RESEARCH PAPER

Whole-cell response to nitrogen deprivation in the diatom
Phaeodactylum tricornutum

Leila Alipanah, Jens Rohloff, Per Winge, Atle M. Bones and Tore Brembu

Department of Biology, Norwegian University of Science and Technology, N-7491 Trondheim, Norway

* To whom correspondence should be addressed. Email: tore.brembu@ntnu.no

Received 1 December 2014; Revised 8 June 2015; Accepted 15 June 2015

Editor: Christine Raines

Abstract

Algal growth is strongly affected by nitrogen (N) availability. Diatoms, an ecologically important group of unicellular algae, have evolved several acclimation mechanisms to cope with N deprivation. In this study, we integrated physiological data with transcriptional and metabolite data to reveal molecular and metabolic modifications in N-deprived conditions in the marine diatom Phaeodactylum tricornutum. Physiological and metabolite measurements indicated that the photosynthetic capacity and chlorophyll content of the cells decreased, while neutral lipids increased in N-deprived cultures. Global gene expression analysis showed that P. tricornutum responded to N deprivation through an increase in N transport, assimilation, and utilization of organic N resources. Following N deprivation, reduced biosynthesis and increased recycling of N compounds like amino acids, proteins, and nucleic acids was observed at the transcript level. The majority of the genes associated with photosynthesis and chlorophyll biosynthesis were also repressed. Carbon metabolism was restructured through downregulation of the Calvin cycle and chrysolaminarin biosynthesis, and co-ordinated upregulation of glycolysis, the tricarboxylic acid cycle, and pyruvate metabolism, leading to funnelling of carbon sources to lipid metabolism. Finally, reallocation of membrane lipids and induction of de novo triacylglycerol biosynthesis directed cells to accumulation of neutral lipids.

Key words: Carbon metabolism, diatom, metabolomics, nitrogen deprivation, Phaeodactylum tricornutum, transcriptome, triacylglycerol.

Introduction

Phytoplankton blooms vary temporally and spatially in accordance with nutrient availability (Brandes et al., 2007; Falkowski and Raven, 2007). Under upwelling conditions, high levels of available nitrate and iron lead to an increase in phytoplankton biomass, which is generally dominated by diatoms (Kudela and Dugdale, 2000; Capone and Hutchins, 2013). Inorganic nitrogen (N) in the form of ammonia or nitrate is utilized by several phytoplankton (Dham et al., 2005); some phytoplankton are also able to use organic forms of nitrogen such as amino acids, nucleic acids, and urea (Baker et al., 2009; Solomon et al., 2010).

Diatoms are a group of unicellular heterokont microalgae believed to include some 200 000 species (Armbrust, 2009). It is estimated that marine diatoms are responsible for about 32% of global phytoplankton primary production (Uitz et al., 2010). Unlike plants and green algae, diatoms and
other heterokonts originate from a serial secondary endosymbiosis event, in which a green alga and subsequently a red alga were engulfed by a heterotrophic eukaryote (Moustafa et al., 2009; Bowler et al., 2010). In addition, a large number of horizontal gene transfer events have further increased the gene repertoire. As a result, diatom genomes contain unique combinations of nutrient assimilation and metabolic pathways that have contributed to their ecological success in the ocean (Prihoda et al., 2012).

The whole-genome sequences of the centric diatom Thalassiosira pseudonana (Armbrust et al., 2004) and the pennate diatom Phaeodactylum tricornutum (Bowler et al., 2008) have provided valuable information on the regulatory and metabolic inventory of these diatoms. The genomes of T. pseudonana and P. tricornutum contain several transporter proteins for uptake of inorganic and organic nitrogen (Rees and Syrett, 1979; Armbrust et al., 2004; Allen, 2005; Hildebrand, 2005; Solomon et al., 2010). Nitrate entering the cell is first reduced to nitrite and ammonium (Allen et al., 2005; Bowler et al., 2010). Ammonium is then assimilated by glutamate synthase/glutamine synthetase to amino acids (Zehr and Falkowski, 1988; Takabayashi et al., 2005). Diatoms possess plastidal glutamine synthetase (GSII) and glutamate synthase (Fd-GOGAT) as well as mitochondrial NAD(P)H-GOGAT and GSIII (Bowler et al., 2010; Allen et al., 2011). Mitochondrial GSIII may catalyse the assimilation of glutamine from ammonium derived from cytosolic catabolic reactions, e.g. deamination and hydrolysis of organic N (Hockin et al., 2012; Kissen et al., 2010; Parker and Armbrust, 2005).

N and carbon metabolism are closely connected to each other. N assimilation and amino acid biosynthesis require reducing equivalents from photosynthesis and carbon skeletons from the tricarboxylic acid (TCA) cycle (Hockin et al., 2012). Moreover, in the photosynthetic apparatus, assimilated N is used for example in ribulose-1,5-bisphosphate carboxylase (Rubisco) and the light-harvesting complex (LHC) (Orellana and Perry, 1995; Foyer et al., 2003; Nunes-Nesi et al., 2010). In response to N deprivation, diatoms reprogram several metabolic pathways. The impact of N deprivation on pigments, photosynthesis, carbon fixation, and N assimilation has been studied in diatoms (Syrett et al., 1986; Kolber et al., 1988; Geider et al., 1993; Granum et al., 2009; Bender et al., 2014). Diatoms store carbon in the form of 1,3-β-D-glucan (chrysosominarin) or lipids (Kroth et al., 2008). Under optimal conditions, chrysosominarin is the major sink of carbon storage in the vacuoles (Granum and Myklestad, 2002). Under several stress conditions, in particular N starvation, diatoms change their carbon storage patterns in favour of neutral lipid accumulation (Eizadora et al., 2009; Norici et al., 2011; Sharma et al., 2012; Valenzuela et al., 2012). Neutral lipids produced from microalgae have been proposed as a sustainable substitute biofuel for fossil fuels (Wijffels and Barbosa, 2010). Other N-containing compounds, such as proteins and nucleic acids, are also affected by a decrease in cellular N content (Olson et al., 1986; Mock and Kroon, 2002; Bertozzini et al., 2013; Mus et al., 2013).

To understand how the oleaginous marine diatom P. tricornutum responds to N deprivation, cells were grown in f/2 medium and in N-free medium, and samplings were conducted at 48 and 72 h after N deprivation. We combined transcriptional and metabolite analyses to monitor the effect of N deprivation at different molecular levels in order to get a better insight into the acclimation strategies employed by P. tricornutum under N deprivation. These data were further complemented by physiological data such as measurements of cell growth, neutral lipids, and other cell chemistry measurements. We use this data to predict metabolic changes in N-deprived cells leading to remodelling of lipid metabolism and triacylglycerol (TAG) accumulation.

Materials and methods

Growth conditions and treatments

Axenic cultures of P. tricornutum clone Pt1 8.6 (CCMP632) were grown in f/2 medium and kept in exponential growth at 15 °C under continuous white fluorescent light (60 μmol photons m⁻² s⁻¹) for 3 weeks. Bacterial contamination was checked regularly by inoculation in peptone-enriched f/2 medium (Andersen et al., 1997). Growth medium (f/2) was made from 0.2 μm-filtered seawater, autoclaved, and enriched with macro- and micronutrients (Guillard, 1975). Three or four replicates of the start culture (6–7 ml) were transferred to 220 ml of medium supplemented with complete f/2 nutrients (replete) or f/2 without added nitrate (deprived). The nitrate concentration in the seawater used for the experiments was measured to 10 μM, which is 1.1% of the f/2 nitrate concentration. Cells were incubated in batch cultures with a starting cell density of 5 × 10⁶ cells ml⁻¹ in sterile culture flasks with a 75 cm² growth area. Cell counting and maximum quantum yield of photosystem II (PSII) (Fm/Fm′) was measured daily using a Bürker–Türk counting chamber and AquaPen-C AP-C 100 (Photon Systems Instruments), respectively. For the other experiments, samples were harvested 48 and 72 h after the beginning of the treatment. Samples for RNA and metabolite analysis were stored at −80 °C, while samples for nutrient and pigment analysis were stored at −23 °C until analysis.

Nutrient analysis

Triplicate cultures for particulate N, carbon, and phosphorus analysis were collected on pre-combusted GF/F filters (particulate C and N analysis) or 0.2 μm GF/F filters (particulate phosphorus), and the flowthrough was used for detection of medium phosphate and nitrate concentration. Triplicate samples for particulate N and carbon analysis along with blank filters were treated with HCl vapour (37%), packed in tin capsules, dried for 2 days at 60 °C, and analysed by an ECS 4010 element analyser (Costech Instruments). All these processes were performed according to Chauton et al. (2013). Inorganic nutrients were measured in the filtrate. NO₃⁻ + NO₂⁻ and PO₄³⁻ were analysed in parallel according to I.O. Analytical cartridge Part A002603 and A002604, respectively, as described by Hansen and Koroleff (1999). Particulate phosphate was first oxidized to PO₄³⁻ according to Norwegian standard NS4725, and then analysed as inorganic PO₄³⁻.

Pigment analysis

Pigment analysis (fg per cell) was performed based on the protocol by Rodríguez et al. (2006). Briefly, 60 ml (N replete) or 100 ml (N deprived) of cultures was collected on GF/F filters. The cells were extracted with 6 ml of 100% ethanol, and extracts were filtered
through Millipore 0.45 μm filters. A volume of 73 μl of the final extracts was mixed with 23 μl of water and injected into a Hewlett-Packard HPLC 1100 Series system. Pigments were separated on a Waters Symmetry C8 column using the high-performance liquid chromatography (HPLC) method of Zapata et al. (2000). Chl $a$ and fucoxanthin were detected by absorbance at 440 nm and identified by a diode array detector ($\lambda$=350–370 nm, 1.2 nm spectral resolution). Standard curves were made by isolating pigments separated by HPLC, verifying their identity and quantifying on a spectrophotometer, and running a dilution series on the HPLC instrument. The specific extinction coefficients (ε: 1 g$^{-1}$ cm$^{-1}$) provided by Egeland et al. (2011) were used for pigment quantification.

Neutral lipid measurement

A volume of 1 ml of culture was stained with 1 μl of 0.1 μg ml$^{-1}$ of BODIPY 505/515 (4,4-difluoro-1,3,5,7-tetramethyl-4-bora-3a,4a-diaza-s-indacene; Life Technologies) dissolved in 2% (w/v) dimethyl sulfoxide and shaken carefully by hand (Govender et al., 2012). After 5 min, 30 μl of culture was transferred to a microscope slide, and a coverslip was placed on top of the culture and sealed using dental wax. At least 20 cells from two replicates were analysed for BODIPY 505/515 fluorescence on a Leica TCS SP5 confocal laser scanning microscope using a ×63 water objective. Z-Sectional images were made using argon laser excitation at 488 nm (17% of maximal intensity), and emission was detected with a spectral detector set from 495 to 550 nm. Non-confocal bright-field images were made simultaneously. A z-stack consisting of 10 scans was made for each cell, encompassing the complete fluorescent part of the cell. The length of the z-stack varied between 4.00 and 5.78 μm; consequently, the z-slice step size varied between 0.44 and 0.64 μm. Laser power, PMT gain, and offset were kept constant for all scans. Image stacks containing the fluorescence channel were imported into ImageJ (Abramoff et al., 2004). To determine the total fluorescence detected in the z-stack, a region was drawn around each cell to be measured, and three regions next to the selected cell that had no fluorescence were used for background subtraction. The corrected total cell fluorescence for each cell was calculated using the following formula (Gavet and Pines, 2010; Potapova et al., 2011):

$$\text{Whole-cell signal corrected} = \text{whole-cell signal} - \left( \frac{\text{area of selected cell} \times \text{mean fluorescence of background}}{\text{area of selected cell}} \right)$$

Background fluorescence, as measured from five z-sectional images of unstained cells, was negligible (<1% of stained N-replete cells). A total of 20–30 cells were analysed for each treatment.

RNA isolation

Depending on cell density, 60–100 ml of cultures was collected on 0.65 μm Durapore membrane filters, washed off the filter using 1 ml of f/2 medium (N-deprived cells were washed with f/2 without nitrate supplement), and centrifuged at 13 000 rpm for 1 min at 4 °C. The supernatant was removed and pellets were flash frozen in liquid N$_2$ and stored at −80 °C. Frozen samples were homogenized using a TissueLyser system (Qiagen) for 2 × 2 min at 25 Hz. The samples were placed in a pre-cooled (~80 °C) adapter set for the first shaking step. Before the second shaking step, the samples were transferred to a room temperature adapter set, and 0.5 ml of lysis buffer (SpectrumTM Plant Total RNA kit; Sigma-Aldrich) was added to each tube. Total RNA was isolated with a SpectrumTM Plant Total RNA kit (Sigma-Aldrich). To eliminate genomic DNA, an on-column digestion was performed using an RNase-free DNase I set (Qiagen). Total RNA was quantified using a NanoDrop ND-1000 Spectrophotometer (NanoDrop Technologies). The RNA quality was verified using formaldehyde gel electrophoresis. In addition, RNA integrity was checked on a 2100 Bioanalyzer (Agilent). All samples had RNA integrity numbers above 7.

cDNA microarray experiments

Total RNA (200 ng) was reverse transcribed, amplified, and labelled according to a Low Input Quick Amp Labeling Kit, One-Color (Agilent Technologies). A total of 1650 ng of cRNA from each sample was fragmented and hybridized with a Gene Expression Hybridization Kit (Agilent Technologies) on 4 × 44K
**P. tricornutum** whole-genome 60-mer oligonucleotide microarrays (Agilent Technologies) in an Agilent G2545A Hybridization Rotary Oven at 10 rpm, 65 °C for 17.5 h. Slides were washed with washing buffer 1 and 2 using a Gene Expression Wash Buffer Kit (Agilent Technologies) and directly scanned using a laser scanner (G2505 B; Agilent Technologies) based on the ‘dynamic range exponent’ option in the scanner software. Images were processed by Agilent Feature Extraction software version 9.5.

Statistical analysis

The Limma package (version 3.20.1) (Smyth, 2005) and R version 3.0.3 were used for statistical analysis and identification of significantly differentially expressed genes. Single-colour feature expression files from the Agilent microarray scans were imported, and spots identified as feature outliers were excluded from the analysis. Weak or undetected spots were given reduced weight. The data were normalized using the quantile method, and no background subtraction was performed. A design matrix was created and pair-wise comparisons between the samples, DN48 (nitrogen-deprived 48 h) and R48 (replete 48 h) and DN72 (nitrogen-deprived 72 h) and R72 (replete 72 h) were performed. The method of Benjamini and Hochberg (1995) was used to estimate the false discovery rate. Genes with an adjusted P value of <0.05 were regarded as significantly differentially expressed and were included in the analysis if all oligonucleotides for each gene had a mean adjusted P value of <0.05. The study is MIAME compliant. Raw data has been deposited in GEO (accession no. GSE58946).

The Gene Ontology (GO) dataset for biological process was downloaded from the **P. tricornutum** database at Joint Genome Institute (http://genome.jgi-psf.org/Phatr2/Phatr2.home.html). GO terms assigned to significantly regulated genes at each time point were listed separately for up- and downregulated genes. Metabolic pathways were analysed using the DiatomCyc database (Fabris et al., 2012).

cDNA synthesis and quantitative real-time PCR

cDNA synthesis was performed using 1 μg of total RNA with a QuantiTect Reverse Transcription Kit (Qiagen) following the manufacturer’s instructions. cDNA samples were diluted five times in ddH₂O before use for quantitative real-time PCR (qRT-PCR) analysis.

Three biological replicates from all treatments were used to perform qRT-PCR on a LightCycler 480 using a LightCycler 480 SYBR Green I Master kit (Roche Applied Science), with a program comprising pre-incubation for 5 min at 95 °C, followed by 50 cycles of amplification consisting of 10 s at 95 °C, 10 s at 55 °C, and 10 s at 72 °C. Primer sequences used in the qRT-PCR experiment are given in Table 1. The microarray dataset was screened for genes that were non-responsive to N deprivation at both time points. Based on this screen, Exportin1 (Phatr2_24186) and Aureochrome1 (Phatr2_8113) were selected as reference genes for the qRT-PCR analysis. PCR efficiencies and C_t values were calculated by linear regression using the LinRegPCR software (Ramakers et al., 2003; Ruijter et al., 2009), and the mean PCR efficiency was calculated for each primer pair. PCR efficiencies and C_t values were used in the REST 2009 software (Pfaffl et al., 2002) to calculate the statistical significance of difference in expression levels in various treatments. The target genes were normalized to the reference genes in the REST 2009 software.

**Results**

Effect of deprived levels of N on physiological responses in *P. tricornutum*

*P. tricornutum* cell growth was monitored daily in both cultures. All N-replete cultures remained in the exponential phase throughout the time course of the experiment. During the experiment period, cell density increased from 5 × 10⁴ to 1.92 × 10⁶ cells ml⁻¹ in replete cultures (Fig. 1A). N-free cultures showed similar cell growth compared with N-replete cultures until 48 h, but significantly lower growth at 72 h, with a cell density of 1.02 × 10⁶ cells ml⁻¹. In order to compare physiological and transcriptional responses before and after the N deprivation started to affect cell growth, the time points of 48 and 72 h were chosen for further physiological and molecular experiments.

Nutrient assays of dissolved inorganic nitrate and phosphate demonstrated that none of the replete cultures encountered any deprivation in dissolved inorganic nitrate or phosphate during the whole experiment (Table 2). An increase in the C:N ratio was observed at both time points in N-deprived cells. In N-deprived cells, C:N deviated from the Redfield ratio (Redfield, 1934). The N:P ratio in replete cells was well below the Redfield ratio, but the cultures were still in exponential phase. Reduction of N:P in deprivd cells coincided with nitrate loss in the medium.

Measurements of chlorophyll a and fucoxanthin (the major carotenoid in diatoms) levels per cell showed that

**Table 1. Genes analysed by real-time qPCR and their respective primers**

<table>
<thead>
<tr>
<th>Phatr2 ID</th>
<th>Accession</th>
<th>Description</th>
<th>Orientation</th>
<th>Sequence (5′→3′)</th>
<th>Amplicon size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>54101</td>
<td>XP_002177983</td>
<td>Nitrate transporter</td>
<td>Forward</td>
<td>GGAATACCTTGCTGTCTCTATGC</td>
<td>58</td>
</tr>
<tr>
<td>34373</td>
<td>XP_002178768</td>
<td>Molybdopterin biosynthesis-like protein CNX5</td>
<td>Reverse</td>
<td>AGGAGAACCTGAGGGTCATCT</td>
<td>100</td>
</tr>
<tr>
<td>17344</td>
<td>XP_002176623</td>
<td>Adenine/guanine permease</td>
<td>Forward</td>
<td>ATGCTCAAGGACCGATGCAAAC</td>
<td>130</td>
</tr>
<tr>
<td>18049</td>
<td>XP_002177871</td>
<td>LHC protein LHC F1</td>
<td>Reverse</td>
<td>CAGCTTGTCTCAGTCAACA</td>
<td>87</td>
</tr>
<tr>
<td>49339</td>
<td>XP_002183906</td>
<td>Pyruvate carboxylase PYC2</td>
<td>Forward</td>
<td>AAATTACGGCTCTTCC</td>
<td>82</td>
</tr>
<tr>
<td>24186</td>
<td>XP_002185483</td>
<td>Exportin 1-like protein XPO1</td>
<td>Reverse</td>
<td>TCCGCTGGTATCATGCAAG</td>
<td>116</td>
</tr>
<tr>
<td>8113</td>
<td>XP_002183783</td>
<td>Aureochrome AUREO1a</td>
<td>Forward</td>
<td>GGTTTCTCAAGTTTAGAACAG</td>
<td>116</td>
</tr>
</tbody>
</table>
the content of these pigments declined progressively in N-deprived cells at both time points, while both pigments were stable in control cells (Table 2). In contrast, the ratio between chlorophyll a and fucoxanthin did not change. We also monitored the effect of N deprivation on the activity of PSII. Maximum quantum yield of PSII ($F_v/F_m$), applied as a proxy measure of photosynthesis, was similar in N-replete and N-deprived cultures after 24 h. A clear drop in $F_v/F_m$ was observed in N-deprived cells ($F_v/F_m=0.33$) after 72 h, while the ratio remained unchanged ($F_v/F_m=0.68$) in the control cultures (Fig. 1B). Quantification of neutral lipids by confocal laser scanning microscopy and BODIPY 505/515 showed that the neutral lipid content increased significantly at 72 h ($t$-test: $P=1.87 \times 10^{-6}$) and was 29.3% higher in N-deprived cells compared with N-replete cells (Fig. 2A). Representative images from the analysis showed that the lipid droplets were larger and more strongly stained by the BODIPY marker in N-deprived cells than in replete cells, implying higher lipid accumulation (Fig. 2B).

Metabolite profiling of the responses to N deprivation was performed using GC-MS. Strong effects on the central metabolism were revealed, with a significant decrease in most of the N-containing metabolites and major fatty acids (Supplementary Table S1). The regulation of biosynthesis and significance of distinct metabolites is further discussed in subsequent sections.

**Gene expression**

Transcriptome responses at 48 and 72 h after N deprivation were analysed using whole-genome oligonucleotide microarrays. The treatment led to strong transcriptome responses: 5279 genes were significantly regulated ($P<0.05$) in N-deprived cultures compared with N-replete cultures 48 h after N deprivation. As expected, the stronger N deprivation at 72 h affected even more genes (6629). Comparison of the N-replete cultures at 48 and 72 h resulted in only 22 significantly regulated genes, probably reflecting higher cell densities (results not shown).

GO analysis was performed on the dataset. As the *P. tricornutum* genome is still poorly annotated, GO terms are assigned to a limited number of genes. The GO analysis still provided an overview of the processes most affected by N deprivation. The most enriched GO terms were similar at 48 h (Supplementary Fig. S1, available at JXB online) and 72 h (Fig. 3); however, there were large differences between GO terms enriched in up- and downregulated genes. The most frequent GO term among the downregulated genes was protein biosynthesis. Other GO terms related to ribosomal assembly and translation were also enriched among downregulated genes, indicating reduced protein biosynthesis. Furthermore, photosynthesis light harvesting was the fourth most used GO term among the downregulated genes, indicating downsizing of the light-harvesting apparatus. In contrast,

---

**Fig. 1.** Physiological responses of *P. tricornutum* to nitrate deprivation. Growth curves (A) and changes in maximum quantum yield ($F_v/F_m$) (B) of *P. tricornutum* in N-replete (f/2 medium) and N-deprived (f/2 medium minus nitrate) cultures. Arrows indicate sampling time points. Values are means±standard deviation of four biological replicates.

**Table 2.** Changes in chemical composition, medium nutrient concentration, and pigment concentration of nitrogen-replete (N+) and nitrogen-deprived (N–) cultures 48 and 72 h after N deprivation (n=4)

<table>
<thead>
<tr>
<th></th>
<th>N+ 48 h</th>
<th>N– 48 h</th>
<th>N+ 72 h</th>
<th>N– 72 h</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cellular nutrient content</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>µg C: µg N</td>
<td>6.03±0.32</td>
<td>9.46±0.73</td>
<td>5.44±0.19</td>
<td>14.65±0.77</td>
</tr>
<tr>
<td>µg N: µg P</td>
<td>4.24±0.26</td>
<td>2.37±0.31</td>
<td>4.89±0.18</td>
<td>1.67±0.1</td>
</tr>
<tr>
<td><strong>Medium concentration</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>µg PO₄³⁻ / l</td>
<td>612.77±2.39</td>
<td>618±2.24</td>
<td>316.99±26.73</td>
<td>576.72±31.38</td>
</tr>
<tr>
<td>µg (NO₃⁻ + NO₂⁻) / l</td>
<td>8797.38±49.23</td>
<td>3.98±1.07</td>
<td>7550.87±108.79</td>
<td>3.8±0.7</td>
</tr>
<tr>
<td><strong>Pigment concentration</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>fg Chl a cell⁻¹</td>
<td>271.4±25.89</td>
<td>153.2±5.96</td>
<td>271.6±26.29</td>
<td>76.9±9.42</td>
</tr>
<tr>
<td>fg fucoxanthin cell⁻¹</td>
<td>100.7±3.21</td>
<td>56.8±2.25</td>
<td>96.2±7.73</td>
<td>32.4±3.98</td>
</tr>
</tbody>
</table>
in the microarray analysis. The qRT-PCR results correlated well with the microarray analysis (Supplementary Fig. S2, available at JXB online).

**Effect of N deprivation on N metabolism**

*N uptake and assimilation*

Transcriptional responses to N deprivation of *P. tricornutum* showed that uptake, assimilation, and scavenging mechanisms were activated (Fig. 5). In our experiment, transcript levels of genes involved in NO\(_3\)\(^-\), NH\(_3\), and urea transport were upregulated (Fig. 4 and Supplementary Dataset S1, available at JXB online). Of four ammonium transporters detected in our microarray data, three were upregulated. The induction of a nitrate transporter (Phatr2_54101) was confirmed by qRT-PCR (Supplementary Fig. S2). Increased transcription of genes encoding nitrate reductase (NR) and both NAD(P)H- and Fd-dependent nitrite reductase was observed at 72 h after deprivation (Figs 4 and 5). Interestingly, two genes encoding molybdopterin biosynthesis proteins were induced (Supplementary Dataset S1). These enzymes might be orthologues of the *Arabidopsis thaliana* cofactor of NR and xanthine dehydrogenase CNX5 and CNX2, respectively (Schwarz and Mendel, 2006). The biosynthesis of molybdenum cofactor (Moco), which forms the active site of molybdenum (Mo) enzymes in eukaryotes, involves six enzymes. The qRT-PCR result also confirmed upregulation of the CNX5 orthologue (Phatr2_34373; Supplementary Fig. S2). None of the genes encoding plastidial GSII/Fd-GOGAT and mitochondrial GSIII (GLNA), which are required for ammonium assimilation, were regulated (*P*<0.05). However, increased transcript levels of two different isoforms of NAD(P) H-dependent glutamate synthase (NADPH-GOGAT, GltD and GltA) were observed in N-deprived cells (Fig. 5). Glutamate dehydrogenase (GDH) is another enzyme that catalyses the reversible conversion of 2-oxoglutarate (2-OG) to glutamate. We observed increased expression of an NADP-GDH (Phatr2_13951; Fig. 5 and Supplementary Dataset S1).

*N scavenging from various organic compounds*

We observed induction of genes encoding two glutamyltRNA(Gln) amidotransferase-like proteins (Phatr2_50401 and Phatr2_45118) and three acetamidase/formamidases (Phatr2_54476, Phatr2_37952, and Phatr2_37719) at both time points in N-deprived cells (Fig. 5). Phylogenetic analyses indicated that Phatr2_54476 is related to FmdA-type formamidases (Supplementary Fig. S3, available at JXB online); the main sub-strate of *Methylophilus methylotropus* FmdA (Wyborn et al., 1994), as well as lupin LaFmd (Rath et al., 2010), is formamide. Phatr2_37952 and Phatr2_37719, which were strongly induced at both time points, encode amidohydrolases belonging to a poorly characterized clade with low similarity to FmdA-type formamidases (Supplementary Fig. S3).

**Purine and pyrimidine biosynthesis and degradation**

Most of the transcripts involved in biosynthesis of purine and pyrimidine were downregulated (Supplementary Dataset S1). Simultaneously, we observed upregulation of several transcripts involved in their catabolic processes, such as urine and pyrimidine deaminases. Furthermore, uracil-xanthine permease
Nitrogen-deprivation response in diatoms

Transcripts encoding Phatr2_16991 and adenine/guanine permease (Phatr2_17344) were upregulated in N-deprived cultures; the response was confirmed by qRT-PCR analysis (Supplementary Fig. S2). Urease (Phatr2_29702), which catalyses the hydrolysis of urea into CO$_2$ and NH$_4^+$, was also transcriptionally induced following N deprivation (Supplementary Dataset S1).

Protein biosynthesis, folding, and degradation

N deprivation influenced both biosynthesis and degradation of proteins and amino acids. Most amino acid biosynthesis pathways were transcriptionally repressed; the strongest downregulation was found for transcripts encoding homoserine dehydrogenase (Phatr2_26813) and N-acetylglutamate kinase (Phatr2_3969) (Fig. 4). Reduction of protein biosynthesis could be observed as a decrease in mRNA levels of many genes encoding aminoacyl-tRNA synthetases, as well as ribosomal subunits and translation elongation factors. The transcription levels of genes encoding 18 peptidylprolyl isomerases and two protein disulfide isomerases that catalyse protein folding were also reduced. In contrast, the transcript levels of genes encoding amino acid degradation enzymes, such as those related to catabolism of branched-chain amino acids, were upregulated (Fig. 4). Five identified autophagy-related genes were upregulated at one or both time points in N-deprived cells (Supplementary Dataset S1). The mRNA levels of several genes involved in ubiquitination were upregulated, but most of the proteasome subcomponents were downregulated.

Effect of N deprivation on photosynthesis and pigment biosynthesis

Glutamate, the main precursor of chlorophyll biosynthesis, declined under N-deprived conditions (Supplementary Table S1). In line with the reduced chlorophyll $a$ and fucoxanthin levels in N-deprived cells, the expression levels of most of the genes encoding enzymes involved in the chlorophyll $a$ and carotenoid biosynthetic pathways were also repressed (Figs 4 and 5, and Supplementary Dataset S1). Of 39 differentially regulated genes encoding LHC proteins, only red algal-like
LHCR10 and two high-light-induced proteins (HLIP1 and HLIP1b) were significantly upregulated at both time points, whereas the LI818-like LHCX4 and LHCR7 showed moderate upregulation 72 h after N deprivation (Fig. 4). Among the downregulated LHCs, repression of LHCF1 was confirmed by qRT-PCR (Supplementary Fig. S2). Similarly, a majority of the transcripts involved in photosynthesis were downregulated (Figs 4 and 5, and Supplementary Dataset S1). Furthermore, a chloroplastic ferredoxin-NADP reductase (Phatr2_23717) was strongly downregulated, indicating that NAD(P)H production through photosynthesis was reduced (Fig. 4).

**Effect of N deprivation on carbon and lipid metabolism**

**Carbon fixation**

Downregulation of several genes connected to the bio-physical carbon-concentrating mechanism was observed in N-deprived cells. Of five carbonic anhydrase (CA) genes...
related to the biophysical carbon-concentrating mechanism that were significantly regulated 72 h after N deprivation (Supplementary Dataset S1), transcript levels of CA-III and two β-CAs (PtCa1 and PtCa2) decreased. A chloroplast bicarbonate transporter (SLC4A_1) was also repressed. In accordance with the β-CA and SLC4A_1 downregulation, a majority of the genes encoding enzymes of the Calvin cycle were downregulated (Figs 4 and 6). Upregulation of several transcripts encoding enzymes involved in the mitochondrial decarboxylation under N-deprived conditions was observed in N-deprived cells (Supplementary Dataset S1). Downregulation of plastid-localized pyruvate carboxylase 2 (PYC2) was confirmed by qRT-PCR analysis (Supplementary Fig. S2).

TCA cycle
Consistent with upregulation of the mitochondrial decarboxylation, TCA cycle transcripts were induced (Figs 5 and 6, and Supplementary Dataset S1). However, genes encoding enzymes towards the end of the TCA cycle were not regulated. Aconitate hydratase (Phatr2_26290) and isocitrate dehydrogenase (Phatr2_14762) transcripts showed the highest level of upregulation. In contrast to the upregulation of TCA transcripts, we observed a decrease in the levels of most of the metabolite intermediates of the TCA cycle (Supplementary Table S1).

Chrysolaminarin biosynthesis and degradation
We observed repression of several genes encoding enzymes potentially involved in gluconeogenesis, as well as chrysolaminarin biosynthesis (Kroth et al., 2008; Chauton et al., 2013), especially 72 h after N deprivation (Fig. 6). Inversely, transcript levels of genes encoding enzymes for chrysolaminarin degradation, such as exo-1,3-β-glucosidases, increased. Chrysolaminarin degradation produces glucose; indeed, glucose levels were higher in N-deprived cells (Fig. 6 and Supplementary Table S1). Consistent with the increased glucose level, cytosolic glucokinase was also induced in our experiment.

Oxidative pentose phosphate pathway (OPPP), glycolysis, and pyruvate metabolism
Surprisingly, all OPPP transcripts were induced (Fig. 6). Transcripts of most of the putatively cytosolic glycolytic enzymes, such as phosphoglycerate mutase (PGAM_7), increased in our experiment, while transcript levels of several plastidial enzymes, such as glyceraldehyde-3-phosphate dehydrogenase (GAPC1), showed the opposite regulation (Fig. 4). In contrast to the transcriptional induction of glycolytic enzymes, metabolite levels of glucose-6-phosphate, fructose-6-phosphate, and pyruvate declined (Supplementary Table S1), which might be the result of their quick conversion to other metabolites. Consistent with the decrease in pyruvate, transcript levels of several genes responsible for pyruvate metabolism were upregulated.

Fatty acid biosynthesis and degradation
Most transcripts related to the chloroplast fatty acid biosynthetic pathway were strongly downregulated (Fig. 4 and Supplementary Dataset S1, available at JXB online); the only upregulated transcript was 3-oxoacyl-[acyl-carrier-protein] synthase (FABFb) (Phatr2_18940). We also observed lower levels of total free fatty acids in N-deprived cells (Supplementary Table S1), which might be a consequence of their incorporation into TAG.

Fig. 6. Cellular pathways and processes affected under N deprivation in *P. tricornutum*. Metabolites detected are indicated by a blue box frame. Red, blue, and black text indicates up-, down-, and no regulation of pathways, genes, or metabolites by N deprivation, respectively. 3-PGA, 3-phosphoglycerate; CA, carbonic anhydrase; ICL, isocitrate lyase; MLS, malate synthase; OPPP, oxidative pentose phosphate pathway; PEP, phosphoenolpyruvate; PK, pyruvate kinase; PPDk, pyruvate orthophosphate dikinase; RuBP, ribulose-1,5-bisphosphate.
Membrane lipid remodelling and TAG biosynthesis
TAG biosynthetic pathways are illustrated in Fig. 7. Of three differentially regulated glycerol-3-phosphate dehydrogenases, transcript levels of Phatr2_36821 increased in N-deprived cells (Fig. 7). This enzyme consumes NAD(P)H to convert dihydroxyacetone phosphate, an intermediate in glycolysis, to glycerol-3-phosphate. Subsequent transfer of two acyl-CoAs to glycerol-3-phosphate by glycerol-3-phosphate acyltransferase and acyl-glycerol-3-phosphate acyltransferase (AGPAT) result in the formation of phosphatidic acid. Of the five detected isoforms of AGPAT, only one putative isoform (Phatr2_45551) displayed increased transcript levels 72 h after deprivation, whereas three others (Phatr2_20460, Phatr2_11916, and LPT1) were suppressed. Phosphatidic acid is dephosphorylated to diacylglycerol, the main precursor of TAG. This process is catalysed by phosphatidic acid phosphatase; a putative PAP (Phatr2_40261) was weakly upregulated. Incorporation of the third fatty acyl-CoA into glycerol-3-phosphate backbone by diacylglycerol O-acyltransferase (DGAT) completes TAG formation. The mRNA levels of two isoforms of DGAT

Fig. 7. Transcriptional changes in genes related to TAG biosynthesis in response to N deprivation. Coloured squares indicate the regulation pattern of genes encoding putative enzymes functioning in the TAG biosynthetic pathway after 48 and 72 h of N deprivation, compared with N-replete cultures. Squares with a diagonal line inside indicate non-significant regulation (P > 0.05). The scale on the right represents gene expression ratio values, which were log2 transformed. Numbers indicate Phatr2 gene IDs. Gene ID 12726 (marked with an asterisk) belongs to the Phatr1 database (http://genome.jgi-psf.org/Phatr1/Phatr1.home.html).
(Phatr2_43469 and Phatr2_9794) were induced in our experiment. Although we could detect TAG accumulation at both the molecular and physiological level, the transcript abundance of several TAG lipases was induced in N-deprived cells.

Upregulation of four isoforms of phospholipase C and two isoforms of phospholipase D indicated that membrane phospholipids are degraded under N deprivation to provide the TAG precursors phosphatidic acid and diacylglycerol (Fig. 7). The highest induction was seen for phospholipase C (Phatr2_48445), with an approximately 4- and 10-fold increase 48 and 72h after deprivation, respectively. Transcript levels of a putative phospholipid:diacylglycerol acyltransferase enzyme (PDAT, Phatr2_8860) were also upregulated at 72h in N-deprived cells (Fig. 7).

### Discussion

Previous studies in diatoms and other microalgae have demonstrated that these organisms undergo dramatic metabolic changes in response to N starvation (Hockin et al., 2012; Valenzuela et al., 2012; Yang et al., 2013). Performing an integrated analysis of the response to N deprivation in *P. tricornutum*, we confirmed these modifications at the physiological, metabolite, and transcriptome levels.

**Reprogramming of N metabolism**

We observed a higher C:N ratio than that suggested by the Redfield ratio (Redfield, 1934) in N-replete and N-deprived cultures, indicating that the C:N:P composition of phytoplankton and marine particulate matter is flexible, especially in nutrient-deprived cells (Geider and La Roche, 2002). Furthermore, an increase in the C:N ratio of N-deprived cultures is probably a result of biomass increase after N exhaustion.

Although transcriptional regulation is indicative and not necessarily directly linked to changes at protein level, some trends are evident. Due to a decrease in the N content of N-deprived cells, many processes connected to N metabolism were affected. A major response to N deprivation was to increase the cellular capacity for N uptake and nitrate reduction to ammonium, as observed at the transcript level. This phenomenon was reported previously in N-deprived *P. tricornutum* and other microalgae at the transcriptome level (Mock et al., 2008; Miller et al., 2010; Valenzuela et al., 2012) (Fig. 7). However, the downregulation of one ammonium transporter (Phatr2_54981) was in agreement with previous reports (Haimovich-Dayan et al., 2013; Krell et al., 2007; Yang et al., 2013). The induction of two molybdopterin biosynthesis genes could be related to an increased need of Moco for NR and xanthine dehydrogenase in *P. tricornutum*. In contrast to our results, a NR gene was repressed under N stress in *Emiliania huxleyi*; the authors postulated that expression of the NR gene is stimulated under high NO₃ concentrations (Bruhn et al., 2010). In addition, they observed no co-ordination between the regulation of nitrate reduction at the transcript and protein levels. Despite the increase in the expression level of genes encoding nitrate-reducing enzymes in N-deficient *T. pseudonana* (Mock et al., 2008), their protein levels were found to decrease in a proteomic study (Hockin et al., 2012); the authors postulated that the levels of these enzymes might be controlled by post-transcriptional modifications. In photosynthetic eukaryotes, ammonium assimilation primarily occurs inside the chloroplast (Bowler et al., 2010). Although we could not detect any regulation of GSII/Fd-GOGAT or mitochondrial GLNA, increased transcript levels of two different NADPH-GOGAT isoforms (GltD and Gltx) indicated that cells increased their N scavenging mechanisms to assimilate more ammonium from other pathways such as protein and amino acid degradation (Fig. 5). In addition, the upregulated levels in N-deprived cells of another ammonium assimilating enzyme, NADP-GDH, might also contribute to ammonium assimilation. GDH is generally believed to act as a catabolic enzyme, catalysing the oxidative deamination of glutamate to 2-OG, and previous studies have implied that GDH is a minor contributor to NH₄⁺ assimilation (Zehr and Falkowski, 1988; Guerra et al., 2013). However, the observed decrease in 2-OG levels might indicate that GDH also can perform the anabolic reaction to scavenge NH₄⁺ under N deprivation. Previous studies have demonstrated that diatoms are able to use other sources of N, such as amides, amines, urea, and amino acids (Shah and Syrett, 1982; Baker et al., 2009). Besides the increase in N-scavenging mechanisms, upregulation of amidases and acidamidase/formamidases indicates that *P. tricornutum* can degrade organic N sources such as amides and formamide from intracellular or possible extracellular sources to produce ammonium when faced with N deprivation, as observed in N-deficient *Aureococcus anophagefferens* (Wurch et al., 2011). In support of our results, a formamidase transcript was induced under N stress in *E. huxleyi* (Bruhn et al., 2010).

Biosynthesis of several amino acids relies on the availability of glutamate, which declined after N deprivation as a consequence of a reduced N pool (Supplementary Table S1). The lower glutamate level compared with the control resulted in a strong drop in levels of other amino acids and repression of amino acid biosynthetic pathways. A decline in the cellular amino acid pool as a result of N deprivation is consistent with results from other diatoms (Granum et al., 2002). Simultaneous with the suppression of amino acid biosynthesis, increased degradation of amino acids through various catabolic pathways was observed at the transcript level, which produced several carbon-containing intermediates that can enter the TCA cycle. Consistent with increased transcripts associated with catabolism of branched-chain amino acids in our experiment, Ge et al. (2014) suggested that branched-chain amino acid degradation directs carbon and energy towards TAG accumulation in N-deprived *P. tricornutum*.

All purine and pyrimidine nitrogen compounds originate from amino acids (glutamine, aspartate, and glycine) (Zrenner et al., 2006). Reduced biosynthesis of purine and pyrimidines is probably a result of decreases in the amino acid pools (Supplementary Table S1) and cell growth (Fig. 1A). At the same time, recycling of purines and pyrimidines provides the cells with an important N source during
N deprivation. Upregulation of the urease gene could also be related to purine degradation. \( \text{NH}_4^+ \) produced from the hydrolysis of urea is hypothesized to be redirected to mitochondria for amino acid biosynthesis via mitochondrial GS/GOGAT (Allen et al., 2011). The strong induction of purine/ pyrimidine permeases might explain the ability of diatoms to import purine and pyrimidine from the environment under limited N conditions, as reported by Allison and Syrett (1987). However, Berg et al. (2008) showed that the purine permease \( \text{AaURA} \) is expressed during growth of \( A. \) anophagefferens on a number of N sources, indicating its role as an important nitrogen source for proliferation of this organism. The high expression of \( \text{AaURA} \) might be related to the growth habitat of \( A. \) anophagefferens in shallow coastal waters, which are in close contact with sediments rich in dissolved organic N.

Eukaryotes utilize autophagy and the ubiquitin–proteasome system for protein degradation (Onodera and Ohsumi, 2004). The ubiquitin–proteasome system is used for rapid degradation of proteins and acts mainly to degrade short-lived proteins such as transcription factors, while the turnover of long-lived proteins, which constitute 99% of cellular proteins, is processed by autophagy (Onodera and Ohsumi, 2004). While N-limited \( A. \) anophagefferens showed moderate downregulation of two autophagy-related genes (Berg et al., 2008), the transcript levels of several autophagy-related genes in our experiment and during N limitation of the green microalga \( \text{Neochloris oleoabundans} \) (Rismani-Yazdi et al., 2012) were induced. Although autophagy generally is not a selective protein degradation process, selective autophagy was stimulated under conditions of nutritional stress, especially N deficiency, in yeast and plants (Onodera and Ohsumi, 2004; Yoshimoto et al., 2004). Therefore, the induction of autophagy-related genes can be explained as a response to N deprivation by selectively degrading excessive proteins into amino acids that are recycled to protein biosynthetic pathways in order to maintain cellular homeostasis. Autophagy is highly regulated at the protein level (Klionsky and Emr, 2000); therefore, it would be worthwhile looking at regulation of autophagy components at the protein level to better understand their role under N deprivation.

Chlorosis is one of the main responses of diatoms to N deprivation. The reduced chlorophyll \( a \) level in N-deprived cells (Table 2) is probably caused at least partly by repression of its biosynthesis. In addition, the co-ordinated decrease in chlorophyll \( a \) and fucoxanthin content under N deprivation suggests that the biosynthesis of these two pigments is co-ordinated under N deprivation; a similar result was also observed in a previous study in \( P. \) tricornutum (Geider et al., 1993). In contrast, N limitation in \( Chaetoceros gracilis \) led to changes in the chlorophyll \( a: \) fucoxanthin ratio (Cleveland and Perry, 1987). The decrease in pigment biosynthesis and LHC proteins corresponds to a reduced photosynthetic apparatus of N-deprived cells and a lower requirement for pigments. These results clearly indicate that N deprivation reduces the photosynthetic efficiency, in agreement with the observed reduction in maximum quantum yield of PSII \( (F_v/F_m) \) (Fig. 1B). Despite the downregulation of LHC proteins, induction of LHCXs was reported under several stresses in diatoms (Zhu and Green, 2008). Induction of \( \text{LHCR10} \) expression by N deprivation was also observed by Yang et al. (2013); furthermore, both LH CX and LHCR-II genes, to which \( \text{LHCR7} \) and \( \text{LHCR10} \) belong, are induced by high light (Nymark et al., 2009, 2013). The increased transcript levels of \( \text{LHCR4} \), along with two LHCRs and two HLIPs, may be related to a photoprotective role during acclimation to low N levels. In summary, N-deprived \( P. \) tricornutum modified N metabolism in order to reduce synthesis of nitrogenous compounds and catabolize excessive N-containing compounds in favour of essential N compounds.

Remodelling of carbon metabolism

The major pathway used by diatoms for carbon fixation is the Calvin cycle. Since \( \beta \)-CAs and \( \text{HCO}_3^- \) transporters are required to concentrate inorganic carbon in the vicinity of Rubisco, their downregulation repressed a majority of the genes encoding enzymes of the Calvin cycle (Fig. 6). Although the Calvin cycle was downregulated, cells might employ other mechanisms such as pyruvate orthophosphate dikinase to dissipate excess energy around the photosystems to reduce the production of reactive oxygen species under N deficiency, as reported by Haimovich-Dayan et al. (2013).

Upregulation of several transcripts encoding enzymes involved in the mitochondrial decarboxylation under N-deprived conditions leads to production of oxaloacetate and pyruvate (Supplementary Dataset S1). Oxaloacetate can replenish \( \text{C}_4 \) acids of the TCA cycle, whereas pyruvate can enter the TCA cycle or fatty acid biosynthesis. The TCA cycle could also be upregulated in response to high levels of protein and amino acid degradation, which generates TCA cycle intermediates and provides precursors for resynthesis of certain amino acids, as observed by Hockin et al. (2012). Furthermore, strong upregulation of the genes encoding aconitate hydratase and isocitrate dehydrogenase leads to production of 2-OG, which acts as a precursor in ammonium assimilation. Malate from the TCA cycle could also be directed to the fatty acid biosynthetic pathway through NADP-dependent malic enzyme (Supplementary Dataset S1). Similar regulation of the TCA cycle was reported for other diatoms under N deprivation (Bender et al., 2014). Thus, a co-ordinated upregulation of the TCA cycle and mitochondrial decarboxylation might shift the flow of carbon skeletons towards fatty acid biosynthesis.

Degradation of chrysolaminarin releases glucose. Glucose cannot enter the metabolic pathway directly and must be converted to glucose-6-phosphate by cytosolic glucokinase. Phosphorylated glucose could further enter the glycolytic pathway and/or OPPP (Fig. 6). Upregulation of OPPP produces NAD(P)H supplying NAD(P)H-dependent pathways like lipid synthesis and nitrogen assimilation. Utilization of glucose-6-phosphate through glycolysis produces energy in the form of ATP and NAD(P)H, as well as the glycolysis end product, pyruvate. Further metabolism of pyruvate produces acetyl-CoA, which can enter the TCA cycle, or \( \text{de novo} \) fatty acid biosynthesis in the chloroplast. Thus, increased degradation of chrysolaminarin, along with induction of OPPP
and glycolysis, could provide N-deprived cells with reducing equivalents to balance reduced NAD(P)H production from photosynthesis, as well as carbon fluxes for the TCA cycle and fatty acid biosynthesis.

The microarray data showed downregulation of de novo fatty acid biosynthesis genes (Fig. 4 and Supplementary Data S1), supporting the observed decrease in free fatty acids (Supplementary Table S1). In contrast, Yang et al. (2013) reported increased total fatty acid levels in N-deprived cells, while RNA-sequencing data from their experiment showed that most of the transcripts involved in chloroplast de novo synthesis of fatty acids were downregulated. We conclude that, under exponential growth, the high cell division rate leads to a high demand for membrane lipids in newly synthesized cells. In contrast, cell division slows or halts in cells faced with N limitation, and there is less need for membrane lipids and fatty acids. Even though de novo synthesis of fatty acids is downregulated, fatty acids are still produced and accumulate in the form of neutral lipids. Moreover, repression of the fatty acid β-oxidation pathway might direct fatty acids from degradation of membrane lipids to TAG production.

Under favourable conditions, fatty acids are incorporated into membrane lipids. The main classes of lipids in diatom chloroplasts are sulphydroxydiacylglycerol, monogalactosyldiacylglycerol, digalactosyldiacylglycerol, phosphatidylglycerol, and phosphatidylcholine; however, phosphoglyceride levels are higher in the endoplasmic and plasma membranes (Hu et al., 2008; Goss and Wilhelm, 2010). Therefore, the downregulation in membrane lipid biosynthesis could be related to a reduced demand for membrane lipids due to the reduced cell growth rate. A major effect of N starvation in many microalgae is the switch in lipid production towards neutral lipid accumulation, mainly TAGs, which act primarily as energy reserves inside the cells (Hu et al., 2008). Increased levels of neutral lipids were also observed in our study (Fig. 2). TAG can be synthesized through different pathways, one of which uses glycerol-3-phosphate as a precursor (Fig. 7) (Hu et al., 2008). The accumulation of TAG was consistent with a decrease in the amount of free glycerol-3-phosphate inside the cells (Supplementary Table S1), implying that this compound might be used as a precursor for TAG biosynthesis. Furthermore, TAG can also accumulate via degradation of glycerophospholipids. The increased expression of phospholipase C and phospholipase D suggests that these enzymes contribute substantially to TAG accumulation through processing of phospholipids. An acyl-CoA-independent mechanism for biosynthesis of TAG from diacylglycerol using phospholipids as acyl donors is catalysed by a putative gene encoding phospholipid:diacylglycerol acyltransferase (PDAT, Phatr2_8860). This could also contribute to TAG accumulation upon N deprivation. The upregulation of several TAG lipases could be related to saturated lipid bodies and could act to recycle previously synthesized TAGs with new TAGs, or they might have been induced to make modifications in TAG structure. Overall, our results suggest that both de novo TAG biosynthesis and remodelling of membrane lipids play important roles in TAG accumulation under N deprivation.

Conclusions

A combined analysis of transcriptional and non-targeted metabolite profiling, along with physiological and biochemical experiments, revealed transcriptional, metabolic, and physiological acclimation in the diatom *P. tricornutum* under conditions of N deprivation. The global expression data suggested that *P. tricornutum* is able to remodel N through catabolism of internal N-containing resources such as amino acids and proteins. N deprivation was also accompanied by a reduction of pigment pools and photosynthetic capacity. We also showed large changes in genes related to carbon and lipid metabolism. Decreased levels of carbon skeletons due to suppression of the Calvin cycle were compensated by breakdown of chrysolaminarin, leading to upregulation of OPPP, cytosolic glycolysis, pyruvate metabolism, and the TCA cycle. These pathways provide precursors for fatty acid biosynthesis. In addition, remodelling of membrane lipids and upregulation of the de novo TAG biosynthetic pathway was further supported by increased levels of neutral lipids, indicating TAG accumulation under N deprivation. Our study provides a detailed picture of *P. tricornutum* acclimation to N deprivation, and can be used as a guide for future metabolic manipulations to increase TAG production.

Supplementary data

Supplementary data are available at *JXB* online.

Supplementary Fig. S1. GO analysis of significantly regulated genes after 48 h of nitrate deprivation.

Supplementary Fig. S2. qRT-PCR analysis of selected genes.

Supplementary Fig. S3. Phylogenetic analysis of FmdA-type amidases/formamidases in *P. tricornutum*.

Supplementary Table S1. Tentatively identified algal metabolites based on GC-MS profiling.

Supplementary Table S2. Genes analysed by qRT-PCR and their respective primers.

Supplementary Dataset S1. Representative mRNA transcripts grouped by cellular pathway.

Acknowledgements

We thank Torfinn Sparstad for excellent technical assistance with transcriptome analyses, Matilde S. Chauton for access to instrumentation and technical advice in particulate nutrient analysis, Kjersti Andresen for help in pigment and nutrient experiments, and Bjørnar Sporsheim for kind guidance in lipid analysis. This work was supported by the Research Council of Norway through grants 184146 and 207794.

References


Mock T, Kroon BMA. 2002. Photosynthetic energy conversion under extreme conditions—I: important role of lipids as structural modulators and energy sink under N-limited growth in Antarctic sea ice diatoms. Phytochemistry 61, 41–51.


