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Molecular and physiological responses of diatoms to variable levels of irradiance and nitrogen availability: Growth of *Skeletonema costatum* in simulated upwelling conditions

G. Jason Smith, Richard C. Zimmerman, and Randall S. Alberte

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**Abstract**

Molecular mechanisms that drive metabolic acclimation to environmental shifts have been poorly characterized in phytoplankton. In this laboratory study, the response of light- and N-limited *Skeletonema costatum* cells to an increase in light and NO$_3^-$ availability was examined. C assimilation was depressed relative to N assimilation early in enrichment, and the photosynthetic quotient (O$_2$ : CO$_2$) increased, consistent with the shunting of reducing equivalents from CO$_2$ fixation to NO$_3^-$ reduction. The concomitant increase in dark respiration was consistent with the increased energetic demand associated with macromolecular synthesis. The accelerations of N-specific rates of NO$_3^-$ uptake and nitrate reductase activity (NRA) over the first 24 h were comparable to observations for coastal upwelling systems. Increases in cell-specific rates of these processes, however, were confined to the first 8 h of enrichment. The abundance of 18s ribosomal ribonucleic acid (rRNA) increased immediately after the environmental shift, followed by increases in levels of NR-specific mRNA that coincided with the acceleration in NO$_3^-$ assimilation. NRA, however, exhibited a diurnal rhythm that did not correspond to changes in NR protein abundance, suggesting that enzyme activity was also regulated by direct modulation of existing NR protein by light and NO$_3^-$ availability.

New production in the sea is driven primarily by NO$_3^-$ assimilation (Dugdale and Goering 1967). NO$_3^-$ is supplied to coastal upwelling systems in periodic pulses of cold, nutrient-rich water driven to the ocean surface by wind forcing (Codispoti 1983). Although upwelling systems produce dense phytoplankton blooms under certain conditions, there is a high degree of temporal and spatial variability in the production cycle, and not all upwelling systems or events...
quent development of phytoplankton blooms depend on the mixing environment, the supply of NO$_3^-$, and the initial algal biomass, which collectively lead to variable rates of NO$_3^-$ depletion as a water mass moves offshore (Zimmerman et al. 1987; Dugdale et al. 1990).

Although some of the metabolic shifts observed in natural upwelling systems may result from shifts in species composition (Chavez et al. 1990), such successional processes might reflect variable metabolic lag responses of individual species to improved conditions, which dictate the ecological success of certain members of the community. For cells exhibiting N deficiency when exposed to the improved growth conditions, the rate of induction of NO$_3^-$ metabolism in particular, may control the dynamics of the phytoplankton bloom.

Environmental shifts that result in improved conditions can induce a suite of molecular and physiological responses that lead to higher growth rates (shift-up, Schaecter 1968). This phenomenon has been used to examine the regulatory pathways involved in the induction of specific enzyme systems and metabolic processes such as protein synthesis (Ingraham et al. 1983). Metabolic induction is of ecological importance because it often produces significant temporal lags between the stimulus (environmental shift) and the ecologically significant response (growth) that can be affected by the nature of the environmental shift, as well as the prior metabolic status of the cell (Ingraham et al. 1983). Time lags in the response of phytoplankton to improved growth conditions have been observed in the field (MacIsaac et al. 1985) and in laboratory culture (Collos 1986). Furthermore, predictions of numerical models indicate that these lags can have significant effect on the timing and intensity of phytoplankton bloom formation (Zimmerman et al. 1987). The physiological processes responsible for the observed lags, however, are poorly understood, and alternative hypotheses have been proposed to explain the field observations (Garside 1991).

Laboratory studies of unialgal cultures subjected to N starvation suggest that the temporal response to N resupply can result in an uncoupling between N uptake and cell division (Collos 1982) and lead to variable lags in the accumulation of cellular metabolites (Sakshaug and Holm-Hansen 1977). The response of *Chlamydomonas reinhardtii* to N resupply entails, at least, differential expression of genes encoding photosynthetic functions (Plumley and Schmidt 1989). Similar regulatory processes acting on N assimilation pathways may also underlie lags in responses of marine phytoplankton to improved growth conditions.

NO$_3^-$ assimilation is subject to some of the most extensive environmental regulation of any metabolic pathway in photosynthetic cells. The rate-limiting enzyme involved in NO$_3^-$ assimilation in higher plants and algae, nitrate reductase (NR, EC 1.6.6.1), has been shown to be nitrate inducible (Cove and Pateman 1969; Beecers and Hageman 1980), light activated (Packard 1973; Aparicio et al. 1976; Martinez et al. 1987), and ammonium repressible (Hipkin and Syrett 1977). Additionally, given the high energetic cost of NO$_3^-$ and NO$_3^-$ reduction, induction of this metabolic pathway can affect the photosynthetic performance of phytoplankton. As a result, modulation of NR gene expression and enzyme activity by environmental factors may affect the time scales of NO$_3^-$ utilization and bloom formation in the sea.

In this study, the metabolic responses of the marine diatom *Skeletonema costatum* to increased light and nutrient availability were measured to determine whether qualitative and quantitative patterns of cellular metabolic adaptation by unialgal cultures were consistent with observations of natural phytoplankton communities, define the molecular levels of control on the induction of NO$_3^-$ metabolism by light and NO$_3^-$ enrichment, and characterize physiological and molecular processes that can serve as useful indices of the response of phytoplankton cells to transient environmental conditions.

**Materials and methods**

**Culture conditions**—Axenic stocks of *S. costatum* (clone SKEL, obtained from the Provasoli-Guillard Center for the Culture of Marine Phytoplankton) were maintained as batch cultures in f/2 medium (Guillard...
1975) at 15°C on a 14:10 L/D cycle at an incident PPF (photosynthetic photon flux) of 30 μmol quanta m⁻² s⁻¹. A 15-liter glass carboy, containing f/2 medium supplemented to 400 μM NO₃⁻, was inoculated with 500 ml of late-log-phase culture yielding an initial density of 2 × 10⁸ cells ml⁻¹. The culture was grown to stationary phase (3.5 × 10⁸ cells ml⁻¹) under a 14:10 L/D cycle at a PPF of 170 μmol quanta m⁻² s⁻¹, then diluted by 50% with fresh f/2 using NH₄⁺ as the sole source of N (final concn, 20 μM) to repress NO₃⁻ metabolism (Fig. 1, point a). The culture reached stationary phase 2 d after NH₄⁺ enrichment (Fig. 1, point b) and the PPF was reduced to 3 μmol quanta m⁻² s⁻¹ with several layers of neutral density screening. After 2 d under low irradiance, the culture was again diluted with f/2 containing NO₃⁻ as the sole N source (final concn, 60 μM). The irradiance was returned to 170 μmol quanta m⁻² s⁻¹ (Fig. 1, point c) ~15 min after exposure to the elevated NO₃⁻ concentration. This last manipulation defined the start of the simulated upwelling treatment.

**Bulk properties**—Samples were collected daily during the light-limited period and every 4 h after the environmental shift. Macronutrient (NO₃⁻, NO₂⁻, NH₄⁺, and PO₄³⁻) concentrations were determined with a flow-injection autoanalyzer (Flow Injection Systems). Cells were removed by centrifugation (1,000 × g, 10 min) and the supernatant was frozen (-20°C) until analyzed for nutrient content.

Cell density was determined from the mean of cell counts in ten hemacytometer fields. Photosynthetic pigments were assayed by duplicate aliquots of the well-mixed culture collected onto Whatman GF/C filters under vacuum (20 mm of Hg). Pigments were extracted by grinding the filters in ice-cold 90% (vol/vol) acetone. Chl a concentrations were determined spectrophotometrically with the equations of Jeffrey and Humphrey (1975).

Particulate organic C (POC) and N (PON) were determined with a CHN analyzer (Control Equipment XA-240) on duplicate samples collected onto precombusted GF/C filters, rinsed with sterile seawater, and dried at 60°C.

![Fig. 1. Changes in Skeletonema costatum cell density during simulated upwelling manipulations. Point a represents time of dilution of stationary-phase, NO₃⁻-grown culture with medium supplemented with NH₄⁺ as sole N source. Point b represents time of low light shift-down. Point c represents time of high light and NO₃⁻ enrichment corresponding to the start of the simulated upwelling event or shift to improved growth conditions. Black bars indicate dark phase of 24-h light-dark cycle.](image)

Protein and nucleic acids were quantified from duplicate samples collected onto GF/C filters, wrapped in aluminum foil, frozen in liquid nitrogen, and stored at -70°C. The frozen filters were homogenized in chilled TBS (Tris-buffered saline: 150 mM NaCl, 20 mM Tris-HCl, pH 7.5) containing 0.1% Nonidet-P40 and centrifuged. Protein content of the supernatant was determined with the bichloro acid technique (Smith et al. 1985) with a BSA standard. Concentrations of ribonucleic acid (RNA) and deoxyribonucleic acid (DNA) in the supernatant were determined with a fluorometric technique (Dorch et al. 1983) modified by the substitution of propidium iodide for ethidium bromide. Relative fluorescence emission was measured at 620 (±5) nm with 552- (±5) nm excitation in an Aminco SPF-500 spectrophotometer. RNA content was determined from the difference in sample fluorescence before and after digestion with 100 μg ml⁻¹ RNase A (DNase-free) for 1 h at 37°C. DNA content was calculated directly from residual fluorescence. Calf thymus DNA and torula yeast RNA were used as fluorometric standards after their concentrations had been calibrated spectrophotometrically (DNA: 1 O.D. 260 nm...
Physiological rate processes: Photosynthesis—Photosynthesis vs. irradiance ($P$ vs. $I$) responses were measured with a polarographic oxygen electrode (Rank Brothers) kept at 15°C. Illumination was provided by a Kodak slide projector (ELH bulb). PPF was adjusted to 15 different irradiances with neutral density filters. Photosynthetic C fixation was measured on samples enriched to 1 μCi ml$^{-1}$ with NaH$^{14}$CO$_3$ (ICN, sp act = 840 μCi μmol$^{-1}$). Samples were incubated for 1 h in glass scintillation vials kept in a seawater-cooled (15°C) photosynthetron. Irradiance was provided by a 300-W quartz-halogen lamp and adjusted for each incubation vial with neutral-density screens. Vials wrapped with black electrical tape were used to determine dark uptake. After incubation, the contents of the vials were collected onto GF/C filters, rinsed with three volumes of sterile seawater, and acidified with 1 N HCl. Incorporation of $^{14}$C was determined by scintillation counting. Quench correction used the sample channels ratio calibrated with internal standards.

Nitrate uptake—$^{15}$NO$_3^\cdot$-saturated uptake was measured at 15 different PPFs in scintillation vials incubated as described above. Each vial (20 ml) was inoculated with distilled water containing 20 mM $^{15}$NO$_3^-$, resulting in a 40 μM enrichment of $^{15}$NO$_3^-$ to the existing culture medium. Cells were collected onto precombusted GF/C filters after a 2-h incubation, rinsed with two volumes of sterile seawater, and dried at 60°C. The $^{15}$N enrichment of the samples was determined with a Roboprep TracerMass mass spectrometer. $^{15}$NO$_3^-$ (PON-specific $^{15}$NO$_3^-$ uptake rate, h$^{-1}$) was calculated according to Dugdale and Wilkerson (1986).

Nitrate reductase activity—Nitrate reductase activity (NRA) was measured with an in vivo assay that measured NO$_2^-$ production by alcohol-permeabilized cells (Brunetti and Hageman 1976). For *S. costatum*, 4% (vol/vol) 1-propanol yielded the highest NRA, and the rate of NO$_2^-$ production was linear over 2 h. Cells were collected by centrifugation (1,000 × g, 5 min), resuspended in incubation medium (4% 1-propanol in sterile distilled water containing 20 mM NaNO$_3$), and incubated in the dark at 15°C on a rotary shaker table. NO$_2^-$ concentrations were measured colorimetrically (Strickland and Parsons 1972) on aliquots (cells removed by centrifugation) taken at 15-min intervals. NRA (μmol NO$_2^-$ liter$^{-1}$ h$^{-1}$) was calculated from the slope of the NO$_2^-$ concentration time series and normalized to the appropriate biomass unit.

Molecular analysis: Determination of RNA transcript abundance—Total cellular RNA was isolated with the single-step acid guanidinium isothiocyanate-phenol-chloroform method (Chomczynski and Sacchi 1987). All reagents used were molecular biology grade from Sigma. Cells were collected by centrifugation (2,000 × g, 5 min, 15°C) and washed by centrifugation through sterile seawater, followed by isotonic TBS containing 5 mM EDTA at 4°C. The cell pellet was extracted with guanidinium isothiocyanate at room temperature for 1 h with intermittent vortexing. The extract was acidified with 0.1 volume of Na-acetate, pH 4.0, further disrupted by grinding in a glass homogenizer, extracted with phenol:chloroform (1:0.2, vol/vol). Aqueous and organic phases were separated by centrifugation. RNA was precipitated from the aqueous phase at −20°C after the addition of an equal volume of cold 100% isopropanol. The RNA pellet was dissolved in guanidinium isothiocyanate buffer, reprecipitated with absolute ethanol at −20°C, dissolved in diethylpyrocarbonate-treated sterile distilled water, quantified spectrophotometrically at 260 nm, and stored at −20°C as an ethanol precipitate until further analysis. RNA yield was 75–90% of the bulk RNA content determined by the fluorometric assay.

Total RNA was size fractionated by electrophoresis on denaturing 1.1% agarose-formaldehyde gels for 2.5 h at 40 V, transferred to NYTRAN membranes (Schleicher and Schuell), and immobilized by baking at 80°C under vacuum (Maniatis et al. 1982). DNA probes were used to quantify the abundance of specific RNA transcripts. The 18S probe (DNA hybridization probe for small subunit rRNA) was generated with universal primers for small subunit rRNA (Sogin 1990) in a polymerase chain reaction (Saiki et al. 1988) amplification of *S. costatum* genomic DNA. The plasmid
pSCNR21 (DNA hybridization probe for NR transcripts) contained an 800 bp cDNA fragment isolated by screening a S. costatum cDNA library with a PCR (polymerase chain reaction) product synthesized with oligonucleotide primers to the NR coding sequence. The design of the NR primers was based on the amino acid sequence of the N-terminal Mo3tRin and C-terminal cytochrome b domains of vascular plant and fungal NR (Smith and Alberte unpubl. data). Probe specificity was determined by hybridization of the northern blots with random-primed $^{32}$P-labeled DNA probes with standard protocols (Feinberg and Vogelstein 1983; Wahl et al. 1987).

Temporal variation in the relative abundances of 18S and pSCNR21 homologous gene transcripts was estimated from dot blots of total RNA prepared with a BIO-DOT (BioRad) microfiltration apparatus. Equal amounts of total RNA (10 µg) from each time point were denatured with glyoxal (Thomas 1983), serially diluted with 1% SDS [sodium dodecyl sulfate (wt/vol)] before application to a NYTRAN membrane saturated with 20 x SSPE (1 x -180 mM NaCl; 10 mM NaH$_2$PO$_4$, pH 7.4; 1 mM EDTA), then baked and hybridized as described above. The same hybridization conditions were used for both probes: 42°C for 24 h with 2 x $10^6$ $^{32}$P cpm ml$^{-1}$ in prehybridization-hybridization buffer [50% formamide, 5 x SSPE, 5 x Denhardt's (1 x = 0.02% Ficoll 400; 0.02% BSA; 0.02% polyvinyl pyrrolidone) and 100 µg ml$^{-1}$ sheared calf thymus DNA]. The washing conditions were: 2 x 15 min at 25°C in 1 x SSPE, 0.1% SDS; 2 x 15 min at 42°C in 0.1 x SSPE, 0.1% SDS; 3 x 30 min at 65°C in 0.1 x SSPE, 0.1% SDS.

Autoradiographs of the dot blots were analyzed on a BioImage VISAGE analytical imaging system with the Whole Band Analysis software package. Relative transcript abundances at each time point were determined from regression analysis of the integrated O.D. of each dot in the dilution series, and sequence abundance was estimated as the change in O.D. per µg total RNA.

In vitro translation—Temporal changes in the composition and abundance of translatable RNA (translational capacity) were characterized by in vitro translation of 5 µg of total RNA from each sample with $^3$H-leucine in a rabbit reticulocyte lysate kit (GIBCO BRL). Protein synthesis was quantified by liquid scintillation counting of 10% (wt/vol) TCA precipitates of the translation reactions and converted to dpm by the channels ratio method calibrated with internal standards. Radiolabeled translation products were separated by SDS-PAGE (Laemmli 1970) and visualized by fluorography (Bonner and Laskey 1974).

Polypeptide composition and western blotting—Changes in cellular polypeptide composition were monitored by SDS-PAGE (polyacrylamide gel electrophoresis) analysis of the same detergent extracts used for protein and nucleic acid quantifications. Subsamples of the extracts were apportioned to equivalent protein content, precipitated on ice with cold (0°C) 80% (vol/vol) acetone, and resolubilized with SDS-PAGE sample buffer (Laemmli 1970). Variation in the relative abundance of NR was determined from western blots of SDS-PAGE gels (Towbin and Gordon 1984) probed with heterologous, polyclonal antibodies to squash-leaf NR (Redinbaugh and Campbell 1983). Immunoreactive polypeptides were localized on nitrocellulose blots with alkaline phosphatase-coupled, goat-antirabbit secondary antibodies. The integrated O.D. of each immunoreactive band was used directly as the index of NR abundance normalized to equivalent protein content.

Results

Cell growth during the simulated upwelling cycle—Cell numbers changed in a predictable pattern throughout the manipulation (Fig. 1). The addition of NH$_4$$^+$ 48 h before the transfer to light-limited conditions (Fig. 1, point a, -88 h) led to a single doubling in cell number. The culture was approaching stationary phase when transferred to a PPF below photosynthetic compensation (Fig. 1, point b, -40 h; see below), during which time cell density declined by 17%. The light and NO$_3$ enrichment (Fig. 1, point c, 0 h) led to another round of cell division that was completed at +30 h. The culture returned to stationary phase after a
Fig. 2. Changes in dissolved nutrient concentrations and biomass indicators of the Skeletonema costatum culture after the shift to improved growth conditions. [a.] Dissolved NO₃⁻, NH₄⁺, and PO₄³⁻ concentrations in the growth medium. [b.] Concentrations of Chl a, DNA, and RNA. [c.] Concentrations of POC and PON. [d.] Variation in the C : N atomic ratio in the particulate fraction.

1.7-fold increase in cell number. About 10% of the cells observed in the stationary-phase culture 48 h after enrichment were spermatogonia, but none were observed before this time.

Nutrient utilization—The NO₃⁻ enrichment (60 μM), relative to the standing stock of cell nitrogen was 0.43 (DIN : PON) and was depleted from the medium after 24 h (Fig. 2a). The residual NH₄⁺ concentration was 2 μM at 0 h, and no net uptake of NH₄⁺ occurred until the NO₃⁻ concentration dropped below 10 μM, ~10 h after enrichment. Orthophosphate remained relatively constant until 28 h after enrichment, then decreased by 60% over the next 4 h following the completion of cell division. NO₃⁻ concentrations remained below 1 μM throughout the experiment. The rate of NO₃⁻ depletion was constant [6.9 (±1.2) μmol NO₃⁻ liter⁻¹ h⁻¹] over the first 8 h after enrichment.

Bulk properties—Concentrations of Chl a, DNA, and RNA in the culture tracked the decline in cell density during the light-limited phase (~40 to 0 h), indicating no net synthesis of these components during this period (data not shown). In contrast, the shift to improved environmental conditions resulted in a 20% increase in the volume-specific content of Chl a between 0 and +20 h and a 32% increase in DNA content between +8 and +12 h (Fig. 2b). Volume-specific levels of RNA exhibited a transient, 3-fold increase between 0 and +12 h, then declined over the subsequent 4 h to a level 50% above the value at 0 h. POC and PON levels also increased over the first 16 h (Fig. 2c). POC accumulated more rapidly than PON after +8 h, reflecting the onset of N depletion. POC content, however, increased an additional 40% during the second light period after enrichment (+20 to +32 h), indicating significant net photosynthesis over this interval. The C : N atomic ratio in the particulate fraction increased from 10.5 at +8 h to 13 at +30 h after enrichment (Fig. 2d), reflecting the rapid depletion of NO₃⁻ from the medium and indicating that the NO₃ pulse was not sufficient to overcome the N deficiency exhibited by the cells at the end of the light-limited phase.

Metabolic rate processes: NO₃⁻ uptake—V¹⁵NO₃⁻ accelerated at a rate of 7.4 x 10⁻⁴ h⁻¹ between +1 and +24 h to a maximum of 0.033 (±0.004) h⁻¹ and then declined immediately after cell division (Table 1; Fig. 3a). Uptake of ¹⁵N was independent of PPF during all of the 2-h incubations (data not shown) and average rates for each time point were calculated by pooling the samples measured at the 15 irradiances. Estimates of V¹⁵NO₃⁻ accounted for only 41% of the observed NO₃⁻ depletion rate in the culture during the first 8 h of the simulated upwelling (Table 2), possibly indicating a loss of ¹⁵NO₃⁻ from cellular pools during sample drying or by DON production.

Nitrate reductase activity—VNRA (PON-specific NRA, h⁻¹), exhibited a pronounced diurnal rhythm (Fig. 3) that peaked in the middle of the light period. Average NRA
growth of S. costatum 995

Table 1. Acceleration of N-specific $^{15}$NO$_3^-$ incorporation, $V^{15}$NO$_3^-$, and in vivo NRA, VNRA, in Skeletonema costatum between 0 h and +24 h of the simulated upwelling. Acceleration of VNRA peak activity was determined from the difference in the peaks in NRA between the first two light periods. Based on data in Fig. 3a.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Acceleration ($\times 10^4$ h$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V^{15}$NO$_3^-$</td>
<td>7.4</td>
</tr>
<tr>
<td>VNRA</td>
<td>2.4</td>
</tr>
<tr>
<td>VNRA, peak activity</td>
<td>0.6</td>
</tr>
</tbody>
</table>

was equivalent to ~50% of the $^{15}$NO$_3^-$ uptake rate during the first and second light periods after enrichment (Table 2, Fig. 3a). No recovery of NRA was observed during the third light period after NO$_3^-$ had been depleted from the medium for 24 h. As with NO$_3^-$ uptake, maximal NRA increased between 0 and +24 h after enrichment (Table 1). Acceleration of NRA over the first 24 h was 65% of the rate for $V^{15}$NO$_3^-$, whereas the daily peak in NRA increased at only 18% of the rate observed for $^{15}$N uptake. The lack of light dependence in $V^{15}$NO$_3^-$ and pronounced light-dependent phasing of VNRA demonstrated that uptake and assimilation were uncoupled.

The patterns of induction observed for $^{15}$NO$_3^-$ uptake and NRA were different when standardized to cell number rather than the N content of the culture (Fig. 3b). Although N-specific and cell-specific rates of $^{15}$NO$_3^-$ uptake exhibited similar increases from 0 to +8 h after enrichment (37 and 47%, respectively), the cell-specific uptake capacity did not continue to increase. Similarly, cell-specific NRA values exhibited only an 18% increase from 0 to +24 h after enrichment, compared to an 85% increase in N-specific activity. Again NO$_3^-$ uptake and assimilation were not in phase. Additionally, cell-specific NRA was lower at +28 h than at +4 h (Fig. 3b), indicating that cell-specific measures were more indicative of NO$_3^-$ depletion from the medium.

Photosynthesis and respiration — Light compensation ($I_c$) and saturation ($I_s$) PPFs for oxygenic photosynthesis did not change after the shift to improved growth conditions, remaining above $[I_c = 7.9 (\pm 2.8) \mu$mol quanta m$^{-2}$ s$^{-1}$, $N = 4$, 0–48 h] and below $[I_s = 73.4 (\pm 18.2) \mu$mol quanta m$^{-2}$ s$^{-1}$, $N = 4$, 0–48 h] the PPFs used for the light-limited and simulated upwelling treatments, respectively. In contrast, maximal Chl a-specific rates of net oxygen evolution and gross carbon ($^{14}$C) assimilation varied dramatically (Table 3). Oxygenic photosynthesis and dark respiration increased by 23 and 78%, respectively, between 0 and +8 h, while C assimilation rates decreased by 43% during the same period. This pattern resulted in a 4-fold reduction in the $P:R$ ratio (Table 3).

Associated with the increase in oxygenic photosynthesis at +8 h was a 2-fold increase in photosynthetic efficiency ($\alpha$), which returned to the pre-enrichment level by +32 h. After the culture reached stationary phase.

Fig. 3. Variation in $^{15}$NO$_3^-$ uptake capacity and in vivo NRA of Skeletonema costatum after shift to improved growth conditions. [a.] Rates normalized to PON content. [b.] Rates normalized to cell number.

Table 2. Comparison of mean PON-specific rates (±1 SD) of NO$_3^-$ depletion ($V$NO$_3$), NO$_3^-$-saturated uptake of $^{15}$NO$_3^-$ ($V^{15}$NO$_3^-$), and NRA ($VNRA$) in Skeletonema costatum between 0 and +24 h of the simulated upwelling (based on data in Figs. 2, 3a).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Rate ($h^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V$NO$_3$</td>
<td>0.046±0.011</td>
</tr>
<tr>
<td>$V^{15}$NO$_3^-$</td>
<td>0.019±0.003</td>
</tr>
<tr>
<td>VNRA</td>
<td>0.010±0.007</td>
</tr>
</tbody>
</table>

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Table 3. Variation in photosynthetic efficiency ($\alpha$), light-saturated photosynthetic rates of gross oxygen evolution ($P_{\text{max}}$, O$_2$), dark respiration ($R_O$), C assimilation ($P_{\text{max}}$,$^{14}$C), photosynthetic quotients (O$_2$ : CO$_2$), and ratios of oxygenic photosynthesis to respiration ($P$ : $R$) during the light and NO$_3$ shift-up of Skeletonema costatum. Daily oxygenic $P$ : $R$ ratios were calculated based on 14 h of light-saturated net photosynthesis and 24 h of dark respiration: \[
\left[\frac{P_{\text{max}} - R_O}{R_O}\right] \times \frac{14}{24}.
\]

<table>
<thead>
<tr>
<th>Time (h)</th>
<th>$n^*$</th>
<th>$P_{\text{max}}$O$_2$</th>
<th>$R_O$</th>
<th>$P_{\text{max}}$,$^{14}$C</th>
<th>O$_2$ : CO$_2$</th>
<th>$P$ : $R$</th>
<th>Daily $P$ : $R$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>3.6(1.0)</td>
<td>331(28)</td>
<td>20(5)</td>
<td>241(18)</td>
<td>1.37</td>
<td>16.4</td>
<td>9.0</td>
</tr>
<tr>
<td>8</td>
<td>7.8(1.6)</td>
<td>408(31)</td>
<td>93(25)</td>
<td>137(7)</td>
<td>2.98</td>
<td>4.4</td>
<td>2.0</td>
</tr>
<tr>
<td>32</td>
<td>3.4(0.2)</td>
<td>283(6)</td>
<td>26(5)</td>
<td>155(7)</td>
<td>1.83</td>
<td>11.0</td>
<td>5.8</td>
</tr>
<tr>
<td>48</td>
<td>4.3(0.4)</td>
<td>278(7)</td>
<td>28(6)</td>
<td>213(5)</td>
<td>1.31</td>
<td>9.8</td>
<td>5.2</td>
</tr>
</tbody>
</table>

* $\mu$mol O$_2$ (mg Chl a)$^{-1} h^{-1}$ s$^{-1}$ (±1 SD).
† $\mu$mol O$_2$ (mg Chl a)$^{-1} h^{-1}$ (±1 SD).
‡ $\mu$mol CO$_2$ (mg Chl a)$^{-1} h^{-1}$ (±1 SD).

(ca. +28 h), oxygenic photosynthesis stabilized, but rates of C assimilation increased. These shifts resulted in a more than doubling of the photosynthetic quotient (PQ = O$_2$ : CO$_2$) between 0 and +8 h, followed by a return to the initial PQ value after the culture reached stationary phase (Fig. 4; Table 3). Calculation of daily $P$ : $R$ ratios indicates that, under conditions of high rates of NO$_3^-$ assimilation (+8 h), daily photosynthetic C gain would only slightly exceed C demand (Table 3).

The ratio of C- and N-specific C : N uptake capacity was 7.7 at the onset of the simulated upwelling treatment but dropped below 5 for the remainder of the experiment (Fig. 4). If we assume that the Redfield value of 6.6 represents balanced uptake capacity, the changes observed here indicate a large increase in the capacity to take up N relative to C immediately after the nutrient pulse. These findings are consistent with the changes in photosynthetic and respiratory energetics and are reflected in the variation in the PQ and $P$ : $R$ ratios described above.

Molecular processes: Macromolecular pools—Cellular content of Chl a, DNA, RNA, and protein in S. costatum declined during the light-limited period (-40 to 0 h; Fig. 5a,b). The shift to improved growth conditions resulted in sequential accumulation of nucleic acids and proteins that preceded cell division. RNA content increased 3.5-fold in the first 12 h after enrichment. This accumulation of RNA, indicative of increased transcription, preceded net protein synthesis by 4 h and DNA synthesis by 8 h. Chl a content increased gradually (~10%) and then decreased by 28% after cell division, most likely reflecting the onset of N depletion. Cellular protein content was 66% higher after cell division whereas total nucleic acid content (RNA + DNA) decreased to pretreatment levels. The RNA content, relative to cellular DNA, increased 1 h after enrichment and declined exponentially after cell division to the pre-enrichment ratio (Fig. 5c), providing further evidence for transcriptional activation by light and NO$_3^-$ enrichment.

Characterization of changes in the RNA transcript population: Translational capacity—The abundance of translatable RNA increased by 69% between 0 and +8 h and was associated with the initial doubling in total RNA content (Fig. 6a). Although total
RNA content continued to increase until the onset of cell division at +20 h, the abundance of translatable RNA declined, indicating that the continued increase in total RNA was associated with the accumulation of ribosomal and other nonmessenger RNAs. A second peak in translatable RNA occurred at +28 h, after the completion of cell division. Translational capacity declined subsequently as the cells entered stationary phase.

Analysis of the in vitro translation products by SDS-PAGE revealed that the initial response to improved growth conditions was associated with the accumulation of transcripts encoding a suite of polypeptides ranging from 20 to 40 kD (kilodaltons) between 0 and +8 h (Fig. 6b). A similar set of polypeptides was synthesized at the time of the second peak (24 h) in translatable RNA content. No measurable synthesis of polypeptides in the 100-120-kD mol wt size class expected for NR (Solomonson and Barber 1990; Y. Gao and G. J. Smith unpubl. data) was observed that corresponded to the variation in NRA however, probably attributable to the low abundance of NR relative to total cellular protein (see below).

Hybridization analysis of the transcript population—Northern analysis of total cellular RNA from S. costatum revealed that the 18S and pSCNR21 probes hybridized to discrete bands with relative lengths of 1.8 kb (kilobase) for the small subunit ribosomal transcript and 5.0 kb for the NR transcript, respectively (Fig. 7). The specificity of these probes made them amenable for quantifying temporal changes in specific transcript abundance by dot-blot analysis.

Abundance of 18S rRNA declined by a factor of 2 under light limitation (−40 to +1 h), then increased by a factor of 2.5 at +4 h (Fig. 8a,c,d). The relative abundance of 18S rRNA increased an additional 3.5-fold between +12 h and +32 h, after the completion of cell division. The two peaks in 18S rRNA abundance were associated with the periods of maximal translational capacity (see Fig. 6a). The first maximum in 18S rRNA abundance preceded acceleration in the rate of cellular protein accumulation and the initial increase in translational capacity (see Figs. 5b, 6a). The second increase in 18S rRNA abundance was greater than the first peak (Fig. 8c) and paralleled the increase in translational capacity between +12 and +24 h (Fig. 6a), although 18S rRNA abundance peaked at +32 h, 8 h after the peak in translational capacity.

The exposure slopes of the pSCNR21 probed dot blot (Fig. 8b) were 10-100-fold lower than those for the 18S probe (Fig. 8a), implying a lower abundance for the transiently expressed NR transcript. The relative abundance of the pSCNR21-positive transcript declined 80% during the light-limited period (−40 to 0 h, Fig. 8d). After the shift to improved growth conditions, NR transcript abundance increased 380% between 0 and +4 h, then declined 50% during the dark period through +16 h (Fig. 8c,d). The abundance of the NR transcript in-
increased gradually during the second light period, reaching a maximum at +32 h, coincident with the second peak in 18S transcript abundance. The abundance of the NR transcript declined over the next dark period to pre-enrichment levels at +48 h, as the culture was in stationary phase in the NO$_3^-$-depleted medium.

Regulation of NRA—The NR protein was detectable in profiles of S. costatum protein only by immunological methods. Western blots of total proteins probed with a het-
Erologous polyclonal NR antiserum revealed an immunoreactive polypeptide in the molecular size range (105-110 kD) observed for higher plant (Solomonson and Barber 1990) and purified S. costatum NR (Y. Gao and G. J. Smith unpubl. data; Fig. 9a). Protein-specific abundance of this polypeptide declined by a factor of 3 during the light-limited phase (-40 to 0 h), increased by a factor of 1.5 in the first hour after enrichment, then declined by a factor of 2 at +8 h (Fig. 9b). Abundance of the NR polypeptide increased by a factor of 3.3 during the dark period prior to the onset of cell division (+8 to +12 h), then declined at the beginning of the second light period after enrichment (+20 to +28 h). NR protein level increased again from +28 to +48 h after enrichment.

Comparison of variation in NRA with NR polypeptide and transcript abundance provides insight into the levels of regulation controlling NRA during transition to improved growth conditions. The cellular content of NR protein and transcripts declined by 80% during the light-limited period, then increased by 30% in the first hour after the light and NO₃⁻ enrichment (Fig. 10). The dramatic increase in NR transcript abundance between 0 and +4 h preceded a similar increase in NR protein through the first half of the dark period between +8 and +16 h. This pattern was observed again between +22 and +48 h after enrichment, but increases were of lower magnitude. Although accumulation of NR transcripts during the light period was correlated with peaks in NRA, the lack of a concomitant increase in NR protein suggests that the diurnal rhythm in NRA resulted from the light activation of existing NR protein.

When expressed as relative levels per cell (Fig. 10), in vivo NRA increased by 280% between 0 and +4 h, although no similar increase in the cellular content of NR protein was observed. The subsequent decline in NRA (37%) between +4 and +8 h was associated with a decline (20%) in NR protein abundance. The low NRA observed during the first dark period, however, was associated with a dramatic increase in NR protein abundance. Cellular NR protein content declined following cell division, in contrast to the pattern for NRA. The increase in NR protein abundance between +32 and +48 h was associated with a loss of NRA. Since changes in NRA and the NR protein pool appear to be uncoupled temporally, diurnal NRA rhythms probably reflect light-induced, posttranslational modulation of existing NR enzyme.

Discussion

Most information on the physiological adaptation of marine phytoplankton comes from steady state cultures. Although these static observations have generated a precise quantitative understanding of balanced growth responses to different environmental conditions (e.g. Bannister 1979; Shuter 1979; Kiefer and Mitchell 1983), the resulting theory explains little of the fine-scale temporal variability observed in the ocean (Cullen 1990). At least part of the problem may stem from the inability of steady state
Growth of *S. costatum* 1001

![Graph showing the growth of *S. costatum*](image)

**Time (h)**

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**Fig. 9.** Analysis of changes in abundance of NR protein in *Skeletonema costatum* cells during simulated upwelling. [a.] Western blot of total cellular protein profiles probed with antisera to squash NR and visualized with alkaline phosphatase-coupled secondary antibodies. Arrow indicates location of the polypeptide band cross-reacting with NR antisera. Equal amounts of protein (50 μg) from sequential samples relative to the start of the enrichment were separated on a 7.5% SDS-PAGE and transferred to a nitrocellulose membrane. The position and size of mol-wt markers (kD) separated on the same gel are indicated on the left. [b.] Temporal variation in cellular protein content (■) (as in Fig. 6b) and the relative abundance of the NR cross-reactive protein (○) determined by densitometry of the alkaline phosphatase-positive bands shown in the top panel. Optical densities of each band were baseline corrected to the density background in each lane.

**Fig. 8.** Characterization of the abundance patterns of the small subunit rRNA (18S) and NR transcripts in total RNA isolated from *Skeletonema costatum* during simulated upwelling. [a, b.] Autoradiographs of dot blots of total RNA isolated from *S. costatum* during the simulated upwelling. Serial dilutions of glyoxal denatured RNA (4 μg to 63 ng, rows) from each time point (columns) were applied to Nytran membranes and hybridized with the 18S rRNA probe (panel a) or the NR probe pSCNR21 (panel b). [c.] Quantification of changes in the abundance of 18S (●) and pSCNR21 (○) hybridizing sequences. Autoradiographs of hybridized dot blots were scanned with a densitometer and the change in exposure intensity (O.D.) with the weight of RNA per dot was determined by linear regression and expressed as the exposure slope. Error bars represent standard errors of the regression slopes. [d.] Changes in the abundance of the 18S rRNA (●) and pSCNR21 (○) transcripts relative to levels observed at the start of the light and NO₃⁻ enrichment (time = +1 h).
Physiological responses to simulated upwelling—The transient responses of this unialgal culture to improved growth conditions qualitatively mimicked the temporal changes in NO\textsubscript{3}\textsuperscript{-} uptake, C uptake, and biochemical composition observed in natural phytoplankton communities from coastal upwelling systems (Barlow 1982; MacIsaac et al. 1985; Dugdale and Wilkerson 1989), although events occurred on a compressed time scale due to the high initial biomass (2.3 \times 10^6 cells ml\textsuperscript{-1}; 700 \mu g Chl a liter\textsuperscript{-1}) and low relative N enrichment (DIN : PON = 0.43) used here. In contrast, freshly upwelled seawater is generally characterized by DIN : PON ratios > 1 and Chl a levels < 1 \mu g Chl a liter\textsuperscript{-1} (Wilkerson and Dugdale 1987; Dugdale and Wilkerson 1989). Prior to the light and NO\textsubscript{3}\textsuperscript{-} enrichment, the cells were N deficient (C : N = 9.9) with respect to the Redfield ratio, and indices of NO\textsubscript{3}\textsuperscript{-} assimilation capacity (V\textsuperscript{15}NO\textsubscript{3}, VNRA) were low. The rates of induction or acceleration of V\textsuperscript{15}NO\textsubscript{3} and VNRA following enrichment were comparable to observations from natural upwelling systems exhibiting lower absolute but higher relative enrichment levels (Wilkerson and Dugdale 1987). Increases in cell-specific 15NO\textsubscript{3} uptake and NRA were confined to the first light period when NO\textsubscript{3}\textsuperscript{-} was present in the media, indicating that NO\textsubscript{3}\textsuperscript{-} plays a direct role in metabolic induction. These observations also suggest that species-specific partitioning of NO\textsubscript{3}\textsuperscript{-} (e.g. storage vs. growth; see Collos 1986) may influence the temporal pattern of induction of NO\textsubscript{3}\textsuperscript{-} metabolism in natural systems.

The effect of improved growth conditions on photosynthetic performance was also dramatic. While rates of CO\textsubscript{2} fixation declined initially, rates of oxygenic photosynthesis increased, suggesting that the increased reducing power generated through enhanced photosynthetic electron transport was being used to drive the assimilation of NO\textsubscript{3}\textsuperscript{-} at the expense of CO\textsubscript{2} fixation. The increase in photosynthetic efficiency (\alpha) at +8 h also indicates a redirection of ab-
sorbed light energy toward \( \text{NO}_3^- \) reduction. The increase in \( \text{NO}_3^- \) assimilation capacity was also associated with a 3-fold increase in dark respiration, which could increase pools of reductant and (or) C skeletons for \( \text{NO}_3^- \) assimilation as has been hypothesized for green algae (Weger and Turpin 1989). Enhanced respiration during this period was also associated with an overall increase in energy-dependent anabolic processes such as nucleic acid and protein synthesis. Thus, although the relative \( \text{NO}_3^- \) enrichment used here was not sufficient to reverse the N-deficient status of the cells (based on C : N ratio), it did induce large and temporally significant transients in rates of photosynthesis and \( \text{NO}_3^- \) assimilation that can affect the biological dynamics of natural upwelling systems.

**Regulation of NRA**—The rate of induction of \( \text{NO}_3^- \) utilization capacity may control the rate of phytoplankton growth in N-limited environments characterized by a pulsed \( \text{NO}_3^- \) supply. The molecular aspects of \( \text{NO}_3^- \) assimilation, particularly the potentially rate-limiting activity of NR, have been well characterized in higher plants, but this information is lacking for phytoplankton (Campbell 1989; Solomonson and Barber 1990). NR expression is known to be influenced by light availability and source of N at the levels of gene transcription, RNA translation, and posttranslational modulation of enzyme activity. This study demonstrated that environmentally induced changes in NR transcript and protein abundance may underlie the adaptive response of phytoplankton to shifts in environmental conditions.

Rapid responses of phytoplankton cells to improved growth conditions may entail activation of existing enzyme pools as well as increased transcription and translation. Changes in the NRA of *S. costatum* cells observed here reflected both increased synthesis of enzyme and activity modulation of existing NR protein. The rapid increase in NR transcript abundance after the environmental shift also provides clear evidence that light and \( \text{NO}_3^- \) availability can induce NR gene expression in diatoms. Dependence of NR transcription on light and availability of N is further suggested by the declines in NR transcript abundance during \( \text{NH}_4^+ \)-enriched and light-limited culture (−60 to 0 h) and during darkness (+12 to +22 h) since increases in NR transcript levels were restricted to periods of photosynthetically saturating PPF (see Fig. 10).

NR transcription and translation appeared to be coupled during the light-limited phase (−40 to 0 h) as NR transcript and protein abundance declined in parallel. These processes became uncoupled, however, after the shift to improved growth conditions, where the rapid increase in NR transcript and activity levels were not immediately associated with a concomitant increase in NR protein levels. In fact, at the cellular level, there was a general lag between the increase in transcript abundance and protein synthesis, presumably reflecting the time required to accumulate biochemical components required for protein synthesis.

NR protein may also be synthesized constitutively during the cell cycle of *S. costatum*, since the doubling in abundance of enzyme at +12 h after enrichment and subsequent reduction to pre-enrichment levels by +24 h was associated with the period of cell division. Accumulation of protein during the dark period is consistent with observations of oceanic phytoplankton in which sustained protein synthesis at night was attributed to the reapportionment of photosynthate (Cuhel et al. 1984).

In vivo NRA exhibited a pronounced diurnal rhythm that did not correspond to changes in NR protein levels, indicating that posttranslational modulation of enzyme activity may be light-dependent. Although in vitro NRA assays have been shown to track NR protein abundance in higher plant leaves and a green alga (Solomonson and Barber 1990; Velasco et al. 1989), our data demonstrate that diel changes in in vivo NRA reflect physiological levels of regulation independent of changes in enzyme abundance in *S. costatum*. The uncoupling of NRA and enzyme abundance, as well as the decline in NRA after \( \text{NO}_3^- \) depletion, demonstrates that both illumination and \( \text{NO}_3^- \) availability directly influence in vivo NRA. In addition, diurnal rhythms in NRA have been observed in marine phytoplankton com-
munities and macrophytic algae with both in vitro and in vivo assays (Martinez et al. 1987; Gao et al. 1992). Thus, caution must be taken to ensure that conclusions drawn regarding environmentally regulated induction of NRA in marine algae are not biased by underlying rhythms.

Under the experimental conditions used here, an absence of diurnal cycles in NO$_3^-$ uptake and photosynthetic capacity of $S$. costatum cells indicates that diurnal modulation of NRA occurs independently of other changes in the physiological status of the cell and most likely reflects a direct role of light in the activation of existing NR protein. Accumulation of NR protein during the dark period after enrichment does suggest, however, that some increase in NR protein synthesis and (or) its stability resulted from the experimental treatment. Although the qualitative pattern is clear from this investigation, the quantitative effects of light and NO$_3^-$ availability on induction of NO$_3^-$ metabolism remain to be determined.

Biochemical markers of responses to transient conditions—A suite of biochemical parameters has been proposed to indicate nitrogen status in phytoplankton, providing information on prior sources of N and an index of growth potential for phytoplankton populations (Dortch and Postel 1989; Flynn 1990). If they are to serve as useful indices of cellular features, biochemical parameters should respond on time scales corresponding to the temporal patterns of environmental change. For example, RNA and protein content have been suggested as suitable markers for N status (Dortch et al. 1983; Dortch and Postel 1989) because, together, they account for upward of 80% of cellular N content (Wheeler 1983). Further, variation in the biochemical marker should be expressed on a cell-specific basis or under field conditions, normalized to a measure of biomass that varies directly with cell number.

In the present study, a very pronounced transient in cellular RNA content was observed that preceded changes in Chl $a$, protein, and DNA levels associated with the phased cell division (see Fig. 5), thus meeting the requirements outlined above. RNA : DNA ratios also provide a clear signal of rapid response to the improved growth conditions in the present study. This ratio returned to baseline levels after a round of cell division and in response to N depletion. Interestingly, the baseline RNA : DNA ratio (wt basis) was similar for both the light- and N-limited phases of the treatment, supporting previous observations that this ratio is a useful marker for transient improvements in phytoplankton growth conditions and metabolic status (Dortch et al. 1983).

Application of standard molecular biology techniques provided sensitive markers of the molecular mechanisms associated with the response of phytoplankton cells to variable environments. Underlying the transient increase in the RNA content of $S$. costatum after enrichment was a rhythmic variation in the translational capacity that was synchronized with the light phase of the photoperiod. Previous studies of green algae and higher plants have documented light-dependent transcription of some nuclear- and chloroplast-encoded genes (Puiseux-Dao 1981; Tobin and Silverthorne 1985). In this study the initial peak in translational capacity was associated with the period of maximal protein accumulation. The second peak in translational capacity, however, occurred after the onset of NO$_3^-$ depletion from the medium and was not associated with any net accumulation of protein, suggesting that NO$_3^-$ depletion inhibited some components of the translational machinery (e.g. amino acid availability). Sensitive measures of the uncoupling between the availability of mRNAs (translational capacity) and realized rates of protein synthesis may elucidate key metabolic steps that dictate lags in phytoplankton growth in response to environmental shifts.

The dynamics of RNA transcript populations can be dissected further through hybridization analyses with specific gene probes. In the present study, the small-subunit rRNA and NR transcripts exhibited different behaviors. Levels of both transcripts increased in the initial response to improved light and NO$_3^-$ conditions for growth and again during the second photoperiod, after NO$_3^-$ had been depleted from the medium, suggesting that transcript accumulation may respond to internal rather
Growth of \textit{S. costatum} 1005

than external levels of N, as has been observed previously for NRA in \textit{S. costatum} (Dortch 1982). The abundance of 18S rRNA tracked the translational capacity of the total RNA more closely than did the abundance of NR transcripts. The dynamics of 18S rRNA, with its primary role in protein synthesis, in addition to ease of detection due to its high abundance in the cell, suggest that 18S rRNA abundance may be a useful index of the growth potential of phytoplankton cells, as has been demonstrated for bacteria (Delong et al. 1989).

The dynamics of specific RNA transcripts can also be sensitive indicators of changes in the metabolic status of phytoplankton cells and may serve as reliable markers of shifts in specific metabolic pathways in response to environmental transients. This approach has real promise for oceanographic applications since bulk measures of RNA and DNA content can be performed routinely on oceanic water samples (Dortch et al. 1983), and nucleic acids of suitable quality for molecular analysis can be obtained with only minor modifications of existing protocols (Giovannoni et al. 1990; G. J. Smith unpubl. data). Northern analysis with taxon-specific gene probes can be used to track functional markers of individual taxa among heterogeneous mixtures of RNAs obtained from natural oceanic samples, as used to examine bacterial diversity in natural systems (Giovannoni et al. 1990).

Environmental regulation of critical metabolic processes, such as NO$_3^-$ assimilation, can be assessed at the molecular level by simultaneously monitoring the abundance of specific transcripts, proteins, and enzyme activities in oceanic samples as reported here for laboratory-grown phytoplankton. Application of these techniques in oceanic environments in conjunction with measurements of water-column biological and physical characteristics are essential for development of a mechanistic understanding of the influence of metabolic lags on temporal and spatial dynamics of phytoplankton bloom formation. Such knowledge is critical to the development of quantitative models that can accommodate the nonlinearity of biological phenomena and yield new insights into oceanic processes.

This laboratory study, based on the concurrent monitoring of whole-cell physiological and subcellular molecular events associated with the transition of light-limited and N-deficient \textit{S. costatum} cells to light and NO$_3^-$ enrichment, has demonstrated that the cellular changes observed here are qualitatively similar to the responses of natural phytoplankton assemblages to improved growth conditions characteristic of coastal upwelling systems. Furthermore, the physiological and molecular responses described here can provide a partial, mechanistic explanation for performance shifts observed in field populations. Cellular lags in responses to environmental transients can be related to an uncoupling of C and N assimilation and the feedback of these physiological features on the molecular processes that govern their expression. It is anticipated that further investigations into the cellular and molecular processes that govern N and C assimilation will not only improve understanding of growth dynamics of marine phytoplankton but advance the ability to predict temporal and spatial patterns of oceanic primary production.

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Growth of S. costatum


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