Genomic Foundation of Starch-to-Lipid Switch in Oleaginous Chlorella spp.¹

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The ability to rapidly switch the intracellular energy storage form from starch to lipids is an advantageous trait for microalgae feedstock. To probe this mechanism, we sequenced the 56.8-Mbp genome of Chlorella pyrenoidosa FACHB-9, an industrial production strain for protein, starch, and lipids. The genome exhibits positive selection and gene family expansion in lipid and carbohydrate metabolism and genes related to cell cycle and stress response. Moreover, 10 lipid metabolism genes might be originated from bacteria via horizontal gene transfer. Transcriptomic dynamics tracked via messenger RNA sequencing over six time points during metabolic switch from starch-rich heterotrophy to lipid-rich photoautotrophy revealed that under heterotrophy, genes most strongly expressed were from the tricarboxylic acid cycle, respiratory chain, oxidative phosphorylation, gluconeogenesis, glyoxylate cycle, and amino acid metabolisms, whereas those most down-regulated were from fatty acid and oxidative pentose phosphate metabolism. The shift from heterotrophy into photoautotrophy highlights up-regulation of genes from carbon fixation, photosynthesis, fatty acid biosynthesis, the oxidative pentose phosphate pathway, and starch catabolism, which resulted in a marked redirection of metabolism, where the primary carbon source of glycine is no longer supplied to cell building blocks by the tricarboxylic acid cycle and gluconeogenesis, whereas carbon skeletons from photosynthesis and starch degradation may be directly channeled into fatty acid and protein biosynthesis. By establishing the first genetic transformation in industrial oleaginous C. pyrenoidosa, we further showed that overexpression of an NAD(H) kinase from Arabidopsis (Arabidopsis thaliana) increased cellular lipid content by 110.4%, yet without reducing growth rate. These findings provide a foundation for exploiting the metabolic switch in microalgae for improved photosynthetic production of food and fuels.

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ago (Heckman et al., 2001). They are among the most widely cultivated microalgae by mankind (Klein-Marcuschamer et al., 2013), with an annual net production of about 4,000 tons biomass worldwide. *Chlorella* spp. have served as a popular food supplement and animal feed in Asia, the United States, and Europe for centuries because of their high protein, lipid, and chlorophyll content (the highest amount among known plants; Lee, 1997; Spolaore et al., 2006). However, strain improvement of industrial *Chlorella* spp. has been hindered by the lack of genomics research models. Moreover, although genetic modification of *Chlorella vulgaris* (Chow and Tung, 1997) and *Chlorella ellipsoidea* (Chen et al., 2001) were achieved, none of the *Chlorella* spp. strains shown to be amenable to transformation are of commercial interest (Tran et al., 2013). Thus transformation of industrial production *Chlorella* spp. strains remains a key goal.

One important feature of *Chlorella* spp. is their ability to support high biomass productivity under both autotrophic and heterotrophic culture modes (Gladue and Maxey, 1994; Radmer and Parker, 1994; Lee, 1997; Doucha and Livansky, 2012). Under photoautotrophic conditions, most *Chlorella* spp. produce abundant proteins and lipids, but 1 to 2 weeks of cultivation is usually required from inoculation to harvest. Such a relatively long production cycle often results in low productivity and vulnerability to changes in the weather (Perez-Garcia et al., 2011). However, when cultivated heterotrophically, they reach much higher biomass productivity (>100 g L⁻¹; Doucha and Livansky, 2012) while accumulating significant levels of starch and much lower levels of lipids and proteins (Sansawa and Endo, 2004; Marsalkova et al., 2010; Choix et al., 2012). Remarkably, when cells are shifted from heterotrophic to photoautotrophic conditions, a short duration (1 to 2 d depending on the weather) of illumination induces a prominent change in the intracellular profile, with marked decreases in starch yet 70% to 120% increases in lipids and proteins (Fan et al., 2012a, 2012b; Han et al., 2012; Li et al., 2014b), resulting in both high target product contents (lipids and proteins) and high biomass productivity (Fig. 1; Supplemental Fig. S1). Such a controlled switch that exploits the advantages of both autotrophy and heterotrophy shows potential in large-scale food and biofuel production. Similar switches have been found in many microalgae and have been exploited in cultivation (e.g., via high-light induction or change of nutritional mode) to improve the productivity of valuable cellular components such as astaxanthin, ketocarotenoid, and α-tocopherol (Ogbonna et al., 1999; Hata et al., 2001; Zhang and Lee, 2001). Nevertheless, the molecular mechanism underlying the switch remains unclear.

In this study, we established an oleaginous industrial *Chlorella pyrenoidosa* strain FACHB-9 as a model to investigate the genomic foundation of heterotrophy-to-photoautotrophy metabolism and the associated metabolic switch. We sequenced and annotated the 56.8-Mbp genome sequence of *C. pyrenoidosa* FACHB-9. Furthermore, we employed replicated deep messenger RNA sequencing (mRNA-seq) to track transcriptome dynamics during the transition from heterotrophy to photoautotrophy at six time points along the complete process of the

![Figure 1](https://www.plant.org)
metabolic switch from starch-centric to lipid-centric production, revealing the underpinning molecular machineries and mechanism. Finally, by demonstrating the first genetic transformation of industrial oleaginous Chlorella spp., we showed that introducing a NAD(H) kinase gene into FACHB-9 increased the lipid content by 45.3% to 110.4% but did not affect the growth rate of the host cells under either heterotrophic or photoautotrophic conditions. These findings provide a foundation for exploiting the metabolic switch in microalgae for improved photosynthetic production of food and fuels.

RESULTS AND DISCUSSION

C. pyrenoidosa as a Research Model for Large-Scale Switchable Production of Starch and Lipids

C. pyrenoidosa is one of the most heavily studied species in the Chlorella genus. Due to its rich protein content, relatively fast growth rate, and tolerance of a wide range of environmental conditions, C. pyrenoidosa has been employed as commercial microalgal feedstock of proteins, lipids, and chlorophyll for both heterotrophic fermentation and outdoor mass cultivation (Running et al., 1994; Li et al., 1998; Mason, 2001). Under a standalone heterotrophic culture mode, C. pyrenoidosa strain FACHB-9 exhibited a competitive growth rate (average 2 g L\(^{-1}\) h\(^{-1}\)) and can reach greater than 150 g L\(^{-1}\) dry cell weight (Supplemental Fig. S2), which is comparable to a commercial yeast (Saccharomyces cerevisiae) fermentation system (approximately 130 g L\(^{-1}\); Chen, 1996). In addition, the maximal biomass of the strain reached 4.2 g L\(^{-1}\) after 13-d photoautotrophic growth using heterotrophic cells as seed, with the intracellular neutral lipid content increasing from 3.5% to 23.6% (1.91 and 1.66 times higher in biomass and lipid productivity, respectively, than with photoautotrophic seed; Fig. 1B).

These observations have allowed us to develop a process for C. pyrenoidosa cultivation called sequential heterotrophy-dilution-photoautotrophy (SHDP; Fan et al., 2012b).

SHDP consists of two phases: first, the heterotrophic culture mode for biomass accumulation, and subsequently, the photoautotrophic culture mode for production of target metabolites (Fig. 1C). In the heterotrophic mode, which lasts several days, Glc is gradually depleted while cell density steadily increased, resulting in accumulation of cellular starch to approximately 40% of dry cell weight (Fig. 1, A and C). This is followed by dilution, which is critical to improve light penetration of high-density cultures (e.g., >5 g L\(^{-1}\)) before moving into the photoautotrophic mode. After the dilution, treatments of heterotrophically growing cells with a sustained high light irradiation of 250 μmol m\(^{-2}\) s\(^{-1}\) for 24 h resulted in a marked increases in lipid, protein, and chlorophyll content but marked decreases in starch (Fig. 1C; Supplemental Fig. S1; Fan et al., 2012b). These SHDP-cultivated cells reach a state in which the metabolite profile is as rich in target products (lipid, protein, and chlorophyll) as that of autotrophically grown cells, yet the cultivation cycle is cut from the typical 1 to 2 weeks under the autotrophic mode to 3 to 5 d under SHDP (Fig. 1A). The high productivity demonstrated capability of commercial-scale cultivation and flexibility to switch cellular metabolites suggests C. pyrenoidosa FACHB-9 as an advantageous research model for the switchable production of starch, protein, and lipids.

General Feature of the Genome

The 56.6-Mbp nuclear genome of C. pyrenoidosa FACHB-9 was assembled from a combination of 15× shotgun and 10× paired-end sequence coverage of Roche 454 Titanium (average read length of 400 bp; Table I; Supplemental Table S1). Half of the assembled genome sequences are contained in nine scaffolds, each longer than 1.39 Mbp. The C. pyrenoidosa nuclear genome is slightly larger than Chlorella variabilis NC64A (46.2 Mbp; Blanc et al., 2010) and considerably larger than Chlorella protothecoides (22.9 Mbp; Gao et al., 2014), with C. pyrenoidosa predicted to contain more genes (10,284 as opposed to 9,791 and 7,039 genes, respectively; Supplemental Table S2), which suggests that great changes have taken place in the size of the genome after the diversification in Chlorella genus. This may be due to the different number of unique genes, multicopy genes, and genome duplication /rearrangements among close relatives (Gao et al., 2014).

To produce a high-quality map of gene structures, total RNA samples from both heterotrophic and autotrophic culture conditions were pooled and sequenced on 454 Titanium (533,300 raw reads produced; Supplemental Table S1) and assembled into 13,018 unique complementary DNA (cDNA) isotigs. Furthermore, to detect rare transcripts, 187,503,403 raw reads of mRNA collected from SHDP culture conditions and six time-points were collected (Supplemental Table S3). All of the cDNA reads that passed quality control were used for transcript-based gene prediction. The vast majority (97%) of the predicted genes were supported by these mRNA-seq reads (fragments per kilobase of exon model per million mapped fragments [FPKM] value > 1; Supplemental Data Set S1), and 89% of the predicted proteins were complete models with both start and stop codons (Table I).

The predicted C. pyrenoidosa protein-coding genes are rich in introns, averaging nearly nine introns per gene (the largest in known chlorophytes; Supplemental Table S2) and only 1.1% intron-free genes (Table I). Repeated sequences (such as retroelements, DNA transposons, simple repeats, and low-complexity sequences) made up merely 3.1% of the whole genome (Supplemental Table S4), suggesting a much lower repeat content than Chlamydomonas reinhardtii (Merchant et al., 2007).
**Table I. General features of C. pyrenoidosa FACHB-9 genome**

<table>
<thead>
<tr>
<th>Assembly Features for Nuclear Genome</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genome size</td>
<td>56.8 Mbp</td>
</tr>
<tr>
<td>Sequencing depth</td>
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<tr>
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<tr>
<td>No. of scaffolds (&gt;2 kb)</td>
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<td>Contig N/L50</td>
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<td>Scaffold N/L50</td>
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<td>Guanine and cytosine content</td>
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<tr>
<td>Gene model quality</td>
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<tr>
<td>Multigene genes</td>
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<tr>
<td>Complete models with both start and stop codons</td>
<td>9,228 (89.7%)</td>
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<tr>
<td>Complete models extend both to 5’ and 3’ untranslated regions</td>
<td>9,159 (89.1%)</td>
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<tr>
<td>Gene model support</td>
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<tr>
<td>Models with cDNA support (454 RNA sequencing [RNA-seq])</td>
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<td>Models with homology support (NCBI nr database)</td>
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<td>Models with Pfam domain alignments</td>
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<tr>
<td>Models with Kyoto Encyclopedia of Genes and Genomes (KEGG)/European Commission assignments</td>
<td>7,799 (75.8%)</td>
</tr>
<tr>
<td>Models with GO assignments</td>
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</tr>
</tbody>
</table>

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**Genomic Features of C. pyrenoidosa That Drive Metabolic Versatility**

Green algae have a complex phylogenetic history and constitute a large resource of genetic and metabolic diversity (Falkowski et al., 2004). A whole-genome phylogeny of Chlorophyta species (consistent with 18S rDNA-based phylogeny; Supplemental Fig. S3) was constructed from the 1,685 single-copy orthologous groups identified from seven sequenced green algal genomes (Fig. 2; Supplemental Results S1). Mapping these sets of single-copy orthologs allowed the selective pressure to be traced at every branch of chlorophyte evolution. At the Chlorophyta level, all 52 functional categories of genes are strongly constrained, with median estimates of $\omega$ (the $K_s/K_a$ ratio of nonsynonymous to synonymous nucleotide divergence) much less than 1 (Fig. 3). The most constrained biological processes include photosynthesis, organelle organization, and protein phosphorylation, whereas the least constrained ones are lipid metabolism, phospholipid biosynthesis, glycolysis, and carbohydrate metabolism. When compared at the genus level (C. pyrenoidosa FACHB-9 and C. variabilis NC64A), the most constrained processes are protein folding, the ubiquitin-dependent protein catalytic process, and cellular component organization, and the least constrained are tRNA aminoacylation for protein translation, the biosynthesis process, and photosynthesis (Supplemental Fig. S4). Most of the functional categories, including lipid and carbohydrate metabolisms, are moderately constrained. Between the two Chlorella species, most genes are under strong negative selection ($\omega < 0.1$); however, 51 genes were found under positive selection with an average $\omega$ greater than 1 (Supplemental Table S5). The largest subclasses are oxidation-reduction process, protein binding, and transport-related proteins. The functional and evolutionary implications of such quickly evolving enzymes appear crucial to driving the diverse metabolic capacity of Chlorella spp.

To probe the evolutionary forces behind the expanded starch and lipid metabolic gene repertoire, phylogenetic analysis was employed to investigate the horizontal gene transfer (HGT) events in C. pyrenoidosa genome. Among the 231 genes related to starch metabolism (49 genes), fatty acid biosynthesis (45 genes), glycerolipid biosynthesis (48 genes), lipid activation (47 genes), and lipidase (42 genes), 10 genes (all were related to the lipid metabolism, accounting for 5.5%) might have originated from bacteria via HGT (Supplemental Figs. S5 and S6).

Three genes encoding fatty acyl-CoA synthase (FACS; g11, g937, and g2922) identified were phylogenetically close to bacterial homologs, suggesting their possible evolutionary origin from bacteria via HGT. From the starting point of phototrophic process, transcriptional levels of all of the three genes exhibited significant up-regulation, while transcriptional levels of the other six long-chain (LC)-FACS genes slightly increased or decreased. The synchronous expression pattern of the HGT candidates FACS, along with the increased production of glycerolipids, indicates their important contribution in converting carbon flux from fatty acids into lipid.

Glycerol-3-phosphate acyltransferase (GPAT) genes usually have stringent one to two copy numbers in microalgae, even in heterokonts Nannochloropsis spp., which have highly expanded gene dose in both fatty acid biosynthesis and triacylglycerol (TAG) assembly pathways (Vieier et al., 2012; Wang et al., 2014). However, in Chlorella spp., in addition to the homologous gene inherited in microalgae, another GPAT gene (g2197) was identified and inferred as being originated from bacteria via HGT. This gene was transcriptionally active and up-regulated in the late heterotrophic process, suggesting its potential roles in lipid metabolism.

In addition to the above four HGT candidates, one $\beta$-ketoacyl-acyl-carrier protein (ACP) reductase gene (g7443) in fatty acid biosynthesis pathway, two enoyl-CoA hydratase genes (g4146, g4040) in fatty acid $\beta$-oxidation pathway in mitochondrion, and three lipase genes (g3730, g7283, and g10256) were inferred as HGT candidates. All of the three $\beta$-oxidation-related HGT candidates were dramatically up-regulated in late heterotrophic process and in the first several hours of phototrophic process. Thus, the diverse evolutionary origin of the Chlorella spp. lipid biosynthesis genes has underlain their massive genetic pools and likely contributed to the extraordinary capacity for lipid accumulation.
Genomic Basis of Carbohydrate and Starch Metabolism in C. pyrenoidosa

Polysaccharides, including starch, are one major sink for photosynthetic carbon in many microalgae. The C. pyrenoidosa genome encodes a distinct functional profile in carbohydrate metabolism (as represented by Gene Ontology [GO] terms) from those of C. reinhardtii, Phaeodactylum tricornutum, and Nannochloropsis oceanica (Supplemental Table S6). In C. pyrenoidosa, GO terms in the carbohydrate metabolic process (GO:0005975) are prominently enriched, including the subcategories chitin metabolic process (GO:0006030), polysaccharide metabolic process (GO:0005976), lipopolysaccharide biosynthetic process (GO:0009103), and amino sugar metabolic process (GO:0006040). These findings suggest that carbohydrate metabolic machinery is a prominent feature of the C. pyrenoidosa genome.

Starch is a common form of polysaccharides and the major storage carbohydrate in plants and green algae (Zeeman et al., 2010). In the C. pyrenoidosa genome, for 12 of the 17 nodes along the starch metabolic pathway, multiple genes were found in each step (Supplemental Table S7). Comparison of the gene doses in each step of the starch metabolic pathways among six sequenced green algal genomes demonstrated that C. pyrenoidosa has a highly specific and enriched starch metabolic gene set (Supplemental Table S7).
algal species and strains revealed that *C. pyrenoidosa* harbors an expanded repertoire of genes involved in starch biosynthesis and degradation (Supplemental Table S8), in which selective steps such as ADP-Glc pyrophosphorylase, glucan-water dikinase, and 1,4-β-glucan branching enzyme were highly elevated in gene dose, while isoamylase and starch phosphorylase were less variable and identical in gene dose to other green algae. This suggests that, in green algae, the evolution of starch synthesis pathways is characterized by conservation of pathway backbones and divergence of gene doses in selected pathway nodes.

Genomic Foundation of Lipid Metabolism

To probe the mechanism behind the robust accumulation of neutral lipids and the rapid starch to lipid switch in *C. pyrenoidosa*, we reconstructed the fatty acid biosynthesis, TAG assembly, and lipid activation/degradation pathways (Supplemental Table S9). In *C. pyrenoidosa*, glycerolipids biosynthesis is presumably carried out by two distinct pathways: the prokaryotic pathway (located in the chloroplast), which is widely distributed in prokaryotes, microalgae, and higher plants, and the eukaryotic pathway, which occurs in the endoplasmic reticulum (Shen et al., 2010). Genes encoding both heteromeric (multisubunit; g2736, g8911, and g9519) and homomeric (multifunctional; g6472 and g6473) acetyl-CoA carboxylase (ACCase) were identified. Six putative LC-FACS genes were identified in the genome. Six copies of diacylglycerol acyltransferase (DGAT) genes were discovered in the *C. pyrenoidosa* genome (Supplemental Table S9). Five of the six DGAT genes encode putative type-2 DGATs, whereas the remaining one encodes type-1 DGAT (g7494). Previous research has indicated that there is an alternate pathway for TAG synthesis in yeast, plants (Dahlqvist et al., 2000), and *Chlamydomonas* spp. (Riekhof and Benning, 2009) that involves phospholipid:diacylglycerol acyltransferase (PDAT) to generate TAG using phospholipids as donors. One PDAT gene was identified in *C. pyrenoidosa* (g2349; Supplemental Table S9).

Comparison of GO term profile in *C. pyrenoidosa* with those in *C. reinhardtii, P. tricornutum*, and *N. oceanica* revealed the overrepresentation of the GO terms for the phospholipid metabolic process (Supplemental Table S6). Moreover, the expansion of several other gene families was apparent, which may contribute to the robust biomass and lipid production phenotypes of *C. pyrenoidosa*. These include genes involved in photosynthesis, responses to stress and other organisms (including bacteria and fungi), phosphorus utilization, and the cell cycle (Supplemental Table S6).

**TFs Involved in Starch and Lipid-Related Metabolic Pathways**

Genome-wide identifications of transcriptional factors (TFs) by computational methods were reported for green algae such as *C. reinhardtii* and *Volvox carteri* (Pérez-Rodríguez et al., 2010; Zhang et al., 2011), red
algal species such as *Galdieria sulphuraria* (Pérez-Rodríguez et al., 2010; Zhang et al., 2011), and Eustigmatophyceae strains such as *Nannochloropsis* spp. (Vieier et al., 2012; Hu et al., 2014). Here, we performed such analysis in *Chlorella* spp. via the characteristic domains of plant TFs using the methods of PlantTFDB (Zhang et al., 2011; Jin et al., 2014). The results revealed 195 TFs in *C. pyrenoidosa* (1.9% of the proteome; Supplemental Data Set S2). Each predicted TF was then assigned into a specific TF family based on its DNA-binding domain. MYB, Cys3His zinc finger domain (C3H), and basic Leucine Zipper (bZIP) are the three largest TFs families, together accounting for 36% of all TFs in *C. pyrenoidosa*. Orthologs in *C. pyrenoidosa* genome of higher plant TFs experimentally shown related to sugar signaling (e.g., WRKY, g3291; Sun et al., 2003) and lipid accumulation (e.g. Dof, g1875; Wang et al., 2007) were identified.

Fourteen and eight TFs were found putatively involved in the regulation of starch and lipid metabolism, respectively (Supplemental Data Set S2; Supplemental Methods S1). In starch-related TF genes, eight TF families that include bZIP, C3H, SQUAMOSA promoter binding proteins, MYB, STERILE APETALA, GATA, B3 (the third basic domain in the VIVIPAROUS1), and ARR-B (a Myb-like DNA binding domain called ARRM [type B]) were identified, whereas in lipid-related TFs, seven families that include Calmodulin binding transcription factors, DNA binding with one finger, Heat stress transcription factors, GATA, MYB-related, CONSTATS-like, and Nodule inception-like were found. Notably, the Dof family of TFs, which contains a single C2C2-type zinc-finger-like motif, was shown to regulate in plants processes that include lipid accumulation and photosynthesis (Papi et al., 2002; Wang et al., 2007; Shaw et al., 2009). One of the family members, Gmdof4, was found involved in lipid synthesis in *Glycine max* by activating the ACCase gene (Wang et al., 2007). Moreover, total fatty acid and lipid content were significantly increased in Gmdof4-overexpressing Arabidopsis (*Arabidopsis thaliana*; Wang et al., 2007) and *C. ellipsoidea* (Zhang et al., 2014). These TFs thus represent potential opportunities for engineering the regulatory network for enhanced starch or oil productivity.

**Transcriptome Dynamics during the Switch Process**

To model the temporal process featuring the switch of the intracellular metabolite profile from one dominated by starch to one dominated by proteins and lipids, we changed the *C. pyrenoidosa* cultivation mode from heterotrophy to photoautotrophy at 72 h (Fan et al., 2012a, 2012b; Han et al., 2012), followed by a 48-h light illumination. At each of the six time points during this SHDP process, three biological replicates of separate culture experiments were each sampled for transcriptome analysis via deep mRNA sequencing and metabolite analysis (Fig. 1C).

A total of 187,503,403 trimmed paired-end reads (50 nucleotides in length after trimming, representing 22× average coverage of the genome at each sample; Supplemental Table S3). Our results suggested that 1,578 and 1,491 nuclear genome-encoded genes were respectively up- and down-regulated greater than 3-fold at least one of the time points (Supplemental Fig. S7; Supplemental Data Sets S3 and S4). A set of 75 transcripts was found whose expression levels together were able to distinguish the cellular state among the six time points in a highly reliable manner, suggesting a transcript-based strategy to quantitatively model the SHDP process (Supplemental Fig. S8).

**Overview of the Switch Process**

To determine the time-dependent differential gene expression when cells were transferred from heterotrophic growth to photoautotrophy, the temporal patterns of the relative transcript abundance over the six time points were grouped by k-means clustering, which revealed 16 distinct groups of protein-coding genes (Fig. 4). During the course of the heterotrophic growth phase, the number of differentially expressed genes steadily increased, suggesting distinct cellular transcriptomic states during growth (Supplemental Fig. S7). The number of genes differentially expressed in the phototrophic process at time point 80 h was significantly higher than the other sampling points (Supplemental Fig. S7). This suggests a vigorous gene regulatory response in photoautotrophy, which can partly explain the rapid and massive changes in intracellular carbon storage forms within the first several hours (Fig. 1C; Fan et al., 2012b).

In the heterotrophic growth process, the genes most strongly up-regulated were those from the citrate cycle and amino acid metabolism, whereas those most down-regulated were from fatty acid biosynthesis and metabolism, pentose phosphate metabolism, and nucleotide metabolism (Supplemental Fig. S9). The shift from heterotrophy into photoautotrophy highlights the increased expression of genes involved in carbon fixation and photosynthesis that would be expected owing to light illumination (Fig. 5). Fatty acid biosynthesis, pentose phosphate pathway, and starch catabolism genes were also transcriptionally up-regulated.

**How Was Starch Synthesized?**

In *C. reinhardtii*, starch biosynthesis and degradation are under circadian clock control that appears uncorrelated with illumination (Rai et al., 2006). However, there is evidence for starch turnover in *Chlorella* spp. during growth in response to light intensity or nutritional status (Rosen et al., 1986; Fan et al., 2012b). In our experiments, the starch content of *C. pyrenoidosa* cells showed a comparable decrease between the exponential and stationary phases of heterotrophic growth (48–72 h; Fig. 1C; Sansawa and Endo, 2004; Fan et al., 2012b), indicating a highly active starch catabolism during the SHDP process (Fig. 6). Three transcripts encoding phosphoglucomutase (g7581) and ADP-Glc pyrophosphorylase (g5866 and
g9503), which define the beginning committed steps of starch biosynthesis, were down-regulated by the end of heterotrophy (72 h). Two granule-bound starch synthase I genes (g2429 and g7634) showed continuous up-regulation during the process of light induction (72–96 h). Transcripts responsible for starch breakdown, including glucan-water dikinase (g5653, g5659, and g7110), β-amylase (g3412 and g9517), phosphoglucan water dikinase (g4469 and g8607), and starch phosphorylase (g4743 and g4758), showed remarkable increases both in the heterotrophic (24–72 h) and phototrophic (74–96 h) phases. The enzyme activities of both ADP-glucose pyrophosphorylase and β-amylase showed the consistent trend of starch metabolism (Supplemental Table S10). Thus, the redirection of carbon flux away from storage carbohydrates in dark-grown cells at 72 h suggests that starch serves as a carbon and energy supply when external supplies of organic carbon are depleted after the end of heterotrophy.

**How Was Lipid Synthesized?**

The heterotrophy-to-photoautotrophy change of cultivation mode increased cellular lipid content but reduced starch content in *C. pyrenoidosa* (Fig. 1C) by an unknown mechanism. Glc was exhausted at the end of the heterotrophic process, and sudden illumination was supplied subsequently. Accordingly, genes encoding enzymes associated with fatty acid and neutral lipid metabolic pathways were differentially expressed (Fig. 7). Of the four genes encoding subunits of heteromeric ACCase, two (biotin carboxylase [g8911] and biotin carboxyl carrier protein [g2736]) decreased...
transcriptionally at the end of heterotrophic phase and then recovered quickly following 2 h of light induction. Enzyme activity of ACCCase confirmed its up-regulation during the process (Supplemental Table S10). Similar patterns were also observed in fatty acid synthetase genes, including MALONYL-COA:ACP TRANSACYLASE, BETA KETOACYL-ACP SYNTHASE, BETA KETOACYL-ACP REDUCTASE, BETA-HYDROXYACYL-ACP DEHYDRATASE, and ENOYL-ACP REDUCTASE (Fig. 7; Supplemental Table S9). Thus, de novo fatty acid synthesis is modulated by light/dark conditions, with synthesis increasing in the light and decreasing in the dark (Sasaki et al., 1997). Transferring the dark-grown cells (72 h) to light conditions (80 h) resulted in a slight increase in transcript abundance of acyl-ACP thioesterase (g1189), suggesting increased fatty acid export from the plastid to the endoplasmic reticulum, where TAG assembly occurs. Among the six putative LC-FACS genes identified in C. pyrenoidosa genome data set, two were almost completely suppressed under either or both growth conditions (FPKM < 5; g2654 and g6978; Supplemental Table S9). Another three genes showed a 4.4- to 17.5-fold decrease in mRNA abundance in the dark. Two of the three, g3983 and g9628, were up-regulated under light illumination; the remaining gene (g6510) showed an increase in transcript abundance at the log phase upon heterotrophic growth. These observations suggest distinct roles for individual LC-FACS genes and the presence of an intricate regulatory mechanism.

Transcriptional levels of TAG assembly pathways, including genes encoding glycerol-3-phosphate dehydrogenase (g2897 and g5566), lysophospholipid acyltransferase (LPAAT; g3070 and g9996), phosphatidic acid phosphatase (g6947), and diacylglycerol acyltransferase (type-2 DGAT; g3280 and g7566) showed a synchronized increase by the end of heterotrophic growth (72 h). In Chlamydomonas and Nannochloropsis spp., transcriptome and genetic analysis revealed contribution of acyltransferases to TAG accumulation and their role as potential targets for manipulating TAG hyperaccumulation (Boyle et al., 2012; Sanjaya et al., 2013; Li et al., 2014a). Several carotenoid biosynthesis genes exhibited similar expression patterns (Supplemental Data Set S3); one exception is that a LPAAT gene (g8553), predicted to function in the chloroplast prokaryotic pathway, was down-regulated at 72 h in dark-grown cells and then recovered by light induction. Thus, TAG accumulation, which usually accompanies secondary carotenoid synthesis (Rabban
et al., 1998; Zhekinisheva et al., 2002), might be regulated by the heterotrophy-to-autotrophy switch, thus protecting the chloroplast from potential photooxidative damage. Therefore, the dynamic modulation of eukaryotic and prokaryotic pathways for glycerolipid synthesis, which accompanies the heterotrophy-to-photoautotrophy switch, may be crucial for forming the various intracellular energy storage compounds.

Figure 6. Genomic and transcriptomic features of central carbon metabolism in C. pyrenoidosa. Metabolic steps are represented by arrows. Dashed lines represent multiple metabolic steps. Boxes indicate those nodes where carbon skeletons from amino acid degradation feed into the pathway. Genes encoding the enzymes of these pathways are labeled in red. Up- or down-regulation of mRNA expression under heterotrophic growth (compared with heterotrophy 0 h, leftward arrows) and the heterotrophy to photoautotrophy transition (compared with heterotrophy 72 h, rightward arrows) based on mRNA-seq data are indicated with red upward arrows and green downward arrows, respectively. The full names of the corresponding genes are given in Supplemental Table S9.
Dynamics of the Central Metabolism That Supports

Central metabolism pathways played a prominent role in the metabolic switch of *C. pyrenoidosa* (Fig. 6). Key enzymes from the tricarboxylic acid (TCA) cycle, respiratory chain, and oxidative phosphorylation were transcriptionally up-regulated at the 24-h time point in the dark (Fig. 5; Supplemental Data Set S3). The energy economy of cell growth is dominated by ATP production and the reducing agent NADH via the TCA cycle, which supports various metabolic processes. Consistent with the enrichment of starch granules at the beginning of the heterotrophic phase (Fan et al., 2012b), up-regulation of gluconeogenesis genes was evident within the first 24 h of heterotrophy, which directed carbon from central carbon metabolism to intracellular carbohydrate stores. One Fru-bisphosphate aldolase (g6760) and two glyceraldehyde-3-P dehydrogenase genes (g4391 and g3826) were up-regulated 3- to 16-fold (Supplemental Data Set S3). Although these enzymes can also catalyze the reversible reactions of glycolysis, transcripts of a phosphoenolpyruvate carboxykinase (g2374), one pyruvate phosphate dikinase (g7054), and three Fru 1, 6-bisphosphatase genes (g5024, g4411, and g3655) also increased markedly.

The glyoxylate cycle serves as a shunt of the TCA cycle and allows cells to utilize simple carbon compounds as carbon source when complex sources such as Glc are unavailable (Lorenz and Fink, 2002). This permits the synthesis of Glc from lipids via acetate generated in fatty acid β-oxidation. Transcripts of isocitrate lyases (g1361 and g4926) and malate synthases (g10080), which are specific to the glyoxylate cycle, showed a 3- to 25-fold increase at the end of the heterotrophic culture conditions (72 h). Accordingly, transcripts of acetyl-CoA synthetase (ACS; g7446, g7414, and g4467) increased 7- to 12-fold at the end of the heterotrophic process. ACS catalyzes acetate to produce acetyl-CoA, a precursor in the glyoxylate cycle, TCA cycle, and fatty acid biosynthesis. In contrast to the initial heterotrophic phase, majority of the transcripts for key enzymes of glycolysis, gluconeogenesis, Calvin and TCA cycles, and the pentose phosphate pathway declined at the end of heterotrophic process (i.e., after 72 h, when Glc became limiting), exhibiting a global down-regulation of central carbon metabolism (Supplemental Data Sets S3 and S4).

In addition, oxidative pentose phosphate (OPP) pathway genes, including 6-phosphoglucononate dehydrogenase (decarboxylating; g4585) and 6-phosphogluconolactonase (g3624), were transcriptionally up-regulated approximately 3-fold under light conditions (80 h). The elevated OPP activity resulted in the generation of reducing equivalents (in the form of NADPH) that drive reductive biosynthesis reactions (e.g., fatty acid synthesis); thus, up-regulation of the OPP pathway may eventually contribute to lipid accumulation.
The pyruvate dehydrogenase complex (PDC) subunits (g2537, g9493, g1999, and g1804) were also up-regulated following 2-h light illumination (Supplemental Data Set S4). The PDC transforms pyruvate into acetyl-CoA, which may then be used in fatty acid biosynthesis or as an input to the TCA cycle to carry out cellular respiration. Five fatty acid biosynthesis genes (g4578, g2736, g8879, g8526, and g8911) were rapidly up-regulated upon transferring cells from dark conditions to photoautotrophy (from 72 – 80 h; Supplemental Data Set S4), suggesting that carbon from photosynthesis or glycolysis might be channeled through acetyl-CoA and then into this pathway. Furthermore, changes of enzymatic activities of the key metabolic notes, including those from glycolysis, TCA cycle, gluconeogenesis, and oxidative pentose phosphate pathway, were consistent with the transcriptome analysis (Supplemental Table S10). Collectively, these findings reveal the complex interactions of the central metabolic pathways that jointly support the switch.

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Reboot of Photosynthetic Pathways Promoted the Switch

Down-regulation of photosynthesis is one response to external shocks in microalgae such as nutrient deficiency, stress, or nutritional alteration (González-Ballester et al., 2010; Miller et al., 2010; Xiong et al., 2010; Msanne et al., 2012; Fan et al., 2014a, 2014b; Schmollinger et al., 2014). Most of the photosystem proteins, light-harvesting complexes, and the cytochrome b6f complex were not down-regulated until 72 h in darkness (Supplemental Data Set S3). In contrast to their inert and decreased expression in heterotrophy (24 and 72 h), most of the photosynthesis-related genes showed a fast response, reaching their maximal expression levels at 2 h after photoautotrophy began (74 h; Supplemental Data Set S4), indicating that C. pyrenoidosa is able to rapidly reboot its photosynthetic apparatus to protect from photo damage after the onset of light.

Notably, carbon fixation genes showed a moderate response during heterotrophy (Supplemental Fig. S9; Supplemental Data Set S3). By contrast, a rapid increase in the transcript abundance of predicted chloroplast Calvin cycle genes was observed at the onset of light induction (74 h; Fig. 5; Supplemental Data Set S4). Moreover, up-regulation was observed for C4 dicarboxylic acid cycle enzymes, including NADP-malate dehydrogenase (NADP-MDH; g4904), carbonic anhydrase (g5902), pyruvate phosphate dikinase (PPDK; g7054), and phosphoenolpyruvate carboxykinase (g2374), following the nutritional conversion from heterotrophy to photoautotrophy (74–96 h; Supplemental Data Set S4). NADP-MDH and PPDK are both reported
to be light-modulated enzymes that function in the C4 pathway of photosynthesis (Hatch and Slack, 1969; Carr et al., 1999; Ocheretina et al., 2000). Thus, nimble response of the carbon fixation pathway genes might underpin rapid adaptation of the microalgae to the phototrophic culture.

Genetic Transformation of *C. pyrenoidosa* FACHB-9 Using Electroporation

To demonstrate amenability of industrial production *Chlorella* spp. strains to genetic engineering, electroporation of FACHB-9 was performed without prior enzymatic treatments and selection on 25 mg L\(^{-1}\) G418. A transformation rate of about 1.07 \(\times\) \(10^{-5}\) per microgram plasmid DNA was achieved (Supplemental Methods S1; Supplemental Table S11). The efficiency of the transformations was strongly affected by the promoter and field strength (Supplemental Table S11), as the highest number of colonies was generated using 3,300 V cm\(^{-1}\) field strength, and the promoter producing the highest number of transformants was *Heat shock protein70* (*Hsp70*; followed by *Tubulin*; Supplemental Table S11). Both genomic PCR and Southern blot confirmed presence and integration of the transgenes in selected colonies (Fig. 8, B and C). Fluorescence microscopy observation analysis confirmed successful stable expression of the exogenous GFP gene in resistant *C. pyrenoidosa* lines (Fig. 8, D and E). Stable integration and expression of the exogenous gene were confirmed after five generations of growth via antibiotic selection in both agar plates and liquid media.
Overexpression of the AtNADK3 Significantly Enhances the Fatty Acids and Lipid Content in C. pyrenoidosa FACHB-9

Our transcriptome analysis indicated the switch from heterotrophy to light environment might lead to generation of reducing energy equivalents (e.g., in the form of NADPH) that drive reductive biosynthesis reactions (e.g., fatty acid synthesis). Thus, increasing the level of the intracellular reducing equivalents may eventually contribute to lipid accumulation. In Arabidopsis, NAD(H) kinase3, a key source of the cellular reductant NADPH, is required for various abiotic stress responses (Chai et al., 2006). The corresponding gene AtNADK3 encodes a NADH kinase that can synthesize NADPH with NADH as the preferred substrate and also utilizes NAD+ (while the other two AtNADK genes can utilize only singular substrate; Chai et al., 2006). Intriguingly, in C. pyrenoidosa FACHB-9 genome, only one endogenous copy of NADK (g6804) was found. Transcriptome analysis showed that the NADK3 native to C. pyrenoidosa was up-regulated by 360% and 41.6% in the heterotrophic and light induction process (Supplemental Data Set S1), suggesting an active role of this transcript in lipid production. To test whether higher dose of this enzyme could lead to NADPH accumulation and thus promote fatty acid synthesis, we introduced into FACHB-9 one more copy of AtNADK3 using the genetic transformation system we developed (Supplemental Methods S1).

Growth curves of the transgenic AtNADK3 strains and wild-type C. pyrenoidosa were similar at either heterotrophy or photoautotrophy (Fig. 9, A and B), suggesting absence of obvious deleterious effects on growth as introduced by the transformation. Because Hsp70 promoter, as a transcriptional enhancer, promoted transgene expression in C. reinhardtii under inducing conditions (Rosales-Mendoza et al., 2012), we compared three culture conditions (heterotrophy, photoinduction, and heat shock) to probe whether the native Hsp70 promoter works as constitutive or inducible type in C. pyrenoidosa. All the three conditions induced AtNADK3 expression, yet with distinct effects (Fig. 9C; Supplemental Fig. S10). Compared with the wild type, all three transgenic strains exhibited significantly increased NADPH and lipid content, by 39.3% to 79.9% (Fig. 9D) and 45.3% to 110.4% (Fig. 9E), respectively. Main fatty acid components of the transgenic strain NADK3-2 and the wild type were similar, as revealed by gas chromatography-mass spectrometry analysis (Fig. 9F). Both light induction and heat shock increased the cellular content of the fatty acids (Fig. 9F) compared with the wild type, yet the relative compositions of total fatty acids were not significantly different from that in the wild type (Fig. 9F). Among the three investigated induction strategies, light was the most effective way (followed by heat shock and then heterotrophy) to induce both gene expression and NADPH accumulation as well as lipid biosynthesis (Supplemental Figs. S10 and S11).

Overexpression of NADK causes perturbation of NADP(H) pool and has positive effects on growth or stress tolerance (Pollak et al., 2007; Panagiotou et al., 2009), e.g., stimulation of photosynthesis metabolism and tolerance of oxidative damages in higher plants (Takahashi et al., 2009; Takahara et al., 2010). Our study is, to our knowledge, the first to show that the NADP(H) pool can be engineered to increase lipid content, by introducing a single NAD(H) kinase gene in microalgae. Thus, these findings suggest a new strategy for strain development for higher lipid productivity in not just industrial microalgae but also higher plants.

CONCLUSION

The green algae Chlorella spp. are among the most widely cultivated microalgae in the world (Klein-Marcuschamer et al., 2013). Genome of the industrial strain C. pyrenoidosa FACHB-9 revealed positive selection and gene family expansion in lipid and carbohydrate metabolism genes, as well as those related to the cell cycle and response to stimuli. HGTS were inferred in several lipid metabolism nodes. Temporal analysis of transcriptomes during the metabolic switch from a starch-rich heterotrophy mode to a protein- and lipid-rich photoautotrophy mode revealed a marked redirection of metabolism in which the primary carbon source, Glc, is no longer supplied to cell building blocks by the TCA cycle and gluconeogenesis, while carbon skeletons from photosynthesis and starch degradation may be channeled directly into fatty acid and protein biosynthesis. By establishing, to our knowledge, the first genetic transformation for industrial Chlorella spp., we show that overexpression of the AtNADK3 significantly enhances the fatty acids and lipid content without affecting its growth. This is, to our knowledge, the first report of increasing microagal fatty acid and lipid content by genetically engineering cellular-reducing equivalents. These findings thus serve as a foundation for engineering the metabolic switch apparatus in this and related algae for optimized and controllable production of food and fuels.

MATERIALS AND METHODS

For a detailed description of all methods, see Supplemental Methods S1.

Strain and Culture Conditions

Chlorella pyrenoidosa FACHB-9 (Freshwater Algae Culture Collection of the Institute of Hydrobiology, Chinese Academy of Sciences) was cultured heterotrophically in Endo medium (Endo et al., 1974) in flasks or in 50- and 500-L fermenters with Glc and urea as the organic carbon and nitrogen source. F-Si medium (Sato et al., 2006) was chosen for photoautotrophic culture lipid production with 2% (v/v) CO2/air supplementation. For SHDP cultivation, the heterotrophically grown cells (in which Glc was already depleted) were resuspended in water at a final cell concentration of about 2 g L-1 and then transferred to 5-d phototrophic cultivation under indoor continuous light at approximately 300 μmol m-2 s-1 at 30°C. For each of
cultivation processes above, samples were taken at specific intervals to determine biomass and cellular component production.

**Genome Sequencing and Assembly**

For *C. pyrenoidosa* FACHB-9 genome sequencing, we employed Roche 454 Titanium to collect single-end and paired-end reads. We first generated a total of 3,730,590 454-Titanium sequence reads (average read length, approximately 400 bp, with pair-end distances of 3, 8, and 20 kb, respectively). Assembly of the single- and pair-ended 454 sequences (1.46 Gbp in total and approximately 26-fold coverage) with Newbler version 2.0 (Roche) resulted in 8,193 contigs that assembled into 1,345 scaffolds longer than 2 kb. The assembled scaffolds were further screened and filtered by searching against all bacterial sequences from the SILVA (Pruesse et al., 2007) and National Center for Biotechnology Information (NCBI) nr database to filter out possible contaminants. The resultant assembly of the *C. pyrenoidosa* nuclear genome contains 1,336 scaffolds totaling 56.8 Mbp in size, with a contig N/L50 of 1,265 kb and a scaffold N/L50 of 9.13 Mbp.

Algal cells grown under different heterotrophic and phototrophic conditions (±Glc, low-/high-light illumination, and logarithmic/stationary phase) were collected and harvested by centrifugation. Each sample was then pooled to prepare libraries of cDNA for mRNA sequencing on 454 Titanium (Roche). A total of 533,300 raw reads were produced using GS-Titanium. All raw reads were trimmed based on quality value before further analysis. All cDNA reads that passed quality control were used for transcript-based gene prediction. About 81% (8,317 genes) of the total number of predicted protein-coding genes were covered by the 454 cDNA sequencing reads.

**Genome Annotation and Analyses**

The assembled draft nuclear genome of *C. pyrenoidosa* was annotated using the QIBEBT-FUNGEA annotation pipeline (accessibility), which combines three different ab initio gene predictors (Augustus version 2.5 [Stanke and Morgenstern, 2005], GeneID v1.2 [Parra et al., 2000], and SNAP [Korf, 2004]) and one evidence-based gene predictor (Cufflinks version 1.1.0 [Trapnell et al., 2010]). The cDNA reads that passed quality control were also used for transcript-based gene prediction. Functional annotation of the predicted protein-coding genes was carried out based on the identification of protein domains using InterProScan (Quevillon et al., 2005) database and BLASTp matches against the NCBI nr, Kyoto Encyclopedia of Genes and Genomes (KEGG) and Conserved Domain Database (CDD) databases, with an E-value < 1 × 10−5. Protein sequences from *C. pyrenoidosa* were used as BLASTp queries against the sequenced alga species, and hits with an E-value < 1 × 10−10 were considered homologous proteins.

For selection pressure analysis of protein-coding genes in green algae, orthologs were used as BLASTp queries against the sequenced alga species, and hits with an E-value < 1 × 10−10 were considered homologous proteins. For selection pressure analysis of protein-coding genes in green algae, orthologs were used as BLASTp queries against the sequenced alga species, and hits with an E-value < 1 × 10−10 were considered homologous proteins.

**RNA-Seq and Differential Expression Analysis**

Total RNA was extracted from *C. pyrenoidosa* cells during the SHDP process. Briefly, the SHDP cultures were performed in 500-mL shaker flasks. Algal cells were first cultivated heterotrophically to achieve high cell density. After Glc was completely consumed, the broth was diluted to approximately 2 g L−1 dry biomass and transferred to a light environment for photosynthetic culture. Three separate flasks (i.e., three biological replicates) were analyzed in parallel (Supplemental Methods S1). As a result, 18 samples in total, with the three biological replicates collected from each of the six time points (0, 24, and 72 h from heterotrophic cells and 2, 8, and 24 h from phototrophic cells; Fig. 1C), were used for mRNA-seq library preparation and then submitted to Sorexia GA-IIX (Illumina) for sequencing (short insert [approximately 400 bp] paired-end, 2 × 100 and 2 × 90 bp). Default parameters were used to screen reads using the Illumina quality control pipeline. The processed sequence files (a total of 187,503,403 reads) were then mapped to the assembled genome by TopHat version 2.0.14 (Trapnell et al., 2009) with a tolerance of up to two mismatches and three indels. For each of the RNA-seq data sets, the transcript relative abundance (TRA) was expressed as the number of aligned reads to annotated gene models using Cufflinks (version 1.1.0; Trapnell et al., 2010) and then normalized to FPKM values. Fold expression changes between different time points were calculated using the TRA log2 ratio. Genes were regarded as differentially expressed if they had at least a 3-fold change and 5% or less false discovery rate. Furthermore, the gene sets with positive TRA log2 ratios that had numerators greater than 50 FPKM were regarded as up-regulated. Down-regulated genes were those with negative TRA log2 ratios that had denominators greater than 50 FPKM.

**Nuclear Transformation by Electroporation**

FACHB-9 cells were cultured up to logarithmic phase, harvested following centrifugation at 3,000 rpm for 10 min, and then processed with a high-exudate buffer containing 0.2 M sorbitol and mannitol on ice. The processed cells were recenterificed at 3,000 rpm and resuspended with a buffer at a concentration of approximately 5 × 10^7 cells mL−1 and then immediately mixed with a final concentration of 30 µg mL−1 plasmid pGreen0029-Uri-eGFP-Nos (freely available by contacting Jianhua Fan), a final concentration of 10 µg mL−1 plasmid pG418, and 25 µg mL−1 salmon sperm DNA. The mixture was placed on ice for 10 min before a 100-µL mixture was supplied into the 2-mm electroporation cuvette for transformation with a Bio-Rad Gene Pulser II Electroporation System. The cells were transformed at a field strength of 1.8 to 4.3 kV cm−1 leading to time constants of 1 to 5 ms. Following electroporation, cells were screened via selective medium (in agar plates with 25 mg L−1 G418; Nichols, 1973) and validated.

**Construction, Validation, and Phenotyping of ATNADK3-Overexpression Strains of C. pyrenoidosa**

To test whether higher dose of this enzyme could lead to NADPH accumulation and thus promote fatty acid synthesis, we introduced into FACHB-9 one more copy of ATNADK3 using the genetic transformation system we developed. The cDNA of ATNADK3 was amplified from the leaves of Arabidopsis (Arabidopsis thaliana) according to the published ATNADK3 cDNA sequence (NCBI accession no. NM_106506), and the fragments were then inserted into the site between Spel and NotI in the plastid pGreen0029-Uri-eGFP-Nos to replace eGFP. Similarly, the promoter Ubiquitini gene was also replaced with endogenous promoter with HiiiD and BpiII to create the pGreen0029-Hsp70-ATNADK3-Nos.

Fatty acid composition of mutant cells was quantitatively determined using a gas chromatography mass spectrometer. For gene expression, the first-strand cDNA synthesis and real-time quantitative PCR were performed using the ReverTra Ace qPCR RT Master Mix with gDNA Remover and SYBR Green Realtime PCR Master Mix (Toyobo), respectively. A pair of specific primers (forward, ATCCGATCCAAATCTGGAT, and reverse, TGGCACAACAAATGTCATCCA) were designed and used for expression-level quantitative detection. The comparative threshold cycle method was used to analyze gene expression relative to control based on the average fold change. The content of NADPH was determined using the coenzyme II NADP (H) content kit (SuZhou Comin).

The *C. pyrenoidosa* whole-genome sequencing project (PRINAI171991) has been deposited at DDBJ/EMBL/GenBank under the accession number ANZC00000000. The version described in this paper is the first version, ANZC01000000. The mRNA-seq raw data are available at the Gene Expression Omnibus database (http://www.ncbi.nlm.nih.gov/geo/) under the accession number GSE40028. The genome and transcriptome data can also be freely accessed through the project Web site (http://www.bioenergychina.org/index/Chlorella_pyrenoidosa_index.html).

**Supplemental Data**

The following supplemental materials are available.

**Supplemental Figure S1.** Confocal microscopy images of Nile red-stained *C. pyrenoidosa* cells during SHDP process and nitrogen deprivation under phototrophic conditions.

**Supplemental Figure S2.** Scale-up the *C. pyrenoidosa* heterotrophic growth in 50 to 500-L fermentors by repeated fed-batch cultivation for seeds supply.

**Supplemental Figure S3.** Phylogenetic position of *C. pyrenoidosa* among algae, plants, and streptomycetes.

**Supplemental Figure S4.** Selection pressure on protein-coding sequences in two *Chlorella* spp.
Supplemental Figure S5. Phylogenetic analysis of lipid synthesis genes that were inferred to have originated from HGT using the neighbor-joining method.

Supplemental Figure S6. Phylogenetic analysis of lipid synthesis genes that were inferred to have originated from HGT, using the maximal-likelihood method.

Supplemental Figure S7. Number of differentially expressed genes identified at each time point for both heterotrophic growth and phototrophic process.

Supplemental Figure S8. Transcriptomic signature analysis of the SHDP process.

Supplemental Figure S9. Overview of metabolic pathways and regulation under heterotrophic growth in C. pyrenoidosa.

Supplemental Figure S10. Expression of AtNADK3 into C. pyrenoidosa cells under different culture conditions.

Supplemental Figure S11. Analyses of ultrastructure for C. pyrenoidosa transformant NADK3-2 expressing AtNADK3 under different culture conditions.

Supplemental Figure S12. Guanine-cytosine content distribution of raw reads from the two 454 sequenced libraries for genome assembly.

Supplemental Figure S13. Saturation curves of transcriptome sequencing reads.

Supplemental Figure S14. Spearman correlation upon biological replicate samples.

Supplemental Figure S15. Comparison of fold difference in RNA abundance during SHDP process estimated from mRNA-seq versus real-time PCR experiments.

Supplemental Figure S16. Quality of reads after filtering via FastQC.

Supplemental Table S1. The genome sequencing data for C. pyrenoidosa using Roche 454.

Supplemental Table S2. Comparison of C. pyrenoidosa genome statistics with those of sequenced chlorophytes genomes.

Supplemental Table S3. Details of the transcriptome sequencing data for C. pyrenoidosa using an Illumina solexa platform.

Supplemental Table S4. Repeat sequences in the C. pyrenoidosa genome.

Supplemental Table S5. Annotation of single-copy orthologs between two Chlorella spp. with an average Ks/Ka value greater than one.

Supplemental Table S6. Selected GO term enrichment and depletion of C. pyrenoidosa proteins compared with other alga.

Supplemental Table S7. Starch metabolic pathway genes in C. pyrenoidosa.

Supplemental Table S8. Starch metabolic pathway gene comparison among green algae.

Supplemental Table S9. Fatty acid and lipid metabolic pathway genes in C. pyrenoidosa.

Supplemental Table S10. Enzyme activities in C. pyrenoidosa grown under the sequential heterotrophy-dilution-photautotrophy model.

Supplemental Table S11. Transformation rates achieved by electroporation of C. pyrenoidosa FACHB-9 cells in the presence of green0029 plasmid with different voltage field strengths and promoters.

Supplemental Data Set S1. BLASTp annotation of C. pyrenoidosa nuclear genome-encoded gene in NCBI nr database.

Supplemental Data Set S2. TEs involved in starch- and lipid-related metabolic pathways.

Supplemental Data Set S3. Expression levels of differentially expressed genes grouped by putative functions under heterotrophic process.

Supplemental Data Set S4. Expression levels of differentially expressed genes grouped by putative functions under photo-induced process.

Supplemental Methods S1. Details for the methods used in this article.

Supplemental Results S1. A more detailed description of the results aside from the main text.
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