**Diatom acclimation to elevated CO2 via cAMP signalling and coordinated gene expression**

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**Diatoms are responsible for** ∼**40% of marine primarycarbon cycle and productivity1, fuelling the oceanic contributing to natural carbon sequestration in the deep ocean2. Diatoms rely on energetically expensive carbon concentrating mechanisms (CCMs) to fix carbon efficiently at modern levels of CO2 (refs 3–5). How diatoms may respond overtheshortandlongtermtorisingatmosphericCO2 remains an open question. Here we use nitrate-limited chemostats to show that the model diatom *Thalassiosira pseudonana* rapidly responds to increasing CO2 by differentially expressing gene clusters that regulate transcription and chromosome folding, and subsequently reduces transcription of photosynthesis and respiration gene clusters under steady-state elevated CO2. These results suggest that exposure to elevated CO2 first causes a shift in regulation, and then a metabolic rearrangement. Genes in one CO2-responsive cluster included CCM and photorespirationgenesthatshareaputativecAMP-responsive *cis*-regulatorysequence,implyingthesegenesareco-regulated in response to CO2, with cAMP as an intermediate messenger. We verified cAMP-induced downregulation of CCM gene** δ**-CA3 in nutrient-replete diatom cultures by inhibiting the hydrolysis of cAMP. These results indicate an important role for cAMP in downregulating CCM and photorespiration genes under elevated CO2 and provide insights into mechanisms of diatom acclimation in response to climate change.**

Burning fossil fuels and land-use change have accelerated CO2 emissions to the atmosphere by a factor ∼100 above natural levels6. About a third of anthropogenic emissions have been absorbed by the oceans7,8, increasing dissolved CO2 and reducing pH (ref. 9). Despite these changes, CO2 concentrations in surface waters remain below half-saturation for most forms of ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco)3, the central enzyme used to fix carbon. Consequently, marine phytoplankton, including diatoms, rely on carbon concentrating mechanisms (CCMs) to ensure adequate delivery of CO2 to the Rubisco active site, minimizing the competitive fixation of oxygen3–5. The required bicarbonate transporters and carbonic anhydrases of these CCMs concentrate CO2 against a gradient, which is energetically costly10. Downregulation of CCMs as part of acclimation to elevated CO2 should result in energy savings to the diatom cell and metabolic rearrangement. Here we use nitrate-limited chemostats to simulate *in situ* nutrient limitation11 while precisely controlling cell biomass and CO2 (ref. 12), allowing us to identify potential signalling pathways triggered either by an abrupt transition to increased CO2, as might occur during coastal upwelling13, or at steady-state exposure to elevated CO2, including 800 µatm predicted for 2100 (ref. 14; Fig. 1a,b).

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Metabolic and regulatory genes were differentially impacted by changes in CO2 (Fig. 1c). The initial response to an abrupt increase in CO2 included upregulation of genes required for transcriptional regulation and kinase activity (Fig. 1c), potentially reflecting CO2specific signalling mechanisms. Once steady-state conditions with elevated CO2 were attained, genes required for energy-producing metabolic pathways, including oxidative phosphorylation, TCA cycle and photosynthesis, were preferentially downregulated, as were genes involved in transcriptional regulation, ion transport, kinase activity and protein degradation (Fig. 1c), suggesting a general reduction in metabolism under high CO2. This corroborates our previous physiological measurements of decreased respiration and photosynthetic oxygen production under elevated CO2 and nitrate limitation12. A significant upregulation of components of the ribosome under steady-state high CO2 (Fig. 1c) may reflect either an increased capacity for protein synthesis or a high turnover of ribosomal proteins, whereas the downregulation of the same gene category in transition suggests disparate modes of regulation between transition and steady-state exposure to elevated CO2.

To identify potential linkages between an initial response and eventual metabolic rearrangements under steady-state conditions, we identified groups of co-expressed genes based on unsupervised hierarchical clustering across a combined data set of 98 RNA sequencing and microarray data sets from a range of conditions15–20, including our chemostat experiments. Of the 400 clusters of coexpressed genes, 160 exhibited significant correlation to CO2 partial pressure during transition or steady-state exposure (Spearman false discovery rate (FDR) *<* 0.0001; Supplementary Table 1). The unsupervised clustering analysis yielded biologically meaningful insights, grouping together functionally related genes associated with energy metabolism, including light harvesting, photosynthesis, and respiration (Fig. 2a,b). These energy metabolism gene clusters were differentially expressed only after acclimation to steady-state elevated CO2, along with other clusters containing genes implicated in cell signalling (Fig. 2c,d), suggesting a suite of gene clusters are probably involved in diatom metabolic rearrangement. Expression of a gene cluster associated with regulation, structural maintenance of chromosomes (SMC), and transcription was correlated with CO2 level during transition and steady-state exposure (Fig. 2e,f). This cluster of regulatory genes, not previously connected to the diatom CO2 response, could be involved in diatom cell regulation vital to sensing and acclimating to rising CO2.

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| **a** Transition **b** Steady-state **c**  Kinases  Regulation of  transcription  Ion transport  Proteasome  Ribosome  Photosynthesis  Citrate cycle  (TCA cycle)  Oxidative  phosphorylation  Transition  Steady-sta  te  Fold enrichment  1  2  3  4  5  0  1  2  3  4  5  Pos. correlated  Neg. correlated  ∗  ∗  ∗  ∗  ∗  ∗  ∗  ∗  300  400  500  600  700  800  900  CO  2    (  μ  atm)  0 1 2 3 4 Low Med High  Time (days) (Acclimated >10 days)  **Figure 1 | Gene set enrichment in transition and steady-state nitrate-limited cultures. a**,**b**, *Thalassiosira pseudonana* exposed to transition from low to high CO2 over four days (**a**) or steady-state acclimation to low, medium or high CO2 (**b**). Error bars indicate 1 s.d. from the mean, *n*=4. **c**, Genes significantly correlated (positively or negatively) with CO2 (Spearman rank correlation *p<*0.05) formed gene sets categorized by KEGG pathway and GO term. The ratios of gene number to expected value (fold enrichment) in each category are plotted on the *y*-axis; dashed line indicates expected value and asterisk indicates significant enrichment in positively or negatively correlated gene sets (hypergeometric test *p<*0.05). |

High external CO2 is expected to reduce the need for the CCM as well as photorespiration, owing to the competitive inhibition of Rubisco’s oxygenase activity by CO2 (ref. 5). Interestingly, a single cluster contained both putative CCM and photorespiration genes and was negatively correlated to CO2 in both transition and steady-state experiments (Supplementary Table 1). This cluster further grouped into two distinct sub-clusters that exhibited different expression patterns across the range of CO2 concentrations examined in our experiments (Fig. 2e,f and Supplementary Fig. 1). One sub-cluster included candidates for participation in the CCM, such as the plastid-localized delta-carbonic anhydrase (δ-CA3; ref. 21) and plastid-targeted transporters. The other sub-cluster included mitochondria-localized photorespiration genes necessary to recycle by-products of the oxygenase activity of Rubisco. The zeta-carbonic anhydrase ζ-CA1 (ref. 21), also grouped with the photorespiration sub-cluster, is the only enzyme known to use cadmium as a cofactor22 and is localized to the outer membrane space21.

Given the strong correlation across our data set of these sub-clusters of genes from separate pathways, we hypothesized they were regulated by a common mechanism. Genes in the CCM/photorespiration cluster all share a short upstream sequence (Supplementary Fig. 1b), TGACGT, recently identified as a *cis*regulatory sequence that downregulates expression of the CO2responsive β-CA in the distantly related diatom *Phaeodactylum tricornutum*23. Furthermore, two transcription factors, one in each sub-cluster, exhibit close homology (Supplementary Fig. 2) and are classified in the same phylogenetic group24 as a cAMP-activated transcription factor in *P. tricornutum* shown to bind the *cis*regulatory element and to contribute to downregulation of the β-CA gene in response to increased CO2 (ref. 23). These lines of evidence suggest that the two transcription factors identified in the CCM/photorespiration cluster may function to downregulate transcription of their own cluster by binding to the common upstream *cis*-regulatory region. This further implies that these transcription factors repress their own transcription, creating a negative feedback on the repression of this cluster, a feature common in biological systems25.

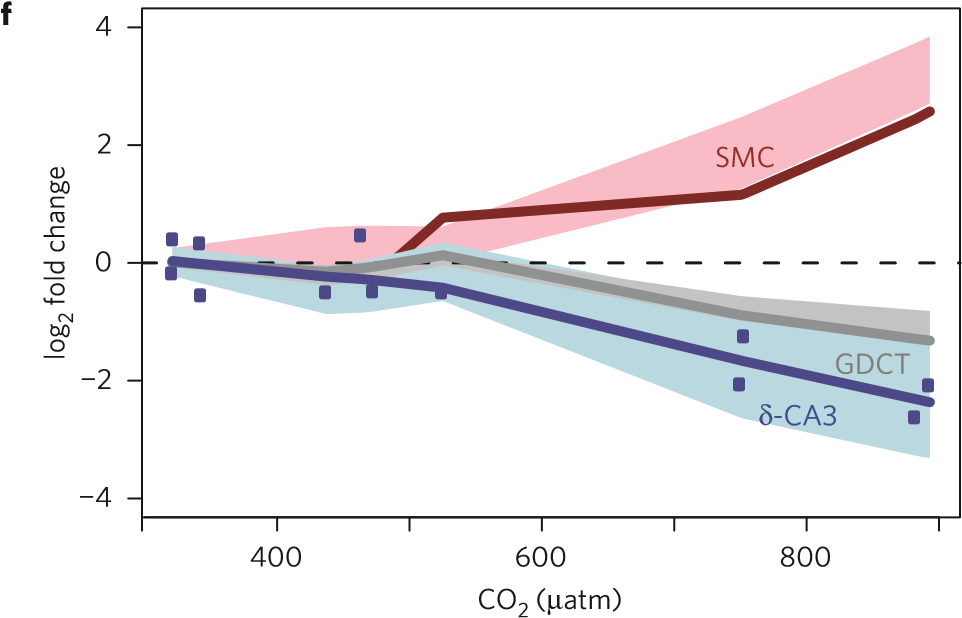
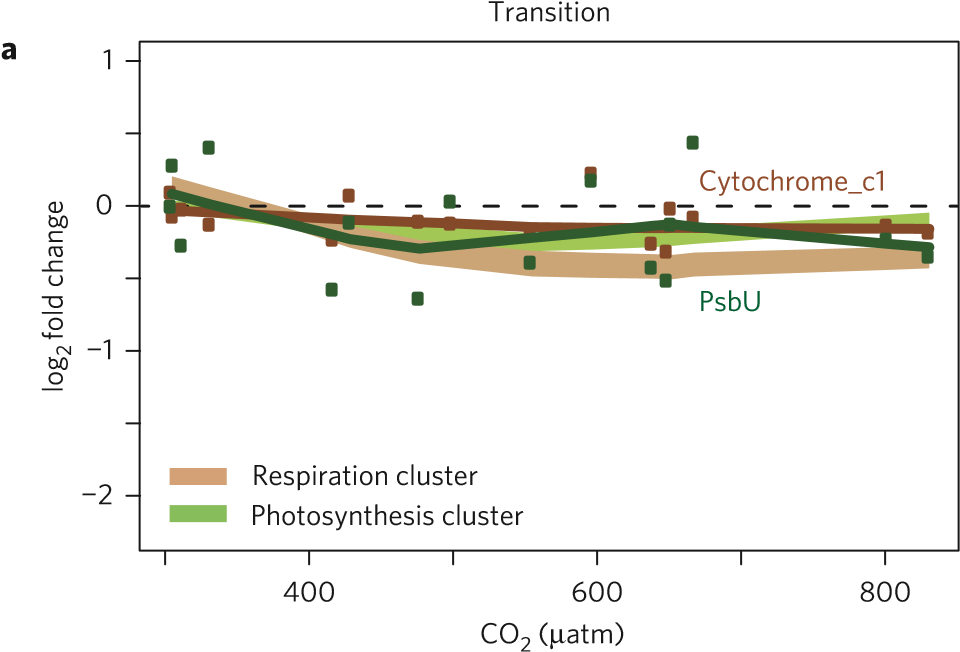
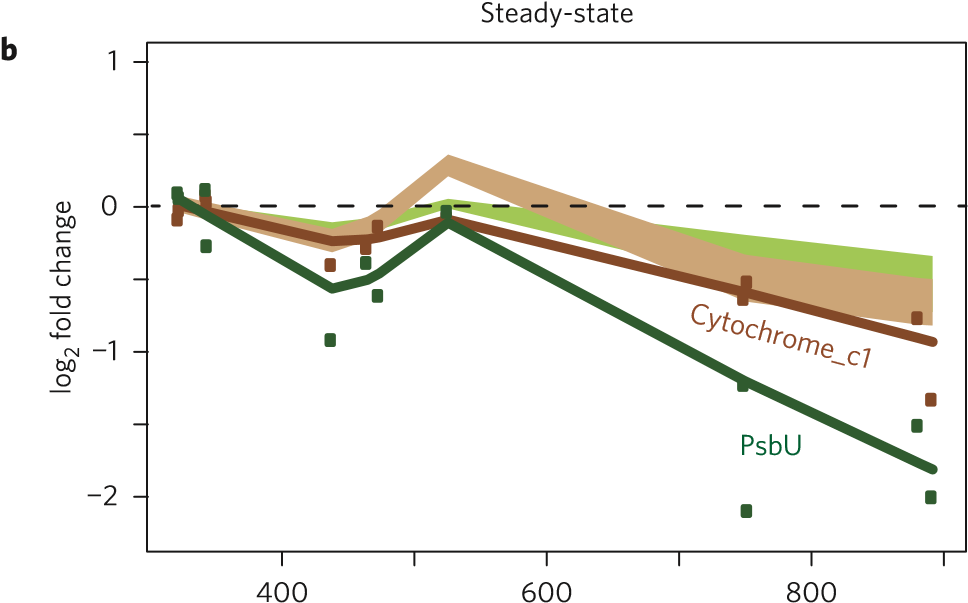
We tested the proposed mechanism of cAMP-regulated gene expression with two indicator genes—carbonic anhydrase δ-CA3 and a putative transporter (pID 262258)—that responded to CO2 in both transition and steady-state experiments with a large fold change (Supplementary Table 2). We grew *T. pseudonana* in nutrient-replete batch cultures and manipulated the CO2 and the cellular level of cAMP by inhibiting its degradation with the phosphodiesterase inhibitor 3-isobutyl-1-methylxanthine (IBMX). The gene encoding δ-CA3 is a member of the CCM gene cluster and contains the upstream TGACGT motif. As predicted, transcript abundanceforthisgenewassignificantly(*p<*0.05,two-wayanalysis of variance (ANOVA)) reduced by both high CO2 and IBMX addition compared to low CO2 (Fig. 3). The gene encoding the putative transporter lacks the cAMP-responsive motif. As predicted, transcript abundance for this gene was increased by elevated CO2, but was not significantly altered by the addition of IBMX (*p>*0.05, *t*-test) (Fig. 3). These data suggest that gene expression patterns in response to CO2 remain consistent independent of nitrate availability and that cAMP selectively downregulates CO2responsive genes possessing the TGACGT motif.

Consistent with our data, mounting evidence23,26 suggests that diatoms sense a change in external CO2 concentrations, in part, through cAMP signalling. In other organisms26,27 the activity of specific adenylyl cyclases are regulated by CO2 or bicarbonate, where increased CO2 stimulates cyclase activity and results in enhanced production of cAMP (ref. 27). This evidence suggests that increasing CO2 concentrations promote a cyclase in *T. pseudonana* to increase production of cAMP, which in turn induces the CCM/photorespiration cluster transcription factors to bind to the TGACGT motif and repress transcription of CCM/photorespiration cluster genes, including their own transcription. The soluble cyclase (sol. CYCc, Fig. 4) and two membrane-bound cyclases (CYCc, Figs 2c,d and 4) are among the candidates for sensing CO2 in *T. pseudonana*. This mechanism is supported by the downregulation of CCM/photorespiration cluster genes, including the transcription factors, by an increase in CO2 and the downregulation of δ-CA3 by addition of IBMX (Supplementary Table 2 and Fig. 3).

Negative feedback mechanisms should result in a decreased repression of the CCM/photorespiration cluster genes over time unless changes in cAMP signalling or transcription factor regulation occur. We observed that cAMP metabolism genes were upregulated at steady-state elevated CO2 (Fig. 4), including a phosphodiesterase, which degrades cAMP by hydrolysis, and two membrane-bound cyclases (CYCc, Fig. 2c,d) that enhance production of cAMP. This indicates a change in cAMP metabolism at steady-state elevated CO2, potentially altering

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400

600

800

400

600

800

CO

2

(

μ

atm)

CO

2

(

μ

atm)

log

2

fold change

log

2

fold change

Regulatory cluster

Photoresp sub-cluster

CCM sub-cluster

δ

CA

3

-

SMC

Signalling cluster

GTPase cluster

CYCc

CYCc

PDE

PDE

Rab

Rab

1

0

−1

−2

−3

3

2

log

2

fold change

1

0

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−2

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2

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0

−2

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4

**c**

**e**

**d**

CO

2

(

μ

atm)

400

600

800

CO

2

(

μ

atm)

GDCT

**Figure 2 | Clusters of co-expressed genes versus CO2 in transition and steady-state experiments. a**–**f**, Fold change (mean ± 2 s.e.m.) of clusters of co-expressed genes normalized to expression under low CO2. Gene clusters involved in energy-producing metabolic pathways (**a**,**b**), sensing and intracellular signal propagation (**c**,**d**) and a regulatory cluster and photorespiration and CCM sub-clusters (**e**,**f**). Expression of selected individual genes is indicated with solid lines representing smoothed mean expression and points representing single transcriptome measurements relative to low CO2. Abbreviations for genes are: Cytochrome\_c1, Cytochrome C1 oxidase; PsbU, Photosystem II subunit; CYCc, adenylate/guanylate cyclase; PDE, phosphodiesterase; Rab, Rab-like small GTPase; SMC, structural maintenance of chromosomes protein; CA, carbonic anhydrase; GDCT, glycine decarboxylase T protein.

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| how signals are propagated in cAMP-responsive pathways, and perhaps explaining why transcript abundance for the δ-CA3 gene is not further reduced under high-CO2 conditions in the presence of IBMX (Fig. 3). Likewise, the CCM sub-cluster transcription factor was no longer significantly repressed at steady-state elevated CO2 (Fig. 4), potentially allowing it to attain sufficient abundance to maintain repression of the CCM and photorespiration pathway.  Placing the CCM and photorespiration gene clusters in the context of the diatom cell, we integrate predicted protein | localization, gene transcription patterns, and metabolic pathways at steady-state elevated CO2 (Fig. 4) to gain insight into potential mechanisms of the diatom CO2 response. A functioning diatom CCM must transport bicarbonate into the plastid to the stroma, where it is converted to CO2 by a carbonic anhydrase10. The CCM sub-cluster contains genes encoding plastid-localized membrane proteins, including bestrophin-like proteins, which are homologous to a family of gated anion-selective channels28 permeable to bicarbonate29 (Supplementary Table 2). These plastid-targeted proteins provide a feasible mechanism for bicarbonate transport to |

0

20

40

60

80

100

a

b

c

c

Transcripts ng

−1

mRNA

Low

Low

High

High

CO

2

IBMX

−

+

−

+

Transporter 2

62258

δ

-

CA

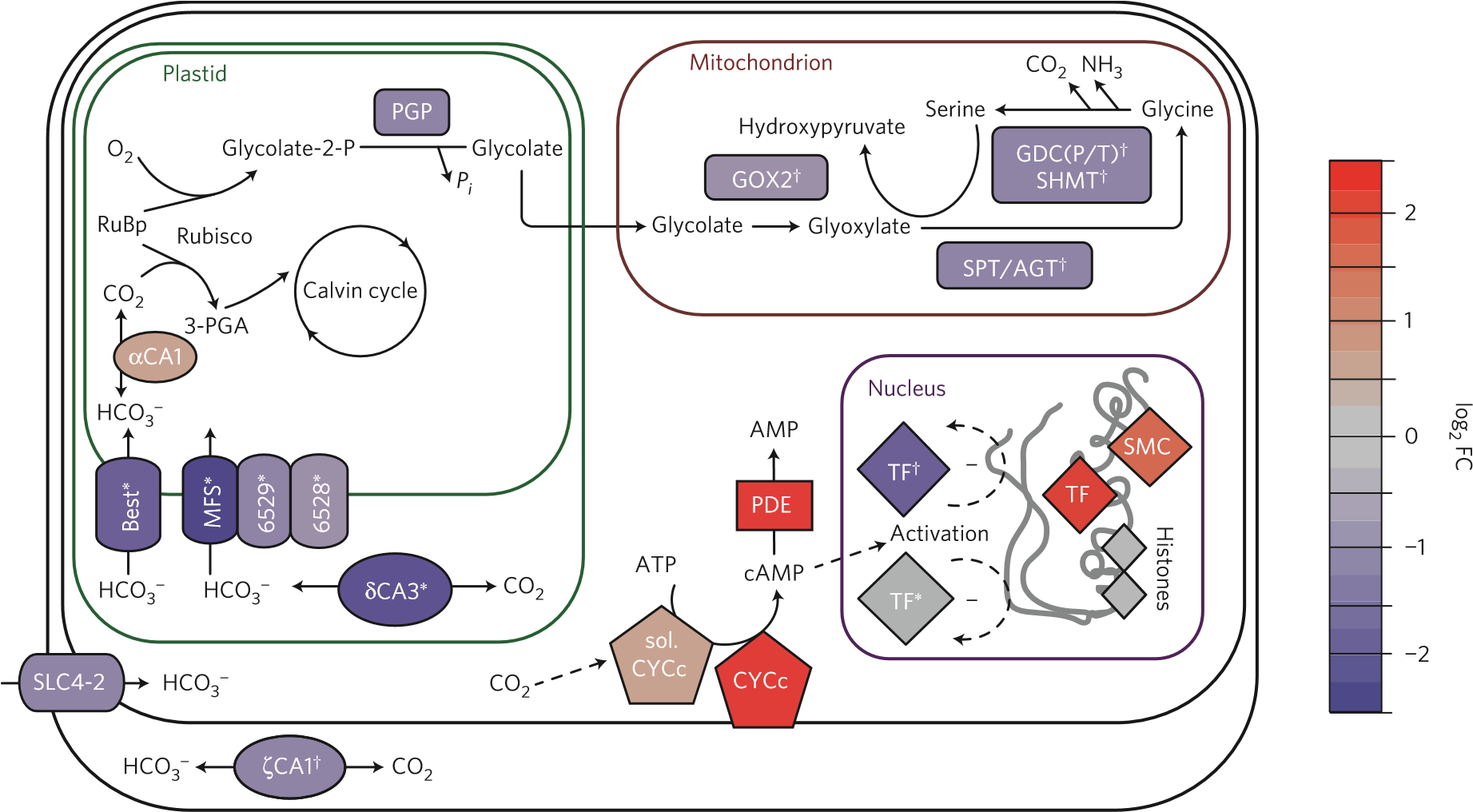
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**Figure 3 | cAMP and CO2 dependence of gene expression in nutrient-replete cultures.** Transcript abundance normalized to messenger RNA (mean ± 1.96 s.e.m.: 95% CI, *n*=3) of genes encoding carbonic anhydrase δ-CA3 (grey) and a putative transport protein pID 262258 (white), in nutrient-replete cultures. *T. pseudonana* cells were exposed to low (240 ± 15 µatm) or high (857 ± 54 µatm) CO2, with or without 1.0 mM of the phosphodiesterase inhibitor 3-isobutyl-1-methylxanthine (IBMX) used to elevate cellular cAMP concentrations. Dashed line indicates the quantification limit of transcript abundance by RT-qPCR and letters indicate significant differences among treatments for δ-CA3 gene expression (*p<*0.05, two-way ANOVA and Tukey HSD).

the stroma (Fig. 4), which has not yet been elucidated in diatoms. Once inside the plastid, bicarbonate can be converted to CO2 by the stroma-localized21, constitutively expressed α-CA1 (Fig. 4). Interestingly, this carbonic anhydrase is not downregulated by CO2, suggesting that, unlike *P. tricornutum*23, expression levels of a stromal-localized CA are not a point of CCM control in *T. pseudonana*. Another essential feature of efficient CCMs is to prevent the passive diffusion of enhanced CO2 concentrations out of the cell in general, or the plastid in particular10. The δ-CA3, localized to the plastid membrane space21, and the ζ-CA1, localized to the outer membrane space21, are well placed to serve this role by converting CO2 into bicarbonate in the periplastid and periplasmic compartments, where it could be selectively transported back into the cytoplasm or stroma. That these genes are rapidly and sustainably downregulated under elevated CO2 suggests that preventing diffusive loss of CO2 is tightly regulated as a point of control for the *T. pseudonana* CCM, in strong agreement with physiology-based model predictions for an efficient diatom

CCM (ref. 10).

Another feature of an efficient diatom CCM is active transport of bicarbonate from the external environment to the cell interior10; the SLC4-2 bicarbonate transporter is the most likely candidate, as it is distantly related to a transporter demonstrated to serve this role in the diatom *P. tricornutum*30, and the gene encoding this transporter was significantly downregulated at steady-state high CO2 (Fig. 4 and Supplementary Table 2). Yet the SLC4-2 gene does not group with the CCM/photorespiration genes, nor share the same upstream motif (Supplementary Table 1). Likewise, the plastid-localized phosphoglycolate phosphatase (PGP) gene, essential to the photorespiration pathway, was downregulated at steady-state (Fig. 4), but does not cluster with the mitochondrial photorespiration genes nor share the upstream region. cAMPsignal-induced regulation cannot account for all the changes



**Figure 4 | Model of cell signalling and metabolite fluxes in *T. pseudonana* after acclimation to high CO2.** Gene expression changes relative to low CO2 in high-CO2 steady-state experiments indicated by the heat map (log2 fold change) overlaid on a model of diatom CO2-responsive signalling and metabolic pathways. Membership in the CCM sub-cluster and photorespiration cluster indicated by ∗ and † respectively. Solid arrows indicate reactions and dashed

arrows indicate regulatory relationships. Abbreviations: SPT/AGT, serine-pyruvate/aspartate aminotransferase; TF, transcription factor; SHMT, serine-glycine hydroxymethyltransferase; GDC(P/T), glycine decarboxylase (P/T) protein; GOX, glycolate oxidase; MFS, major facilitator superfamily; Best, bestrophin superfamily transporter; PGP, phosphoglycolate phosphatase; SLC4, solute carrier family 4 bicarbonate transporter; 3-PGA, 3-phosphoglyceric acid; RuBP, ribulose-1,5-bisphosphate; others as in Fig. 2.

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in gene expression, because these and other known CCM and photorespiration genes were not downregulated in lock-step (Fig. 4) nor do they all contain the same upstream region. Transcription of a SMC gene, co-expressed transcription factors, and histone proteins were significantly affected by an abrupt increase in CO2 (Supplementary Table 2). These proteins function to modify chromosome structure to alter gene expression (Fig. 4), and may explain shifts in CO2-responsive genes not regulated by the cAMPinduced transcription factors.

The metabolic rearrangement we observed under steady-state elevated CO2 and nitrate limitation resulted from downregulation of the energy-consuming processes of CCM and photorespiration because cells were unable to increase growth rate owing to nitrate limitation, a common limiting factor in the oceans11. Diatoms are also limited by other factors, such as iron or light, in vast regions of the ocean11, and may exhibit different metabolic rearrangements or growth rate enhancement in response to elevated CO2 under these other conditions. Our analysis discovered genes that exhibit CO2 responses regardless of nitrogen status, and provide insights into the mechanisms of CO2 sensing, signalling and metabolic rearrangement necessary to predict how diatoms will acclimate in future oceans. Our approach, examining the behaviour of coexpressed gene clusters in both transition and after steady-state acclimation, is a powerful tool for beginning to untangle the mechanisms of sensing and responding to rising CO2. Many of the CO2-responsive genes identified by our clustering analysis were not previously connected to *T. pseudonana* CCM or CCM-regulation, allowing us to investigate new genes and mechanisms involved in regulating diatom physiology and biogeochemical cycles in the face of climate change.

## Methods

Methods and any associated references are available in the [online version of the paper.](http://dx.doi.org/10.1038/nclimate2683)

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## Author contributions

G.M.M.H., R.L.M. and R.D.G. carried ot RNA sample preparation, sequencing and

RT-qPCR. G.M.M.H., J.A. and C.B carried out bioinformatics and statistics. G.M.M.H., J.A., M.V.O., N.S.B. and E.V.A. carried out experimental design. The manuscript was prepared by G.M.M.H., J.A., R.D.G. and E.V.A. All authors contributed to discussion of results and comments on the manuscript.

## Additional information

Supplementary information is available in the [online version of the paper.](http://dx.doi.org/10.1038/nclimate2683) Reprints and permissions information is available online at [www.nature.com/reprints.](http://www.nature.com/reprints) Correspondence and requests for materials should be addressed to G.M.M.H. or E.V.A.

**Competing financial interests**

The authors declare no competing financial interests.

## Methods

**Chemostat cultures.** For a full description of chemostat culturing methods see ref. 12. Briefly, axenic *T. pseudonana* cells in four biological replicates (duplicate chemostats × 2 experimental runs) were acclimated to nitrate limitation at 70% (1.5 day−1) of maximum growth rate for more than ten days (*>*15 generations) under a continuous light level of 80 µmol photons m−2 s−1. Cell biomass was maintained at ∼2×105 cells ml−1 by 10 µM nitrate, carbonate chemistry stabilized to 300, 475 or 800 µatm CO2, verified by calculating31 *f*CO2 from pH and dissolved inorganic carbon (DIC) measurements. After steady-state acclimation, ∼1×108 cells were harvested on 0.2 µm polycarbonate filters by gentle vacuum filtration and then flash frozen. Transition samples and carbonate chemistry were collected daily from chemostat cultures as CO2 levels were increased from ∼300–800 µatm at a rate ≤ 0.2 µatm min−1 over four consecutive days (six generations) after pre-acclimation to 300 µatm CO2 and nitrate limitation (Fig. 1a,b). During transition, ∼1.5×107 cells were harvested daily on 0.2 µm polycarbonate filters by gentle vacuum filtration and then flash frozen.

**Nutrient-replete cultures.** Triplicate cultures of axenic *T. pseudonana* were grown in f/2 medium under continuous illumination (300 µmol photons m−2 s−1) at 20 ◦C with constant aeration. Exponentially growing cultures were harvested with a cell density of ∼4.2×105 cells ml−1 at 240±19µatm or 857±54µatm CO2, verified by DIC and pH measurements (Supplementary Table 3). To inhibit the hydrolysis of cAMP, 1.0 mM 3-isobutyl-1-methylxanthine (IBMX) was added to the cultures, as described in previous work with the diatom *Phaeodactylum tricornutum*32. Before and after a 100 min exposure to 1.0 mM IBMX, ∼4×108 cells were harvested by gentle vacuum filtration on a 0.8 µm filter and then flash frozen for RT-qPCR analysis.

**SOLiD libraries.** A total of 28 barcoded transcriptome libraries were prepared from chemostat culture samples. RNA was extracted from filtered cells using the ToTALLY RNA kit (Life Technologies). Messenger RNA was selectively amplified using MessageAmp II aRNA Amplification kit (Life Technologies) and used to prepare SOLiD barcoded libraries (SOLiD Total RNA-seq kit, Life Technologies). Libraries were sequenced on a SOLiD 5500XL sequencer in two runs: one containing steady-state barcoded libraries and one containing transition barcoded libraries.

***In silico* read processing.** Reads were quality controlled (using a cutoff of *p*=0.99 and minimum length = 30), trimmed, and aligned to *T. pseudonana* gene models using the Burrows–Wheeler Alignment tool and the SEAStAR tool [(https://github.com/armbrustlab/SEAStAR)](https://github.com/armbrustlab/SEAStAR). The aligned reads were counted for each gene model (Joint Genome Institute: Thaps3 extended models) using the SEAStAR tool. RNA sequences and analysis products were deposited in NCBI’s Gene Expression Omnibus and are accessible through GEO series accession number [GSE67971.](http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE67971)

**Gene set enrichment analysis.** Read counts from transition and steady-state transcriptomes were first normalized by the trimmed mean of M-values method (TMM: R package, edgeR; ref. 33), then each gene was normalized to its own average expression level at low CO2 (*<*312 µatm for transition samples and *<*350 µatm for steady-state samples) and log2 transformed. The normalized gene expression data were tested for correlation to CO2 using the Spearman rank correlation test (cor. test, R stats). Genes with significant (*p<*0.05) correlation to CO2 in transition and steady-state transcriptomes were tested for significant gene set enrichment with the hypergeometric test (phyper, R stats) in categories defined by Kyoto Encyclopedia of Genes and Genomes (KEGG 58.1, 1 June 2011), Gene Ontology (GO 2009).

**Gene clustering analysis.** A data set of 98 microarray and RNA-seq samples representing ten experiments15–20 was used to identify co-expressed gene clusters. All RNA-seq data were TMM normalized, and log2 ratios versus triplicate control samples were combined with microarray log2 ratios. To estimate the co-expression of transcripts, Pearson pairwise correlation distances were computed across all samples, following normalization such that all within-sample standard deviations were equal to one. Hierarchical clustering of these distances using Ward’s method (fastcluster)34 identified a hierarchy of co-expressed genes, from which 400 co-expressed groups were selected using arbitrary cut height. Quality controls required a cluster to have at least 15 genes expressed with normalized mean square residual35 *<*0.6 across the CO2 experiments and to be significantly correlated with CO2 in transition or steady-state experiments (FDR *<* 0.0001) to be considered for further analysis. Multiscale bootstrap resampling was performed to estimate the significance and reproducibility of sub-clusters among this hierarchy. Candidate *cis*-regulatory regions were identified using MEME (ref. 36) from 0 to 800 base pairs upstream of gene start sites, and TOMTOM (ref. 37) was used to assess similarity of candidate *cis*-regulatory motifs to previously characterized *cis*-regulatory elements.

**Homology and alignments.** Searches for homologous sequences were performed by a hidden Markov model search tool (HMMsearch; ref. 38) across the gene models. Multiple alignments were performed with MAFFT (ref. 39) and visualized in Jalview40.

**RT-qPCR.** To quantify gene expression we used primers (Supplementary Table 4) to amplify genes for δ-CA3 (ref. 41) and a putative transporter (pID 262258) from poly(A) selected RNA (MicroPoly(A)Purist kit, Life Technologies) on a StepOnePlus instrument with the Power SYBR Green RNA-to-CT 1-Step kit (Life Technologies). The gene copies were quantified in technical triplicate by a standard curve generated from the gene amplicon in a linearized 2.1 TOPO vector (Life Technologies) and normalized to mRNA in each sample. A two-way ANOVA (R stats, aov) was performed on δ-CA3 gene expression with a Tukey HSD *post hoc* test (R, stats, TukeyHSD) to determine significant differences between treatments and groups respectively. A Student’s *t*-test (R, stats, *t*-test) was performed on gene expression of the putative transporter at high CO2 only, because the low-CO2 gene expression was below the quantification limit of the standard curve.

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**NATURE CLIMATE CHANGE** |

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