Remodeling of intermediate metabolism in the diatom *Phaeodactylum tricornutum* under nitrogen stress

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Contributed by Paul G. Falkowski, December 1, 2014 (sent for review October 15, 2014; reviewed by Angela Falciatore and Thomas Mock)

Diatoms are unicellular algae that accumulate significant amounts of triacylglycerols as storage lipids when their growth is limited by nutrients. Using biochemical, physiological, bioinformatic, and reverse genetic approaches, we analyzed how the flux of carbon into lipids is influenced by nitrogen stress in a model diatom, *Phaeodactylum tricornutum*. Our results reveal that the accumulation of lipids is a consequence of remodeling of intermediate metabolism, especially reactions in the tricarboxylic acid and the urea cycles. Specifically, approximately one-half of the cellular proteins are cannibalized; whereas the nitrogen is scavenged by the urea and glutamine synthetase/glutamate 2-oxoglutarate aminotransferase pathways and redirected to the de novo synthesis of nitrogen assimilation machinery, simultaneously, the photobiological flux of carbon and reductants is used to synthesize lipids. To further examine how nitrogen stress triggers the remodeling process, we knocked down the gene encoding for nitrate reductase, a key enzyme required for the assimilation of nitrate. The strain exhibits 40–50% of the mRNA copy numbers, protein content, and enzymatic activity of the wild type, concomitant with a 43% increase in cellular lipid content. We suggest a negative feedback sensor that couples photosynthetic carbon fixation to lipid biosynthesis and is regulated by the nitrogen assimilation pathway. This metabolic feedback enables diatoms to rapidly respond to fluctuations in environmental nitrogen availability.

**Significance**

When starved for nutrients, diatoms redirect carbon toward biosynthesis of storage lipids, triacylglycerols (TAGs). We examined how this modification is achieved in the diatom *Phaeodactylum tricornutum*. Under nitrogen stress, the cells cannibalized their photosynthetic apparatus while recycling intracellular nitrogen and redirecting it to synthesize nitrogen assimilation enzymes. Simultaneously, they allocated newly fixed carbon toward lipids. In contrast, a nitrate reductase knocked-down strain shunted ∼40% more carbon toward TAGs than the wild type without losing photosynthetic capacity. Our results show that diatoms can remodel their intermediate metabolism on environmental cues and reveal that a key signal in this remodeling is associated with nitrogen assimilation. This insight informs a strategy of developing a much more efficient pathway to produce algal-based biofuels.


**Reviews:** A.F., CNRS, and T.M., University of East Anglia.

The authors declare no conflict of interest.

Data deposition: The DESeqs output for all 10,402 genes reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, www.ncbi.nlm.nih.gov/geo (accession no. GSE56346), and all reads reported in this paper have been deposited in the National Center for Biotechnology Information's Short Read Archive (accession no. SRP010732).

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1419818112/-/DCSupplemental.
and its resupply. Our results reveal how carbon is redirected toward lipid biosynthesis under nitrogen stress in *P. tricornutum*.

**Results**

**Physiological Characteristics.** Under optimal growth conditions, when nitrate (as the sole nitrogen source) and other nutrients are not limiting and light is saturating for growth, ~45% of the photosynthetically fixed carbon was incorporated into protein (Fig. 1). Under these conditions, only ~9% of the cellular carbon was allocated into lipids, of which ~1% was allocated for storage as TAGs. However, when being nitrogen-stressed, the fraction of the carbon that was shunted toward cellular protein decreased by 50%, whereas the carbon allocated to TAGs increased by an order or magnitude.

Biophysical analysis of the kinetics of variable chlorophyll fluorescence on the microsecond timescale (17) revealed that nitrogen-stressed cells had ~50% lower photosynthetic energy conversion efficiency [determined as the ratio of variable fluorescence (Fv) to maximal fluorescence (Fm)]. Fv/Fm in photosystem II (PSII), but the effective absorption cross-section of PSII reaction center (εPSII) was 17% ± 1%.

Although the abundance of total protein decreased by ~50% under nitrogen stress, the losses were disproportionally distributed. Proteins involved in photosynthetic pathways decreased markedly. Western blot analysis revealed that the plastid-encoded proteins ribulose-1,5-bisphosphate carboxylase/oxygenase (RuBisCo) and the D1 protein (PsbA), a core subunit of PSII, decreased by 60% and 20%, respectively. Despite the massive increase in carbon allocated to TAGs (Fig. 1 and Tables S1 and S2), DGAT type II (DGAT2D), a key rate-limiting enzyme associated with TAGs biosynthesis, decreased by 70% (Fig. 2). In contrast, nitrate reductase (NR), an enzyme essential for the assimilation of nitrate, increased by almost fourfold under nitrogen stress (Fig. 2).

Nitrogen stress also induced a change in the lipid profile; in conjunction with the marked increase in TAGs, nitrogen stress also led to threelfold decrease in intact polar lipids (IPLs), which primarily are associated with membranes (Fig. S1 and Tables S1 and S2); consequently, the cellular IPL/TAG ratio was 25-fold higher in nitrogen-replete cells. In nitrogen-stressed cells, lipids associated with thylakoid membrane lipids, such as monogalactosyldiacylglycerol (MGDG), digalactosyldiacylglycerol (DGDG), and sulfoquinovosyldiacylglycerol (SQDG), accounted for >90% of the cellular IPLs and were dominated by DGDG, which can increase the structural stability of thylakoid membranes in diatoms (18) (Fig. S2 and Tables S1 and S2).

**Transcriptome Analysis.** Analysis of the transcriptomes of axenic cultures in the two growth conditions yielded 10,234 sequences corresponding to 98% of the annotated genes (genome.jgi-psf.org/Phatr2), of which 5,620 genes were differentially expressed (DE) (Materials and Methods). Of the DE genes, 49% were up-regulated, whereas 51% were down-regulated in the nitrogen-stressed cells compared with the control. We further manually assigned genes related to metabolic pathways and functions to 1 of 16 categories and calculated the percentage of down-regulated, up-regulated, and non-DE (NDE) genes in each category (Fig. S2 and Dataset S1). This analysis revealed that genes related to nitrogen assimilation and the urea and TCA cycles were mostly down-regulated (Figs. S2–S4).

The most up-regulated genes in the transcriptome were associated with nitrate and urea uptake and assimilation [including the glutamine synthetase/glutamine 2-oxoglutarate aminotransferase (GOGAT) pathway], the pyruvate dehydrogenase complex, and the TCA cycle (Fig. 3). Genes related to chlorophyll biosynthesis and fucocyanin chlorophyll a/c binding proteins were the most down-regulated; fucocyanin chlorophyll a/c binding protein transcripts decreased by >300-fold. The expression of genes associated with photosynthetic carbon assimilation decreased markedly and was accompanied by down-regulation of the genes involved in both glycolytic and gluconeogenesis pathways. In addition, genes related to the pentose phosphate pathway and aerobic respiration exhibited a very high fraction of the NDE genes (Figs. S2 and S3 and Dataset S1).

About one-third of all the known genes related to lipids biosynthesis were down-regulated, including acetyl-CoA carboxylase, which represents a major branch point in intermediate metabolism that commits carbon to lipid biosynthesis (Fig. S4 and Dataset S1). Of the rest, 62% were NDE, and only 8% were up-regulated (Fig. S2 and Dataset S1). The genes responsible for TAGs metabolism in *P. tricornutum* are not well-annotated; however, based on the gene assignments, the expressions of genes related to TAGs biosynthesis were mostly NDE. Of five known DGAT type II, only *dgat2D* (43469) was up-regulated (Fig. S4B and Dataset S1). Fifty-seven percent of the genes related to biosynthesis of IPIs were NDE as were all genes related to biosynthesis of phospholipid, DGDG, and MGDG biosynthesis. The two genes related to SQDG biosynthesis were either NDE or down-regulated (Dataset S1).

**Prediction of Metabolic fluxes.** Based on an assumption of steady-state growth under the two conditions and the population of mRNAs of each of the genes in the transcriptomes, we calculated predicted fluxes of the intermediate metabolites of the cells. The analysis predicted that the fluxes of 92% of the reactions involving intermediate metabolites were lower than in nitrogen-replete conditions. The model further predicts that fluxes related to the production of glutamate, glutamine, and 2-oxoglutarate as well as those associated with the urea and TCA cycles increased significantly (Fig. 4).

Overall, this computational analysis suggests that nitrogen stress leads to a remodeling of intermediate metabolism centered around glutamate. This metabolic hub involves transfer of...
Changes in transcript abundance of genes involved in carbon and nitrogen metabolism pathways between nitrogen-stressed and -replete conditions.

**Fig. 3.** Changes in transcript abundance of genes involved in carbon and nitrogen metabolism pathways between nitrogen-stressed and -replete conditions. A full description of the genes, exact fold-change values, and false detection rate can be found in Dataset S1. GOGAT, glutamine oxoglutarate aminotransferase; GS, glutamine synthetase; NH$_4^+$, ammonium; NO$_3^-$, nitrate; PDC, pyruvate dehydrogenase complex.

An amino group to form glutamine and subsequent reactions involving both glutamic dehydrogenase and GOGAT (Fig. 4A) and seems to critically conserve intracellular nitrogen by recycling protein degradation products through the urea cycle (Fig. 4B). Our metabolic flux analysis further predicts that this hub is coupled to the intermediate metabolism of carbon in the TCA cycle. Specifically, the analysis predicts an increase in the oxidation of malate to oxaloacetate, resulting in an increased flux of catabolically produced reductant (Fig. 4C). In contrast, the central hub involving pyruvate, phosphoenolpyruvate, and oxaloacetate is down-regulated in nitrogen-stressed cells, and this intermediate metabolic pathway is unlikely to be a significant source of carbon for lipid biosynthesis.

**Characterization of an NR Knock-Down Strain.** To further examine the cellular responses of diatoms to nitrogen stress, we generated an NR knock-down (KD) transformant strain, NR21. This transformant had ~40% of the NR mRNA copy numbers, 50% of the NR protein amount, and 50% of the NR activity rates compared with the wild type (WT) under nutrient-replete conditions (Fig. 5A and B). The change in NR is specific; the transformant had the same amount of total protein as the WT and the amounts of RubisCo for both cell lines that were unchanged between the strains (Fig. 5A). Moreover, the strain's Fv/Fm was identical to that of the WT. However, the strain grew slower with maximal growth rates that were 70% of the WT under nutrient-replete conditions (Fig. 5C). In effect, strain NR21 is a nitrogen-deficient phenotype, which is reflected by a 43% increase in the amount of lipids produced under nutrient-replete conditions (Fig. 5D).

**Discussion**

The remodeling of intermediate metabolism resulting from nitrogen stress in the marine diatom, *P. tricornutum*, is analogous to physiological changes associated with the developmentally programmed process of grain filling in terrestrial grasses. During grain filling in wheat, for example, nitrogen stored in the photosynthetic proteins in the leaf flag is catabolized and redirected to the seeds as the plant matures (19, 20). Similar to the loss of nitrogen and impaired photosynthetic abilities of the wheat flag leaf (21), *P. tricornutum* becomes chlorotic under N stress and cannibalizes and remobilizes plastid proteins and polar lipids toward energy storage, primarily in the form of TAGs. In effect, diatom plastids function like an intracellular flag leaf under nitrogen-stressed conditions. Diatoms are able to accumulate lipids as a storage product, albeit within a single cell rather than a fruiting body (Fig. 1 and Tables S1 and S2). However, the remodeling of intermediate metabolism in nitrogen-stressed diatoms is not developmentally programmed. How is it executed in these single-celled algae?

Nitrogen stress sets a global constraint on protein translation by limiting amino acid biosynthesis (22). Thus, on nitrogen stress,
Tables S1 and S2 reporting the up-regulation of Predicted metabolic fluxes of intermediate metabolites related to NR Activity. Error bars represent Levitan et al. Protein. P. tricornutum =| NR21 glutamine synthetase; maCoA, malonyl CoA; NH$^+$ argininosuccinate; asp, aspartate; carbam-P, carbamoyl-phosphate; citrul, citrulline; pyr, pyruvate; suc-CoA, succinyl CoA. color-coded according to their pathway: black, nitrogen uptake and assimilation; red, TCA cycle. acCoA, acetyl-CoA; arg, arginine; arg-suc, arginosuccinate; asp, aspartate; carbam-P, carbamoyl-phosphate; citrul, citrulline; DHG, glutamate dehydrogenase; gin, glutamine; glu, glutamate; GS, glutamine synthetase; maCoA, malonyl CoA; NH$^+$, ammonium; NO$^-$, nitrite; NO$_2^-$, nitrate; oxa, oxaloacetate; 2-oxoglutarate; PEP, phosphoenolpyruvate; pyr, pyruvate; suc-CoA, succinyl CoA.

Like in all other eukaryotic microalgae, in diatoms, the intermediate metabolism of carbon and nitrogen metabolism is closely coupled with a hub centered around glutamate and 2-oxoglutarate (Fig. 4). Our results suggest that this hub operates to redirect intracellular nitrogen derived from the catabolism of enzymes that are temporarily incapable of supporting growth toward a set of enzymes that confers selective advantage under nitrogen stress. The latter is primarily associated with nitrogen uptake and assimilation (including NR and GS/GOGAT) degradation of proteins, including the urea cycle and glutamate dehydrogenase, and energy-generating/anaplarotic reactions, such as the TCA cycle. This observation is consistent with other studies with WT P. tricornutum reporting the up-regulation of TCA cycle genes (24, 25) and the nitrogen uptake and assimilation

**Fig. 4.** Predicted metabolic fluxes of intermediate metabolites related to intermediate carbon and nitrogen metabolic pathways under nitrogen-replete and -stressed conditions. The flux line width represents the relative volume of the flux as calculated by the model. (A) Fluxes that represent recycling of internal nitrogen recycling. Dark gray, glutamate dehydrogenase- and aspartate aminotransferase-related fluxes; light gray, GS/GOGAT pathway; black, nitrogen uptake and assimilation related fluxes. (B) Fluxes that are mainly related to the TCA and the urea cycles. Fluxes are color-coded according to their pathway: black, nitrogen uptake and assimilation and shunting carbon toward FA metabolism; blue, urea cycle; green, pyruvate-related; red, TCA cycle. acCoA, acetyl-CoA; arg, arginine; arg-suc, arginosuccinate; asp, aspartate; carbam-P, carbamoyl-phosphate; citrul, citrulline; DHG, glutamate dehydrogenase; gin, glutamine; glu, glutamate; GS, glutamine synthetase; maCoA, malonyl CoA; NH$^+$, ammonium; NO$^-$, nitrite; NO$_2^-$, nitrate; oxa, oxaloacetate; 2-oxoglutarate; PEP, phosphoenolpyruvate; pyr, pyruvate; suc-CoA, succinyl CoA.

P. tricornutum cells redirect their photosynthetically fixed carbon toward lipids, while concurrently, shunting their internal pools of nitrogen toward the nitrogen assimilation machinery (Fig. 2, Fig. S1, and Tables S1 and S2). The net effect of this intracellular reorganization is that cells can quickly respond to changes in external nitrogen availability, thereby affording a great competitive advantage in many aquatic environments where nitrogen can be introduced into the environment rapidly through turbulence (23). Thus, instead of filling grains with proteins and lipids, diatoms redirect intracellular nitrogen to enhance their nitrogen assimilation capacity at the expense of photosynthetic carbon assimilation (Fig. 2). Choreographing this intermediate metabolic dance requires one or more signal transduction pathways that lead to changes in both nuclear and plastid gene expression. Indeed, our transcriptomic data and computational flux model reveal that several central metabolic pathways, such as lipid biosynthesis, are down-regulated, whereas others, such as the TCA and nitrogen assimilatory cycles, are markedly up-regulated. Because internal cellular nitrogen is not shuttled into synthesis of more lipid biosynthetic machinery, we propose a mechanism that leads to increased lipid accumulation in nitrogen-stressed cells.

**Fig. 5.** Physiological and NR-related characteristics of exponentially grown WT P. tricornutum and the NR21 strain under nitrogen-replete conditions with nitrate as a sole nitrogen source. (A) NR and RbcL protein abundance. (B) NR characteristics: mRNA copies, protein abundance, and enzyme activity. (C) Fatty acid methyl esters (FAMEs). (D) Cell growth. (Inset) Maximal growth rates. n = 3. Error bars represent ±1 SD.
genes under nitrogen stress (25). We hypothesize that the selection for the up-regulated pathways is so fundamental to the cell's survival that suppressing a diatom's ability to incorporate nitrogen must influence the cell's carbon allocation. This hypothesis is supported by the phenotype of the NR21 KD transformant, where the restriction of nitrogen assimilation leads to massive accumulation of lipids without causing chlorosis or a significant change in photosynthetically capacity.

The most noteworthy change in the lipid profile of nitrogen-stressed *P. tricornutum* was the increase in TAGs accumulation and diversity paralleled by a decrease in IPLs (Fig. S1 and Tables S1 and S2). Regardless of whether TAGs biosynthesis is a result of the catabolism of IPLs or de novo synthesis of FAs, the final step is catalyzed by DGATs, which convert diacylglycerols to TAGs (26). Previously, it was reported that a DGAT gene could be up-regulated under nitrogen stress (24). In our study, only one of five DGAT paralogs, dgat2D (ProID 43469), had a significant increase in transcript abundance (Fig. S4A), but its relative protein amount was 70% lower under N stress (Fig. 2). This apparent paradox can be simply explained by the decreased overall metabolic flux of carbon in the cell caused by a higher quantum requirement for growth and lipid biosynthesis (11). Thus, based on our results and previously published studies, we suggest that the subsequent accumulation of TAGs under nitrogen stress seems to be a consequence of allocation of carbon and reductant, rather than an up-regulation of lipid biosynthesis genes (Figs. 1 and 4 and Figs. S1–S4) (24, 25, 27).

Our results strongly suggest that the remodeling of intermediate metabolism due to nitrogen stress is a result of at least two different processes. The first process is related to shuffling of preformed nitrogen as reflected by the loss of photosynthetic machinery. The second process is related to redirecting photosynthetically fixed carbon from amino acids to other sinks, especially lipids, similar to the findings reported by Ge et al. (27). Although nitrogen-stressed cells recycle the nitrogen from preexisting photosynthetically fixed carbon toward nitrogen-deficient storage molecules, they continuously synthesize carbon skeletons, primarily in the forms of 2-oxoglutarate, fumarate, and malate (Figs. 3 and 4). These relatively oxidized intermediate metabolites become potential sinks for photosynthetically produced reductant, leading to the formation of lipids.

There is a clear difference in the allocation of reductant between the nitrogen-stressed WT and NR21. Although NR21 shifts only 4% of its cellular reductant from nitrate toward carbon reductant, we calculate a 70% increase in the fraction of NADPH that is used to reduce carbon to FA (Tables S3–S5) (28). In contrast, nitrogen-stressed cells shift 16% of their cellular reductants from nitrate to carbon reduction and increase the amount of reductant that is used for FA biosynthesis by 160%. This significant increase strongly supports a massive accumulation of cellular reductant; because reductants are primarily synthesized in the plastid, it is highly likely that the retrograde signal alters the pattern of nuclear gene expression associated with nitrogen stress (29). Indeed, the phenotype of the NR21 KD strain clearly indicates that lipid biosynthesis can be uncoupled from protein shuffling (Fig. 5). Analysis of the transcriptome in the NR21 strain grown under nitrogen-replete conditions provides additional evidence that the increased lipid accumulation is not accompanied by up-regulation of the glutamate and 2-oxoglutarate hub gene. We further hypothesize that NR itself is a signal that is involved in the allocation of reductant toward lipids, whereas the plastid controls the shuffling of intercellular nitrogen. In addition, *P. tricornutum* operates a redox-dependent posttranslational modification that can support rapid responses to variable environmental conditions (30). Specifically, it was suggested that the redox regulation of nitrogen assimilation enzymes, including NR, could act as a feedback mechanism; the chloroplast redox state is monitored by those enzymes, and posttranslational modification of the enzyme plays a key role in regulating intermediate metabolism. The combined retrograde and NR feedback signals potentially allow the cell to rapidly respond to changes in nutrient availability in the environment (23). The physiological phenomenon has been observed in several times in shift-up metabolic experiments (31), where diatoms rapidly assimilate and horde inorganic N under a boom and bust strategy. This strategy has allowed diatoms to become extremely successful in highly turbulent environments (32, 33), where survivor cells become the seeds of new populations. However, the underlying mechanisms responsible for the responses have never been explained. Finally, the results presented here suggest that redirecting carbon toward lipid biosynthesis can be achieved genetically by altering the expression of a relatively small number of genes not directly involved with lipid biosynthesis.

**Materials and Methods**

**Cultivation and Experimental Planning.** Cultivation of the *P. tricornutum* strain (accession Pt1 8.6, the Provasoli–Guillard National Center for Culture of Marine Phytoplankton) (34), the NR21 transformant, and nitrogen stress experiments were done following our previously described experiments (11) and as described in SI Materials and Methods. For characterizing the NR21 strain vs. the WT, optically thin cultures were sampled during exponential growth under nitrogen-replete conditions with nitrate as their sole nitrogen source.

**Analytical Methods.** Exponentially growing cultures were analyzed for their IPLs and FAMEs using previously described methods (11, 12, 35–39). The percentage of carbon allocated to each biosynthetic compound was calculated given the following information: 16.5 and 9.4 pg C per cell, 13.6 and 3.6 pg protein per cell, and 7.7 and 5.5 pg carbohydrate per cell for the nitrogen-replete and -stressed cultures, respectively (11, 12); TAG and IPL percentages are calculated based on the data provided in Tables S1 and S2. Total protein determination was performed as previously described (11). The abundance of four selected proteins (PbA, RbcL, NR, and DGAT2D) was determined by Western blots. PSI biophysical characteristics were measured on a custom-built fluorescence induction and relaxation instrument (17, 40). The kinetics of the single-turnover saturating flash were analyzed to obtain the maximum quantum efficiency of photochemistry (Fv/Fm) and the functional absorption cross-section of PSI (PSII). Samples for quantitative PCR were collected, extracted, processed, and analyzed as previously described (11). The NR activity was measured following the method described by Eppley et al. (41) with modifications (3). Additional details are available in SI Materials and Methods.

**RNA-Seq.** Samples for RNA-Seq were harvested and extracted from both the nitrogen-replete and -stressed cultures as described for the quantitative PCR samples. TruSeq RNA (Illumina) was used to prepare mRNA libraries for each of the six samples according to the manufacturer’s instructions. The S0- and S1-ended libraries were multiplexed and sequenced on an Illumina MiSeq platform. The raw reads were trimmed for adapter and low-quality sequences and then aligned to *P. tricornutum*’s version 2.0 set of 10,402 filtered gene models (genome.gji.doe.gov/Phatr2/Phatr2.info.html) using CLC Genomics Workbench (v6.02) (42). Functional metabolic assignment for the different gene models were done according to KEGG database (43), Diatomyc database (www.diatomyc.org), and published literature (44, 45). Additional details on the data analysis can be found in SI Materials and Methods.

**Computational Metabolic Flux Prediction.** Computational prediction of metabolic fluxes was done using an extension of flux balance analysis that infers a metabolic flux distribution from transcriptomic data (46–48). In our analysis, the minimum and maximum reaction rates of the flux balance analysis were set based on the expression level of the genes associated with each reaction. In addition, based on a given limited translational efficiency and a limited accumulation of enzyme over the time, we set the objective function to maximize the correlation between the flux vector and a vector of corresponding gene expression data. Additional details are available in SI Materials and Methods.

**Construction of an Inverted Repeat Vector for Silencing NR, Genetic Transformation, and Screening.** An inverted repeat construct for silencing of the NR gene was generated using standard molecular cloning methods for
RNAi based on the pKS-5h ble-FA plasmid (49). The inverted repeat sequence was designed to match the C terminus of the NADH binding domain (50). The pKS-5h ble nfrA-FA vector was inserted into P. tricornutum using a PDS-1000/He Particle Delivery System (Bio-Rad) as previously described (51). The insertion was verified by PCR of the antibiotic-resistant marker. To select for the best transformants, growth curves were obtained by growing the transformants on 17 mM chlorate (ClKO

Levitan et al. 48(2):185

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www.pnas.org/cgi/doi/10.1073/pnas.1419818112

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Acknowledgments. We thank Angela Falciatore for the gift of the pKS-5h ble-FA plasmid, Andrew Allen for providing a nitrate reductase antibody for our initial work, and Helen Fredricks (Woods Hole Oceanographic Institution) for assistance with lipid analysis. We also thank Benjamin Baillieu (University of Liege) for constructive discussions and Ayye Harel (Rutgers University) for advice on visualizing metabolic fluxes. This research was supported by the US Department of Energy Consortium of Algal Biofuels Commercialization Program, a gift from James G. Gibson (to P.G.F.), the Bennett L. Smith Endowment, and the Rutgers Energy Institute. L.T.G. was supported by a doctoral fellowship from the Portuguese Foundation for Science and Technology (FCT-MCTES) (SFRH/BD/61387/2009).

6 of 6 | www.pnas.org/cgi/doi/10.1073/pnas.1419818112

Levitin et al.