Transcriptome analysis reveals the genetic foundation for the dynamics of starch and lipid production in *Ettlia oleoabundans*

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\textbf{A B S T R A C T}

The oleaginous microalga *Ettlia oleoabundans* accumulates both starch and lipids to high levels under stress conditions such as nitrogen starvation (N−). To steer biosynthesis towards starch or lipids only, it is important to understand the regulatory mechanisms involved. Here physiological and transcriptional changes under nitrogen starvation were analysed in controlled flat-panel photobioreactors at both short and long time-scales. Starch accumulation was transient and occurred rapidly within 24 h upon starvation, while lipid accumulation was gradual and reached a maximum after 4 days. The major fraction of accumulated lipids was composed of de novo synthesized neutral lipids - triacylglycerides (TAG) - and was characterized by a decreased composition of the polyunsaturated fatty acids (PUFAs) C18:3 and C16:3 and an increased composition of the mono-unsaturated (MUFA)s and saturated (SFA)s fatty acids C18:1/C16:1 and C18:0/C16:0, respectively. RNA-sequencing revealed that starch biosynthesis and degradation genes show different expression dynamics from lipid biosynthesis ones. An immediate rapid increase in starch synthetic transcripts was followed by an increase in starch degrading transcripts and a decrease in the starch synthetic ones. In contrast, increased gene expression for fatty acid and TAG synthesis was initiated later and occurred more gradually. Expression of several fatty acid desaturase (FAD) genes was decreased upon starvation, which corresponds to the observed changes to higher levels of MUFAs and SFAs. Moreover, several homologs of transcription regulators that were implicated in controlling starch and lipid metabolism in other microalgae showed differential gene expression and might be key regulators of starch and lipid metabolism in *E. oleoabundans* as well. Our data provide insights into the genetic foundation of starch and lipid metabolism in *E. oleoabundans* under nitrogen starvation and should facilitate metabolic engineering towards tailored strains with desired storage compound composition.

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multiple genes or complete pathways [8]. The selected gene targets for metabolic engineering have been identified by transcriptomics or were based on detailed biochemical and genetic information from other microalgal or plant models. Most of these studies focused on microalgae that accumulate only lipids or starch under stress conditions such as nitrogen or phosphorus starvation, while only a few have focused on transcriptomics and metabolic engineering of so-called “hybrid” producers: oleaginous microalgae that are capable of simultaneous accumulation of starch and neutral lipids such as triacylglycerides (TAG). The oleaginous microalgae *Ettlia oleoabundans* and *Acutodesmus obliquus* and several *Chlorella* species are known hybrid producers of industrial relevance that accumulate high levels of both starch and lipids under nitrogen starvation [9–11]. To achieve economically viable production of these hybrid producers multiple-product biorefineries and cultivation optimization are important factors, but for many industrial applications it is also desired that carbon-partitioning can be altered to produce mainly TAG or starch and to genetically control the switch between starch and lipid metabolism. The genetic foundation for this dual accumulation strategy however is unknown in most of these microalgae, due to a lack of well-annotated genome sequences and a limited number of transcriptome studies. It is also plausible that the regulatory mechanisms for storage compound accumulation in hybrid producers are more complex than in microalgae mainly accumulating a single storage compound. *E. oleoabundans* is a hybrid producer with several properties that make it a very suitable candidate for industrial production. It can accumulate up to 56% of its cell dry weight as lipids under nitrogen starvation, mainly in the form of TAG, and while isolated as a freshwater strain it can also grow and accumulate storage compounds under saline conditions and high pH as well [12–15]. Furthermore, continuous TAG production in continuously growing cells has been reported [16]. However, the genomic foundation for these favourable industrial traits of *E. oleoabundans* currently is not yet established. The few studies that addressed the transcriptional changes upon nitrogen starvation in this species, have analysed either a single timepoint at an early-stage (< 24 h) [15] or a late-stage (11 days) during batch nitrogen starvation [17] or investigated transcriptomes during nitrogen limitation under turbidostat cultivation [18]. For a full understanding of the cellular and transcriptional changes upon N-starvation it is necessary to track the temporal dynamics of metabolite concentrations, physiological parameters and transcript abundances over both a short and long timespan, to identify key regulatory genes controlling metabolic switch-points. Some recent studies in e.g. *Chlamydomonas reinhardtii*, *Chlorella* spp., *Nannochloropsis* spp. and *Monoraphidium neglectum* have performed this correlation analysis of storage compound accumulation and transcriptome patterns, which indicated possible transcription factors and metabolic nodes that are involved in or even control the “switch” between starch and lipid metabolism [19–24]. In addition, integration of transcriptomics with genome-scale metabolic models could aid in pinpointing the critical metabolic nodes that should be targeted in metabolic engineering.

In this study, we therefore set out to perform an in-depth analysis of the temporal dynamics of storage compound accumulation and transcriptome changes during nitrogen starvation in *E. oleoabundans*, with the aim to identify the metabolic genes and transcriptional factors that are involved in the switch between starch and lipid metabolism in this microalgae. Our results revealed the differences in temporal dynamics of gene expression for the starch and lipid pathways upon nitrogen starvation, and identified transcriptional regulators that might be involved. These insights are valuable for rational metabolic engineering in this important industrial microalgae.

2. Materials & methods

2.1. Culture conditions and nitrogen starvation experiment

*Ettlia oleoabundans* UTEX 1185 (culture collection of Algae, University of Texas, Austin) was grown in Bold’s Basal Medium (BBM) at pH 7.5 and 25 °C, with 25.2 mM KNO₃ (N-replete conditions) or without nitrate (N-starvation). Culture purity during experiments was monitored by microscopical observation and analysis on a Multisizer™ 3 Coulter Counter® (Beckman Coulter). Experiments were performed in flat panel air-lift photobioreactors (Labfors 5 Lux, Infors HT, Switzerland) with a working volume of 1.7 L, at pH 7.5 and 25 °C, and sparged with 2% CO₂ at an air flow of 1.2 L min⁻¹ and photon flux density (PFD) of 800 μmol s⁻¹ m⁻² at a 12:12 light:dark cycle.

The pH was maintained at pH 7.5 by the on-demand supply of 0.1 M HCl. A nitrogen-replete batch pre-culture of *E. oleoabundans* (250 mL) in exponential growth phase was used to inoculate all three photobioreactors, and microalgae were further batch cultivated until a biomass concentration of 2 g cell dry weight (CDW)/L was reached. Subsequently, continuous nitrogen-replete cultivation (N⁺) was continued for several days until a stable biomass concentration was maintained. For the nitrogen starvation experiment biomass from the nitrogen-replete continuous culture was collected, spun down, rinsed and resuspended in nitrogen-free BBM medium. The reactors were rinsed with distilled water, filled with nitrogen-free BBM medium and inoculated with the same volume of the collected microalgae. During the nitrogen-depleted phase (N⁻), the reactors were run in non-steady state continuous mode. Medium was fed continuously at 800 mL/d to compensate for the volume-loss due to sampling. Samples for analysis of biomass and cell parameters from the nitrogen-replete continuous phase (N⁺) were collected at four timepoints in the light-phase preceding nitrogen starvation (−25, −23, −19 and −15 h; Fig. 1). Samples for biomass, cell parameter and molecular analysis from the onset of nitrogen starvation (N⁻) were collected at time-points at the start, middle and end of each light-phase and at the end of each dark-phase over a 4-day period (0, 2, 6, 10, 23, 29, 33, 46, 53, 71, 77 and 99 h; Fig. 1). Samples were centrifuged, and cell pellets snap frozen in liquid nitrogen and stored at −80 °C. The experiments were performed as three biological replicates.

2.2. Biomass analysis

2.2.1. Cell dry weight, cell count and cell parameters

Cell dry weight (CDW) concentrations were determined as described by Kliphuis et al. [25], by filtering culture broth (around 10 mg of biomass) through pre-dried (100°C overnight) and pre-weight Whatman glass fibre filter paper (GF/F; Whatman International Ltd., Maidstone, UK). The filter was washed twice with filtered demineralized water to remove adhering inorganic salts and trace elements and subsequently dried overnight at 100°C before weighing. Cell numbers were determined using a Multisizer™ 3 Coulter Counter® (Beckman Coulter). Quantum yield (QY) was determined using an Aquapen-C (Photon Systems Instruments, Czech Republic).

2.2.2. Total fatty acids

Extraction and quantification of total fatty acids (TFA) were adapted from Breuer et al. [26]. Around 20 mg of pellet was transferred to bead beating tubes (Lysing Matrix E; MP Biomedicals, Santa Ana, CA, USA) and lyophilized overnight. Freeze-dried cells were disrupted by a bead beating step in a Precellys® 24 bead beater (Bertin Technologies) for 2× (3×60 s with 120 s pause in between at 2500 rpm), followed by 1× (2×60 s with 120 s pause in between at 2500 rpm) in the presence of a chlorormform:methanol mixture (1:1.25) to extract the lipids from the biomass. The internal standards C15:0 (tripentadecanoin - T4257; Sigma-Aldrich, St Louis, MO, USA) and C19:0 (trinonadecanoin - T4632; Sigma-Aldrich, St Louis, MO, USA) were added to the extraction mixture to enable fatty acid quantification. Methylation of the fatty acids to fatty acid methyl esters (FAMEs) and the quantification of the FAMEs by GC/MS analysis were performed as described by Breuer et al. [26]. TFA concentration was calculated as the sum of triacylglycerides (TAG) and polar lipids (PL).
2.2.3. Triacylglycerides and polar lipids

Quantification of TAG was similar to the TFA analysis method with some modifications. After TFA extraction, the chloroform-methanol mixture was evaporated under N₂ gas and the TFA fraction dissolved in 1 mL hexane and separated based on polarity using a Sep-Pak Vac silica cartridge (6 cm³, 1000 mg; Waters, Milford, MA, USA) equilibrated with hexane. The neutral TAG fraction was eluted with 5 column volumes (10 mL) of hexane-diethyl ether (7:1 v/v). Subsequently the polar lipid fraction containing glycolipids and phospholipids was eluted from the silica cartridge with 5 column volumes (10 mL) of hexane:acetone:water (2:2:1 v/v/v). All solvents were then evaporated from the fractions under N₂ gas. TAG and PL fractions were then methylated and analysed as described in the TFA analysis section.

2.2.4. Total carbohydrates

Total carbohydrates were extracted and analysed by a colorimetric assay according to DuBois et al. [27]. A biomass dry-weight of ~20 mg was hydrolysed in 5 mL 2.5 M HCl in a heat block at 100 °C for 3 h. Samples were neutralized using 5 mL of 2.5 M NaOH. Then 0.5 mL of 5% w/w phenol and 2.5 mL of concentrated sulfuric acid was added to 0.5 mL of hydrolysed sample. The samples were incubated at 35 °C for 30 min before reading of the absorbance at 485 nm against a blank of 0.5 mL 5% w/w phenol, 2.5 mL concentrated sulfuric acid and 0.5 mL of deionized water. Glucose was used as a standard.

2.2.5. Starch

Starch content was analysed by enzymatic degradation of starch to glucose using the thermostable α-amylase and amyloglucosidase enzymes from the Total Starch Assay Kit (AA/AMG) (Megazyme International, Ireland) using the protocol described by de Winter et al. [28]. Around 10 mg of freeze-dried cells were disrupted by bead beating in bead beating tubes (Lysing Matrix E; MP Biomedicals, Santa Ana, CA, USA) for 1 × 10 s at 5500 rpm in a Precellys® 24 bead beater (Bertin Technologies), in the presence of RLT buffer with β-mercaptoethanol (Qiagen). Subsequently, tubes were placed on ice and spun down for 2 min at 13,000 × g. RNA was further cleaned using the RNasey Plant Mini Kit (Qiagen). In short, supernatant was transferred to a Qiashredder column, and columns centrifuged for 2 min at 13,000 × g. Supernatant was transferred to a 2 mL Eppendorf vial, 0.5 V
of 96% ethanol was added and mixed. The mixture was then processed on Qiagen RNeasy columns, including on-column DNaseI treatment, according to the manufacturers’ instructions. Further preparation of RNA and RNA sequencing was performed by BaseClear (Leiden, The Netherlands).

RNA quality control and quantification were performed using a BioAnalyzer and an Illumina TruSeq strand-specific mRNA library was prepared (average mRNA library size of 345 bp) for the construction of the reference transcriptome using 125 bp paired-end sequencing (PE 125) from a pool of all RNA samples (N-replete and N-deplete). Individual sample libraries were prepared by 50 bp single-end read sequencing (SR 50). Paired-end and single-end sequencing reads were generated using the Illumina HiSeq2500 instrument. FASTQ sequencing files were generated using the Illumina CASAVA software and internally quality filtered based on Illumina’s chastity filtering procedure, then PF reads (raw reads) were used for further analysis. Time series data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus [29] and are accessible through GEO Series accession number GSE104807 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE104807).

2.4. De-novo transcriptome assembly and functional annotation

RNA-seq raw reads were quality controlled using FASTQC v.0.11.3 and Trimmomatic [30] (automatically invoked by Trinity software [31]). The reference transcriptome was assembled with Trinity v.2.0.6 using the following parameters: -seqType fq –SS_lib_type FR –trimmomatic –quality_trimming_params SLIDINGWINDOW:20:25 MINLEN:80 and have been deposited at DDBJ/EMBL/GenBank under the accession GFWX00000000.

Trinitrate v.2.0.2 (https://trinitrate.github.io) was used for transcriptome annotation. Transcripts and their predicted ORFs were queried with BLASTx and BLASTp against both the SwissProt [32] and UniRef90 [33] databases and only the top-scoring hit was retained. HMMER [34] was used to search for conserved protein domains in predicted ORFs against the pfam-A database [35]. BLAST homologs and Pfam domain entries were loaded into the pre-formatted Trinotate SQLite database which contained UniProt-associated annotation information. The subcellular localizations of each protein were predicted using a standalone version of TargetP v1.1 [36] in plant mode and without cut-offs (winner takes it all).

2.5. Transcript abundance and differential expression analysis

Scripts bundled with Trinity software were mainly adopted to quantify the transcripts and find the differentially expressed subset. Transcript-level rather than unigene-level expression was investigated to retain isoform-specific information. Reads from individual libraries were aligned to the reference transcriptome with bowtie v.1.1.2 [37] and quantified using RSEM v.1.2.19 [38]. A table of TMM-normalized FPKM expression matrix and a separate table of raw fragment counts were generated for further analysis and visualization. Differentially expressed (DE) transcripts were identified from raw counts with the Bioconductor package EdgeR v.3.1.3 [39]. Three biological replicates for each condition were provided. The most significant differentially expressed transcripts (FDR < 0.001 and FC > 4) were extracted for further analysis. A hierarchically clustered heatmap was generated from the Pearson correlation matrix of pairwise sample comparisons based on the most significant DE subset. The most significant DE subset was also partitioned into expression clusters by cutting hierarchical clustering trees of transcripts using MeV v.4.9.0 (https://sourceforge.net/projects/mev-tm4/).

2.6. Gene ontology analysis

GO terms were extracted from transcriptome annotation using scripts from Trinotate and visualized using BGGs WEGO [40]. GO functional enrichment tests on differentially expressed transcripts were performed with scripts from Trinity v2.4.0 that wrapped the Bioconductor package Goseq [41]. Blast2GO v.4.0.7 [42] was used to map GO terms to plant GO Slim [43] terms in order to obtain a broad overview of the transcriptome. For each time point, enriched GO terms were visualized using GOpplot v1.0.2 [44]. Most significant over- and under-represented GO terms (top 10 for each time point, then combined) were extracted using custom scripts and visualized using the GOpplot package (http://wenccke.github.io/).

2.7. KEGG pathway analysis

A standalone version of KOBAS 2.1.0 [45] was used to functional annotate the transcriptome with KO terms, the corresponding pathways and enzymes were extracted from information retrieved using KEGG API. The TMM-normalized FPKM expression matrix was filtered by removing low expressed transcripts (average expression value below 1) and transcripts without proper KO terms, then used as input for Pathview ([46]; online version) to visualize significant pathways (internally use GAGE [47] for pathway enrichment analysis) and some manually selected pathways. Pathways with q-values smaller than 0.05 were combined and shown as heatmaps.

3. Results

E. oleoabundans was grown in continuous mode in controlled flat-panel photobioreactors and subjected to full nitrogen-starvation during a 4-day period. Before and during this starvation period, culture parameters were monitored and physiological and transcriptional changes in culture samples measured to determine the temporal dynamics of the nitrogen-stress response. For the pre-cultures grown in nitrogen-replete conditions (+N) samples were taken at four timepoints in the light-phase preceding nitrogen starvation, while from the onset of nitrogen starvation (~N), time-series samples were taken at the start, middle and end of the light-phase and at the end of the dark-phase over a 4-day period (Fig. 1).

3.1. Biomass dynamics under nitrogen starvation

Upon nitrogen starvation, E. oleoabundans could sustain an increase in biomass concentration (cell dry weight) during the light phase in the first two days (Fig. 2A). Overall, the biomass concentration gradually decreased, which coincided with a decreased biomass productivity over time (Fig. 2D). The protein content and quantum yield (QY) also showed a steady decline throughout the starvation period (Fig. 2D and Supplementary Fig. A.1).

Nitrogen starvation induced the fast accumulation of carbohydrates, increasing from 15% at the start to a maximum content of 33% of the cell dry weight after 10 h starvation. Within the same time frame and with the same accumulation pattern, starch content increased from 8% at the start to a maximum content of 28% of the cell dry weight (Fig. 2B). This indicated that the increase in total carbohydrates was mainly due to starch accumulation. The starch content subsequently decreased again to N-replete levels of 18% in the light phase at day 4. In contrast, accumulation of total fatty acids (TFA) was more gradual from a content of 16% at the start to a maximum of 41% after 4 days. The increase in lipid content was mainly due to TAG accumulation, which increased from a content of 0.4% to 30% after 4 days, while the polar lipid (PL) content was stable around 13% (Fig. 2C). This indicates that TAG biosynthesis occurred mainly de novo, although some interconversion from starch to TAG or from membrane lipids to TAG cannot be excluded. During the dark phases starch was consumed and decreased to 15% content, while a decrease of fatty acids (indicating fatty acid consumption), was not observed. In the hybrid producer E. oleoabundans starch therefore is the primary storage compound that is
accumulated in the light phase and that serves as a transitory energy source that is respired in the dark phase.

Concomitant with the accumulation of storage compounds the fatty acid composition of the different lipid fractions also changed during nitrogen starvation. A clear change from polyunsaturated fatty acids (PUFAs) towards mono-unsaturated and saturated fatty acids was observed in all fractions (Fig. 3). Under N-replete conditions a large part of all lipid fractions was composed of C18:3 and C16:3 fatty acids (approximately 30% and 15%, respectively). These species decreased during N-starvation (to 5% and 2.5%, respectively), while there was an increase in the (mono-)unsaturated fatty acids C18:1, C16:2 and C16:1 and the saturated fatty acid C18:0 respectively. The biggest change was observed for C18:1 that increased from 5% to almost 40% in the lipid fraction. PUFAs with higher carbon chain lengths (e.g. C20:3 and C20:5) were not detected in any of the lipid fractions in N-replete and N-starvation conditions.

Production rates for total carbohydrates (TCH) and starch showed an identical rapid increase during the first day of N-starvation. This indicates that increased TCH levels in the first day can mainly be ascribed to starch production. Subsequently, both TCH and starch production rates decreased to levels below those for N-replete growth, although more severe for TCH. This means that production rates of carbohydrates other than starch were reduced much stronger. In the lipid fraction TAG productivity increased slowly in the first 2 days following N-starvation, and then decreased again to N-replete rates. The total lipid productivity remained stable on the first day of N-starvation and then started to decrease (Fig. 2D).

3.2. Differential gene expression dynamics under nitrogen starvation

A total of 25.6 million high-quality RNA-seq clean reads were de novo assembled into 61,010 transcripts belonging to 42,338 “genes” with a N50 length of 1838 bp and average GC content of 58.9% (Table 1). Functional annotation of the transcriptome was carried out with the Trinotate pipeline, and 44,789 transcripts (73.4%) were annotated using UniRef90 database (Table 1). The high-quality reference transcriptome served as a solid foundation for subsequent gene expression analysis.
Fig. 3. Fatty acid composition of (A) total fatty acids, (B) TAG and (C) polar lipids fractions of *E. oleoabundans* grown in N-replete (N+) and N-starvation (N−) conditions in air-lift flat-panel photobioreactors. Mean values (*n* = 3) are shown. Onset of N-starvation was at *t* = 0 h.
3.2.1. Global gene expression analysis

Time series experiments were performed (7 time-points, 3 replicates; GEO accession GSE104807), reads from each RNA-seq library were individually aligned back to the reference transcriptome and the abundance of each transcript was determined. The sequencing depth for individual samples ranged from 8.5 to 19 million reads, and over 97% of the reads could be mapped to the reference transcriptome for all the samples. The transcriptome coverage, defined here as the percentage of genes with TPM (Transcripts Per Kilobase Million) abundance bigger than one, was 68.2–80.5% (Supplementary Table 5). The hierarchical clustering of transcripts from samples obtained from triplicate photo-bioreactors indicated a good experimental reproducibility (Supplementary Fig. A.2). Differentially expressed transcripts were then determined from raw count values for each time point from the onset of N-starvation. A total of 12,975 transcripts were either up-regulated or down-regulated compared to control samples (0 h) and the number varied between different time points (2014-6589) (Supplementary Table 1).

Hierarchical clustering of expression profiles of each transcript resulted in 23 distinct expression clusters (Fig. 4 and Supplementary Table 1). The largest clusters were cluster 21, 15, 11 and 4, which contained 2938, 1834, 1145 and 1081 differentially expressed gene isoforms respectively. Many transcripts related to fatty acid, TAG and starch metabolism belong to these clusters. The diverse expression clusters to which the genes from a metabolic route belong to, indicates that different genes from these metabolic pathways are most probably under the control of different transcriptional regulators and therefore do not follow simultaneous activation. Some transcriptional regulators group into these clusters as well and might play a regulatory role in the nitrogen-starvation activated response of fatty acid and starch biosynthesis genes. These include transcriptional regulators such as the 5′-AMP-activated and SNF1-related protein kinases (TR3188|c0_g1 and g2, and TR24203|c0_g1), a dual specificity tyrosine-phosphorylation-regulated kinase (DYRK3: TR935|c0_g2_11), PHR1-LIKE 1 protein (PHL1: TR19823|c0_g1) and a nitrogen regulatory protein P-II homolog (GLNB: TR228|c0_g1 and TR228|c0_g3). Clusters 1–3, 12, 19 and 20 showed very distinct expression peaks and interestingly several transcripts related to reactive oxygen stress grouped to the clusters 12, 19 and 20. Detailed description of the transcriptional response to nitrogen starvation is discussed below.

The GO terms for upregulated transcripts were enriched with terms related to fatty acid and lipid metabolism as well as carbohydrate and starch metabolism, as is expected in view of the observed biomass changes (Supplementary Fig. A.4 and Supplementary Table 2). In addition, GO terms for ammonium uptake and incorporation were enriched (direct response to absence of nitrogen) as well as GO terms related to reactive oxygen species (ROS) stress, indicating that N-starvation might cause the production of ROS. The GO terms for down-regulated transcripts were enriched with terms related to the chloroplast and photosynthesis, which makes sense in view of the breakdown of pigments (cell bleaching) and reduced photosynthetic capacity (e.g. quantum yield) due to the lack of nitrogen (Supplementary Fig. A.4 and Supplementary Table 3). Taken together, GO analysis indicates that most of the transcripts can be grouped into a small number of broad yet distinct functional categories that correlate well to the observed biomass and physiological changes. GOBubble plots based on plant GO Slim terms (Supplementary Fig. A.4) showed that the highest number of increased and decreased GO Slim terms was found after 6 to 10 h of nitrogen starvation. This indicates that most transcriptional changes upon nitrogen starvation occur relatively rapid. The broader GO Slim terms indicated that “carbohydrate metabolic process” (GO:0005975) was the only GO Slim term that for all time points significantly increased compared to T = 0 (negative z-score H0 versus other time points). Other GO Slim terms like “catabolic process” (GO:0009056) and “response to stress” (GO:0006950) also increased for an extended period. In contrast, the GO Slim terms like “photosynthesis” (GO:0015979), “plastid” (GO:0009536) and “thylakoid” (GO:0009579) were clearly decreased from 6 to 29 h (positive z-score H0 versus other time points), indicating a reduction in the photosynthetic capacity.

KEGG pathway enrichment was performed using GAGE (Supplementary Table 4) and data were mapped on KEGG pathways to generate graphs showing detailed temporal changes (Supplementary Fig. A.5). The most striking observations from the KEGG pathway enrichment analysis was an up-regulation of the TCA cycle (ko00020), Proteasome (ko03050), Protein processing in endoplasmic reticulum (ko04141) and Phagosome (ko04145) pathways, and a down-regulation of the Photosynthesis (ko01195) pathway (Supplementary Table 4). This indicates increased protein and organelle degradation and correlates with the increased GO Slim terms such as “catabolic process” (GO:0009056), “generation of precursor metabolites and energy” (GO:0006901) and “protein metabolic process” (GO:0019538), as well as the decreased photosynthesis, plastid and thylakoid GO Slim terms mentioned above (Supplementary Fig. A.4) and decrease in quantum yield (Supplementary Fig. A.1).

3.2.2. Transcriptional changes in starch metabolism

The transient and rapid increase in starch productivity and accumulation in the first 10 h of N-starvation coincided with increased expression of several genes involved in starch biosynthesis (Fig. 5 and Supplementary Table 1). In particular, starch biosynthesis genes encoding phosphoglucomutase (conversion of glucose-6-phosphate to glucose-1-phosphate) showed a strong up-regulation in expression from the onset of N-starvation, while the ADP-glucose phosphorylase (conversion of glucose-1-phosphate to ADP-glucose) was only up-regulated from 10 h on. Multiple starch synthase genes showed different responses. While the soluble starch synthase S3Y and the granule-bound starch synthases SSG1 and SSG2 showed a fast and transient up-regulation from 2 to 6h from the start, the soluble starch synthase S3Y showed a strong transient increase in expression from 6 to 10h. Likewise, the 1,4-alpha-glucan-branching enzyme (GLGB1) that modifies amylose to starch was only up-regulated in the first 6h. Among the starch degradation genes, gene expression of the alpha-glucan water dikinase (GWD1) was increased from 6h on. This enzyme increases the phosphorylation-level of starch and thereby stimulates the degradation of starch in higher plants and Chlamydomonas [48]. Starch breakdown involves several amylases and we found that multiple genes for alpha-amylase (AMY1 and AMY3), beta-amylase (AMYB, AM1, BAM1) and isoamylase (ISOA1 and ISOA3) showed fluctuating expression responses. Overall though, they exhibited down-regulation at the start of starch accumulation and up-regulation in the later stages, from 6 to 10h on. Hexokinase performs the first step in glycolysis, the conversion of glucose to glucose-6-phosphate, and hexokinase genes were upregulated from 10h starvation on. This indicates breakdown of glucose, which for a large part could be starch-derived glucose, as starch

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**Table 1**

**Summary of de novo transcriptome assembly and annotation for *E. oleo-bundans***

<table>
<thead>
<tr>
<th>Features</th>
<th>Results</th>
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degradation increases from this time on as well. Overall, these results suggest that the increased production and accumulation of starch can mainly be attributed to the strong up-regulation of the starch biosynthesis genes and less to a decrease in starch degradation. The subsequent decrease in starch productivity and accumulation from 29 h on concurred with decreased gene expression for the soluble starch synthase SSY2 as well, which supports this hypothesis.

3.2.3. Transcriptional changes in fatty acid and lipid metabolism

Transcriptional changes for genes related to lipid biosynthesis (Fig. 6 and Supplementary Table 1) also correlated with the observed increase in productivity and accumulation of TFA and TAG (Fig. 2). Several genes required for fatty acid biosynthesis showed increased expression within 2 to 10 h from N-starvation. The first step in fatty acid biosynthesis requires the precursor acetyl-CoA, which can be derived from different biochemical conversions. A first route is via the pyruvate dehydrogenase complex (PDH) that converts pyruvate to acetyl-CoA. A second route is via acetyl-CoA synthetase (ACS), that converts acetate to acetyl-CoA. Finally, a third route is via the TCA cycle where ATP citrate lyase (ACLY) can convert citrate to acetyl-CoA and oxaloacetate.

Fig. 4. Hierarchical clusters of gene expression profiles during nitrogen starvation. Expression levels are shown as median-centered log2(fpk + 1) values for the time series H0-H99 and the number of transcript isoforms in each cluster are indicated.
Gene expression of the PDH subunits ODPA1 and ODPB1 (TR1574|c0.g2 and TR13897|c0.g2), several acetyl-CoA synthetases (TR16088|c0.g1|g2, TR18695|c0.g1, TR2994|c0.g1|g2|g3 and TR26776|c0.g1) and ATP citrate lyases (TR25574|c0.g2 and TR26430|c0.g2) clearly increased within 2 to 10 h from N-starvation and remained so until 99 h of starvation. Increased expression of PDH, ACS and ACLY genes can result in a higher supply of acetyl-CoA, the final precursor for fatty acid biosynthesis, and thereby increase the carbon flux towards lipid synthesis (Fig. 6). The subsequent steps in fatty acid biosynthesis involve acetyl-CoA carboxylase (ACCase) and malonyl-CoA-ACP transacylase (FABD), and gene expression for these enzymes also increased in the first 10 h from N-starvation. The fatty acid synthesis (FAS) complex genes were up-regulated as well, including the ones encoding chloroplastic 3-oxoacyl-ACP synthase I (KASC1), 3-oxoacyl-ACP reductase (FABG), NADPH-dependent enoyl-ACP reductase (FABL) and 3-hydroxyacyl-ACP dehydratase (FABZ), as well as the acyl-CoA thioesterase (FATA). Overall however, this response was more delayed than that for starch biosynthesis.

For subsequent biosynthesis of triacylglycerides (TAG), transcripts encoding glycerol kinase (GK), glycerol-3-phosphate dehydrogenase (GPDA), two glycerol-3-phosphate O-acyltransferases (GPAT) and the phosphatic acid phosphatase (PAP) were strongly up-regulated from 2 h after the onset of N-starvation, while gene expression of lysophosphatic acid acyltransferase (LPAT) - which is responsible for the intermediate conversion of lysophosphatic acid to phosphatic acid - only increased from 29 h on. These genes perform biosynthesis of the lipid backbone glycerol-3-phosphate (G3P) and the subsequent incorporation of free fatty acids up to diacylglycerol (DAG). The diacylglycerol O-acyltransferases (DGAT) genes required for the final conversion of diacylglycerol (DAG) to triacylglycerol (TAG) also showed an initial decrease in expression, but were clearly up-regulated only from 29 h of N-starvation on, indicating that the DAG pool is built up before the final DAG to TAG conversion. Both a DGAT-1 and a DGAT-2 gene were up-regulated and appear to be responsible for the final conversion of DAG to TAG. The transcriptome annotation predicted a single DGAT-1 gene and five DGAT-2 gene sequences, and our data indicate that only one of the five DGAT-2 genes responds to N-starvation. The subcellular locations of these DGAT enzymes and the corresponding TAG biosynthesis pathways in *E. oleoabundans* are still to be determined. Interestingly, a gene encoding a putative lipid body-associated caleosin (peroxigenase) protein (TR11980|c0.g1.i1) was also up-regulated from 29 h on, indicating that increased lipid body formation is concomitant with DAG to TAG biosynthesis. Incorporation of free long-chain fatty acids into lipids requires the activation to their acyl-CoA form, which is usually performed by long-chain acyl-CoA synthetases (LACS). A gene isoform encoding a chloroplastic long-chain-fatty-acid CoA ligase (TR19322|c0.g1.i1) showed up-regulation from 53 h on and might serve this role.

TAG lipase activity negatively affects TAG accumulation, while phospholipases are involved in liberation of free fatty acids from (plastid) membrane lipids and interconversion into TAG. Transcript levels of two TAG lipase genes (SDP1: TR9756|c0.g3 and TR22634|c0.g2.i1) were decreased in the first 10 h of N-starvation, while at the end of N-starvation a marked increase was observed (Supplementary Table 1). The late upregulation of TAG lipases follows the earlier up-regulation of TAG biosynthesis genes, which could explain the initial steady increase in TAG accumulation and final leveling-off in TAG productivity. Besides de novo TAG synthesis, interconversion from membrane lipids to TAG is a common mechanism under stress conditions in microalgae [49–52], and the enzyme phospholipid:diacylglycerol acyltransferase (PDAT) plays an important role in this [51,53]. A PDAT homolog was not identified in our reference transcriptome, but several chloroplastic phospholipase A1 genes (DAD1 homologs) did show increased expression at different timepoints during N-starvation (Supplementary Table 1). Phospholipase activity results in the conversion of phospholipids to lysophospholipids and free fatty acids, which can be further incorporated into TAG upon N-starvation.
acids. Lysophospholipase subsequently converts lysophospholipids further to free fatty acids and glycerol-phosphate derivates [54,55]. A lysophospholipase gene (TR21369|c0_g1_i5) showed increased expression from 29 h on. Reacylation of lysophospholipids to phospholipids is performed by lysophospholipid transferases, and two lysophospholipid transferase genes (LPT1: TR19784 and TR9894|c0_g2_i1) were downregulated within the first 10 h. The gene expression profiles of the (lyso)phospholipases and lysophospholipid transferases could indicate that free fatty acids are liberated from membrane phospholipids upon nitrogen starvation, which subsequently can be used in TAG biosynthesis. The decrease in chloroplast-related GO Slim terms in the period before the clear increase in DGAT expression at 29 h also supports that TAG biosynthesis initially was not only de novo, but might also be the result of some interconversion from membrane lipids to TAG. Altogether, there are several indications that in the earlier stages of N-starvation there is active recycling of plastidial membrane phospholipids to TAG. Therefore incorporation of membrane-derived fatty acids into TAG cannot be excluded and awaits further investigation, e.g. by using carbon radiolabelling studies. Expression of several chloroplast-specific fatty acid desaturase (FAD) genes was repressed from 6 h on (Fig. 7), which explains the increased incorporation of mono-unsaturated (C18:1 and C16:1) and saturated (C18:0) fatty acids in the different lipid fractions and decrease of C18:3 and C16:3 fatty acids. The combined reduction in expression of FAD6C (C16:1 → C16:2 and C18:1 → C18:2) and FAD3C (C16:2 → C16:3 and C18:2 → C18:3) supports the increase in the C16:1, C16:2 and C18:1 fatty acid fractions. Concomitant with this, expression of the elongase-complex genes 3-ketoacyl-CoA synthase (KSC1), very-long-chain (3R)-3-hydroxyacyl-CoA dehydratase (HACD) and very-long-chain enoyl-CoA reductase (ECR) showed increased expression. This could result in higher C16:0 to C18:0 conversion, and higher C18:0 levels could thereby increase C18:1 production as well. Both increased C18:0 and C18:1 levels were observed, though C16:0 levels were not clearly reduced. This indicates that most probably the observed increase in C18:1 and C18:0 is mainly caused by the reduction in fatty acid desaturases gene expression. In support of this we did not observe significant differential gene expression of two stearoyl-ACP desaturase genes (SAD3 and SAD5) in our reference transcriptome, which are responsible for desaturation of C16:0 and C18:0. From the start of N-starvation a chloroplastic FAD4 desaturase gene expression was also reduced (TR22599|c0_g2_i1), which converts [sn-2-C16:0]-containing glycerophospholipids into [sn2-C16:1]-containing glycerophospholipids, while at the same time expression of a chloroplastic
palmitoyl-monogalactosyldiacylglycerol delta-7 desaturase (ADS3, TR16760[c0_g1_i1]) was upregulated in the early phases of N-starvation. This indicates decreased C16:0 to C16:1 conversion in the glycerophospholipid fraction, while in the glycolipid fraction (specifically MGDG) there was an initial increased conversion of C16:0 to C16:1. Overall, the combined effect of changes in expression of several desaturase genes involved in C16:0 to C16:1 conversion, led to an increased C16:1 content in the lipid fractions. Finally, expression of a palmitoyl-protein thioesterase gene (TR646[c0_g3_i1]) was upregulated in the late stages of N-starvation. This could supply additional free palmitate (C18:0) derived from lipid-containing proteins for incorporation into e.g. TAG.

3.2.4. Transcriptional changes in photosynthesis, carbon fixation and oxidative stress

Expression of genes for the photosystems proteins initially increased (chlorophyll a/b binding proteins), which coincided with a decreased expression of chlorophyllase (Supplementary Table 1). This could indicate that as N-starved cells cannot synthesize novel (N-containing) photosystems due to the lack of nitrogen, cells try to maintain the level of active photosystems by reducing chlorophyll degradation and maintaining its incorporation chlorophyll into existing photosystem proteins. However, at 10 to 29 h post-starvation this can no longer be sustained, indicating a decrease in photosystem biosynthesis and (most probably) photosynthetic efficiency (as can also be observed from the decreased quantum yield). The observed decrease of chloroplast-related GO Slim terms in this period is in concordance with this.

Many genes involved in CO2 fixation (RuBisCo, PGK, PEP carboxylase, PEPC kinase) and CO2 concentrating mechanisms (carbonic anhydrases) show a rapid up-regulation in expression in the first 2 h, a decrease afterwards and increase again after 99 h (Supplementary Table 1). This correlates to the initial sustained expression of photosystem genes and indicates that N-starved cells initially try to maintain photosynthesis and carbon fixation. Ultimately, when because of N-starvation cells cannot maintain active photosystems they can experience oxidative stress by the production of reactive oxygen species (ROS) such as hydrogen peroxide (H2O2) and superoxide (O2− ) that can cause cell damage, as has been observed in e.g. Chlorella and Acutodesmus species [59–62]. In response to N-starvation, E. oleoabundans showed increased transcript abundance for superoxide dismutase (SOD), catalase, peroxiredoxin and several peroxidases which indicates that these ROS were produced in this species as well (Supplementary Table 1).

3.2.5. Transcriptional changes in nitrogen uptake and incorporation

As expected, N-starvation induced changes in expression of genes involved in nitrogen metabolism. The depletion of nitrate led to reduced gene expression for nitrate transporters, nitrate reductase and ferredoxin-nitrite reductase. At the same time, increased expression started for ammonium transporter genes and the glutamine synthetase (GS) and NADH-dependent glutamate synthase (GOGAT) pathways. This indicates that cells switch to an alternative mode of nitrogen incorporation via ammonium uptake (Supplementary Table 1).

3.3. Transcriptional regulators of starch and lipid accumulation

The patterns in starch and lipid accumulation suggest the existence of a regulatory mechanism for redirecting carbon-partitioning (switching) between the associated metabolic pathways. Identification of the associated “switch” regulators would be of great interest to improve existing metabolic engineering strategies that currently are often based on increased expression of a specific metabolic gene or pathway, or introduction of heterologous transcription regulators from higher plants [6].

In this study, we identified several transcriptional regulators with significantly altered gene expression upon N-starvation, some of which have been implicated in the nitrogen-starvation triggered lipid accumulation in microalgae (Supplementary Table 1). Genes encoding 5′-AMP-activated and SNF1-related protein kinases (TR3188[c0_g1_i1] and g2, and TR24203[c0_g1_i1], respectively) clearly showed increased expression from 10 h N-starvation on, and very strongly from 53 h on. This is in line with a previous transcriptome study in E. oleoabundans [17] where after 11d of N-starvation the 5′-AMP-activated protein kinase SnRK1 showed up-regulation. SnRK1 is a global regulator of carbon metabolism in plants [63,64].

In Nannochloropsis, bZIP-type regulators have been implied to control several steps in the synthesis of TAG [22]. We identified two such bZIP regulators with opposite expression profiles in our transcriptome set: TR2010[c0_g1_i1] and TR14421[c0_g1_i1]. TR14421[c0_g1_i1] expression was activated 2 h after N-starvation, but showed down-regulation from 29 h...
on, while TR2010|c0.g1 was down-regulated until 53 h, after which the gene was activated. We also identified a transcript (TR935|c0.g2|1) with homology to the dual specificity tyrosine-phosphorylation-regulated kinase (DYRKPI) from Chlamydomonas, a negative regulator of carbon storage that inhibits both starch and lipid accumulation and photosynthetic efficiency under N-starvation in this algae [65]. Expression of this transcript was markedly decreased from the start but increased from 29 h starvation on. Differentially expressed transcripts encoding two homologs of PHR1-like regulators from higher plants were also found (TR19823|c0.g1 and TR25002|c0.g1). PHR1 and its homolog in Chlamydomonas PSR1 are Phosphate Starvation Response transcription factors, and PSR1 was found to be essential for lipid and starch accumulation during phosphate starvation in Chlamydomonas [66]. A homolog of the nitrogen regulatory protein P-II (GLNB: TR228|c0.g1 and g3) was upregulated from 6 h starvation on. The P-II regulator from Chlamydomonas was found to negatively control TAG accumulation in lipid bodies during acclimation to nitrogen starvation in this green alga [67].

If and how these regulators affect carbon partitioning for the accumulation of starch and lipids remains to be investigated through functional analysis.

4. Conclusions and discussion

Metabolic engineering of microalgae towards increased lipid or starch productivity requires deeper insight into the genetic details of the metabolic pathways involved and the regulatory mechanisms controlling these pathways. For the oleaginous green microalgae E. oleoabundans this genetic information is still very limited, therefore we set out to study the biomass and transcriptional changes that occur upon nitrogen starvation, to find genetic triggers of starch and lipid accumulation in this industrially relevant microalga. Starch and lipid accumulation patterns showed different dynamics over the applied 4-day starvation period. Whereas the starch content increased strongly and reached its peak within the first day upon nitrogen starvation, lipids (mainly TAG) accumulated more gradually (Supplementary Fig. A.2). A pattern where starch accumulates rapidly ahead of lipid accumulation is observed in other hybrid oleaginous microalgae [9–11]. Starch seems to be the preferred primary storage compound in these species, which is partially due to the fact that its synthesis is biochemically and energetically more favourable than fatty acid biosynthesis [10,68]. On the other hand, lipids have a higher energy density and therefore for the longer term might be the preferred storage compound. Several studies indicate that starch and lipid accumulation are both dependent on de novo biosynthesis and therefore competition for common carbon precursors affects carbon-partitioning between starch and lipids. Increasing the carbon precursor supply can thereby be a strategy to influence carbon-partitioning and improve lipid accumulation in some cases [69,70]. Nitrogen starvation also affected the fatty acid composition of the lipid fractions, with a distinctive decrease of the polyunsaturated fatty acids C18:3 and C16:3, concomitant with a large increase of the mono-unsaturated fatty acid C18:1, and small increases of the mono-unsaturated fatty acid C16:1 and the saturated fatty acids C18:0 and C16:0. These changes in fatty acid profile correspond to observations from earlier studies on N-starvation in E. oleoabundans [71,72], and was observed in other hybrid oleaginous microalgae as well [9,73]. Besides these metabolic factors, several biosynthetic genes and transcriptional factors have been identified that play a role in accumulation and carbon-partitioning of starch and lipids [6,20,22,53,65]. We therefore determined the transcriptional changes upon N-starvation through RNA-sequencing and observed that gene transcripts for starch- and lipid-related metabolic pathways correlated well with the observed biomass changes. The accumulation of starch in the first day could mainly be ascribed to a general increase in transcript abundance for several starch biosynthesis genes. The peak in starch accumulation correlated particularly well with the rapid increase and subsequent decrease in expression of starch synthase genes (SSY2, SSY3, SGG1, SGG2), while the ultimate return to N-replete starch levels also appears to be caused by an increase in transcript levels of different amylase genes (α-, β- and isoamylases) in the later stages.

Expression of fatty acid biosynthesis genes and TAG biosynthesis genes was in general more delayed compared to those for starch biosynthesis and remained at increased levels until 4 days of starvation. Transcripts for the enzymatic reactions performed by PDH, ACS, ACLY and ACCase, which supply the fatty acid biosynthesis precursors acetyl-CoA and malonyl-CoA respectively, were clearly increased, but especially PDH was constitutively upregulated from 6 to 99 h. Likewise, for TAG biosynthesis the genes for G3PDH and glycerol kinase (glycerol-3-phosphate synthesis) and GPAT (incorporation of first acyl chain to glycerol-3-phosphate backbone) were in general up-regulated constitutively from 2 h starvation on, while other TAG biosynthesis genes had a more limited time-frame where they showed upregulation. In both pathways (fatty acid and TAG biosynthesis) there appears to be a trend where continuous supply of pathway precursors is important for the gradual accumulation of lipids/TAG. Metabolic engineering of microalgae for increased lipid production has focused on early and late genes in both pathways, either based on the hypothesis that the supply of fatty acid precursors is key for an increased flux through the FA biosynthesis or on the hypothesis that the ultimate conversion(s) to TAG are rate-limiting. Effective and ineffective metabolic engineering for increased production of lipids/TAG has been reported for both the first (e.g. [6,74]) and the second approach (e.g., [6,75]), yet it still remains unclear whether the key metabolic nodes for increasing lipid/TAG production are conserved among the plethora of oleaginous microalgae. In this respect, a recent study has demonstrated that overexpression of a native DGAT-2 gene in E. oleoabundans can more than double the lipid and TAG content and productivity, as well as alter the saturated fatty acid composition of cells [76]. This indicates that metabolic engineering focused on the final conversion of DAG to TAG is a successful strategy for increased lipid production in E. oleoabundans. The DGAT-2 gene that was used for overexpression is not the one showing differential gene expression in our study (TR20062|c0.g1), but another one of the four DGAT-2 genes found in the reference transcriptome (TR12237|c0.g1, g2 and g3). On the other hand, the location of de novo TAG biosynthesis remains unclear at this moment. While most of the fatty acid biosynthesis transcripts were predicted to be located in the chloroplast, the subcellular location of the TAG biosynthesis transcripts was variable or unspecified (Supplementary Table 1). Whether TAG biosynthesis occurs both in the chloroplast and on the endoplasmic reticulum in E. oleoabundans, as is the case in many microalgae [77], therefore needs to be verified by further biochemical experimentation.

Several transcriptional regulators have been implicated so far to play a central role in lipid accumulation in microalgae. These include the bZIP-type regulators that are suggested to control several steps of TAG synthesis in Nannochloropsis [22], and the nitrogen-responsive regulator (NRR1) and phosphorus stress response regulator (PSR1) in Chlamydomonas [20,53]. While these putative switch regulators were identified through transcriptomics and in silico studies, the direct role of the PSR1 gene in lipid accumulation was confirmed by mutagenesis and overexpression. The kinase DYRKPI has been implicated to repress both starch and lipid accumulation in Chlamydomonas and to result in sustained photosynthetic efficiency under N-starvation [65]. From these described transcriptional regulators, we identified several homologs in E. oleoabundans that showed significant differential gene expression upon nitrogen starvation. The actual role of these regulators in starch and lipid accumulation and carbon partitioning now needs to be further analysed through e.g. gene inactivation and gene overexpression approaches.

Finally, it remains unclear whether accumulation of these storage compounds in E. oleoabundans is controlled by genetic switches only (i.e. via specific activating or repressing regulators) or that
accumulation and carbon partitioning can also be regulated at the metabolic level, e.g. via changes in enzyme quantity or enzyme activity at specific key metabolic nodes. To test these hypotheses, a high-quality genome sequence of *E. oleoabundans* and efficient tools for reverse genomics, such as those for targeted gene knockdown [78] and gene editing [79] in *Nannochloris oceanica* will be highly valuable.

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**Author contributions**

MHJS and JH conceived, designed and performed all experiments, sample processing and biomass analysis. MHJS, YG and DW performed data processing, analysis and interpretation. JX and RHW acquired funding and supervised the work. MHJS and YG wrote the manuscript. All authors revised the manuscript. All authors read and approved the final manuscript.

**References**


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