) were examined. Cellular Cd concentrations [~0.85 nmol/optical density measurement at 750 nm (OD₇₅₀)] were equivalent in ST- and LT-treated cells and no Cd was detected in cells both in Ctrl and after recovery (Figure 1E). Compared to Ctrl, ST, and LT exposed cells shared a similar ionome profile with higher Cu, Zn, and Mn, but lower Ca and Mg concentrations (Figure 1F). Cd exposure only slightly increased Fe cellular concentration, whereas K concentrations were either lower (ST) or higher (LT) than in Ctrl (Figure 1F). The effects of Cd on the ionome were almost reversed upon recovery, with only K remaining high and Ca readjusting to a higher level.

Some differences in the ionome of ST and LT exposed cells were also observed. Although Fe was only moderately affected, ST-exposed cells contained significantly lower Ca, K, Mg, Zn, and Cu, but higher Mn. This dynamic suggested that the response initiated at ST progressed after LT exposure reflecting either an aggravation of Cd-induced deleterious effects, or an acclimation.

Cd deeply remodels the transcriptome

To examine the molecular mechanisms underlying the phenotypic responses to Cd, three biological replicate samples for each of the four growth conditions were subjected to a transcriptomic analysis using RNA-Seq. A principal component analysis (PCA) of gene expression profiles in each condition showed a clear segregation of the samples (Figure 2A). PC1 and PC2 explained 59% and 30% of the gene expression variance by separating conditions according to (1) the presence or absence of Cd in the culture medium (PC1) or (2) whether or not the cells were pre-acclimated to Cd (PC2), respectively.

Then, differentially expressed genes (DEGs; \log_2 fold-change >1 or less than -1; false discovery rate (FDR) adjusted P < 0.05) were identified in four specific comparisons (Figure 2B; Supplemental Dataset S1 and Supplemental Figure S4): between ST (1) or LT (2) exposed cells and Ctrl, characterizing the ST and LT responses to Cd exposure, respectively, and between 3 d recovered cells and LT exposed (3) or Ctrl (4) cells, characterizing the recovery response to Cd exposure or the residual response to Cd exposure following recovery, respectively.

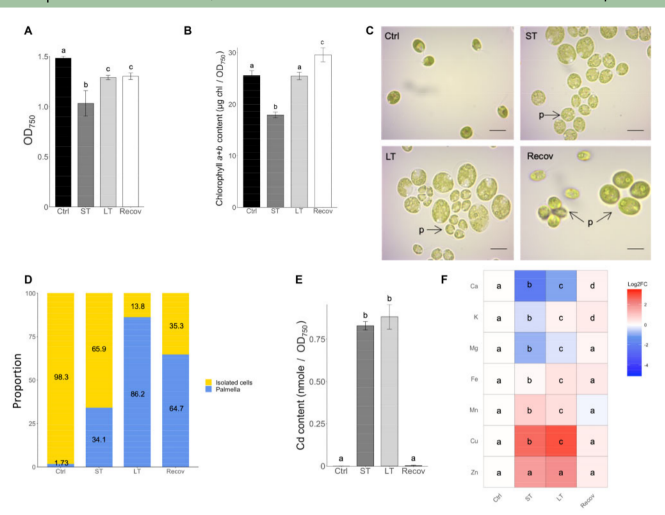
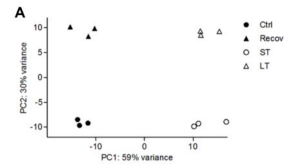


Figure 1 Phenotypic characterization of the control and Cd-exposed cultures. Control cultures grown for 6 months without Cd (Ctrl), ST (3 d) Cd-exposed cultures, LT (6 months) Cd-exposed cultures and recovery cultures (6 months Cd-exposed cells grown for 3 d without Cd) were characterized. A, Biomass estimation and (B) Chl content after 3 d of treatment. Values are means \pm so of three biological replicates. Significant differences (P < 0.05) according to one-way ANOVA followed by Tukey's test are shown by different letters. C, Microscopic observations of each culture after 3 d of treatment, palmella (p) are indicated by an arrow. Scale bars = 15 μ m. The pictures are representative of three biological replicates. D, Proportion of palmella and isolated cells in each condition after 3 d of treatment. Values are from the observation of 10 fields at the microscope (n > 900). E, Cell Cd concentrations after 3 d of treatment. F, Heatmap of the log₂ fold-change of mineral element concentrations in ST, LT, and Recovery cultures relative to the Ctrl. Values are means (\pm so in E) of three biological replicates and statistical differences are shown with different letters, according to one-way ANOVA followed by Tukey's test.

Interestingly, shared DEGs were identified among the different responses allowing their classification into seven specific groups of genes (Figure 2C, Gp 1-7). Groups 1 and 2 represented DEGs characterizing the ST and LT responses to Cd exposure, respectively, whereas group 3 contained DEGs common to both ST and LT responses. The latter characterized the core acclimation response to Cd exposure, while group 2, containing nearly half of all identified DEGs, represented genes progressively regulated by Cd. DEGs of group 4 were specific to the recovery response, being regulated by the transition from LT exposure to an absence of Cd. Group 5 genes were common to ST and recovery response, possibly related to the acclimation to ST changes in culture conditions. Group 6 corresponded to genes specifically regulated upon Cd removal, whereas group 7 included genes, which remained differentially regulated compared to Ctrl 3 d after recovery.

To gain further insight into the mechanisms involved in each of the described responses, a Gene Ontology (GO) enrichment analysis was carried out (Supplemental Dataset S2), completed by a detailed manual examination of the DEGs (Table 1). Special attention was paid to functions related to the phenotypic traits shown above to be impacted by the treatments (Figure 1).

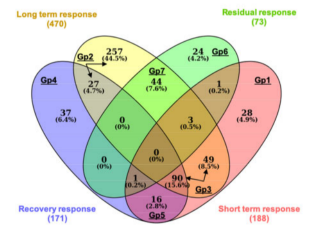
Within the 188 DEGs of the ST response, a notable enrichment was found for metal ion transport (P=0.0003), Fe ion transport (P=0.01), superoxide metabolic process (P=0.04), and calmodulin-dependent protein kinase activity (P=0.0001; Supplemental Dataset S2). Moreover, six genes involved in cell-wall synthesis, eight genes involved in chloroplast- and photosynthesis-related processes, six genes responsible for Fe homeostasis and many transporters were highlighted by the manual analysis (Table 1). Nine genes



В

	Up	Down	Total
ST	103	85	188
LT	184	286	470
Recovery	82	89	171
Residual	37	36	73

C



Gp1: ST response Gp2: LT response

Gp3: Cd acclimation
Gp4: Recovery response

Gp5: Response to habitat modification

Gp6: Response to Cd removal

Gp7: Remaining response after recovery

Figure 2 Transcriptomic analysis of the Cd ST, LT, and recovery responses. A, PCA of RNA-Seq gene expression data of control (Ctrl) cultures without Cd, ST (3 d) Cd-exposed cultures, LT (6 months) Cd-exposed cultures, and recovery cultures (6 month Cd-exposed cells grown for 3 days without Cd). B, Number of up-, downregulated, and DEGs (\log_2 fold-change >1 or less than -1; FDR adjusted P < 0.05) in ST (control versus 3 d Cd-exposed cells), LT (control versus 6 month Cd-exposed cells), Recovery (6 month Cd-exposed cells versus 6 month Cd-exposed cells grown for 3 d without Cd) responses. C, Venn diagram of the identified DEGs for the four responses described in (B), and defining seven sub-groups (Gp) of genes and their potential contribution in those responses. Numbers are numbers of DEGs for each response.

related to P homeostasis were also identified [six transporters, two phosphatases, and one transcription factor (TF)] and signal transduction was well represented with 15 DEGs (Table 1).

The LT response analysis revealed a persistence of metal ion transport (P = 0.005), Fe ion transport (P = 0.03), and calmodulin-dependent protein kinase activity (P = 0.0008) functional categories, as well as the emergence of other categories, including Cu ion transport (P = 0.003), Fe homeostasis (P = 0.03), porphyrin-containing compound biosynthesis (P = 0.03), and cell-wall catabolism (P = 0.04; Supplemental)Dataset S2). Functions revealed by the manual analysis were shared with the ST response (Table 1). Among the key functions common to ST and LT responses (Gp3, Figure 2C) were cell-wall synthesis, chloroplast, and photosynthesis, Fe homeostasis, ion transport, P homeostasis, and signal transduction (Table 1). The LT response mobilized more genes for these functions, together with a diversification of the functions, illustrating progressive and deepened acclimation to Cd with time, as highlighted by the comparison of Gp1, Gp2, and Gp3 in Table 1.

As more than half of the 171 DEGs of the recovery response were shared with the ST and LT responses, a large overlap of enriched functional categories was observed (Table 1; Supplemental Dataset S2). Interestingly, light harvesting and chloroplastic translation were identified among the 37 Recovery-specific DEGs (Gp4, Figure 2C) by both GO enrichment (P = 0.0004 and P = 0.001, respectively; Supplemental Dataset S2) and manual analyses (Table 1).

Finally, no GO enrichment was observed in the residual response, with the manual analysis revealing 10 genes related to cell-wall synthesis as regulated in this group.

Overall, the transcriptomic analysis pinpointed functions (e.g. Fe homeostasis, photosynthesis, cell wall) that were directly related to the phenotypes observed in the four growth conditions. It also uncovered additional functions (e.g. P metabolism) not investigated so far in our analysis. These functions were then examined in detail to further understand acclimation to Cd exposure in Chlamydomonas.

Chloroplast and photosynthesis are major targets of Cd toxicity

Several DEGs involved in photosynthesis and other chloroplast-related functions were identified in all four studied responses. Only a few photosynthesis-related genes were impacted, mostly downregulated, upon ST Cd exposure (Figure 3A). Among those, only *FNR1* was directly related to the photosynthetic machinery and displayed significantly diminished expression (Figure 3A). *FNR1* encodes a ferredoxin-NADP⁺ reductase, which reoxidizes the photosystem I (PSI) direct acceptor, ferredoxin (FDX1). This was concomitant with a decrease in several photosynthesis-related parameters: the amount of ChI (Figure 1B), the relative amount of active photosystems (PSI and PSII) per ChI (Figure 3B), and the relative electron transport rate (rETR) of PSII (Figure 3C).

Table 1 Manual annotation of the functional categories involved in the Cd ST, LT, and recovery responses

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Functional categories	ST	ST response	LT	LT response	Recovery	Recovery response	Residual	Residual response	Ср1	_	Gp2	Gp3		Gp4	Gp5	J	Срб	Ū	Gp7
	ΛD	Down	Up	Down	η	Down	η	Down	Up D	Down L	Up Down	υρ Down	vn Up	Down	Up Down	Up	Down	ηD	Down
Cell wall																			
Synthesis	2	-	9	12	4	9	4	10	2	-	2 8	2	_	-	1	_	9	_	4
Degradation			2	3				_		-	2 2								_
Chloroplast and photosynthesis																			
Light harvesting				4	7						4		4						
NPQ			-				-											_	
Electron transfer		2	2		2	_					1	_	_		_				
Rubisco activation				-							_								
Chl synthesis				9	_						9								
ChI degradation	-	-	-	_	-	1						1 1							
Carotenoid synthesis				2	-						2								
Apparatus assembly		-		-	_							1							
Prot transport to chloroplast			-		_						_								
Chaperone	_		-		_							2							
Translation		-		-	9						-		5		-				
Other		-			-										-				
Fe homeostasis	9		∞			9			_		1	2							
Transport																			
. Ca			-				_											_	
¥		2		-						_		_							
z				3	-						3								
۵	-	5	-	9	2						-	1 5							
Fe	2		4			4					1	5							
Cu	-		3			2					2	_							
Zn		_	-	_	_						_	_							
Mn	_		_			_						_							
Heavy metals			-								1								
Other	7	3	4	7		_				5	2 6	2 1							
Phosphatase		2		3	2			-				2							_
Transcription	2	3	2	11	3	5		2	-		3 6	3 3		-	1		-		_
Translation		_	-	3			2				1 2	1				2			
Amino acid metabolism	4	2	5	6		2					1 7	4 2							
Metal chelation, phytochelatin	_		-			1						-							
Oxidative stress response	_	_	_	2	2	_					_	1	_						
Carbohydrates																			
Anabolism	_			2	3				_		2		2						
Catabolism	_		2	8	3	_		2			1 7	_		-			_		_
Lipids																			
Anabolism	2	_	7	4	3	_					4	1	_		1				
Catabolism	_	_		8		_	_				7	1		-		_			
Signal transduction	7	8	9	24	2	4	-	2	3	•	3 15	3 8		2			_		_

The numbers of upregulated and downregulated genes belonging to each functional category in the ST (control versus 3 d Cd-exposed cells), LT (control versus 6 month Cd-exposed cells), Recovery (6 month Cd-exposed cells versus 6 month Cd-exposed cells grown for 3 d without Cd), and Residual (control versus 6 month Cd-exposed cells grown for 3 d without Cd) responses, as well as in each sub-group defined in Figure 2C, are indicated. The sign (up or down) of regulation for genes present in groups overlapping with the Recovery response (Gp2, 3, and 5) is indicated according to the ST and LT responses: the upregulated genes in these two responses are downregulated in Recovery.

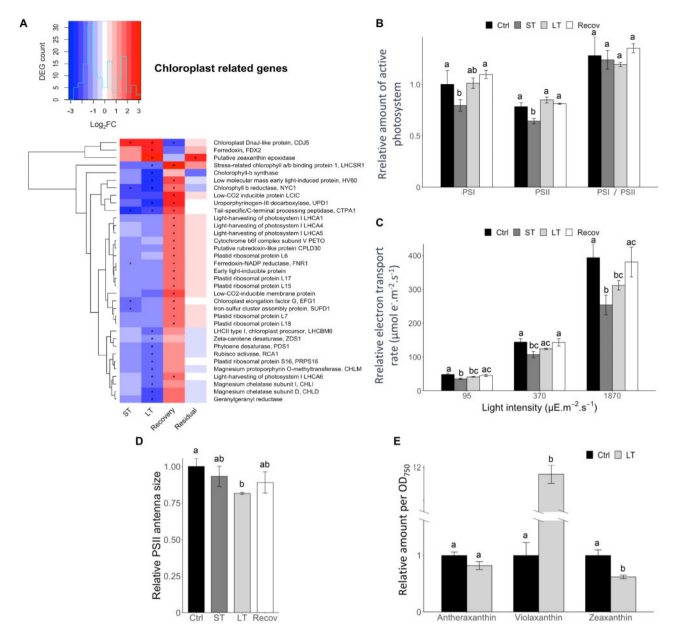


Figure 3 Impact of Cd on the expression of chloroplast-related genes and on photosynthesis. A, Heatmap of chloroplast-related gene expression (Log₂ fold-change) in ST (control versus 3 d Cd-exposed cells), LT (control versus 6 month Cd-exposed cells), Recovery (6 month Cd-exposed cells versus 6 month Cd-exposed cells grown for 3 d without Cd), and Residual (control versus 6 month Cd-exposed cells grown for 3 d without Cd) responses. Significant differences are marked by asterisks (detailed in Supplemental Dataset 1). B, Relative amount of PSI, PSII, and PSI/PSII ratio in control (Ctrl) cultures without Cd, ST (3 d) Cd-exposed cultures, LT (6 months) Cd-exposed cultures and recovery cultures (6 months Cd-exposed cells grown for 3 d without Cd). C, rETR of PSII and (D) relative PSII antenna size in Ctrl, ST, LT, and Recovery cultures exposed to three different PPFD (95, 370, 1,870 μE m^{-2} s⁻¹). E, Relative concentration of xanthophyll cycle pigments in Ctrl and LT cultures after 7 d of Cd treatment. The values are normalized relative to the Ctrl (0 μM Cd). The Y axis is log scaled. Values are mean \pm so of three biological replicates. Significant differences (P < 0.05) according to one-way ANOVA followed by Tukey's test are shown by different letters.

In contrast, following LT exposure, a larger set of genes was differentially expressed. The expression of five genes coding for enzymes involved in different steps of the Chl or carotenoid biosynthesis pathways (CHLD, CHLI, CHLM, ZDS, PDS) was significantly decreased, together with three genes encoding light-harvesting complex proteins (LHCA6, LHCSR1, and LHCBM6; Figure 3A; Supplemental Dataset S1). The latter was in agreement with slightly decreased (20%) PSII

antenna size upon LT Cd exposure (Figure 3D). However, surprisingly, the amount of photosystems (Figure 3B) and the overall PSII activity (Figure 3C) were restored in this condition, which may explain the better fitness of the cells compared to the ST exposure (Figure 1A).

Moreover, a gene encoding a putative zeaxanthin epoxidase, which plays an important role in the xanthophyll cycle allowing nonphotochemical quenching and protection from

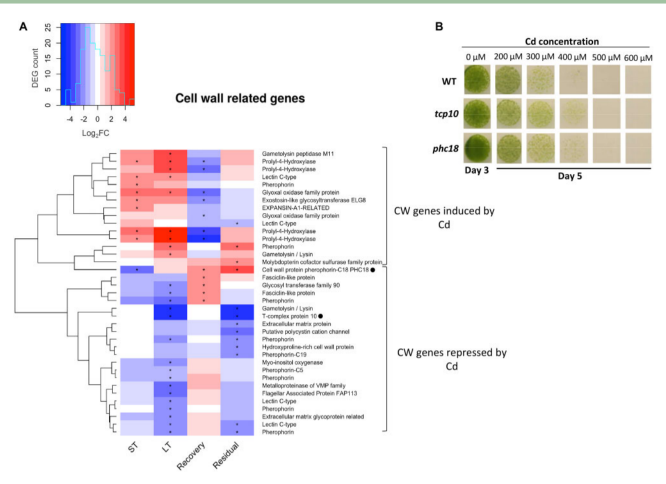


Figure 4 Impact of Cd on the expression of cell-wall-related genes. A, Heatmap of cell-wall-related gene expression (Log₂ fold change) in ST (control versus 3 d Cd-exposed cells), LT (control versus 6 month Cd-exposed cells), Recovery (6 month Cd-exposed cells versus 6 month Cd-exposed cells grown for 3 d without Cd), and Residual (control versus 6 month Cd-exposed cells grown for 3 d without Cd) responses. Significant differences are marked by asterisks (detailed in Supplemental Dataset S1). Two sub-groups of genes that up- or downregulated by Cd are indicated. B, Cd tolerance test on agar plates for tcp10 and phc18 mutants. WT, tcp10, and phc18 cells (10⁵ cell/ml) were spotted in three replicates on agar plates supplemented with different Cd concentrations (0, 200, 300, 400, 500, and 600 μM, respectively). Pictures were taken after 3 and 5 d of growth and are representative of the three biological replicates.

photooxidative damage (Havaux and Niyogi, 1999; Baroli et al., 2003; Havaux et al., 2007), was upregulated in LT conditions. This enzyme ensures the production of violaxanthin from zeaxanthin through antheraxanthin. While zeaxanthin content was slightly lower in LT exposed cells than in Ctrl cells, violaxanthin content was 10 times higher (Figure 3E). This suggested that an increased pool of xanthophyll pigments plays a significant role in Cd acclimation in LT conditions.

After the recovery phase, an almost complete reversal of the Cd-induced responses at ST and LT was observed, with genes downregulated in the presence of Cd being upregulated in the recovery response (i.e. returning to a control state) and vice versa (Figure 3A). Accordingly, neither PSII antenna size (Figure 3D), the amount of PS (Figure 3B) nor the overall PSII activity (Figure 3C) was different from the Ctrl.

The cell wall is important for Cd tolerance

Many genes involved in cell wall synthesis and degradation were differentially regulated in all four experimental conditions (Figure 4A), with a more or less equal number of upand downregulated genes (Table 1). More precisely, a set of genes was upregulated after ST Cd exposure, and this response was maintained at LT, whereas another set of genes was downregulated after LT exposure (Figure 4A). This suggested that cell-wall remodeling plays a role in Cd tolerance in Chlamydomonas as previously shown in other plant species (Krzesłowska, 2011).

A large majority of these genes were differentially expressed in the LT and Residual responses (groups 2, 6, and 7 in Figure 2C), accounting for 79% of cell wall-related DEGs (Table 1). This suggested an important role of the cell wall in LT tolerance to Cd. Moreover, many of these genes induced in presence of Cd were repressed after the recovery phase, and reciprocally (Figure 4A).

To further investigate the role of the cell wall in Cd tolerance in Chlamydomonas, mutants for two candidate genes were obtained from the Chlamydomonas Stock Center (Li et al., 2016b): (1) TCP10 (T-Complex Protein 10, Cre06.g303200), which was downregulated in LT and Residual responses and (2) PHC18 (pherophorin C-18,

Cre24.g755997), which was downregulated in ST response and upregulated in Recovery and Residual responses. These genes were selected based on their distinct regulation profile and their putative role as constituents of the cell wall. In particular, PHC18, a pherophorin belonging to a glycoprotein family specific to the *Chlamydomonadales*, is an essential component of the complex extracellular matrix (ECM) in *Volvox carteri*, which is modulated in response to environmental changes (Hallmann, 2006). TCP10 has not been characterized in detail, but was shown to be differentially regulated in a cell cycle mutant (Tulin and Cross, 2015).

Exposure to 50 μ M Cd for 3 d in liquid cultures had no impact on the growth, Chl content and ionome profile of the mutants compared to Ctrl conditions and to the WT. However, both mutants displayed a slight but reproducible increased tolerance to 300 and 400 μ M Cd on agar plates, compared to the WT (Figure 4B; Supplemental Figure S5), consistent with the downregulation of their gene expression by Cd (Figure 4A). This suggests that both cell-wall genes play a role in Cd acclimation in Chlamydomonas.

P homeostasis-related genes are downregulated upon Cd exposure

Several P homeostasis-related genes were differentially regulated in ST, LT, and Recovery conditions (Supplemental

Dataset 1; Figure 5A). Following ST exposure, the gene encoding the P deficiency master regulator Phosphate Starvation Response 1 (PSR1; Wykoff et al., 1999; Rubio et al., 2001; Moseley et al., 2006) was downregulated, as well as those encoding the intracellular PHO1 phosphatase, four Na⁺/Pi symporters (including PTB2/8/12 animal-like high-affinity Pi transporters), and a periplasmic phosphate binding component of a phosphate ABC transporter, while the plant-like high affinity H⁺/Pi cotransporter PTA3 was upregulated. This gene regulation profile was maintained after LT exposure, with in addition the downregulation of the genes encoding the PTB4 Na⁺/Pi symporter and PHOX, an extracellular alkaline phosphatase, and was mostly reversed upon recovery. This strong downregulation of P uptake-related genes would supposedly result in a decrease of P cellular content in response to Cd exposure. Surprisingly, the total P content was only significantly lower in ST-exposed cells compared to Ctrl cells, whereas it was significantly higher after the recovery phase (Figure 5B). However, these genes were identified based on their homology with the yeast Pi uptake system (Moseley et al., 2006) and since no cellular localization or functional characterization was carried out for PTA or PTB genes in Chlamydomonas, the interpretation of their regulation is difficult.

It is interesting to note that the Chlamydomonas PSR1 TF is homologous to the Arabidopsis PHR1 (Phosphate Starvation

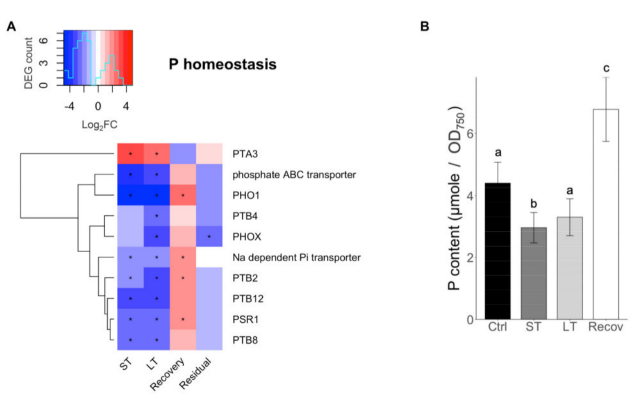


Figure 5 Impact of Cd on phosphorus homeostasis. A, Heatmap of P homeostasis-related gene expression (Log₂ fold change) in ST (control versus 3 d Cd-exposed cells), LT (control versus 6 month Cd-exposed cells), Recovery (6 month Cd-exposed cells versus 6 month Cd-exposed cells grown for 3 d without Cd), and Residual (control versus 6 month Cd-exposed cells grown for 3 d without Cd) responses. Significant differences are marked by asterisks (detailed in Supplemental Dataset S1). B, P content in control (Ctrl) cultures without Cd, ST (3 d) Cd-exposed cultures, LT (6 months) Cd-exposed cultures, and recovery cultures (6 month Cd-exposed cells grown for 3 d without Cd) Values are mean \pm sp of three biological replicates. Significant differences (P < 0.05) according to one-way ANOVA followed by Tukey's test are shown by different letters.

Response 1) TF (Rubio et al., 2001) and that both TFs share well-conserved mechanisms to mediate P homeostasis. As PHR1 is suspected to play a role in metal homeostasis in Arabidopsis (Khan et al., 2014; Briat et al., 2015), this function

may also be conserved for the Chlamydomonas PSR1. Since no modification in cellular P content was observed upon Cd exposure, the downregulation of PSR1 could be related to an involvement of this TF in metal homeostasis.

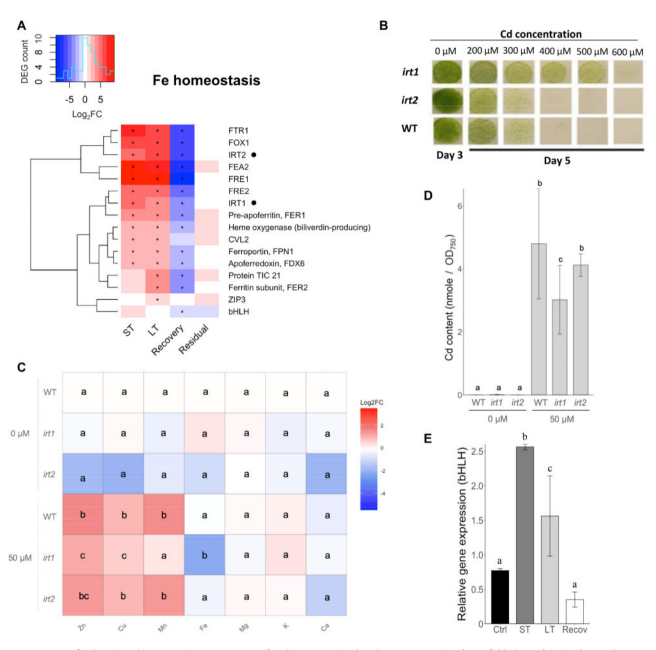


Figure 6 Impact of Cd on iron homeostasis. A, Heatmap of Fe homeostasis-related gene expression (Log₂ fold change) in ST (control versus 3 d Cd-exposed cells), LT (control versus 6 month Cd-exposed cells), Recovery (6 months exposed cells versus 6 month Cd-exposed cells grown for 3 d without Cd), and Residual (control versus 6 month Cd-exposed cells grown for 3 d without Cd) responses. Significant differences are marked by asterisks (detailed in Supplemental Dataset S1). B, Cd tolerance test on agar plates for *irt1* and *irt2* mutants. WT, *irt1*, and *irt2* cells (10^5 cell/mL) were spotted in three replicates on agar plates supplemented with different Cd concentrations ($0 \mu M$, $200 \mu M$, $300 \mu M$, $400 \mu M$, $500 \mu M$, and $600 \mu M$, respectively). Pictures were taken after 3 and 5 d of growth and are representative of the three biological replicates. C, Heatmap of the 10^6 log conditions of mineral element concentrations in the *irt1* and *irt2* mutants in Ctrl ($0 \mu M$ Cd) as well as in WT and the *irt1* and *irt2* mutants in ST (10^6 d exposure) conditions relative to the WT in Ctrl (10^6 μM Cd). Statistical differences are shown by different letters (10^6 concentrations. E, Relative expression level of the Creo5.g241636 gene encoding a bHLH TF in control (Ctrl) cultures without Cd, ST (10^6 d) Cd-exposed cultures, LT (10^6 months) Cd-exposed cultures and recovery cultures (10^6 month Cd-exposed cells grown for 3 d without Cd). D and E, Values are mean 10^6 so 10^6 so

Fe deficiency-responsive genes are upregulated upon Cd exposure

After ST Cd exposure, a large set of genes involved in Fe homeostasis was upregulated. This included genes involved in both low-affinity Fe²⁺ (IRT1 and IRT2; Hanikenne et al., 2009; Urzica et al., 2012) and high-affinity Fe³⁺ uptake systems (FRE1, FOX1, FTR1, and FEA2; Herbik et al., 2002; La Fontaine et al., 2002; Merchant et al., 2006; Allen et al., 2007a; Hanikenne et al., 2009; Narayanan et al., 2011; Glaesener et al., 2013), which are known to be induced upon Fe deficiency, as well as genes involved in intracellular Fe trafficking and storage (FRE2, FER1, CVL2, and FPN1; Singh et al., 2007; Jeong et al., 2008; Urzica et al., 2012; Figure 6A). The same set of genes was upregulated after LT exposure, with in addition the genes encoding the second plastid localized ferritin subunit (FER2) and the putative Zn/Fe transporter ZIP3 (Figure 6A), and all were downregulated upon recovery as for the chloroplast- and cell wall-related genes described above.

As IRT1 and IRT2 are putative Fe²⁺ transporters, possibly also able to transport Cd (as Cd²⁺) into the cell, Cd tolerance was examined in irt1 and irt2 mutants upon Cd exposure (200-600 μM) for 5 d on agar plates. The irt2 mutant displayed a tolerance level similar to the WT, whereas the irt1 mutant displayed strongly increased Cd tolerance (Figure 6B). lonome profiling further revealed that no significant differences were observed between the WT and the irt1 and irt2 mutants in control conditions (Figure 6C). In contrast, upon 3 d of growth in liquid culture in the presence of 50 μ M Cd, the irt1 mutant accumulated lower amounts of Cd (Figure 6D), possibly accounting for its increased Cd tolerance. The accumulation of Cu, Zn, and Mn increased in all three genotypes upon Cd exposure, but did significantly less so in the irt1 mutant (Figure 6C). Fe accumulation was also reduced in the irt1 mutant in Cd condition.

Finally, a basic Helix-Loop-Helix (bHLH) TF (Cre05.g241636), previously shown to be induced in Fe deficiency condition (Urzica et al., 2012), was identified as moderately regulated by Cd in the RNA-Seq data (Figure 6A). Interestingly, this bHLH TF is related to the Arabidopsis group IVc of bHLH, which is involved in Fe homeostasis regulation (Zhang et al., 2015; Li et al., 2016a; Liang et al., 2017; Naranjo-Arcos et al., 2017). The Cre05.g241636 expression profile was further examined by RT-qPCR, which revealed a strong upregulation of the gene upon ST and, to a lower extent, upon LT, Cd exposure, and its downregulation to control levels upon recovery (Figure 6E).

The PSR1 and Cre05.g241636 bHLH TFs are part of a regulatory network

A potential involvement of the PSR1 and Cre05.g241636 bHLH TFs in metal homeostasis and in the response to Cd exposure was further examined. Motifs corresponding to the Transcription Factor Binding Sites (TFBSs) of their Arabidopsis. homologs were retrieved from PlantTFDB (Jin et al., 2017) and used to search the regulatory sequences (1,000-bp upstream the transcription start site) of the recovery response DEGs. This RNA-Seq dataset was selected as

both TFs were found to be differentially expressed in these conditions (Supplemental Dataset S1). Out of the 171 DEGs, 89 and 63 genes were putative targets of Cre05.g241636 or PSR1, respectively, and, among them, 33 were putative targets of both TFs forming a putative co-regulation network (Supplemental Dataset S3; Figure 7).

This in silico analysis identified eight Fe homeostasis genes as putative targets of (1) Cre05.g241636 only (FTR1 and FRE1), (2) PSR1 only (FRE2 and FER1), and (3), interestingly, both TFs (FER2, FOX1, FEA2, and IRT1). Similarly, three out of four P homeostasis-related recovery DEGs were also putative targets of both TFs (PHO1, PTB2, and a Na⁺/Pi symporter encoding genes), whereas the fourth gene (encoding a protein phosphatase 2C, PP2C) was only targeted by Cre05.g241636. Five genes related to metal homeostasis (other than Fe) were linked to one of the two TFs: CTR1 (Cu transporter) and PCS (phytochelatin synthase, PCS) were putative targets of PSR1, whereas CTR2 (Cu transporter), ZTR1 (Zn transporter), and NRAMP1 (Fe/Mn transporter) were among the putative targets of Cre05.g241636. Interestingly, the FSD1 gene encoding the chloroplastic FeSOD isoform, whose activity is reduced under Fe deficiency (Allen et al., 2007b), was also putatively targeted by Cre05.g241636.

Finally, 21 genes with chloroplast-related functions were also identified as targets of PSR1 and/or Cre05.g241636. These genes have diverse functions, including protein transport into the chloroplast, electron transfer, but mainly light harvesting and chloroplastic translation (Supplemental Dataset S3; Figure 7). These genes were mainly specific to the Recovery response (Gp 4 in Figure 2C).

These data indicate that PSR1 and Cre05.g241636 are possibly part of a putative regulatory network, with both TFs controlling overlapping sets of P and metal homeostasis targets. The data also suggest that Cd exposure alters the functioning of the network in Chlamydomonas.

As P and Fe are of critical importance for the development and functioning of photosynthetic cells, we next examined whether such a network has been conserved during the evolution to land plants using publically available transcriptomic data from Arabidopsis (Bustos et al., 2010; Mai et al., 2016). First, we collected expression data for (1) the phr1phl1 Arabidopsis double mutant exposed to P starvation; PHR1 and PHL1 (PHR1-like) are two key regulators of the P starvation response in Arabidopsis and are homologous of the Chlamydomonas PSR1 TF (Rubio et al., 2001; Bustos et al., 2010) and (2) the fit Arabidopsis mutant exposed to Fe sufficiency and deficiency; FIT (FER-like Irondeficiency-induced TF) is, among multiple bHLH TFs involved in Fe homeostasis in Arabidopsis, a major regulator of the Fe deficiency response (Colangelo and Guerinot, 2004; Brumbarova et al., 2015; Mai et al., 2016). Among all 5,588 mis-regulated genes in the Arabidopsis mutants compared to the WT (Supplemental Dataset S4A), 779 were common to both mutants (Supplemental Dataset S4B). Second, we focused our analysis on root expression data, as it was the

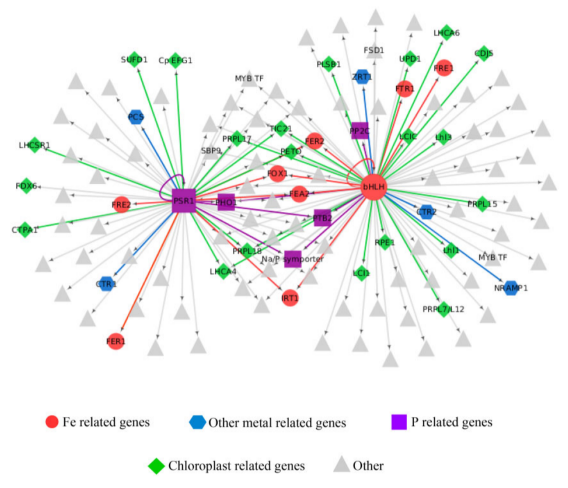


Figure 7 A PSR1 and Cre05.g241636 bHLH transcriptional network. The regulatory network of the PSR1 and Cre05.g241636 bHLH TFs is based on an analysis of TF binding sites in the promoters of genes differentially expressed during the Recovery response (see Figure 2C and "Materials and Methods"). Each node corresponds to a gene and putative regulatory interactions between the TFs and the target genes are symbolized by arrows. Nodes have a specific color and shape based on the functional category of the corresponding genes. Nodes of interest are labeled by the corresponding gene names.

only organ common to both studies and identified 279 overlapping genes (Supplemental Dataset S4C). Interestingly, genes specific to both P and Fe starvation responses in Arabidopsis were included among those 279 overlapping genes suggesting that PHR1, PHL1, and FIT are possibly involved in both responses. Third, to examine this hypothesis in more details, nine Arabidopsis genes were initially selected for promoter analysis (highlighted in yellow in Supplemental Dataset S4C): IRT1, MTP3, AHA7, VTL5, and UGT74E2 are part of the Fe deficiency response in Arabidopsis; PHT1;4, MGD2, and PS2 are linked to P homeostasis in Arabidopsis; and FLA15, whose putative ortholog was found in the PSR1 regulon in Chlamydomonas (Supplemental Dataset S3). For four of those Arabidopsis nine genes (VTL5, MGD2, PHT1;4, and FLA15), a Chlamydomonas putative ortholog was differentially expressed in the Cd dataset (Supplemental Dataset S1). The promoter sequences (-1,500/+100 bases) of those nine Arabidopsis genes were searched as above for occurrences of the P1BS motif, specific of the P deprivation response, and the E-box motif, specific of bHLH TFs. All nine

promoters contained at least one occurrence of both motifs (Supplemental Figure S6; Supplemental Dataset S4, D and E). The data for FIT and the Fe-deficiency genes IRT1, MTP3, and AHA7 are in agreement with previously published data (Colangelo and Guerinot, 2004). Fourth, to generalize this observation, we collected promoter sequences of all genes (2,407) mis-regulated in roots of the phr1phl1 or fit mutants exposed to P or Fe deficiency, respectively (Supplemental Dataset S4A), sorted them in three groups, phr1phl1-specific (940 genes), fit-specific (1,188 genes), or overlapping (279 genes) promoters, and probed those groups of promoters for P1BS or E-box enrichment. An enrichment of P1BS motif in the promoters of the overlapping genes was found relative to fit-specific promoters (exact Fisher's test, P = 0.01), whereas no difference was observed relative to the phr1phl1-specific promoters, suggesting that a P/Fe regulatory network does exists in Arabidopsis. As positive control, the P1BS was also enriched in the phr1phl1-specific promoters relative to the fit-specific promoters (exact Fisher's test, P = 2.83e-5). The reciprocal analysis was less conclusive.

No enrichment was observed for the E-box in the overlapping group of promoters of the two specific groups. It may stem from the fact that the E-box motif is short and is the core binding site of all bHLH TFs, i.e. of more than 120 bHLH TFs in Arabidopsis involved in very different functions (Heim et al., 2003; Toledo-Ortiz et al., 2003) including P starvation response (Chen et al., 2007), as opposed to the P1BS motif which is longer and specific to the P response. However, because the genes investigated here (*fit*-specific and *fit/phr1phl1* overlap) are mis-regulated in a *fit* mutant, it is very likely that they are under the control of Fe starvation response-related bHLH TFs.

Altogether, this analysis suggests that the regulatory network mediated by PSR1 and a bHLH TF to coordinate P and Fe homeostasis in Chlamydomonas is conserved and possibly even more complex in Arabidopsis.

PSR1 and Cre05.g241636 have a role in Fe homeostasis and chloroplast function regulation

Based on these in silico predictions (Figure 7), mutant lines for the *PSR1* and Cre05.g241636 bHLH genes were obtained from the Chlamydomonas Library Project (CLiP) collection. The two mutants were crossed back with a WT strain and the progenies displayed a ~1:1 segregation of par^R and par^S, confirming single insertion of the mutagenic cassette in the genome (Supplemental Table S1A). Furthermore, the reduced growth of the *psr1* mutant on a medium with reduced P level co-segregated with the presence of the Par^R cassette in the genome and thus the *psr1* mutation (Supplemental Table S1B). This also further confirmed that despite residual expression of *PSR1* in the *psr1* background (Figure 8A), the mutant displays typical defects in P homeostasis (Wykoff et al., 1999; Moseley et al., 2006).

The two mutants were further phenotyped in detail. First, reverse transcription-quantitative polymerase chain reaction (RT-qPCR) analysis confirmed that, in control conditions, the expression of the PSR1 and bHLH Cre05.g241636 genes was strongly reduced in the psr1 and bhlh mutant lines, respectively (Figure 8A), indicating that those two mutants are knockdown strains. Although the two TFs were interconnected in a putative regulatory network, the bHLH Cre05.g241636 gene expression was not significantly different in the psr1 mutant, and reciprocally (Figure 8A). Moreover, expression analysis of a selection of putative targets (Figure 7) confirmed that the two TFs are involved in the transcriptional regulation of these genes, at least in control conditions. Indeed, the expression of FRE1 and FSD1, putative targets of the bHLH Cre05.g241636, was significantly upregulated in the bhlh mutant whereas the expression of CTR1, putative target of PSR1, was downregulated in the psr1 mutant (Figure 8A). As predicted, the expression of IRT1, FEA2, and PHO1 was altered in both mutants. Bulked segregant analysis of pools of 20 Par^R and 20 Par^S progenies from a psr1 \times WT backcross showed that the PSR1, PHO1, IRT1, and FEA2 genes displayed lower expression levels in the Par^R strains compared to Par^S strains, indicating that their de-regulation co-segregated with the *psr1* mutation (Supplemental Figure S7). Finally, the analysis also revealed that the strong induction of Fe deficiency genes, *FRE1*, *IRT1*, and *FEA2*, upon ST Cd exposure was similar in WT and both *psr1* and *bhlh* mutants, suggesting that additional regulators are involved in the response to Cd.

Second, ionome profiling revealed lower P concentration, as well as higher Cu and Fe concentrations, in the psr1 mutant in control conditions compared to the WT (Figure 8B), whereas no major alteration was observed in the bhlh mutant (Figure 8B). Upon ST Cd exposure, both mutants displayed reduced Cd accumulation compared to the WT, with a significantly lower accumulation in bhlh compared to psr1 (Figure 8C). The Cd treatment resulted in reduced (P, Ca, K, Fe) or increased (Cu, Zn, Mn) accumulation in the WT and the two mutants. However, the amplitude of these changes were different among genotypes: (1) the P and Ca reductions were more pronounced in the psr1 and bhlh mutants, which both displayed an additional reduction of Mg concentration and (2) the increased accumulation of Cu (bhlh), Zn (psr1, bhlh), and Mn (psr1, bhlh) was of a lower scale in the mutants compared to the WT. The data overall suggested a Cd-induced systemic alteration of the ionome in both mutant backgrounds.

Third, although the *bhlh* and *psr1* mutants had no major growth phenotypes upon Cd exposure (Figure 8D), the *bhlh* mutant displayed distinct Cd sensitivity of the rETR of PSII (Figure 8E; Supplemental Table S2): the *bhlh* mutant maintained a higher rETR than the WT. This phenotype co-segregated with the paromomycin resistance in the progeny of a $bhlh \times WT$ backcross, indicating that it is linked to the bhlh mutation (Supplemental Table S1C). These observations are consistent with the presence of several photosynthesis-related genes among the putative targets of this bHLH TF.

Discussion

Although important progress has been achieved in the last decade in our understanding of the molecular mechanisms underlying Cd uptake and accumulation (Clemens and Ma, 2016; Clemens, 2019), knowledge about the effect of LT, chronic, Cd exposure on photosynthetic organisms remains limited (Yu et al., 2018, 2020). Here, we assessed the response to short (3 d) and long (6 months) Cd exposure in Chlamydomonas, as well as the reversibility of this response. We showed that LT acclimation to Cd relied on molecular pathways similar to the response to short Cd exposure, but mobilized three times more DEGs (Figure 2 and Table 1; Supplemental Figure S4). The CW, adjustment of metal homeostasis (Fe in particular) and photosynthesis, as well as a regulatory network involving the main P homeostasis regulator PSR1 (Rubio et al., 2001) and a bHLH TF (Urzica et al., 2012; Hidayati et al., 2019) contributed to Cd acclimation. These profound physiological alterations enabled restored growth (Figure 1, A and B) and photosynthetic function (Figure 3, B and C). Nevertheless, this response was almost fully reversed within 3 d upon growth in Cd-free medium

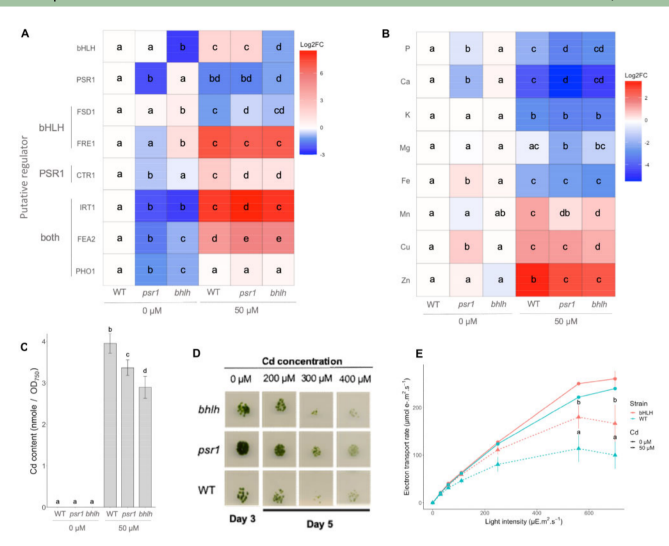


Figure 8 Impact of Cd on *psr1* and Cre05.g241636 *bhlh* mutants. A, B, Heatmaps of the \log_2 Fold Change of the relative expression levels of a selection of PSR1 and/or Cre05.g241636 bhLH TF putative target genes (A) and of mineral element concentrations (B) in WT as well as in the *psr1* and *bhlh* mutants in Ctrl (0 μM Cd) and ST (3 d 50 μM Cd exposure) conditions. All \log_2 fold changes are relative to the WT in Ctrl (0 μM Cd). Statistical differences are shown by different letters (P < 0.05) according to one-way ANOVA followed by Tukey's test. C, Cd content of WT, *psr1*, and *bhlh* mutants in Ctrl (0 μM Cd) and ST (3 d 50 μM Cd-exposure) conditions. Values are mean ± sp (n = three biological replicates). Significant differences (P < 0.05) according to one-way ANOVA followed by Tukey's test are shown by different letters. D, Cd tolerance test on agar plates for *psr1* and *bhlh* mutants. WT, *psr1*, and *bhlh* cells (10⁴ cell/mL) were spotted in three replicates on agar plates supplemented with different Cd concentrations (0 μM, 200 μM, 300 μM, and 400 μM, respectively). Pictures were taken after 3 and 5 d of growth and are representative of the three biological replicates. E, rETR of PSII in WT and *bhlh* strains exposed to a range of light intensities (30 – 700 μE m⁻² s⁻¹) upon growth in control conditions (0 μM Cd) or in the presence of 50 μM Cd for 3 d. Values are mean ± sp of three biological replicates. Significant differences (P < 0.05) according to one-way ANOVA followed by Tukey's test were observed at all light intensities (Supplemental Table S2) but are shown here by different letters for the two highest light intensities only due to lack of space.

(Figure 2; Supplemental Figure S4), concomitantly to a sharp decrease of Cd concentration (Figure 1E), highlighting a major plasticity of metabolic processes in Chlamydomonas. Although new mutations likely appeared in the cultures during the LT Cd exposure, rapid and almost fully reversible response to Cd suggests that acclimation, rather than adaptation, processes are still controlling the cell physiology. Finally, the residual response was mainly dominated by genes involved in CW synthesis and to a lesser extent in signal transduction, translation, and transcription, suggesting that CW remodelling is still perturbed after the recovery phase.

This is in line with the observation of palmelloid structures in the corresponding cultures.

Cell-wall remodeling and multicellularity-like structures as a first protection

The Chlamydomonas CW contains pectins and glycoproteins, which have strong binding affinity for metal ions through their carboxylate groups (Haynes, 1980; Collard and Matagne, 1990; Crist et al., 1994), and constitutes a physical barrier providing an efficient protection against toxic metal ions such as Cd (Cain and Allen, 1980; Macfie et al., 1994;

Penen et al., 2017). Consistently, most CW-related DEGs of the ST Cd response were upregulated and involved in CW maturation and synthesis (Figure 4A and Table 1), possibly increasing the CW buffering capacity. This interpretation has however to be mitigated as a CW-less Chlamydomonas strain was shown to internalize less Cd than the WT (Macfie and Welbourn, 2000), suggesting that the CW binding capacity could facilitate Cd accumulation, possibly by making metal ions available to nonspecific divalent cation transporters located at the cell surface (see below).

So far, the role of the CW following Cd chronic exposure has been under-documented. After LT Cd exposure, the ST response, with mostly upregulated genes, was maintained but was complemented by a strong downregulation of another set of CW-related genes, including pherophorin-encoding genes (Figure 4A and Table 1). This observation is consistent with the formation of palmelloids upon Cd exposure (Figure 1, C and D; Supplemental Figure S3B). Palmelloids are groups of growing and dividing daughter cells within a unique CW (Nakamura et al., 1975). Palmelloid formation is a typical stress response in Chlamydomonas (Iwasa and Murakami, 1969; Lurling and Beekman, 2006; Jamers and Coen, 2009; Boyd et al., 2018) and was recently reported for Cd exposure (Samadani and Dewez, 2018; Cheloni and Slaveykova, 2021). Cd is indeed known to disturb the cell cycle (Cao et al., 2018). Unlike other responses to LT Cd exposure, palmelloids were maintained upon recovery (Figure 1, C and D), consistently with the presence of many lowly expressed pherophorin genes in the residual response (de Carpentier et al., 2019; Figure 4A, bottom). This suggests that reversion of the palmelloid formation is a slow process or that recovery from a LT acclimation to Cd is a stressful event itself.

In addition to palmelloids, larger aggregates of up to several tens of cells surrounded by a matrix (as defined by de Carpentier et al., 2019) were also observed. Such biofilm-like structures could enable Cd tolerance by preventing Cd access to the cell wall and therefore entering into the cell, as suggested in bacteria (Wu et al., 2015). Noticeably, the Chlamydomonas close relative Volvox carteri forms small spherical multicellular colonies (Olson and Nedelcu, 2016) surrounded by a complex ECM, which is modulated in response to environmental changes (Hallmann, Domozych and Domozych, 2014). Pherophorins are essential components of this ECM (Hallmann, 2006). Interestingly, a gene encoding a volvocine algal regA like (VARL) TF (VARL12) was among downregulated genes in ST and LT Cd conditions (Supplemental Dataset S1). The VARL protein family is related to RegA (Duncan et al., 2007), a repressor of germ cell fate in V. carteri (Kirk et al., 1999), and was only identified in Chlamydomonadales and the liverwort Marchantia polymorpha (Sharma et al., 2013). Although no function was identified for VARL TFs in Chlamydomonas, the downregulation of VARL12 in conditions where biofilmlike structures and palmelloids were identified is an interesting feature to further investigate, as it might represent premises towards a form of multicellular organization, which could protect the cells. In this context, cells included in palmelloid structures maintained higher photosynthetic efficiency compared to isolated cells, when exposed to Cd (Supplemental Figure S8).

The key contribution of CW remodeling to Cd acclimation was further confirmed in this study by the analysis of the *phc18* and *tcp10* mutant lines, which both displayed slightly increased Cd tolerance, in agreement with the downregulation of the corresponding genes upon Cd exposure (Figure 4, A and B; Supplemental Figure S5). Further studies will be required to determine how these genes contribute to CW reorganization, palmelloid formation, and control of Cd and nutrient uptake.

Finally, in addition to CW alterations, Chlamydomonas cells displayed other structural changes upon Cd exposure, including motility loss, which is a common reaction under various stressful conditions (Hessen et al., 1995; Khona et al., 2016) and increased cell volume (Figure 1C). The latter was also reported upon chronic Cd exposure in *C. bullosa* (Visviki and Rachlin, 1994), and may result from Cd-induced abnormal cell division.

The uptake and trafficking machinery as a mean to balance Cd-induced metal homeostasis perturbations

As a divalent cation, Cd enters and is distributed within the plant cells via divalent metal transporters with pervasive specificity (for instance, see Vert et al., 2002; Cailliatte et al., 2009; Wong and Cobbett, 2009; Clemens, 2019). Transporters from the same protein families are found in the Chlamydomonas genome (Hanikenne et al., 2005a, 2009; Merchant et al., 2006; Blaby-Haas and Merchant, 2012), and likely also contribute to Cd transport. Therefore, Cd can compete for uptake with other metal nutrients and alters their cellular homeostasis. Consistently, the ionome of Chlamydomonas was strongly impacted upon both ST and LT Cd exposure (Figure 1F), with Cd-induced decreased Ca, K, and Mg, and increased Cu, Zn, and Mn concentrations, respectively (Figure 1F).

Reflecting the need to adjust the homeostasis of those elements, several genes encoding metal transporters were upregulated upon ST and/or LT Cd exposure, including CTR1 and CTR2 for Cu (Page et al., 2009), ZIP3 for Zn and/or Fe (Merchant et al., 2006; Hanikenne et al., 2009) and NRAMP1 for Mn (Allen et al., 2007b) for instance (Supplemental Dataset S1). Cu, Zn, and Mn levels were down to Ctrl levels after three days of recovery, which is consistent with the downregulation of almost all transporters (Supplemental Dataset S1). Moreover, the Cd-induced reduction of Ca in cells is in line with a previous study (Mosulén et al., 2003) showing that Cd uptake in Chlamydomonas is diminished in the presence of additional Ca in the medium, suggesting competition for the same transporters, as suggested in several species (Clemens et al., 1998; Baliardini et al., 2015; Ahmadi et al., 2018). The same study also highlighted a

decrease of N uptake upon Cd exposure, which is in agreement with the downregulation of one nitrate and two ammonium transporters following LT Cd exposure (Supplemental Dataset S1).

Those ionome alterations may represent direct responses to Cd (e.g. a modified uptake to out-compete Cd) or may be an indirect effect of the induction of broad specificity transporters. For instance, Cd is known to alter Fe uptake (Lombi et al., 2002; Lešková et al., 2017; Hanikenne et al., 2021). Fe is a key element in respiratory and photosynthetic electron transport as well as for Chl synthesis (Nouet et al., 2011). In Arabidopsis, increased Fe in the medium allowed a decrease of Cd uptake (He et al., 2017), and the main Fe²⁺ uptake transporter in roots, the ZIP protein IRT1, was shown to represent a major source of entry for Cd and other divalent cations upon Fe deficiency (Rogers et al., 2000; Vert et al., 2002). In Chlamydomonas, both ST and LT Cd exposure had a major impact on Fe homeostasis, with the induction of 10 and 12 Fe-responsive genes, respectively (Figure 6A). This response involved both Fe³⁺ (FRE1, FOX1, FTR1; Page et al., 2009) and Fe2+ (IRT1, IRT2) uptake systems as well as an accessory gene (FEA2; Hanikenne et al., 2009; Narayanan et al., 2011; Glaesener et al., 2013), and genes encoding proteins (possibly) involved in intracellular Fe trafficking and storage (FER1, FER2, CVL2, ZIP3, and FPN1; Singh et al., 2007; Jeong et al., 2008; Urzica et al., 2012).

Moreover, the MSD3 gene, encoding a Mn superoxide dismutase, was upregulated following ST and LT exposure and this gene was shown to be induced upon Fe deficiency in Chlamydomonas to compensate the loss of FeSOD activity in the chloroplast (Allen et al., 2007b). Although no Fe deficiency was observed in our experiment, the MSD3 upregulation may contribute to Fe saving (Page et al., 2012) and is in line with the induction of all other Fe deficiency-related genes. The observed increase in Mn uptake in the presence of Cd (Figure 1F) might provide additional Mn for superoxide dismutases.

This Cd-dependent expression dynamics enabled maintenance of Fe levels in cells after both ST and LT exposure, in contrast to most other nutrient cations (Figure 1F). This response was very rapidly and fully reverted upon Cd removal from LT cultures (Figure 6A). Chlamydomonas thus evolved very dynamic regulatory mechanisms to maintain Fe homeostasis, further illustrating the critical importance of Fe for the cell functioning in the alga.

Among Cd-induced Fe uptake transporters, IRT1 and IRT2 were good candidates for a direct contribution to Cd uptake by the Chlamydomonas cells. Although they belong to another ZIP subfamily (Hanikenne et al., 2005a; Blaby-Haas and Merchant, 2012), IRT1 and IRT2 may indeed play a similar role as their IRT1 counterpart in plants (Vert et al., 2001). While an *irt2* Chlamydomonas mutant had no clear Cd tolerance or ionome phenotypes upon Cd exposure, an *irt1* mutant displayed reduced Cd accumulation upon Cd exposure (Figure 6D), as well as increased Cd tolerance (Figure 6B). The *irt1* mutant also accumulated less Zn, Cu, and Mn

upon Cd exposure, suggesting a broad specificity of IRT1. Our data thus strongly suggest that Cd enters the cells via IRT1, which is induced by a Cd-triggered Fe deficiency. Paradoxically, this upregulation may contribute to increased Cd uptake. Proper Fe supply is however maintained thanks to the induction of the Fe³⁺ uptake system that possibly compensates the Fe uptake imbalance.

Finally, our study also highlights TIC21 as another gene induced by Cd exposure and possibly involved in Fe distribution into the cell (Figure 6A). The homologous gene to TIC21 in Arabidopsis, PIC1 (for permease in chloroplasts 1), was suggested to mediate plastid Fe-uptake (Duy et al., 2007, 2011). Although this function remains controversial (Shi and Theg, 2013) and was not demonstrated in algae, TIC21 displayed an expression profile similar to Fe homeostasis-related genes as it was upregulated upon Cd exposure and rapidly downregulated following the recovery phase (Figure 6A). Interestingly, chloroplastic Fe accumulation triggers the recovery of Cd induced photosynthesis inhibition in plants (Solti et al., 2016). Here, the induction of TIC21 in LT-exposed cells was concomitant with the restoration of photosynthesis efficiency (Figure 3, B and C). Further work will be needed to determine if TIC21 is involved in the adjustment of Fe supply or protein import into the chloroplast in response to Cd, but its genic expression profile supports an involvement in the Fe deficiency response.

Altogether, as no difference of Cd concentration was observed between ST and LT Cd-exposed cells, the reported adjustments of metal ion uptake seem more important to maintain metal homeostasis than to decrease Cd uptake. Consequently, the improved fitness of LT exposed cells (Figure 1A) may rather be related to a decrease of the harmful effects of Cd.

Photosynthetic adjustments to cope with LT Cd exposure

Once in Chlamydomonas cells, Cd is mainly accumulated in vacuoles and in the chloroplast (Nagel et al., 1996; Penen et al., 2017, 2020). Metal cofactors are critical for the photosynthetic electron transport chain (Nouet et al., 2011) and the Cd-induced metal homeostasis disturbance in the chloroplast directly impacts photosynthesis. This is illustrated by the fact that the photosynthetic capacity was negatively impacted upon ST exposure to Cd (Figure 3, B and C), which presumably accounts for the marked decrease in biomass (Figure 1A). However, the negative impact on the cell photosynthetic capacity does not seem to result from a massive downregulation of the related genes to acclimate to ST Cd exposure (Figure 3A), but more likely results from a physiological impact of Cd. Overall, this ST impact of Cd is in good agreement with the known effects of Cd on photosynthesis (Faller et al., 2005; Gillet et al., 2006; Perreault et al., 2011; Parmar et al., 2013). One of these effects is the degradation of PSI active sites, as illustrated in ST exposed cells (Figure 3B). In this respect, the upregulation of the plastid localized ferritin subunit FER1 could play a role in buffering Fe released by this PSI degradation, as it was suggested for Fe deficiency (Busch et al., 2008; Long et al., 2008).

LT-exposed cells displayed a paradoxical relation between the downregulation of photosynthesis-related gene expression and maintained cell photosynthetic capacity. Although the number of active PSI and PSII sites (Figure 3B), the rETR (Figure 3C), and the Chl content (Figure 1B) were equivalent to the Ctrl conditions, pigment synthesis- and light harvesting complex-related genes were down-regulated upon LT Cd exposure (Figure 3A). In this respect, impairment of LHCBM6 accumulation has been previously shown to lead to an overall decrease in PSII antenna size, but also to a diminished photoprotection capacity governed by LHCSR1 (Peers et al., 2009; Girolomoni et al., 2017). The downregulation of LHCBM6 in LT-exposed cells is in line with the reduction by 20% of the functional LHCII antenna size found here (Figure 3D). Because these changes in light harvesting machinery-related gene expression at LT (1) fully recovered upon Cd removal (Figure 3A) and (2) were already initiated at ST, they were likely triggered by Cd presence, and presumably as a response to the negative impact of Cd exposure. These changes probably aim to limit the production of ROS that takes place in the chloroplast when light absorption is larger than the calvin cycle capacity; the RUBISCO protein level and CO₂ fixation were shown to be massively decreased upon Cd exposure (Gillet et al., 2006; Penen et al., 2020). A large increase in Violaxanthin (Vx) content was also found at LT, which is linked to the upregulation of zeaxanthin epoxidase (Figure 3, A-E). Vx acts as antioxidant in the lipid phase of the thylakoid membrane or the protein/ lipid interface (Havaux and Niyogi, 1999; Baroli et al., 2003; Havaux et al., 2007).

The apparent paradoxical observations made here at LT (no impact on growth and photochemistry; decrease in the expression of genes involved in pigment-protein light harvesting complexes; high content in antioxidant Vx), strongly indicates that the cells reached a new equilibrium where damages due to Cd exposure are under control.

P and metal homeostasis are interconnected in Chlamydomonas

Cd exposure yielded another paradoxical observation, with the downregulation of multiple P homeostasis-related genes upon Cd exposure without effect on P cellular concentration, which remained similar to control conditions (Figure 5B). This may result from the fact that among 10 type B and 4 type A Pi transporter genes identified in the Chlamydomonas genome (Moseley and Grossman, 2009), only four and five of those genes were downregulated upon ST and LT Cd exposure, respectively. The *PTA3* gene, encoding a Mn/Pi co-transporter (Fristedt et al., 1999; Jensen et al., 2003), was in contrast upregulated in ST and LT conditions (Figure 5A). As this gene is regulated by Mn deficiency, and not by P deprivation (Allen et al., 2007b), the PTA3 transporter may rather play a role in Mn over-accumulation occurring after ST and LT Cd exposure (Figure 1F).

Several lines of evidence exist that P and metal homeostasis are interconnected in photosynthetic cells. In Chlamydomonas, Cd tolerance was previously shown to depend on P concentration in the medium (Wang and Dei, 2006) and a large part of the Cd accumulated in vacuolar Ca polyphosphate granules (Penen et al., 2017). Fe, K, and Zn uptake were also influenced by P availability, whereas Cu, Fe, and Zn over-accumulated in presence of Cd under P limitation (Webster et al., 2011). The differential expression of P homeostasis-related genes under Cd exposure could therefore indirectly be the result of Cd-induced alterations of metal homeostasis. This connection was observed in Arabidopsis where Fe deficiency- and P starvationregulated genes showed a significant overlap (Li and Lan, 2015), in a similar trend as observed for Chlamydomonas in this study: to an upregulation of Fe deficiency genes upon Fe starvation corresponded a downregulation of P genes (Figures 5, A and 6, A).

In Arabidopsis, the P starvation response is mainly under the control the of PHR1 and PHR1-like (PHL1) TFs (Bustos et al., 2010). Furthermore, some Fe deficiency-responsive genes repressed under P starvation in WT plants were misregulated in the *phr1phl1* double mutant, suggesting an involvement of these TFs in Fe deficiency-responsive gene expression (Briat et al., 2015; Hanikenne et al., 2021). PHR1 also plays a role in Zn and P homeostasis coordination (Khan et al., 2014), suggesting that this TF is involved in the homeostasis of various nutrients.

In Chlamydomonas, the PSR1 TF is homologous to PHR1 (Rubio et al., 2001) and its role in P starvation response is well conserved (Wykoff et al., 1999; Moseley et al., 2006). Our study shows that the regulatory function of these TFs in metal homeostasis is also conserved since nine metal homeostasis-related genes are part of the PSR1 regulatory network in Chlamydomonas (Figure 7). This is further supported by the apparently conserved regulatory link between (1) PHR1 and the chloroplastic ferritin encoding gene FER1 in Arabidopsis (Bournier et al., 2013) and (2) PSR1 and both genes (FER1 and FER2) encoding the two chloroplastic ferritin subunits in Chlamydomonas (Figure 7). Moreover, a psr1 mutant displayed altered metal accumulation, both in control conditions and upon Cd exposure (Figure 8B), and reduced expression of putative metal homeostasis gene targets (Figure 8A), further highlighting the function of PSR1 in metal homeostasis. In particular, the psr1 mutant internalized less Cd (Figure 8C), which is in agreement with the downregulation of PSR1 upon ST and LT Cd exposure that could contribute to cell preservation (Figure 5A).

A bHLH TF is involved in Fe homeostasis and the response to Cd in Chlamydomonas

With the noticeable exception of CRR1 and its key role in Cu homeostasis (Kropat et al., 2005), metal homeostasis regulation by TFs remains poorly described in Chlamydomonas, especially for Fe, although the involvement of a bHLH TF (Cre05.g241636) was proposed based on its upregulation

upon Fe deficiency and its homology to Arabidopsis group IVc of bHLH (Urzica et al., 2012; Zhang et al., 2015; Li et al., 2016a; Liang et al., 2017; Naranjo-Arcos et al., 2017). This hypothesis is supported by (1) the observation that seven out of the nine Fe homeostasis genes included in our promoter analysis are putative targets of this bHLH TF (Figure 7), including all genes of the high affinity Fe³⁺ uptake system (see above) and (2) by the mis-regulation of Fe-related genes in a bhlh mutant in Ctrl condition (Figure 8A). Interestingly, FSD1 (encoding an FeSOD), whose transcript level and corresponding enzymatic activity are decreased under Fe deficiency (Allen et al., 2007b), as well as the genes encoding the Cu transporter CTR2, the Zn transporter ZRT1, and the Mn transporter NRAMP1, are all putatively targeted by this bHLH TF, suggesting that it may play a more general role in metal homeostasis. The reduced accumulation of Cu, Zn, and Mn in a bhlh mutant exposed to Cd supports this hypothesis (Figure 8B). The mutant also accumulated less Cd and displayed higher photosynthesis efficiency (Figure 8, C and E), together with maintained growth, in the presence of Cd. As for IRT1, this is a paradox: the induction of the Cre05.g241636 gene by Cd might per se be detrimental for the cells by possibly sustaining Cd uptake. However, this negative side effect may be the cost of enabling maintained Fe, and possibly other metal, homeostasis, which is critical for the cells, in the presence of Cd.

If the Cre05.g241636 bHLH TF is important to control metal homeostasis and is induced upon Cd exposure, it is not essential for the induction of Fe deficiency genes (Figure 8A) or to maintain Fe cellular levels in the presence of Cd (Figure 8B) and putative targets are induced in the bhlh mutant (FRE1, FSD1, Figure 8A) in control conditions possibly compensating the mutation, all suggesting that additional regulators are likely to be involved in Fe homeostasis regulation in Chlamydomonas. In Arabidopsis, Fe homeostasis regulation is orchestrated by multiple bHLH TFs such as FIT, POPEYE (PYE), and subgroup Ib bHLHs (bHLH38/39/100/ 101; Colangelo and Guerinot, 2004; Long et al., 2010; Wang et al., 2013). Acting upstream of bHLH Ib and FIT TFs, the bHLH IVc TFs are functionally redundant and play a primary role in Fe homeostasis regulation (Zhang et al., 2015; Li et al., 2016a; Liang et al., 2017; Naranjo-Arcos et al., 2017), whereas the MYB10 and MYB72 TFs act downstream of the bHLH TFs and are important for Fe allocation in Arabidopsis (Palmer et al., 2013). MYB72 was shown to be induced by Cd exposure (Lešková et al., 2017). However, in Chlamydomonas, Cre05.g241636 is the only bHLH TF related to the Arabidopsis group IVc of bHLHs and no other bHLH was found to be responsive to Fe or other metal deficiency (Urzica et al., 2012; Urzica, 2017). Thus, the induction of Fedeficiency genes in the bhlh mutant in response to Cd must be controlled by other TFs. Interestingly, two MYB TFs were found to be putative targets of Cre05.g241636 (Figure 7). The Cre10.g421021 MYB gene was upregulated upon ST exposure and downregulated in Recovery, whereas the Cre14.g633789 MYB gene had an expression dynamic similar to the Cre05.g241636 bHLH (upregulated in ST and LT but downregulated in Recovery). In addition, the GCN5-related N-acetyltransferases NAT30 (Cre13.g581150) previously shown to be induced by Fe deficiency (Urzica et al., 2012) was upregulated in the ST/LT conditions but downregulated in Recovery, as Cre07.g331475, another GCN5-related N-acetyltransferases coding gene (Supplemental Dataset S1). Since these proteins are involved in gene expression regulation through histone acetylation, and as homologs were shown to control Fe homeostasis in Arabidopsis (Xing et al., 2015), such epigenetic modifications may also play a role in metal homeostasis regulation in Chlamydomonas.

The PSR1 and Cre05.g241636 bHLH TFs are interconnected in a conserved regulatory network

regulator of P homeostasis main Chlamydomonas (Wykoff et al., 1999; Moseley et al., 2006), may also be involved in the regulation of Fe and metal homeostasis genes. We showed here that it forms a putative regulatory network with the Cre05.g241636 bHLH TF (Figure 7). This is in agreement with the recent report showing that the two proteins interact in a regulatory complex controlling homeostasis and triacylglycerol metabolism genes (Hidayati et al., 2019). Through mutant analysis, we confirmed here that both TFs control the expression of several metal homeostasis genes, including IRT1 and FEA2 (Figure 8A). Our in silico analysis of mis-regulated genes in fit and phr1phl1 Arabidopsis mutants supports that such a regulatory network was at least partly conserved, then diversified, during the evolution of land plants (Supplemental Dataset S4 and Supplemental Figure S5). Not only are deficiency responses for Cu, Fe, or P relying on conserved TFs and signaling processes (Wykoff et al., 1999; Rubio et al., 2001; Kropat et al., 2005; Bustos et al., 2010; Bernal et al., 2012; Urzica et al., 2012), but our work also indicates that a coordinated regulation of these responses is conserved over a large evolutionary time in the plant lineage.

Overall, this network effectively links the homeostasis of different metals (Fe, Cu, Zn, Mn) and P in Chlamydomonas. It is mobilized and is required during LT acclimation to Cd. Further work will be required to dissect in detail these interactions, in the absence and presence of Cd. Our work dramatically expands our molecular understanding of (1) the response to chronic exposure to Cd, a highly toxic and persistent pollutant posing a major environmental threat and (2) the integrated regulatory control of nutrient homeostasis in photosynthetic cells.

Materials and methods

Strain and growth conditions

Chlamydomonas (*C. reinhardtii*) strain CC-4403 (Chlamydomonas Resource Center, Duke University) was cultivated in 25-mL culture in modified TAP medium (Harris et al., 2009) where inorganic phosphate was replaced by β -glycerophosphate (TAgP) to avoid Cd-P complex formation (Collard and Matagne, 1990), buffered to pH 7.0 with glacial

acetic acid and sterilized by autoclaving. The required amount of $CdCl_2$ was then added to the Erlenmeyer flasks from a stock solution (1 mM, filtered through a 0.22 μ m Millipore filter). Cultures were grown under continuous photosynthetic photon flux density (100 μ E m⁻² s⁻¹), at 25°C and placed on an agitator platform (110 rpm).

All Chlamydomonas knockout mutants (Supplemental Table S3) were provided by the CLiP (https://www.chlamyli brary.org/; Li et al., 2016b) and were genotyped by PCR (Supplemental Table S4) to confirm the presence of the insertion cassette (ClB1). The culture conditions were the same as above and the background strain CC-5325 (cw15 mt⁻) was used as control. The psr1 and bhlh mutants were back-crossed with the 137c WT strain (CC-125, mt⁺). The crosses, maturation of zygotes, and random analysis of meiotic products were carried out as described (Hanikenne et al., 2001).

Experimental design

Starting from pre-culture obtained from a single colony, three control cultures were set up in TAP liquid medium (control) and three cultures were set up with addition of Cd (50 μ M CdCl₂). This Cd concentration was selected as the best compromise between an effective toxicity (25% growth reduction) and ensuring the survival of the cells over a long period of time. Each culture was transferred into fresh medium with a starting OD₇₅₀ of \sim 0.005 (corresponding to \sim 5 \times 10⁴ cells/mL) every 7 d for 6 months.

After this period, two transition conditions were tested in addition to the control and LT Cd exposure (Supplemental Figure S1): (1) the three control cultures were duplicated and one duplicate of each culture was exposed to 50 μ M Cd for 3 d and (2) the three Cd-treated cultures were duplicated and one duplicate of each culture was grown for 3 d in TAgP medium free of Cd. After three days of growth, the four triplicated conditions (control, exposed for 3 d, exposed for 6 months, and grown without Cd for 3 d after a 6-month exposure) were sampled for further analysis.

Cell density and Chl quantification

The cell density was estimated by OD_{750} in a spectrophotometer (ThermoScientific, Genesys 20). The Chls were quantified using optical density measurements as follows. A sample of 200 μ L of culture was centrifuged 1 min at 2,500g and the pigments were extracted by adding 800 μ L of 100% ethanol. The mix was centrifuged at maximum speed (>12,000g) for 1 min and 400 μ L of supernatant were used to measure the OD_{650} on a 96-well plate on a fluorimeter (Fluorometer Perkin Elmer 1420, Multilabel Counter Victor 3). The Chl a+b (μ g/mL) content was then calculated by multiplying the OD_{650} by 25.11 (correction factor) and normalized according to the culture cell density.

Tolerance test on agar plates

Two-day-old liquid cultures (\sim 5 \times 10⁶ cells/ml) were diluted to 10⁶, 10⁵, and 10⁴ cells/mL in fresh TAgP medium.

Drops of 10 μ L or 15 μ L of each cell concentration were spotted on TAgP 1.5% (w/v) agar (agar M, Sigma-Aldrich) plates containing 0 μ M, 200 μ M, 300 μ M, 400 μ M, 500 μ M, and 600 μ M Cd, respectively. The plates were then exposed to 100 μ E m⁻² s⁻¹ at 25°C for 10 d. Pictures of the plates were regularly captured.

Metal quantification by inductively coupled plasma atomic emission spectroscopy

Ten milliliter of 3-d-old cell culture (OD₇₅₀ \sim 0.7, corresponding to \sim 7 \times 10⁶ cells/ml) grown in TAgP liquid medium in the presence or absence of 50 μM Cd were sampled and centrifuged at 2,000 \times g for 5 min. The supernatant was then discarded and the pellet sequentially washed with 5 ml of 1 mM EDTA (twice) and 5 ml of distilled water. After each washing step, the cells were resuspended by manual shaking and centrifuged for 5 min at 2,500 g and the supernatant discarded. A cell digestion was then realized by the addition of 3 ml of 65% HNO₃ to the pellet, which was then resuspended by manual shaking and stored for 2 d at 4°C; 7 ml of distilled water and 200 µl of 65% HNO₃ were then added to the tubes and the mixture was used for mineral element profiling by inductively coupled plasma atomic emission spectroscopy as described (Nouet et al., 2015).

Pigment quantification by high pressure liquid chromatography

Pigments were extracted on ice from 1 mL of 7-d-old cell culture (OD₇₅₀ \sim 1.9, corresponding to \sim 2 \times 10⁷ cells/mL). Samples were centrifuged for 2 min at 2,000g and the pellet was resuspended in 90% methanol and frozen at -80° C.

The microtubes containing the pigments were centrifuged for 2 min at 13,000g and the supernatant was filtered using a 0.2 µm PTFE filter (Acrodisc CR, PALL). Then, 200 µL of filtered supernatant were transferred to high pressure liquid chromatography (HPLC) vial. The analysis was realized on a column C-18 (Nova-Pack, silica-based, particle of 4 µm, length of 150 mm and inner diameter of 3.9 mm, Waters, WATO36975). The equipment (Shimadzu) consisted of an injector (SIL-20AC), a pump (LC-20AT), a degassing unit (DGC-20A5R), an oven (CTO-10ASVP) in which the column was located, a holder with a photodiode array (SPD-M20A) and a communication module (CBM-20A). The Empower software (Waters, version 6.1.2154.917) was used to control the HPLC, collect, and analyze the data. The analyses were carried out at 25°C with a flow of 1 mL/min. The mobile phase composition varied during the analysis as three solutions were used following a gradient mode: (1) solution A (90% methanol + ammonium acetate 100 mM); (2) solution B (90% acetonitrile); and (3) solution C (100% ethyl acetate). The quantification was based on peak area at 430 nm after having averaged measures of standard solutions (DHI Labo Products).

Photosynthesis measurements

In vivo, Chl fluorescence and time-resolved absorption change measurements were performed at room temperature (25°C) using a JTS-10 spectrophotometer (Bio-logic, France) or a SpeedZen Fluorescence imaging system (BeamBio, Uani, France) and actinic light was provided by light sources peaking at 640 nm or 520 nm, respectively. For all experiments, cells grown in low photosynthetic photon flux density (PPFD) (50 μ E photons m⁻² s⁻¹) were harvested during exponential growth phase (OD₇₅₀ approximately 0.3–0.5, (corresponding to approximately 3–5 \times 10⁶ cells/mL) and concentrated to a concentration of 10 μ g Chls/mL in fresh TAP medium.

The apparent photochemical yield of PSII (Φ PSII) was calculated as $(F_{\rm m'}-F_{\rm s})/F_{\rm m'}$, where $F_{\rm s}$ is the fluorescence level excited by actinic light ($I_{\rm lum}$), and $F_{\rm m'}$ is the maximum fluorescence emission level induced by a 150 ms superimposed saturating pulse (>3,500 μ E m⁻² s⁻¹). The relative electron transfer rate was obtained as rETR_{II} = Φ PSII \times $I_{\rm lum}$ (Genty et al., 1989).

The ratio between active PSI and PSII centers was estimated from the amplitude of the fast phase (<1 ms) of the electrochromic shift (ECS) signal (at 520–546 nm) after illumination with a single-turnover laser flash. The contribution of PSII was calculated from the decrease in the ECS amplitude after the flash upon the addition of the PSII inhibitors 3-(3,4-dichlorophenyl)-1,1-dimethylurea (DCMU, 20 μ M) and hydroxylamine (1 mM), whereas the contribution of PSI corresponded to the amplitude of the ECS that was insensitive to these inhibitors (Roberty et al., 2014).

PSI relative content was also assessed by measuring maximal P_{700} absorption change in the presence of DCMU (20 μ M) with a probing light peaking at 705 nm (6 nm full width at half maximum). In order to remove nonspecific contributions to the signal at 705 nm, absorption changes measured at 740 nm (10 nm full width at half maximum) were subtracted (Godaux et al., 2015).

Functional PSII antenna size was measured as the rate of Chl fluorescence induction from open (F_0) to closed PSII centers (F_m) in the presence of DCMU (20 μ M), to block reoxidation of the PSII electron acceptor QA (Joliot and Joliot, 2006).

RNA extraction, library preparation, and illumina sequencing

A volume equivalent to 5×10^7 cells was sampled and centrifuged 5 min at 2,000g. The pellet was then resuspended in 2 mL of TAgP, transferred in an eppendorf microtube and centrifuged for 5 min at 2,500g. The supernatant was discarded and the pellet immediately frozen in liquid nitrogen and stored at -80° C.

First, cells were mechanically disrupted using glass beads for one minute at 25 Hz in a grinder (Retsch, MM 200). Total RNA was then extracted using the NucleoSpin RNA Plant (Macherey-Nagel). The quantity and quality were assessed using a Nanodrop (Nanovue plus spectrophotometer).

Libraries for RNA-Seq were prepared from 1 μ g of total RNAs with the TruSeq Stranded mRNA Library Prep Kit (Illumina, San Diego, CA, USA), multiplexed and sequenced with an Illumina NextSeq500 device (high throughput mode; 75-bp single-end reads, \sim 20 million reads/sample).

Transcriptomic analysis

Read quality was assessed using FASTQC (v.0.10.1, http://www.bioinformatics.babraham.ac.uk/projects/fastqc/).

Quality trimming and removal of adapters were performed using TRIMMOMATIC (v.0.32; Bolger et al., 2014), with the following parameters: trim bases with quality score lower than Q26 in 5' and 3' of reads; remove any reads with Q < 26 in any sliding window of 10 bases; crop 1 base in 3' of all reads, and discard reads shorter than 70 bases, and resulted in the loss of \sim 8% of the reads. The resulting high quality reads were then aligned on Chlamydomonas strain CC-503 reference genome V5, downloaded from Phytozome (https://phytozome.jgi.doe.gov) on September 16, 2016, using TopHat (v2.1.1; Trapnell et al., 2009), with the following parameters: -read-mismatches 2; -min-intron-length 40; max-intron-length 2,000; -report-secondary-alignments; no-novel-juncs and providing an indexed genome annotation file. The aligned reads were counted using htseq-count (v0.6.1p1; Anders et al., 2015).

DEGs were identified by pairwise comparisons with the DESeq2 package (v.1.12.3; Love et al., 2014). Genes were retained as differentially expressed when the \log_2 fold-change was > 1 or < -1, with a FDR (Benjamini–Hochberg) adjusted P < 0.05.

GO enrichments were carried out using GOseq as implemented in the run_GOseq.pl perl script of the Trinity suite (Grabherr et al., 2011).

The consistency of the RNA-Seq data analysis was confirmed by RT-qPCR applied to a selection of genes (Supplemental Figure S9).

Promoter scanning for TFBSs

The promoter sequence (1,000-bp upstream from transcription start site) of each DEG of the Recovery response was extracted from Chlamydomonas reference genome V5 using bedtools (Quinlan and Hall, 2010). This dataset was searched for the presence of motifs corresponding to the TFBS of the PSR1 and Cre05.g241636 TFs A. thaliana homologs retrieved from PlantTFDB (Jin et al., 2017). The analysis was carried out using the FIMO tool (Grant et al., 2011) from the MEME suite (Bailey et al., 2009). A regulatory link was inferred based on the presence of a TFBS in the promoter sequence of a gene (P < 0.05) and the resulting network was visualized in Cytoscape (Shannon et al., 2003). The promoter sequences (1,500-bp upstream to 100-bp downstream from transcription start site) of the Arabidopsis genes analysed in Supplemental Dataset S4 were extracted from the TAIR10 (Lamesch et al., 2012) assembly using Araport11 (Cheng et al., 2017) annotation and bedtools (Quinlan and Hall, 2010). The motif enrichment analysis in these promoters was carried out using the AME tool (McLeay and Bailey, 2010) from the MEME suite and the TFBS from PlantTFDB as above.

RT-qPCR analysis

One microgram of total RNA extracted as described above was used for cDNA synthesis with Oligo dT and RevertAid H Minus Kit (Thermo Fisher Scientific, Belgium). RT-qPCR experiments were carried out on a QuantStudio 5 machine (Applied Biosystems, Foster City, CA, USA) using Takyon qPCR Kits for Probe Assay (Eurogentec, Belgium). Three technical replicates were run for each sample/primer combination (Supplemental Table S5) as housekeeping genes and relative gene expression levels relative to two housekeeping genes (SSA9 and CBPL, Supplemental Table S5, Idoine et al., 2014; Lekeux et al., 2018) were calculated by the $2^{-\Delta\Delta Ct}$ method as described (Nouet et al., 2015).

Statistical analysis

The normality of distribution and homogeneity of variance of the data were assessed using the Shapiro-Wilk test and the Levene test as implemented in R, respectively. If both conditions were not fulfilled, a pairwise t test was carried out to evaluate significant differences between treatments. Otherwise, one-way ANOVA (Analysis of variance) was used, followed by a multirange Tukey's test if significant differences were detected at a 95% level of confidence.

Accession numbers

PHC18 (Cre24.g755997), TCP10 (Cre06.g303200), IRT1 (Cre12.g530400), IRT2 (Cre12.g530350), bHLH (Cre05.g241636), PSR₁ (Cre12.g495100), FRE1 (Cre04.g227400), FSD1 (Cre10.g436050), CTR1 (Cre13.g570600), FEA2 (Cre12.g546600), (Cre08.g359300), PTB2 (Cre07.g325741), (AT3G52370), AHA7 (AT3G60330), IRT1 (AT4G19690), MTP3 (AT3G58810), MGD2 (AT5G20410), PS2 (AT1G73010), PHT1;4 (AT2G38940), UGT74E2 (AT1G05680), VTL5 (AT3G25190), PHR1 (AT4G28610), PHL1 (AT5G29000), FIT (AT2G28160). The RNA-Seq reads have been deposited in the National Center for Biotechnology Information (NCBI) Sequence Read Archive (SRA) Database with BioProject identification number PRINA608616.

Supplemental data

The following materials are available in the online version of this article.

Supplemental Figure S1. Scheme of the experimental design.

Supplemental Figure S2. Impact of Cd on the Chlamydomonas cultures.

Supplemental Figure S3. Microscopic observation illustrating multicellular structures forming under Cd exposure.

Supplemental Figure S4. Heatmap of all 577 significantly differentially expressed genes.

Supplemental Figure S5. Replicated Cd tolerance test on agar plates for *tcp10* and *phc18* mutants.

Supplemental Figure S6. Promoter analysis in nine Arabidopsis genes mis-regulated in roots of both *phr1phl1* and *fit* mutants compared to the WT.

Supplemental Figure S7. Bulked segregant analysis of the expression of predicted PSR1 target genes in the progeny resulting from a $psr1 \times WT$ backcross.

Supplemental Figure S8. Cd tolerance in palmelloids.

Supplemental Figure S9. Validation of RNA-Seq data.

Supplemental Table S1. Genetic analysis of the *psr1* and *bHLH* mutants.

Supplemental Table S2. Statistics for pairwise comparisons of electron transport rate of PSII between WT and *bhlh* in control conditions and upon Cd exposure (50 μ M) presented in Figure 8E.

Supplemental Table S3. List of CLiP mutants used in the study.

Supplemental Table S4. List of genotyping primers for the CLiP mutants.

Supplemental Table S5. List of RT-qPCR primers.

Supplemental Dataset S1. Lists of differentially expressed genes from all pairwise comparisons described in Figure 2.

Supplemental Dataset S2. Gene ontology enrichment analysis for pairwise comparisons and sub-groups described in Figure 2.

Supplemental Dataset S3. List of putative PSR1 and Cre05.g241636 bHLH targets in the recovery response, as described in Figures 2 and 7.

Supplemental Dataset S4. Analysis of the conservation of a P/Fe coordinated regulation in Arabidopsis.

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