Divergent gene expression among phytoplankton taxa in response to upwelling

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Summary

Frequent blooms of phytoplankton occur in coastal upwelling zones creating hotspots of biological productivity in the ocean. As cold, nutrient-rich water is brought up to sunlit layers from depth, phytoplankton are also transported upwards to seed surface blooms that are often dominated by diatoms. The physiological response of phytoplankton to this process,

commonly referred to as shift-up, is characterized by increases in nitrate assimilation and rapid growth rates. To examine the molecular underpinnings behind this phenomenon, metatranscriptomics was applied to a simulated upwelling experiment using natural phytoplankton communities from the California Upwelling Zone. An increase in diatom growth following 5 days of incubation was attributed to the genera Chaetoceros and Pseudo-nitzschia. Here, we show that certain bloom-forming diatoms exhibit a distinct transcriptional response that coordinates shift-up where diatoms exhibited the greatest transcriptional change following upwelling; however, comparison of co-expressed genes exposed overrepresentation of distinct sets within each of the dominant phytoplankton groups. The analysis revealed that diatoms frontload genes involved in nitrogen assimilation likely in order to outcompete other groups for available nitrogen during upwelling events. We speculate that the evolutionary success of diatoms may be due, in part, to this proactive response to frequently encountered changes in their environment.

Introduction

Wind-driven coastal upwelling associated with eastern boundary currents delivers rich supplies of nutrients to illuminated surface waters. This phenomenon provides ideal conditions for blooms of phytoplankton that render coastal upwelling regimes centres of new production, even though their relative ocean area is small (Capone and Hutchins, 2013). Typically dominated by large chainforming diatoms, phytoplankton blooms in upwelling zones rapidly sequester carbon dioxide and are the base of short, efficient food chains that comprise a significant percentage of the global fish catch (Ryther, 1969; Estrada and Blasco, 1985; Lassiter *et al.*, 2006; Lachkar and Gruber, 2013).

The phytoplankton community in upwelling zones is postulated to undergo a 'conveyer belt cycle' in which viable cells are upwelled into sunlit waters to seed a surface bloom. The community is then advected away from the upwelled source, and some cells eventually sink

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out of the photic zone. Surviving cells at depth and positioned in future upwelled waters are able to act as seed stock once winds are favourable for upwelling (Wilkerson and Dugdale, 1987, 2008). This continuity between a subsurface population and surface bloom during an upwelling event has been observed through a combination of glider and remote sensing techniques (Seegers *et al.*, 2015).

As a result of this vertical transport to a higher light and nutrient-rich environment, phytoplankton exhibit a physiological response, termed shift-up, that includes an acceleration of processes such as nitrate assimilation and growth (MacIsaac *et al.*, 1985; Wilkerson and Dugdale, 1987). An increase in nitrate uptake rates has been repeatedly observed in simulated upwelling mesocosm experiments (Dugdale and Wilkerson, 1989; Fawcett and Ward, 2011) and in a laboratory experiment on the diatom *Skeletonema costatum* (Smith *et al.*, 1992). Shift-up, as expressed through rapid nitrate assimilation, is hypothesized to be linked to the success of diatoms in upwelling regions; it is believed that diatoms respond quickest to available nitrogen once conditions are optimal (Fawcett and Ward, 2011).

Diatoms likely exhibit a unique cellular response that orchestrates this rapid increase in nitrogen assimilation; however, molecular characterization of the shift-up response in general is currently lacking. Only upregulation of the nitrogen assimilation gene, nitrate reductase, has been observed in *Skeletonema costatum* under labsimulated upwelling conditions, indicating that there is a molecular basis for the shift-up response (Smith *et al.*, 1992). To characterize the phytoplankton community's response and investigate the molecular basis for shift-up, we applied comparative metatranscriptomics to a simulated upwelling event in a shipboard incubation experiment. Metatranscriptomics is increasingly being utilized with eukaryotic phytoplankton communities to provide a deeper understanding of molecular responses among resident phytoplankton groups (Caron *et al.*, 2017). With the growing availability of reference transcriptomes and genomes of eukaryotic phytoplankton, unprecedented levels and confidence in gene annotation are being obtained from environmental sequences (Keeling *et al.*, 2014; Alexander *et al.*, 2015b). Our results indicate that phytoplankton functional groups exhibit a highly distinct transcriptional response to being upwelled in which diatoms constitutively express genes involved in nitrogen assimilation. This strategy possibly allows diatoms to outcompete other groups for available nitrogen once physical conditions are optimal for growth.

Results and discussion

Experimental overview and physiological observations

Upwelling was simulated by collecting seawater from the 10°C isotherm (96 m) at a site lacking natural upwelling along the California coast (Fig. 1). Satellite-derived sea surface temperature and shipboard wind data indicate that upwelling-favourable conditions were not present for 13 days prior to the incubations, and viable phytoplankton cells were detected at the sampling depth (Supporting Information Figs S1 and S2). Although multiple factors contribute to the residence time of cells at depth, these data suggest an upper limit of 13 days prior to sampling. Seawater was incubated onboard the ship for up to 5 days to simulate vertical transport of upwelled waters, although the simulation included a temperature increase that would have likely been more rapid than that of natural upwelling (Supporting Information Fig. S3).

Results from the experiment indicate that a bloom of large phytoplankton (> 5 μ m) was induced with observations of shift-up, i.e., acceleration, of nitrate uptake and primary production resulting in faster growth rates within

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Fig. 1. Experimental site.

A. Satellite-derived sea surface temperature (SST) of the area on the day the incubations began. White areas indicate no data as a result of cloud cover. Water was collected at the point marked with a +.

B. Conductivity-temperature-depth (CTD) measurements for potential temperature (°C), density (σ_t ; kg m⁻³), fluorescence (raw fluorescence units) and photosynthetically active radiation (PAR; µmol photons m⁻² s⁻¹) on the afternoon prior to the start of incubations. The dashed line denotes the depth at which water was collected (96 m).

these large cells. Macronutrient concentrations in the upwelled waters remained high throughout the incubations; however, significant growth in the large (> 5 µm) phytoplankton community was observed (P < 0.05; Fig. 2A and Supporting Information Fig. S4A). The initial dissolved iron concentration was approximately 1.28 nmol l⁻¹, which is marginally higher than the typical values (< 1 nmol l⁻¹) observed in the region. For complete drawdown of nitrate, an iron-to-nitrate ratio of 8 nmol l⁻¹:20 µmol l⁻¹ is generally required (Bruland *et al.*, 2001). The initial ratio of 1.28 nmol l⁻¹:21.86 µmol l⁻¹ in the incubations therefore indicates that iron had the potential to be a limiting nutrient, which resulted in 15 µmol l⁻¹ of unused nitrate after 120 h.

This success of large phytoplankton is consistent with previous studies showing phytoplankton from large size fractions as the significant contributors to growth and new production during upwelling. Large phytoplankton consistently have greater increases in biomass and appear to outcompete small cells for nutrients during a bloom although reduced grazing pressure may also have an effect (Wilkerson *et al.*, 2000; Fawcett and Ward, 2011). Early on in the incubations, grazers of large phytoplankton may not have been abundant enough to control growth compared to those of small phytoplankton (de Baar *et al.*, 2005).

Nevertheless, the large phytoplankton community also exhibited clear physiological responses to being

upwelled. Maximum photochemical yields (F_v:F_m) of the whole community increased from 0.25 to 0.51 within the first 72 h (Supporting Information Fig. S4B). Dissolved inorganic carbon and nitrate uptake in the large cells increased throughout the experiment and was significantly higher than the small cells (Fig. 2B and C). The particulate carbon-to-nitrogen ratio (C:N) was initially 31.5:1 in the > 5 μ m size fraction but decreased to approach the expected elemental composition of 6.6:1 (Redfield et al., 1963), while C:N values remained fairly constant and above the Redfield ratio for the small size fraction (Fig. 2D). This return to Redfield stoichiometry for the larger phytoplankton cells was coupled with increasing biomass-specific NO_3^- uptake rates (V_{NO_3}) that were approximately double that of biomass-specific carbon uptake rates (V_{DIC}) (Fig. 2E and F).

For the larger phytoplankton, these data indicate a positive response once released from light limitation or a possible resting stage. The initial low F_v : F_m signifies that the community was in a quiescent state but able to return to higher photosynthetic efficiencies within 72 h. A high initial C:N ratio that approaches the Redfield-predicted value has also been observed in similar mesocosm experiments (Kudela and Dugdale, 2000; Fawcett and Ward, 2011). These studies suggest that the initial high C:N ratio indicates severe N limitation, which likely occurred as the phytoplankton cells in aged upwelling water began to sink to depth. Once released from light



Fig. 2. Measurements from the initial upwelled water and incubations at 72 and 120 h for the community > 5 µm (red) and < 5 µm (blue).

A. Chlorophyll a (closed circles), particulate nitrogen (open circles) and biogenic silica (green).

B. Primary productivity expressed as inorganic carbon uptake (μ mol C I⁻¹ day⁻¹).

C. Absolute nitrate (NO₃⁻) uptake rates (ρ , NO₃⁻ taken up per unit time).

D. Ratios of particulate carbon to particulate nitrogen.

E. Biomass-specific dissolved inorganic carbon uptake rates (V_{DIC}), i.e., inorganic carbon uptake normalized to biomass as particulate carbon (µmol C I⁻¹ day⁻¹/µmol C I⁻¹ or day⁻¹).

F. Biomass-specific nitrate uptake rates (V_{NO_3}), i.e., nitrate uptake rates normalized to biomass as particulate nitrogen (µmol N I⁻¹ day⁻¹/µmol N I⁻¹ or day⁻¹). Error bars indicate standard deviation of the mean (n = 3).

limitation, the community is able to stabilize with large phytoplankton controlling the total C:N as time progresses. It is also possible that there was C-rich detrital material elevating the initial measurement (Fawcett and Ward, 2011) but acceleration of nitrate uptake, especially in relation to carbon uptake, to drive the phytoplankton community towards balanced growth appears to play a role. The larger cells seem to be able to take advantage of nitrate as conditions become optimal and dominate the community since they uptake nitrate at higher rates than the smaller cells.

Taxonomic composition

Large phytoplankton typically dominate blooms during upwelling as observed in this experiment (Estrada and Blasco, 1985; Lassiter *et al.*, 2006). These phytoplankton are typically chain-forming colonial diatoms such as *Chaetoceros* spp. and *Pseudo-nitzschia* spp. In addition to chlorophyll *a*, the upwelling simulation produced significant increases in biogenic silica signifying that the phytoplankton growth can mostly be attributed to diatoms (P < 0.05; Fig. 2A). Microscopic cell counts indicated that chlorophytes along with dinoflagellates were most abundant whereas initial cell abundances of diatoms were low. Following incubation, diatoms quickly became the dominant phytoplankton group, particularly those members of the genera *Chaetoceros* and *Pseudo-nitzschia* (Table 1).

Obtaining taxonomically annotated mRNA read counts also allows inferences of the relative taxonomic compositions. Metatranscriptome assembly resulted in 3.1 million contigs with levels of annotation similar to previous studies utilizing KEGG and reference transcriptomes from the Marine Microbial Eukaryote Transcriptome Sequencing Project (MMETSP; Supporting Information Table S1) (Keeling *et al.*, 2014; Alexander *et al.*, 2015b; Cohen

Table 1. Cell abundances from microscopic counts $(10^3 \times \text{cells } \text{I}^{-1} \pm 1 \text{ standard deviation of the mean of triplicate samples) of four different phytoplankton groups (diatoms, Bacillariophyceae; dinoflagellates, Dinophyceae; green algae, Chlorophyceae; haptophytes, Haptophyceae) at the initial time point and following 72 and 120 h of incubation.$

Class	To	T ₇₂	T120
		12	- 120
Bacillariophyceae			
Total	$\textbf{5.8} \pm \textbf{2.5}$	250 ± 55	2500 ± 170
Chaetoceros spp.	0.44 ± 0.63	89 ± 38	980 ± 760
Pseudo-nitzschia spp.	$\textbf{0.5} \pm \textbf{0.71}$	66 ± 55	1200 ± 710
Dinophyceae	$\textbf{6.1} \pm \textbf{3.8}$	9.7 ± 4.6	17 ± 24
Chlorophyceae	40 ± 13	250 ± 45	310 ± 85
Haptophyceae	$\textbf{0.90} \pm \textbf{0.14}$	bd	bd

Diatom abundances are shown for the total diatom assemblage (Total) and for members of the genera *Chaetoceros* and *Pseudo-nitzschia.* bd = below detection limit.

et al., 2017). These relative abundances of transcripts show that the initial community was relatively diverse and that dinoflagellates were the dominant phytoplankton group in the pre-upwelled subsurface community (Fig. 3A). Although microscopic cell counts suggest chlorophytes may have been more abundant (Table 1), dinoflagellates may have been exhibiting mixotrophy allowing them to be more transcriptionally active. By 72 and 120 h following incubation, there was an overwhelming increase in the abundance of mRNA reads attributable to diatoms, which is consistent with the bulk measurements, microscope counts and previous studies (Estrada and Blasco, 1985): diatoms were unequivocally the dominant group within the simulated upwelling event.

The taxonomic composition of diatoms followed a similar trend as the whole community with an initially more diverse diatom community that transitioned into one dominated by just two genera: Chaetoceros and Pseudonitzschia (Fig. 3B and Table 1). Chaetoceros appeared to make rapid early gains but the community became mostly Pseudo-nitzschia by 120 h. These two genera were also dominant within a previous mesocosm experiment examining shift-up at a nearby coastal California site (Kudela and Dugdale, 2000). Chaetoceros spp. were found as resting spores and may quickly germinate following upwelling to make early gains in cell abundance (Pitcher, 1990). Although a resting stage for Pseudonitzschia spp. is not known (Lelong et al., 2012), they are significant members of the phytoplankton community throughout the upwelling cycle and dominated after 120 h consistent with a peak in the Pseudo-nitzschia produced toxin, domoic acid (Lelong et al., 2012) (Supporting Information Fig. S4C). The average domoic acid concentration was 1.34 μ g l⁻¹ by 120 h, and although this concentration is lower than peak concentrations observed during blooms in California coastal waters (Schnetzer et al., 2013), it nevertheless supports that Pseudonitzschia spp. were abundant within the incubations by 120 h. This presence of Pseudo-nitzschia is unsurprising considering the reports that members of this genus often dominate subsurface chlorophyll maxima (Ryan et al., 2005), thin layers (Rines et al., 2002; McManus et al., 2008) and upwelled communities (Seegers et al., 2015) that often result in harmful algal blooms in this region.

Comparative gene expression of phytoplankton groups

Examining shifts in the total transcript pool provides a broad depiction of the responsiveness of different groups. By comparing expression levels at 0 and 72 h, these shifts reveal the initial whole transcriptome responses to simulated upwelling by the main detected groups of phytoplankton (Fig. 4A). Diatoms had a high proportion of overrepresented genes after upwelling compared to other



Fig. 3. Average taxonomic distribution by mapped reads from each time point.

A. Percentage of reads from the whole community.

B. Percentage of reads for diatom genera within all reads assigned as diatoms.

groups, over 950 (20%) of which were significantly overrepresented (*P*-value < 0.05). Dinoflagellates showed an opposite pattern with gene expression skewed towards overrepresentation in the preupwelled condition while haptophytes had an even distribution of overrepresented genes under both conditions. Interestingly, chlorophytes also had a higher number of significantly overrepresented genes postupwelling and they were able to maintain their relative proportion of the overall transcript pool unlike the dinoflagellates and haptophytes.

Smaller changes across all four groups were observed when examining shifts from 72 to 120 h (Supporting Information Fig. S5). Relatively minor changes in the whole transcript pool and a less pronounced taxonomic shift from 72 to 120 h indicates that most of the activity in relation to diatom dominance likely occurred in the first 72 h. This timing and slowing of response also corresponds to field observations that predict a 5–7 day window for cells to achieve balanced growth and transition from shift-up to a low nutrient shift-down (Dugdale and Wilkerson, 1989; Wilkerson *et al.*, 2006). It has been speculated that these shifts, or variable transcript allocation, are a reflection of r- and K-type growth strategies (Alexander *et al.*, 2015b). Our observations appear to follow this paradigm with diatoms exhibiting r-type growth and the highest transcript reallocation in terms of gene count.

Analysis of the expression of genes with shared KEGG Ortholog (KO) annotation allows for direct comparisons between taxonomic groups as orthologs normally retain the same function throughout evolutionary history. Similar or different expression of a gene among groups may signify correspondingly similar or different investments in



Fig. 4. KEGG Ortholog (KO) gene expression comparison among the four main detected phytoplankton groups: diatoms (blue), dinoflagellates (red), chlorophytes (green) and haptophytes (orange).

A. Histograms of KO counts binned by \log_2 fold change intervals of 1 for 0 and 72 h and fitted with a Gaussian curve. Dashed vertical lines indicate a \log_2 fold change of -1 or 1. The number and percentage of only the significantly (*P*-value < 0.05) overrepresented genes at 0 h (preupwelling; left) and 72 h (postupwelling; right) are annotated on each plot.

B. Venn diagram of expressed KOs at 72 and 0 h for each group.

C. Heatmap for the 1476 commonly expressed KOs at 0 and 72 h. Each row indicates an expressed KO with darker red (positive fold change) indicating overrepresentation at 72 h and darker blue (negative fold change) indicating overrepresentation at 0 h.

cellular processes at given time points. We detected 1476 orthologous genes expressed by all four taxonomic groups at 0 or 72 h (Fig. 4B and C). Only 18 genes were binned as highly overrepresented at 72 h by all four phytoplankton groups, of which many were related to chlorophyll synthesis. Over 550 genes had low absolute fold change values, many of them positive, across all four groups. These included more photosynthesis-related genes such as photosystem II constituents, photosynthesis electron transport proteins, light-harvesting chlorophyll protein complex proteins and most of the genes associated with the Calvin cycle. The shared expression of these genes across groups is unsurprising considering the community is transitioning from a deep and dark environment to a sunlit environment and would benefit from investing in photosynthetic machinery. Other genes that were highly expressed but showed little change in expression across all four groups were associated with other predictable cellular functions such as ribosomal proteins, translation initiation factors and all constituents of the citric acid cycle.

Of particular interest is the clear overrepresentation at 72 h of approximately 200 genes per taxonomic group that show little or negative fold change in the other three groups (Fig. 4C and Supporting Information Data Set S1). It is important to note that although differences in shifts in the total transcript pool were observed (Fig. 4A), all groups are still responding and highly increasing their expression of a distinct set of genes compared to the other groups. This pattern continues to hold when examining the genes that were shared between diatoms and just one or two of the other groups (Supporting Information Fig. S6). The genes highly expressed by each group appear to be of diverse function as they do not cluster into certain categories or modules but can be broadly interpreted as investments in different metabolic processes (Supporting Information Fig. S7). These unique responses may reflect fundamental differences in life strategies and ecological traits among functional groups.

To further explain the dominance of diatoms in these systems, expression of diatom annotated genes was investigated. A total of 1132 KOs were found solely in diatoms, likely due to the abundance of diatoms in our samples resulting in an improved metatranscriptome assembly for that group (Fig. 4B). However, only 173 of these KOs were significantly overrepresented at either 0 or 72 h. It is difficult to determine the importance of the remaining genes as most were expressed in low abundances.

Diatom taxa, however, were not found to respond equivalently to being upwelled; clear differences were noted between *Chaetoceros*, *Pseudo-nitzschia* and other diatoms (Fig. 5A). Expression of 2807 orthologs was detected in the genera *Chaetoceros*, *Pseudo-nitzschia* and all other diatom genera combined mostly consisting of *Thalassiosira*. Similar to what was observed for major taxonomic groups, there was large overrepresentation of distinct sets of genes, particularly in *Chaetoceros* spp., also potentially reflecting transcriptional investments in different processes at different times.

Ninety-nine significantly differentially expressed genes showed opposing fold-changes in *Chaetoceros* and *Pseudo-nitzschia* when compared with other diatoms (Supporting Information Data Set S2). This highlights that gene expression may not be as accurately assessed by combining genes at high-level taxonomic groupings. The high expression of a gene at one time point or treatment by one group may be cancelled out by another group with opposing expression leading to the incorrect conclusion for the group as a whole. Additionally, one genus could be driving expression of many genes rather than being distributed across the entire group.

Molecular characterization of the nitrogen assimilation response

Gene expression was assessed among specific diatom genera and other phytoplankton groups to investigate nitrogen assimilation and utilization, which underlies the shift-up response. Querying nitrogen-related genes for these groups reveals differences in gene expression for the diatoms compared to other phytoplankton (Fig. 5B and Supporting Information Fig. S8). The genes that were most highly expressed both pre- and post-simulated upwelling, referred to here as frontloaded, were almost all from diatoms and related to nitrogen assimilation: nitrate transporter, nitrate reductase, nitrite reductase and ammonium transporter (Fig. 5C). These frontloaded genes also showed similar expression in diatoms (absolute log_2 fold change < 1) when comparing the initial and iron-amended treatment following 72 h, which could be expected as iron was likely not limiting at his time point (Supporting Information Fig. S9).

The only similarly expressed nitrogen-related gene by other groups was glutamine synthetase within dinoflagellates and haptophytes. Within diatoms, the nitrate assimilation genes all had a positive fold change suggesting slightly greater abundance of these genes post-upwelling when compared to pre-upwelling matching our observations of increased nitrate uptake at 72 and 120 h (Fig. 2C and F). The change in expression in nitrate reductase was very low which contrasts a simulated upwelling experiment with a *Skeletonema* species (Smith *et al.*, 1992). *Skeletonema*, however, was not found to be an abundant genus within this study, and this variation further highlights potential genera-specific differences in response to upwelling.



Fig. 5. Diatom gene expression.

A. Heatmap for the 2807 commonly expressed KOs at 0 and 72 h for *Chaetoceros*, *Pseudo-nitzschia* and all other diatoms. Each row indicates an expressed KO with darker red (positive fold change) indicating overrepresentation at 72 h and darker blue (negative fold change) indicating overrepresentation at 0 h.

B. Cell schematic representation depicting frontloading of genes associated with nitrogen assimilation and utilization for diatoms. The model is based on Alexander and colleagues (2015a) and utilizes the same KO numbers. Colour indicates the average abundance of the genes (log₂ CPM) minus the absolute value of the log₂ fold change to highlight the most abundant, lowest fold change (i.e., frontloaded) genes. Genes are abbreviated as follows: NRT, nitrate transporter; GLNA, glutamine synthetase; NR, nitrate reductase; AMT, ammonium transporter; NIRA, nitrite reductase (ferredoxin); ASSY, arginosuccinate synthase; CPS, carbamoyl-phosphate synthetase; OTC, ornithine carbamoyltransferase; ASL, argininosuccinate lysase; URE, urease, ARG, arginase; GLT1, glutamate synthase (NADP/NADH); GLT2, glutamate synthase (ferredoxin); GLTD, glutamate synthase; small chain; GDHA/GDH2, glutamate dehydrogenase.

C. MA plot of nitrogen-related genes for the four main detected phytoplankton groups: diatoms (blue), dinoflagellates (red), chlorophytes (green) and haptophytes (orange). Clusters of these N-related genes are shown using *k*-means clustering (k = 8) with confidence ellipses at the 90% level to aid visualization of frontloading. Selected genes are labelled within clusters that represent the most highly frontloaded genes (black ellipse) and additional frontloaded genes (medium grey ellipse). Genes are abbreviated in the same manner as Fig. 5B.

D. Differential transcript abundance between 0 and 72 h for *Chaetoceros* (red) and *Pseudo-nitzschia* (blue) for expressed KEGG Orthologs (KOs). Each pie represents a KO and increases in size with absolute values of its coordinates to optimize visibility. Gene circles that are shaded and have grey borders are not significantly represented in either library (*P*-value \geq 0.05). Select gene names discussed in the text are labelled as follows: CHSY, chondroitin sulfate synthase; FTN, ferritin; GUCY1B, guanylate cyclase soluble subunit beta; HERC1, ubiquitin-protein ligase; MRT4; mRNA turnover protein, POT1, protection of telomeres protein 1; PRKG1, cGMP-dependent protein kinase 1; SOD, superoxide dismutases.

The urea cycle is believed to facilitate recovery from prolonged nitrogen limitation for diatoms (Allen *et al.*, 2011) but may also be important for the shift-up response. The urea cycle genes carbamoyl-phosphate synthetase and argininosuccinate synthase were also frontloaded by diatoms (Fig. 5B and C and Supporting Information Fig. S8). Several others were significantly overrepresented post-upwelling including ornithine carbamoyltransferase, arginosuccinate lyase and urease. The exception in diatoms was arginase, the final enzyme in the urea cycle that was significantly overrepresented pre-upwelling by diatoms not including *Chaetoceros* (Supporting Information Fig. S8). Low expression of arginase post-upwelling is similar to the diatom response to iron enrichment (Marchetti *et al.*, 2012) and may suggest that in both of these scenarios, there are alternative fates for urea cycle intermediates such as nitrogen storage or silica precipitation (Kröger *et al.*, 2001; Llácer *et al.*, 2008).

High relative expression of almost all of these nitrogenrelated genes in diatoms compared to most of the other phytoplankton groups at both 0 and 72 h suggests that investing in nitrogen assimilation and utilization is a priority even when conditions are not optimal for growth. Previous studies indicate that these genes are not constitutively expressed in diatoms and expression can decline with nitrogen and iron availability (Allen *et al.*, 2008, 2011; Cohen *et al.*, 2017). It is likely that expression of these assimilation genes declines if the community were to face nitrogen limitation at the surface and, at some point at depth, expression returns to high levels.

This transcriptional investment, particularly in the primary genes for nitrate assimilation such as nitrate transporters and nitrate reductase pre-upwelling, may contribute to the rapid response of diatoms as part of their shift-up process. By maintaining elevated pools of nitrogen-related gene transcripts or expressed proteins, upwelled cells are set up to rapidly assimilate available nitrogen whereas other phytoplankton groups only upregulate these genes after being upwelled into the euphotic zone. These results further support the hypothesis that one reason diatoms dominate upwelling regions is because they have the ability to take up and assimilate nitrate more quickly than other phytoplankton groups (Fawcett and Ward, 2011).

Employing this transcriptionally proactive approach to abiotic changes, or 'frontloading', has also been characterized with environmental stress response genes in coral and yeast (Berry and Gasch, 2008; Barshis et al., 2013). Additionally, it is similar to what has been observed in diatoms within a previous metatranscriptomic study in relation to iron stress. Iron-enrichment experiments in the northeastern Pacific Ocean demonstrated oceanic diatoms continued expressing genes encoding for iron-free photosynthetic proteins rather than substituting genes encoding for iron-containing functionally equivalent proteins which was different from other phytoplankton groups (Marchetti et al., 2012; Cohen et al., 2017). This strategy is speculated to provide oceanic diatoms with the ability to rapidly acclimate to the inevitable return to iron-limited conditions just as our observations show a strategy that provides certain diatoms with the ability to rapidly take up nitrogen following upwelling. Constitutive frontloading is suggested to provide organisms with resilience to such stressors (Barshis et al., 2013). Along similar lines, bloom-forming diatoms such as Chaetoceros and Pseudo-nitzschia may have evolved to frontload transcripts of particular genes depending on frequently encountered environmental fluctuations such as conditions associated with the upwelling conveyor belt cycle rather than simply reactively responding to these changes.

Chaetoceros and Pseudo-nitzschia expressed genes

Analysing genes assigned to two of the most dominant diatom genera, *Chaetoceros* and *Pseudo-nitzschia*, provides further insight into the molecular mechanisms these

genera use at depth and as part of their shift-up response. From KOs with module annotations, it is evident that the significantly overrepresented genes at both time points fall into a diverse set of functional categories even at a high level grouping (Supporting Information Fig. S10). To obtain finer resolution, gene expression among all KOs for these genera was examined.

Many overrepresented genes from Chaetoceros and Pseudo-nitzschia in the pre-upwelled condition indicate a stress response (Fig. 5D). For example, several proteins encoded by highly expressed genes promote proteasome and ubiquitin activity suggesting that the cells are degrading unneeded or damaged proteins. RNA turnover was also likely increased with high expression of exosomerelated transcripts, while expression of protection of telomeres protein promoted the stabilization of DNA (Mivoshi et al., 2008). One highly expressed gene was chondroitin sulfate synthase, which is potentially related to transparent exopolymer particle (TEP) production (Passow, 2002a; Passow, 2002b). TEP is found to be generated by Chaetoceros within the stationary phase, which may have contributed to the aggregation and sinking of these cells as well as the high particulate carbon-to-nitrogen ratios observed in the initial community (Fig. 1D).

Pseudo-nitzchia in particular expressed a set of distinctive genes as part of its shift-up response. Ferritin was highly expressed post-upwelling, possibly providing a method of storing the essential micronutrient iron (Marchetti *et al.*, 2009). As iron availability in the California upwelling regime can be sporadic and potentially growth limiting, ferritin may provide an advantage to *Pseudo-nitzschia* by concentrating iron for longer-term storage (Bruland *et al.*, 2001), although it may also be used for iron homeostasis (Pfaffen *et al.*, 2015).

At 72 h, Pseudo-nitzschia also uniquely and highly expressed a subunit of soluble quanylate cyclase (sGC or GUCY1B, Fig. 5D). sGC is the only proven receptor of nitric oxide (Denninger and Marletta, 1999) and synthesizes cyclic guanosine monophosphate (cGMP), a second messenger related to many physiological responses (Delledonne, 2005). cGMP activates protein kinase G (PRKG1), which was also significantly expressed (Fig. 5D). Although nitric oxide has been hypothesized to be an infochemical for intercellular signalling and monitoring of stress in diatoms (Vardi, 2008; Amin et al., 2012), Pseudo-nitzschia are generally not believed to have a nitric oxide synthase gene as a putative sequence was detected in only one species, P. multistriata (Di Dato et al., 2015). Pseudo-nitzschia may be utilizing sGC to monitor exhibition of stress from other genera which could allow them to rapidly adapt to changing conditions or respond to sexual cues (Basu et al., 2017). Nitric oxide is also produced by the activation of nitrate reductase (Sakihama et al., 2002). As increased nitrate reductase activity occurs as a part of the shift-up response, sGC may be used to monitor the continuation of that response and promote certain cellular functions such as gliding of pennates or binary fission (Thompson *et al.*, 2008). Inhibition of sGC prevents the germination of *Leptocyclindrus danicus* resting spores suggesting that it may be involved in transitioning from a resting stage in certain diatoms (Shikata *et al.*, 2011).

Examination of this gene in our reference database reveals that it is highly conserved among *Pseudo-nitzschia* spp. but not ubiquitously present among diatoms (Supporting Information Fig. S11). From this high expression of sGC as part of the upwelling response, evolutionary conservation of this gene, and potential to act as an important signalling device among *Pseudo-nitzschia* spp., we speculate that it may play an important role in shift-up. Additionally, uncharacterized genes that have been previously observed to also increase in expression in comparable microarray studies (Ashworth *et al.*, 2013; Nymark *et al.*, 2013) may elucidate additional mechanisms for the shift-up response in *Chaetoceros* and *Pseudo-nitzschia* (Supporting Information Data Set S3).

Conclusions

Our simulated upwelling experiment in the California Upwelling Zone is consistent with previous physiological observations of the shift-up response in upwelled phytoplankton: growth of large chain-forming diatoms and increased nitrate assimilation rates. The application of metatranscriptomics to the entire phytoplankton community highlights the divergent transcriptional response of major phytoplankton groups and diatom genera, potentially reflecting variations in their life history strategies. By frontloading nitrogen-related genes, diatoms exhibit the potential for possessing an abundant transcript and/or protein pool allowing them to respond to available nitrate more rapidly than other phytoplankton. This trait is not unlike the response of oceanic diatoms to iron enrichment and may indicate that diatoms have evolved to frontload transcripts in response to frequently encountered changes in their environment. Although the characterization of the shift-up response has largely been focused on nitrogen-related pathways, it is likely that other uncharacterized genes and pathways are also important for diatom success.

Experimental procedures

Sample collection

On 17 July 2014, upwelling conditions were not present at a site within the California Upwelling Zone $(35^{\circ}56.071'N,$

121°44.022′W; Fig. 1A and Supporting Information Figs S1 and S2). At 05:00 PDT (12:00 GMT) at the same location, viable phytoplankton cells were detected via imaging flow microscopy (FlowCAM; Fluid Imaging Technologies Inc., Scarborough, ME, USA) at 96 m although the initial phytoplankton assemblage was small (a total of ~ 75 cells ml⁻¹ greater than 5 μ m in size were found). The sampling depth corresponded to the 10°C isotherm (Fig. 1B).

Seawater from this depth was processed immediately for the initial time point. To simulate upwelling, additional seawater from the same depth was filled into a large acidrinsed HDPE barrel for homogenization, dispensed to triplicate 10 I Cubitainers[®] (Hedwin Corporation, Newark, DE) and incubated in an on-deck plexiglass incubator with flow-through seawater at 33% incident irradiance. Seawater collection and incubations followed trace metal clean techniques as they were conducted as part of a larger study to examine diatom responses to iron addition or removal (R. H. Lampe, unpublished). Temperature and on deck irradiance values throughout the incubation are provided in Supporting Information Fig. S3.

Based on macronutrient drawdown, triplicate cubitainers were harvested following 72 and 120 h of incubation. Subsamples from each cubitainer were preserved or measured for chlorophyll *a*, taxonomic composition (by microscopy and FlowCAM), biogenic silica, F_v : F_m , domoic acid, dissolved nutrients, particulate carbon and nitrogen, carbon and nitrogen uptake and RNA. Chlorophyll *a*, particulate carbon and nitrogen and nitrate uptake rates were size fractionated using a series filter cascade, while carbon uptake rates were size fractionated using a mesh spacer. Additional methods are described in the Supporting Information.

CTD, satellite and meteorological data

Potential temperature, density, photosynthetically active radiation and fluorescence were obtained from sensors mounted on a 24-bottle rosette onboard the R/V Melville (Seabird 911+ conductivity-temperature-depth sensor). Satellite-derived sea surface temperature data on a 0.0125° grid is from the NOAA POES AVHRR satellite courtesy of the NOAA/NESDIS Center for Satellite Applications Research. These data were downloaded from the NOAA CoastWatch Browser and were plotted with matplotlib (Hunter, 2007) for Python v2.7. Wind speed and direction were obtained from the shipboard meteorological system (MetAcq) on the R/V Melville.

Chlorophyll

A total of 400 ml of seawater was gravity-filtered through a 5 μ m polycarbonate filter (47 mm) followed by a GF/F filter (25 mm) under gentle vacuum pressure (< 100 mmHg).

Filters were rinsed with 0.45 μ m filtered seawater and immediately frozen at -80°C until analysis. Chlorophyll *a* extraction was performed using 90% acetone at -20°C for 24 h and measured via *in vitro* fluorometry on a 10-AU fluorometer (Turner Designs, San Jose, CA) using the acidification method (Parsons *et al.*, 1984).

Biogenic silica

Biogenic silica was determined via filtration of 335 ml onto 1.2 μ m polycarbonate filters (45 mm). Concentrations were measured using a NaOH digestion in teflon tubes (Krause *et al.*, 2009) and a colorimetric ammonium molybdate method (Brzezinski and Nelson, 1995).

Particulate carbon, particulate nitrogen and nitrate uptake

To assess nitrate (NO₃⁻) uptake rates, 618 ml of subsamples were spiked with Na¹⁵NO₃ at no more than 10% of ambient nitrate concentration and incubated for 8 h in the flow-through plexiglass incubator. Following incubation, seawater filtration commenced immediately and was performed by gravity through a 5 μ m polycarbonate filter (47 mm) and with an in-line vacuum (< 100 mmHg) onto a precombusted (450°C for 5 h) GF/F filter (25 mm). Cells on the 5 μ m polycarbonate filter were then rinsed onto an additional precombusted GF/F filter (25 mm) using an artificial saline solution. Filters were then stored at –20°C.

Prior to analysis, filters were dried at 50°C for 24 h then encapsulated in tin. Particulate nitrogen (PN), particulate carbon (PC) and atom percentage of ¹⁵N were then quantified from this filter using an elemental analyser paired with an isotope ratio mass spectrometer (EA-IRMS). Samples were not acidified to remove particulate inorganic carbon. Absolute uptake rates (ρ , NO₃ taken up per unit time) were calculated using a constant transport model (Eq. 3 from Dugdale and Wilkerson, 1986). Biomass-specific NO₃ uptake rates (V, NO₃ taken up per unit PN per unit time) were also calculated according to the constant specific uptake model (Eq. 6; Dugdale and Wilkerson, 1986). ¹⁵NO₃ uptake rates were not corrected for possible losses of ¹⁵N in the form of dissolved organic nitrogen (Bronk et al., 1994); therefore, the reported values are considered conservative estimates or net uptake.

Dissolved inorganic carbon uptake

A total of 60 ml of samples from each cubitainer were distributed into light and dark bottles cleaned with 1.2 mol l⁻¹ HCl. For each bottle, 1.2 μ Ci of NaH¹⁴CO₃ was added and mixed. A 1 ml subsample was taken and added to vials containing 6 mol l⁻¹ NaOH to trap and validate the

initial inorganic H¹⁴CO₃ quantities. The light and dark bottles were incubated on-deck for 6.5–8 h. Samples were filtered onto stacked polycarbonate filters (5 and 1 µm) separated with a mesh spacer. Blank control bottles also containing 1.2 µCi of NaH¹⁴CO₃ were filtered onto a GF/F filter after 5 min and had counts similar to dark bottles. Filters were vacuumed dried, placed in scintillation vials with 0.5 ml of 6 mol I⁻¹ HCI, permitted to degas for 24 h and counted using a Beckman Coulter LS 6500 scintillation counter. Reported values are light bottles minus dark bottles. Biomass-specific dissolved inorganic carbon (DIC) uptake rates (V_{DIC}) were calculated by normalizing DIC uptake to PC.

Cell abundances and flow microscopy (FlowCAM)

Phytoplankton cell abundances and species composition were determined by microscopic examination. For microscopic counts, 50 ml of samples were preserved in 2% Lugol's lodine and settled for > 24 h in Utermöhl chambers (Utermöhl, 1958). Counts were performed at 100×, 200× and 400× using a Leica DMIL inverted microscope on a minimum of 400 total cells in at least five fields of view.

The viability of cells from the initial seawater and throughout the incubations was monitored through imaging flow microscopy (FlowCAM, model VS4; prosilica colour camera; C71 syringe pump) with Visual Spreadsheets v3.1. The FlowCAM was operated in trigger mode with a 532 nm, 5 mW laser. In this mode, image acquisition by the camera was triggered by chlorophyll or phycoerthrin fluorescence with a minimum threshold of 400 (background noise had a setting of 150–200).

Samples were drawn from the cubitainers into 50 ml Falcon tubes and stored at 4°C in the dark pending processing (typically within 3 h of collection). At least 5 ml of sample was filtered through a 300 µm nitex mesh screen and passed through the system at a flow rate of 0.2–-0.3 ml min⁻¹ using a syringe pump equipped with a 5 ml glass syringe. Both a 300 µm flow cell (10× objective) and 100 µm flow cell (20× objective) were used. A digital size filter was applied so that only cells > 5 µm were captured in images. The flow cell and tubing were well flushed with Milli-Q water and 70% ethanol between each sample run to avoid cross-contamination.

RNA extraction and sequencing

Seawater was filtered onto 0.8 μ m Pall Supor filters (142 mm) using a peristaltic pump, immediately flash frozen in liquid nitrogen, then stored in either liquid nitrogen or at -80°C until extraction. RNA was extracted using the ToTALLY RNA Total RNA Isolation Kit and treated with DNase 1 (Life Technologies, Grand Island, NY). The extraction procedure was followed according to the manufacturer's instructions with additional first step of glass bead addition to assist with organic matter disruption. RNA quantity and purity were assessed prior to sequencing on an Agilent Bioanalyzer 2100. Total RNA from the triplicate samples for the initial time point (T0) and the first time point (T72) were pooled into one sample due to low RNA yields. Triplicate samples were maintained for the second time point (T120). Library prep was conducted with the Illumina TruSeq Stranded mRNA Library Preparation Kit and HiSeq v4 reagents. Sequencing of barcoded samples was performed on an Illumina HiSeq 2000 (125 bp, paired-end).

Metatranscriptome assembly, annotation and read quantification

Reads were trimmed for quality and adapter removal using Trimmomatic v0.32 (paired-end mode, adaptive quality trim with 40 bp target length and strictness of 0.6, minimum length of 36 bp) (Bolger et al., 2014). Trimmed paired reads that overlap were merged into single reads with BBMerge v8.0. Merged pairs and non-overlapping paired-end reads from all samples were then used to assemble contias using ABvSS v1.5.2 with varied k-mer sizes (32, 55, 78 and 102)(Birol et al., 2009). Assemblies for each k-mer size were merged using Trans-ABySS v1.5.3 to remove redundant contigs (Robertson et al., 2010), and those shorter than 125 bp were discarded. Read counts were obtained by mapping trimmed reads to contigs with Bowtie2 v2.2.6 (Langmead and Salzberg, 2012) and filtered by mapping quality (MAPQ \geq 10) with SAMtools v1.2 (Li et al., 2009). Mapping percentages are provided in the Supporting Information Table S3.

Annotation was assigned by best homology (lowest E-value) to protein databases using BLASTX v2.2.31 (*E*-value $\leq 10^{-5}$). For taxonomic identification, Marine-RefII, a custom reference database was used. MarineRefll contains predicted protein sequences of marine microbial eukaryotes and prokaroytes including all sequenced transcriptomes from the Marine Microbial Eukaryote Transcriptome Sequencing Project (Keeling et al., 2014). MarineRefII was supplemented with transcriptomes of isolated phytoplankton from these incubations adding increased confidence in the taxonomic annotation of some contigs (Supporting Information Table S4). To assign gene function to contigs, the same methodology with the Kyoto Encyclopaedia of Genes and Genomes (KEGG; Release 75) was used (Kanehisa et al., 2017). The best hit with a KEGG Ortholog (KO) number from the top 10 hits was chosen. Similarly, analysis of module annotations (MO) was conducted by selecting the top BLASTX hit with a KEGG MO number from the top 10 hits. A summary of annotation results is provided in Supporting Information Table S2.

Differential expression analysis

Differential expression was assessed by summing read counts of contigs within a taxonomic group (phylumbased or genus for only the diatoms, Chaetoceros and Pseudo-nitzschia) by KEGG Gene Definition or KO annotation. edgeR v3.12.0 was used to calculate normalized fold change and counts-per-million (CPM) from pairwise comparisons within each taxonomic group using the exactTest function (Robinson and Smyth, 2008; Robinson et al., 2010). By normalizing within each taxonomic group, shifts in relative abundances are accounted for although reduced sequencing depth for proportionally lower groups may influence fold change estimations for genes detected in one sample but not another. Significance (*P*-value < 0.05) was calculated by using edgeR's estimate of tagwise dispersions utilizing the available replication within each taxonomic group (Supporting Information Table S2 and Supporting Information Fig. S12) (Chen et al., 2014). ExactTest output in combination with the taxonomic distributions per gene was plotted using a custom plotting function available at https:// github.com/marchettilab/mantaPlot.

Shared expression of gene was considered when a gene was detected in at least one of the libraries under comparison for each taxonomic group. For binning of genes displayed in heatmaps (Figs 4C and 5A), a positive or negative fold change, variance greater than the number of taxonomic groups and fold change greater than or less than all other groups were used. Genes with a log₂ fold change greater than 2 or less than -2 but had a variance less than the number of taxonomic groups were considered similarly overrepresented by all groups. Otherwise, the expression level was considered similar on the basis of fold change. These data were visualized with pheatmap v1.0.8. For KEGG module-based differential expression, quantitative metabolic fingerprinting was used (Alexander et al., 2015a). Briefly, read counts annotated for each KEGG module category were summed and then normalized by the total number of reads for the time point per functional grouping. These data were also visualized with pheatmap v1.0.8. Gene expression ratios for microarray studies and orthologous genes were obtained from The Diatom Portal (Ashworth et al., 2016).

Statistical procedures

One-way ANOVAs followed by Dunn's multiple comparison test were performed on the biological and chemical properties of the seawater (non-gene expression data) in Graphpad PRISM v7.03.

Data deposition

The data reported in this article have been deposited in the National Center for Biotechnology (NCBI) sequence read archive under the accession no SRP074302 (BioProject accession no PRJNA320398). Assembled contigs, read counts and annotations are available on Zenodo (https://doi.org/10.5281/zenodo.1256894). Isolate 18S rDNA sequences, transcriptome raw reads, assemblies and predicted peptide sequences are deposited in Cyverse (http://www.cyverse.org) under the project name unc_phyto_isolates (Supporting Information). Isolate 18S rDNA sequences are also deposited in Genbank (accession nos KX229684–KX229691).

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher's web-site:

Appendix S1. Supporting Information

Data Set S1. Uniquely overrepresented genes displayed in Fig. 4C for each taxonomic group.

Data Set S2. *Chaetoceros* and *Pseudo-nitzschia* genes that show significantly opposing expression to other diatoms.

Data Set S3. Genes of unknown function significantly expressed in our study and with positive fold-changes in similar studies on *T. pseudonana* (tps) and *P. tricornutum* (pti). The study with *T. pseudonana* shows genes with positive fold-changes during exponential growth and light after three days. The dataset with *P. tricornutum* shows genes expressed after 48 hours of darkness followed by 24 hours of re-exposure to light.