Engineering of oleaginous organisms for lipid production
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Phototrophs are attractive candidates for commercial lipid production. Lipid biosynthetic pathways in these organisms have been largely characterized but the mechanisms partitioning resources toward storage lipids are poorly understood. One promising strategy to study and enhance biomass lipid bioproduction in oleaginous microorganisms is to combine genome-scale metabolic modeling and genetic and metabolic engineering. Here we describe recent advances in in vitro, in vivo, and in silico manipulations of phototrophic metabolism that increase total lipid content or redirect lipid production toward more favorable products such as polyunsaturated fatty acids used as nutritional supplements or in biofuel production.

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Introduction
Excessive demands by modern society have been depleting nature’s resources over the past centuries. Exploring and developing new sustainable resources to counter increasing consumption has therefore been the focus of current research efforts in the academic and private sector. The emphasis has partly been on investigating the use of phototrophic organisms that are able to fix carbon dioxide by utilizing light energy and produce energy-dense products. Lipids derived from photosynthetic organisms have therefore been explored to augment existing sources for energy and nutrition [1,2]. Efforts for generating lipid-derived products at the industrial scale have been predominantly centered around microalgae and peripherally around oleaginous yeast. While the biosynthetic pathways for fatty acid and lipid metabolism in these organisms are fairly well understood, how organisms prioritize carbon and energy toward those pathways is still being explored. As a result, there has been a surge in the attempts to understand and subsequently manipulate the cellular processes responsible for allocating resources for the production of lipid biomass. Carbon partitioning is the natural method by which an organism directs resources into various metabolic processes. In phototrophs, this process is often at odds with the accumulation of engineered targets, that is desired lipids, with a bias toward non-desirable carbohydrate biosynthesis [3]. Strategies for maximizing lipid biomass include engineering the cellular metabolism toward a desired biomass composition and taking advantage of natural processes of enhancing carbon toward lipid biomass such as unfavorable growth conditions. These conditions lead to environmental and nutrient stress induced growth arrests in photosynthetic microorganisms which usually result in an accumulation of storage metabolites, including lipids [4]. However, the organism’s inherent partitioning bias between lipids and carbohydrates remains intact. Thus, large scale screens to identify organisms with inherently favorable partitioning have been a common method of choice [5,6]. Increasing the total resource accumulation is another strategy that has been explored since excess nutrient supply is generally diverted to storage molecules. Mixotrophic cultures, where a CO2-fixing phototroph is provided additional resources in the form of an organic carbon source, often results in higher biomass yields [7]. However, the benefit in high yield has to outweigh the cost of providing additional carbon. To circumvent this challenge, designing symbiotic co-cultures with organic carbon-producing phototrophs and organic carbon-utilizing oleaginous heterotrophs is a promising strategy for advanced lipid biomass production [8]. Finally, genome-scale modeling of cellular metabolism can be used as a framework for understanding intracellular carbon partitioning and to create optimized, at times non-intuitive, strategies for maximizing target compounds.

Challenges in lipid-producing microorganisms
Organisms currently being investigated for the production of biofuels and lipids include plants, unicellular phototrophs, and oleaginous heterotrophs. The use of land plants for the production of bioethanol either directly from the plant storage sugars or via processing agricultural waste are options that have been aggressively investigated [9]. For lipid-derived sources, edible and non-edible oilseed crops are being explored [10]. The challenges associated with these sources are centered on the food versus fuel conundrum [11] and the environmental impact [12].
These challenges are constantly cited as the driver for choosing a microbial-based platform for lipid products. Unicellular phototrophs, such as bacteria and microalgae, exhibit high photosynthetic efficiency, can be grown on non-arable land with minimal inputs (CO₂, light, and vitamins), and naturally produce desired products, such as nutritional omega-3 fatty acids. However, most industrial-scale applications for biofuels are currently limited by the challenge of generating biomass along with the economics of scale: bioreactors are inefficient and open (outdoor) cultures are complex [13]. The economics become more feasible with the concurrent generation of high-value products such as omega-3 nutritional supplements, carotenoid-based antioxidants, as well as pigments [14–16]. The success of microbial-based lipid biomass platforms, however, is highly dependent on our understanding of metabolic and regulatory bottlenecks.

**Lipid biosynthetic pathways**

Fatty acid and lipid biosynthetic pathways in plants, unicellular phototrophs, and oleaginous yeast have been well studied (for recent reviews see for example [17–20]). A visual overview of these pathways can be found in Figure 1. Acetyl-CoA carboxylase (ACCase) catalyzed generation of malonyl-CoA is the first committed step of fatty acid biosynthesis; Huerliman and Heimann have published a comprehensive review of algal ACCases [21]. Fatty acid de novo biosynthesis occurs in the chloroplast of phototrophs or in the cytosol of yeast with the acyl-activating moiety differing between the compartments (acyl carrier protein versus CoA respectively). The fatty chain length is extended two carbons at a time involving four enzymes that make up the fatty acid synthase complex. In phototrophs, a cytosolic set of fatty acid synthase enzymes can extend the chain length to form very long (C20 — C24) saturated or monounsaturated fatty acids. While polyunsaturated fatty acids (PUFAs) of the C16 and C18 variety are synthesized in the chloroplast, very long chain omega-3 PUFAs are synthesized in the endoplasmic reticulum and the cytosol. Acyl-editing, a well orchestrated cycle of fatty acid elongation and desaturation, synthesizes high-value omega-3 fatty acids and is largely responsible for their incorporation into triacylglycerols (TAGs) [22]. Lipid biogenesis begins by siphoning triose phosphate away from central carbon metabolism, reducing the oxo functional group, and acylation with activated fatty acids to form phosphatidic acid. Phosphatidic acid is the starting point for the biogenesis of the vast array of membrane lipids (phosphatidylethanolamine, diacylglycerol, monogalactosyldiacylglycerol, etc.) as well as TAGs. Photosynthetic membrane biogenesis is a chloroplastic process with specific fatty acid requirements. Non-photosynthetic membrane synthesis occurs in the cytosol and endoplasmic reticulum, requiring chloroplast generated fatty acids in phototrophs or cytosol generated fatty acids in heterotrophs. The metabolic input required for TAG biosynthesis requires either an excess of carbon and energy or the conversion of existing resources. For the latter, remodeling of existing membrane lipids into TAGs accompanies various types of physiological changes or stress including changes in light, pH, temperature, and nutrient [23–28]. While the biosynthetic pathways have been largely characterized, the regulatory mechanisms that partition resources toward storage lipids are still poorly understood, thus presenting an unexplored resource for design strategies.

**Strategies for lipid production**

Taking advantage of our current knowledge about lipid biosynthetic pathways, several strategies have been exploited to optimize lipid production in photosynthetic organisms ranging from exposure to environmental and nutrient stress to metabolic engineering using transgene expression as well as co-culturing and in silico investigations of flux distributions. Figure 2 gives a schematic overview of the discussed strategies for enhanced lipid production in phototrophs.

**Inherent carbon partitioning**

Oleaginous microorganisms naturally exhibit differential carbon partitioning between carbohydrates and lipids. Liang and Jiang reviewed the potential of oleaginous microorganisms for biodiesel production including bacteria, fungi, microalgae, and yeast and collected the total biomass and biomass lipid content for selected microalgae [29]. On the basis of this data, *Chaetoceros gracilis* produces the highest lipids per biomass followed by *Tetraselmis tetratehe*, *Nannochloropsis oculata*, *Chaetoceros muelleri*, and *Chlorella protothecoides* as shown in Figure 3. Their lipid content per dry weight ranges from 23% for *Nannochloropsis* to 60% for *Chaetoceros*. Among the other investigated organisms are microalgae with higher lipid content but lower total biomass resulting in lower lipid productivity such as *Chlorella* sp. with a lipid content of 52% and a biomass yield 15 times lower compared to *Chaetoceros*, *Thalassiosira weissflogii* and *Cylotella cryptica* showed favorable lipid accumulation in a comprehensive screen of marine diatoms [6]. A screen of 69 strains of oleaginous yeast grown under various culture conditions identified a variety of candidates with lipid content comparable to that of photosynthetic microorganisms [30°]. The most productive oleaginous yeast genus (*Rhodosporidium*) had a biomass yield 1 — 2 orders of magnitude higher than the most attractive diatom candidates with similar total lipid composition [30°]. The dependence of growth rate and carbon fixation on light in phototrophs is probably the source of this disparity. Yet, supplementing phototrophs with organic carbon increased the biomass yield by approximately threefold in different *Chlorella* strains and *Nannochloropsis* [7]. In a separate study, *Chlorella sorokiniana* cultured under mixotrophic and pure heterotrophic conditions were able to achieve biomass yields comparable to yeast [31]. To offset the additional cost of organic
carbon supplementation, wastewater and biodiesel waste (i.e. glycerol) have been successfully used in microalgae [32,33] and oleaginous yeast cultivation [34–36].

Environmental or nutrient stress in microalgae induces a shift toward nutrient storage [4]. Lipid accumulation and fatty acid redistribution during nutrient deprivation has been extensively studied in diatoms [37–40,41*,42**], microalgae [43,44**,45,46] and oleaginous yeast [35]. Studies in *Chlamydomonas reinhardtii* [44,46] and *Phaeodactylum tricornutum* [42**] showed degradation of the photosynthetic apparatus accompanies stress-induced lipid accumulation in phototrophs. This effect can also be artificially initiated at faster time scales using small molecules as shown in *C. reinhardtii* [47]. Thus, the accumulation of TAGs during stress is a reallocation of existing protein and lipid biomass allocated to the photosynthetic apparatus; biomass that is also partitioned into carbohydrates on the basis of the organism’s carbon allocation biases. Metabolic flux analysis of *C. protothecoides* [48] and *Trichosporon cutaneum* [49] experimentally determined the carbon partitioning of a microalga and oleaginous yeast respectively.

**Manipulation of lipid metabolism using metabolic engineering**

Recent advances in metabolic engineering tools for photosynthetic organisms demonstrated that they are attractive candidates for industrially relevant lipid production. Transgenic expression of high-efficiency desaturases and elongases was, for example, applied to enhance the production of PUFAs. Hamilton *et al.* optimized the omega-3 content in *P. tricornutum* by expressing a Δ5-elongase gene from the green alga *Ostreococcus tauri* [50]. Co-expression of microalgal Δ5-elongase and Δ6-desaturase into the oleaginous yeast *Pichia pastoris* led to docosahexaenoic acid production in the transgenic yeast which naturally cannot produce PUFAs longer than C18 linoleic and α-linolenic acid [51]. Because of the fact that fatty acid synthesis is tightly regulated in most organisms, overexpression of genes
Strategies employed to enhance lipid production in photosynthetic organisms. Lipid biomass production can be enhanced using different strategies such as redirecting carbon toward lipid production using nutrient or environmental stress. Genome manipulations promise a favorable lipid biomass production while not slowing down growth rate. Because of the fact that both partners contribute to the biomass content, co-culturing can be used to enhance lipid productivity. Genome-scale models can facilitate and guide efforts in all shown strategies.

involved in fatty acid production can lead to reduced flux toward TAG production. To minimize regulatory effects such as feedback inhibition, Tai and Stephanopoulos co-overexpressed the genes corresponding to the first and last step in the lipid synthesis pathway in the oleaginous yeast *Yarrowia lipolytica* [52]. The biomass lipid content was increased more than fourfold to about 41% with a slightly changed fatty acid profile. Overexpression can also be used to change the fatty acid profile as shown in *P. tricornutum* [53]. The overexpression of glycerol-3-phosphate dehydrogenase resulted in a shift toward neutral fatty acid production. Using antisense and interfering RNA, Trentacoste and coworkers knocked-down genes involved in lipid catabolism in the diatom *Thalassiosira pseudonana* to increase lipid accumulation without compromising growth [54]. Another important step toward the exploitation of diatoms as biofuel producers is a targeted genome modification using, for example, meganucleases and transcription activator-like effector nucleases (TALENs) [55]. The authors were able to engineer seven genes in the lipid metabolism of *P. tricornutum* resulting in a 45% increase in TAG production. An alternative, RNA-dependent targeted genome engineering technology is the CRISPR/Cas9 system that has been shown to be efficient in a wide range of organisms [56], including industrial relevant *Saccharomyces cerevisiae* strains [57] as well as *Escherichia coli* [58]. Compared to TALENs, the CRISPR/Cas system promises to be easier to design and use but has not been implemented in microalgae yet. However, we anticipate this technology to be applied for lipid metabolism engineering due to its promising features.

Recently, Karas and coworkers demonstrated the transfer of genetic material from *E. coli* to the diatoms *P. tricornutum* and *T. pseudonana* using conjugation [59].

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**Figure 2**

Environmental shock
- Growth under nutrient or environmental stress
- Effects on carbon partitioning

Genome-scale models
- *In silico* predictions of flux distributions
- *In silico* strain design

Photosynthetic organisms of interest
- Lipid biomass enhancement
- Biofuels
- Antioxidants
- PUFAs
- Pigments

Metabolic engineering
- Transgene expression
- RNAi
- TALEN

Co-culturing
- Both partners contribute to biomass
- Phototroph produces carbon

High-value products

**Figure 3**

Lipid biosynthesis in phototrophs. Lipid productivity (mg/L/day) against biomass productivity (g/L/day) of selected microalgae on the basis of data from Ref. [29]. Circle sizes are scaled by lipid content. Lipid content and biomass productivity corresponding to the higher lipid productivity were selected in case a range was given.
method can provide for the efficient transfer of large amounts of genetic material such as entire pathways and will facilitate and accelerate genome manipulations in these organisms.

Often, the aforementioned metabolic engineering approaches result in strains unable to dynamically regulate gene expression [60]. Xu and coworkers significantly improved fatty acids production by rewiring an existing transcriptional regulator to dynamically control expressions of genes involved in synthesis and degradation of malonyl-CoA in E. coli. Because of lacking knowledge about the regulatory networks of oleaginous microorganisms this engineering approach was not much exploited yet.

**Enhanced lipid production through co-culturing**

Another strategy to improve lipid production involves the exploitation of symbiotic relationships between microorganisms with at least one photosynthetic species. So far, organisms exploited for these applications include fungi, microalgae, and phototrophic and heterotrophic bacteria [8]. Compared to monocultures, co-cultures can demonstrate enhanced growth rates as well as higher total biomass and lipid content since both partners contribute to biomass production and composition. Thus, the choice of the partners influences the overall fatty acid composition. Lipids obtained from a co-culture of *Trichosporonoides spatulata* and *Chlorella vulgaris*, for example, were mainly composed of palmitic, oleic, and stearic acid and showed a significantly higher content of saturated fatty acids than those of the monocultures [61].

When partnering up yeast and microalgae, yeast provides additional CO₂ to the algae which in return generates O₂ for the yeast [61]. Partnering microalgae and fungi can also enhance the harvesting process which requires a high amount of energy and can be very costly [62]. Wrede and co-authors studied fungal-assisted flocculation for different algae, including marine microalgae, for this purpose. Another possibility for facilitating the harvesting process is the encapsulation of the co-cultured organisms in alginate gel beads as demonstrated for *T. spatulata* and *C. vulgaris* [61]. Although biomass and lipid production slightly decreased compared to free cell cultivation, harvesting was facilitated and less costly. Most studies focus on artificial co-cultures although native communities have been shown to be viable feedstocks for biofuels and bioproducts too [63,64].

**Modeling**

Genome-scale metabolic models have proven to be powerful tools to understand and predict genotype — phenotype relationships. Flux balance analysis [65] can be used to predict intracellular flux distributions with good accuracy [66] allowing for *in silico* predictions of carbon partitioning. Elucidation of intracellular flux *in silico* gives insight into carbon partitioning and can be used to generate targets for metabolic engineering. This has successfully been deployed by using a genome-scale reconstruction of the oleaginous yeast *Mortierella alpina* to optimize the industrial production process of PUFAs [67].

Several mathematical optimization strategies are available that facilitate the identification of targeted genetic manipulation strategies *in silico*. These tools allow for the redesign of metabolism while simulating the impact on a specified objective. Additionally, computational approaches reveal non-obvious redesign strategies by suggesting manipulations in pathways distant from the studied target [68]. Different genetic manipulations can be identified using the available computational approaches. Some of these procedures, including OptKnock, can be used to identify gene deletions resulting in the overproduction of a desired metabolite while ensuring biomass production [69]. Another class of computational approaches for metabolic engineering can be used to identify a combination of gene deletions, down regulations and overexpression, for example OptForce [70]. This method was successfully used to optimize the production of fatty acids with a chain length of six to 16 carbons in *E. coli* [68]. Other available approaches for metabolic engineering can be used for strain design and identify non-native pathway additions. For example, Ip and coworkers applied *in silico* metabolic engineering to predict the effects of gene overexpression and insertion of heterologous pathways on free fatty acid production and experimentally validated their findings [71*]. Currently available procedures were recently reviewed in [72,73].

Genome-scale metabolic modeling can also be exploited to understand the complex interactions between microorganisms in natural and artificial communities. So far, community modeling has not been utilized to optimize the lipid content and composition in co-cultures but was successfully applied to simulate other aspects of microbial communities, such as the interactions between the partners [74].

**Conclusion**

Microalgae and oleaginous yeast accumulate a large part of their dry weight in lipids. However, to use these organisms for industrial applications, lipid biomass yield and productivity need to be further optimized. In this review, we provided an overview on promising strategies to enhance lipid production. Unfavorable growth conditions such as environmental and nutrient stress enhance lipid production while slowing down the growth rate. Recent advances in genetic engineering and manipulation of photosynthetic organisms promise to aid the construction of biofuel production strains without compromising growth. Genome-scale modeling promises
to aid the identification of potential phototroph lipid producers but also to facilitate and speed up the metabolic engineering of these organisms. Additionally, genome-scale modeling facilitates gaining a more detailed understanding of the phototrophs metabolic capabilities and regulation which is essential for engineering these organisms.

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References and recommended reading

Papers of particular interest, published within the period of review, have been highlighted as:

* of special interest
** of outstanding interest

Large-scale screen of inherent carbon partitioning in oлигейн yeast. Identified species with high biomass yield and a large lipid fraction. Valuable insight into the biodiversity of carbon partitioning in yeast.


In depth lipid analysis of nitrogen and phosphorus starvation. Elucidated a two step transition in phosphorus limitation as well as characterization of the fatty acid and major lipid class distributions.


Comprehensive characterization of proteome and lipid remodeling following nitrogen starvation. Revealed almost half the proteome is degraded and re-partitioned to the nitrogen assimilation machinery and storage compounds.


Systems-level analysis of the transition to nitrogen starvation at the transcript and protein level. A nitrogen conservation mechanism is initiated that reduces expression of nitrogen rich proteins; mainly elements of photosynthesis.


Targeted knockdown of genes involved in lipid catabolism using RNA interference can enhance lipid accumulation without compromising growth in the diatom *Thalassiosira pseudonana*.


A highly efficient method for targeted genome editing in diatoms on the basis of meganuclease and TALE nucleases was developed. The method was used to construct modified *Phaeodactylum tricornutum* strains with 45-fold enhanced TAG production.


Highly efficient conjugation-based method for genetic information transfer from *Escherichia coli* to diatoms. This system facilitates research in diatoms not only by enabling the characterization of genes but also by allowing introduction of entire pathways into the organism.


A constraint-based modeling approach for simulating the effect of a heterologous gene insertion is presented and, together with FBA-based approaches for strain optimization, applied to predict engineering strategies for overproduction of free fatty acids in *Escherichia coli*.

