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## Enhancing PUFA-rich polar lipids in *Tisochrysis lutea* using adaptive laboratory evolution (ALE) with oscillating thermal stress

Gachelin Manon <sup>1,\*</sup>, Boutoute Marc <sup>1</sup>, Carrier Gregory <sup>2</sup>, Talec Amélie <sup>1</sup>, Pruvost Eric <sup>1</sup>, Guihéneuf Freddy <sup>1,3</sup>, Bernard Olivier <sup>4</sup>, Sciandra Antoine <sup>1</sup>

<sup>1</sup> Laboratoire d'Océanographie de Villefranche (LOV, UMR 7093), Sorbonne Universités, CNRS, Station zoologique, 181 Chemin du Lazaret, 06230, Villefranche-sur-Mer, France

<sup>2</sup> Laboratoire Physiologie et Biotechnologie des Algues (PBA), IFREMER, Nantes, France

<sup>3</sup> SAS Inalve, 181 chemin du lazaret, 06230, Villefranche-sur-Mer, France

<sup>4</sup> Biocore, INRIA Sophia Antipolis Méditerranée, Valbonne, France

\* Corresponding author : Manon Gachelin, email address : [manon.gachelin@obs-vlfr.fr](mailto:manon.gachelin@obs-vlfr.fr)

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### Abstract :

Adaptive laboratory evolution is a powerful tool for microorganism improvement likely to produce enhanced microalgae better tailored to their industrial uses. In this work, 12 wild-type strains of *Tisochrysis lutea* were co-cultivated under increasing thermal stress for 6 months. Indeed, temperature was oscillating daily between a high and a low temperature, with increasing amplitude along the experiment. The goal was to enhance the polyunsaturated fatty acid content of the polar lipids. Samples were taken throughout the evolution experiment and cultivated in standardized conditions to analyze the evolution of the lipid profile. Genomic analysis of the final population shows that two strains survived. The lipid content doubled, impacting all lipid classes. The fatty acid analyses show a decrease in SFAs correlated with an increase in monounsaturated fatty acids (MUFAs), while changes in polyunsaturated fatty acid (PUFAs) vary between both photobioreactors. Hence, the proportion of C18-MUFAs (18:1 n-9) and most C18-PUFAs (18:2 n-6, 18:3 n-3, and 18:4 n-3) increased, suggesting their potential role in adjusting membrane fluidity to temperature shifts. Of particular interest, DHA in polar lipids tripled in the final population while the growth rate was not affected

### Key points

► Adaptive laboratory evolution on a mix of 12 *T. lutea* strains led to survival of 2 ► Thermal stress impacted cell size, total lipid cell content, and all lipid classes ► DHA cell content partitioned to polar lipids tripled throughout the experiment

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**Keywords** : *Tisochrysis lutea*, Polar lipids, Adaptive laboratory evolution (ALE), Temperature, DHA



# Introduction

Microalgae are known for their diverse and original metabolisms, leading to an ability to produce many bioactive compounds depending on species and cultivation strategies. They therefore offer potential applications in food and feed, cosmetics, and pharmaceuticals (Spolaore et al. 2006). Microalgae have an interesting potential for food and feed, as natural resources become more and more limited (FAO 2017). The phenotypic plasticity of their metabolisms allows to cope with environmental fluctuations. It can also be enhanced to better fit their cultivation conditions and optimize their industrial productivities. Adaptive laboratory evolution (ALE) is widely used for strain improvement of yeast and bacteria (Dragosits and Mattanovich 2013; Wang et al. 2014). Few studies have been published, which tested evolution protocols on microalgae, but they obtained promising results to enhance different metabolic pathways. Bonnefond et al. (2017) enhanced lipid productivity. Fu et al. (2012, 2013)

increased carotenoid content. Perrineau et al. (2013) observed higher tolerance to high salinity. Li et al. (2015) obtained cells resistant to high CO<sub>2</sub> concentration. Wang et al. (2016) enhanced the phenol degradation capacity.

Fish oils are often the lipid sources in feed, resulting in pressuring wild fish stocks. Moreover, fish oil supply is unsecured, with fluctuating quality and quantity along the year. There is therefore room for another source of lipids (Naylor et al. 2009). Some microalgae species can accumulate 20 to 50% lipid in dry biomass and even up to 80% under certain conditions (Santos-Sanchez et al. 2016; Bellou et al. 2014). Their fatty acid (FA) profiles can be rich in specific essential fatty acids,  $\omega$ 3-, and  $\omega$ 6-long-chain polyunsaturated fatty acids (LC-PUFAs) (over 20% of total fatty acids). Temperature is a factor that strongly influence growth rate and lipid content (Roleda et al. 2013; Zhu et al. 2016) with strong species-specific effects (Zhu et al. 2016; Renaud et al. 1995; Thompson et al. 1992; Renaud et al. 2002; Bonnefond et al. 2017). Temperature also affects the saturated/unsaturated FA ratio and profile

(Renaud et al. 1995; Thompson et al. 1992). It has been hypothesized that this acclimation contributes to maintain membrane fluidity (Ackman et al. 1968; Mortensen et al. 1998).

Temperature has already been used as a selection factor in long-term adaptive experiments to improve lipid productivity. A 6-month experiments with *Tisochrysis lutea* consisting in applying daily fluctuating temperatures of increasing amplitude (Bonnefond et al. 2017) contributed to enlarge thermal niche (+ 16%) together with the maximal growth rate (+ 9%). It also resulted in modifying the FA profile. The key idea of this evolution experiment was to maintain a dynamic selection pressure to force the cells to adapt and therefore survive to their environment.

Here, we use ALE for enhancing *Tisochrysis lutea* in PUFA-rich polar lipids. This oleaginous microalga is commonly used in aquaculture (Rico-Villa et al. 2006) especially for its significant content of docosahexaenoic acid (DHA) (Renaud et al. 1995; Thompson et al. 1992; Liu et al. 2013). Twelve wild-type strains were equi-proportionally mixed in the inoculum to increase initial genomic diversity. This mix of strains was continuously cultivated for 6 months under thermal stress. The ALE experiment consisted in daily fluctuating temperature with increasing amplitude. The stress intensity (amplitude of the fluctuations) was increased weekly, balancing between a reasonable growth rate and the induction of durable modifications in the fatty acid profile of polar lipids. Genomics analyses were performed at the end of each selection cycle to determine the remaining strains and track genomic modifications.

## Material and methods

### Microalgal strains

The 12 strains of the photoautotrophic microalga *Tisochrysis lutea* used in this study (supplementary data, table S1) originated from culture collections (CCAP927/14, CCMP463, RCC179, RCC1344, RCC3691, RCC3692, RCC3693, RCC3699, NIVA 4-91, IMFG, and Argenton 1998). The strain Tiso S-5 (maintained at PBA laboratory, IFREMER, Nantes, France) was derived from the original strain CCAP927/14 after a sequential selection procedure (1% lowest lipid content).

### Cultivation device and protocol

#### The adaptive laboratory evolution device

The microalgae were grown under computer-controlled conditions in duplicate flat panel photobioreactors (TL1 and TL2), 6 cm culture thickness. These home-designed water-jacketed 1.8 L photobioreactors (called “selectiostats,” see Fig. 1) were specially designed to maintain and survey continuous cultures for several months. Bio-fouling was prevented as much as possible by bubbling 0.2  $\mu\text{m}$  filtered air along the transparent walls of the selectiostats, and their simple geometry makes the optical measurements of PAR and turbidity reliable.

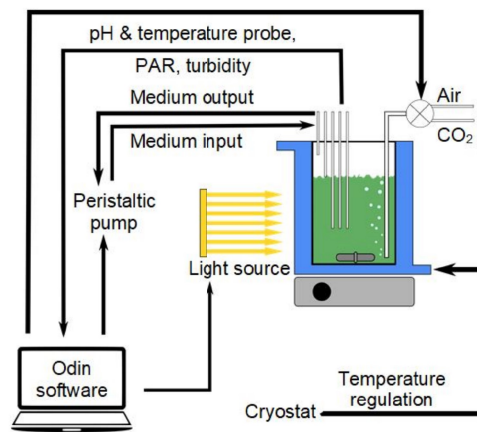


Fig. 1 Simplified diagram of a selectiostat and its controlled parameters

Cultures were grown in turbidostat mode, i.e., dilution rate was automatically regulated to maintain the turbidity and thus cell density between 3.5 and 4 million cells/mL. Turbidity was estimated from the beam attenuation of a light source at 750 nm through the thickness of the culture. It was kept sufficiently low to avoid self-shading and nutrient limitation. Continuous light was provided by a LED panel (Nichia NVSL219BT 2700°K) placed on one side of the photobioreactor. Intensity of incident light in the visible spectrum (PAR; photosynthetically active radiation) was set at 300  $\mu\text{mol photons/m}^2/\text{s}$ , and occasionally controlled with a spherical Biosphere Instrument QSL-100 probe. PAR was also continuously measured with a flat Skye PAR Quantum Sensor probe positioned against the transparent wall on the other side of the photobioreactor. Measured PAR behind the photobioreactor was oscillating between 150 and 200  $\mu\text{mol photons/m}^2/\text{s}$ . Cultures were homogenized through magnetic stirring and bubbling. pH was continuously measured and regulated at 8.2 by controlled micro-injections of CO<sub>2</sub> in the bubbled air. The renewing medium was prepared in 20-L Nalgen tanks with autoclaved (121 °C, 30 min), 0.2- $\mu\text{m}$  filtered natural seawater, enriched with sterile Conway-Walne medium (Walne 1974). Medium was introduced in the selectiostats with a peristaltic pump (Gilson Minipuls 3), through a 0.22- $\mu\text{m}$  Whatman filter. The flow-rate of the peristaltic pumps was automatically regulated and regularly checked.

As temperature was the stressing factor, special attention has been paid to its control. In particular, the stainless-steel frame of the selectiostats were designed to reduce thermal inertia of cultures as much as possible that allowed a rapid circulation of fresh water whose temperature was regulated with a programmable cryostat (Lauda Proline RP845). Temperature regime imposed to selectiostats (Table 1) followed a succession of cycles, each of them consisting in a diel alternation of 16 and 8 h at high (HT) and low (LT) temperatures, respectively. The low and high temperatures were chosen in such a way that the mean daily temperature remained constant throughout the whole experiment (around 24 °C), except for the last cycle. The decision to shift from one cycle to the next one depended on the culture fitness (estimated from the growth rate), according to the “ratchet” protocol experienced by Bonnefond et al. (2017).

Table 1: Temperature stress cycles implemented in the selectiostats during the 150 days of ALE experiment. \*As the cultures collapsed on the second day of S7 cycle, they were restarted with less stressful conditions, identical to those of S5 cycle

Stress cycle	HT (°C)	LT (°C)	Cycle duration (days)	Mean temperature (°C)
S1	26	21	15	24.33
S2	27	19	7	24.33
S3	28	17	7	24.33
S4	29	16	7	24.67
S5	30	14	15	24.67
S6	31	12	15	24.67
S7	32	10	2*	24.67
S8	30	14	24	24.67
S9	31	12	7	24.67
S10	32	10	7	24.67
S11	33	8	7	24.67
S12	34	6	15	24.67
S13	35	8	7	26.00

A stress indicator (Fig. 2) cumulating the pressure that the cells have faced in the selectiostat from initial experimental time to current time  $t$  was calculated as the cumulated difference from the reference temperature:

$$\varphi(T) = \int_0^t |T - T_{ref}| dt$$

$T$  being the temperature at time  $t$  and  $T_{ref} = 24$  °C, the nonstressful temperature chosen as reference.

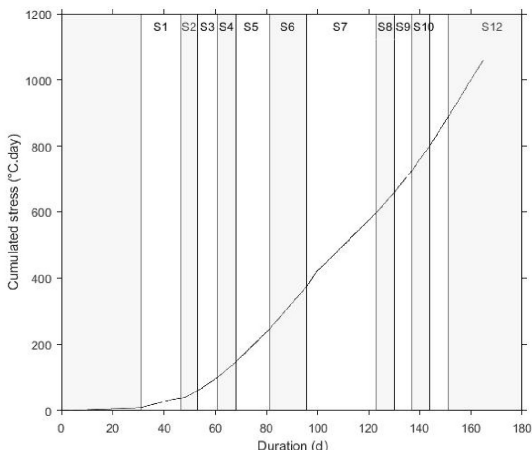


Fig. 2 Calculated stress indicator representing the cumulated stressed faced by the cells in the selectiostats during the experiment

Before use, selectiostats were sterilized with 10% HCl for 20 min and rinsed with sterile culture medium (note that they had to be cleaned similarly 3 times during the experiment to eliminate biofouling). They were inoculated with an assemblage of the 12 strain cultures at the same concentration (15,000 cells/mL). The ODIN software (developed by Inria) allowed automatic control of the selectiostats, by recording at high frequency the parameters measured online (PAR, turbidity, pH, temperature, speed of dilution pump), and by controlling the pump speed to maintain constant turbidity (see Fig. 1).

Cell density was estimated, using a relationship established between the continuous acquisitions of turbidity and cell concentrations measured twice a month with a Multisizer 3 Beckman coulter. Cell density in continuous cultures varies accordingly to the following equation:

$$dx/dt = \mu x - Dx$$

with  $x$ , the cell concentration;  $\mu$ , the net growth rate; and  $D$ , the dilution rate (medium flow rate divided by culture volume). When the turbidostat is at equilibrium ( $dx/dt = 0$ ), the net growth rate equals the dilution rate.

### Standardized measurement of the overall mixed population properties

At the end of each stress cycle, a benchmark experiment in standardized conditions was performed to measure the properties of the overall mixed population. Triplicate batch cultures of 85 mL were inoculated with samples taken in each selectiostat. Duplicate batch cultures inoculated with the initial strain *T. lutea* CCAP 927/14 was systematically cultivated in parallel and served as reference. Indeed, CCAP 927/14 is the strain of *T. lutea* most used in aquaculture. All the cultures were controlled with the Multi-Cultivator MC 1000-OD culturing device (Photon Systems Instruments, PSI). Growth conditions were the same as in the selectiostats, except that T°C was maintained constant at 20 °C, and pH was not regulated. Optical density (OD) was measured continuously at 720 nm in each flask culture to follow the growth in real time. The maximum growth rate was calculated during the exponential phase, using OD measured after 24 h post-inoculation and whose values did not exceed 0.3 (i.e., the value below which OD is linearly correlated with cell density).

Culture samples of 2 mL were filtered in duplicate on precombusted glass-fiber filters (Whatman GF/C, threshold 1.2  $\mu$ m) and kept at 75 °C until particulate carbon and nitrogen analyses made with a CHN analyzer (2400 Series II CHNS/O, Perkin-Elmer).

As the biomass reached in each batch culture of the PSI Multi-Cultivator was not sufficient for accurate lipid estimation, triplicate cultures were pooled together to provide one large sample. Two sub-samples of 100 mL and 50 mL, centrifuged at 3000 rpm during 15 min (10 °C) after addition of 50  $\mu$ L of FeCl<sub>3</sub> (50 mg/mL) to facilitate sedimentation and formation of cell pellets, were stored at -80 °C until extraction.

## Lipid class separation

Total lipids were extracted following the method previously described by Bligh and Dyer (1959). The total lipids were then dried using a Büchi R-200 rotavapor and then freeze-dried to remove solvent traces, then determined gravimetrically and stored at  $-80\text{ }^{\circ}\text{C}$  under nitrogen atmosphere.

The lipid classes (i.e., neutral lipids (NL), glycolipids (GL), and phospholipids (PL)) were obtained by solid-phase extraction (SPE) fractionation on silica columns (Interchim, SI-S-500/6 columns) with 6 volumes of chloroform (NL), 4 volumes of acetone (GL), 6 volumes of methanol, and 2 of methanol/10% ammonia (PL) (Vorbeck and Marinetti 1965), respectively. Each class was then dried under vacuum, determined gravimetrically, and stored at  $-80\text{ }^{\circ}\text{C}$  under nitrogen atmosphere.

## Fatty acids analysis

To determine the fatty acid (FA) composition and content, the lipid class samples were methylated following the protocol described by Morrison and Smith (1964), with 7% boron trifluoride in methanol and toluene as extraction solvent for the fatty acid methyl esters (FAME). Each sample was washed 3 times to ensure maximal FAME recovery. The FAME profiles for both phospholipids and glycolipids were obtained separately and analyzed quantitatively (fg FAME/cell) and qualitatively (% total fatty acids, TFA). FAME were analyzed by gas chromatography (GC) on a Perkin-Elmer XL Autolab GC equipped with a flame ionization detector (FID) and a (Restek, 30 m length, 0.32 mm internal diameter). The column was operated isothermally at  $182\text{ }^{\circ}\text{C}$  while injector and detector were maintained at  $250\text{ }^{\circ}\text{C}$ . Helium was used as carrier gas at 7 psig. The variability was 3% for major components, < 9% for intermediate ones, and 25% for minor components (< 0.5% of total FAME); commercial nonadecanoic acid (19:0) in chloroform was co-injected with FAME sample for FA quantification.

## DNA extraction and sequencing

The cell pellets of samples TL1 and TL2 stress cycles S1, S8, and final population S12 were stored at  $-80\text{ }^{\circ}\text{C}$  prior

to DNA extraction, using a phenol-chloroform method described by Hu et al. (2004). Sequencing was carried out with an Illumina HiSeq3000, with 20 to 30  $\mu\text{g}$  of DNA, by the genomic platform GeT-PlaGe, INRA (Toulouse, France). The libraries were realized in TrueSeq Illumina kit ( $2 \times 150$  pairedend sequenced for a DNA fragment of 430 pb on average). Raw data was filtered to first remove all Illumina residual adapters, and to finish keep the reads with sufficient quality. It was then treated with softwares TrimGalore and home-made script to discard all reads with a mean sequencing score under Q30 or a length shorter than 120 pb. After filtering, more than 80% of the total reads were kept for each sample. The version 2.0 of reference genome of *Tisochrysis lutea* (approx. 85 Mb) (Bertheliet et al. 2018) was used to map the samples' genomes with BWA-MEM software (run with default parameters). For all samples, more than 86% of the pairreads were aligned properly and in average 225 depth of sequencing genome were obtained for each sample. A search for single-nucleotide polymorphism (SNPs) was performed with Freebayes software (minimal alleles' depth 20, minimal frequency in total reads 5%, pooled-continuous option) (Garrison and Marth 2012). The results were edited using a home-made script.

To identify the strains coexisting in the cultivated population during the ALE experiment, each of the 12 initial strains were sequenced independently using the same protocol. Some specific SNPs were identified for each of them. These SNPs were used to allow the identification of the strains actually represented in the cultures during the experiment with a cross-comparison of the SNPs in each strain and the two cultivated populations. A presence index was calculated for each initial strain, by dividing its number of SNPs with the total number of SNPs.

## Results

### Overall evolution in the selectiostat

An example of the monitoring of selectiostat TL1 over 1 week is presented in Fig. 3.

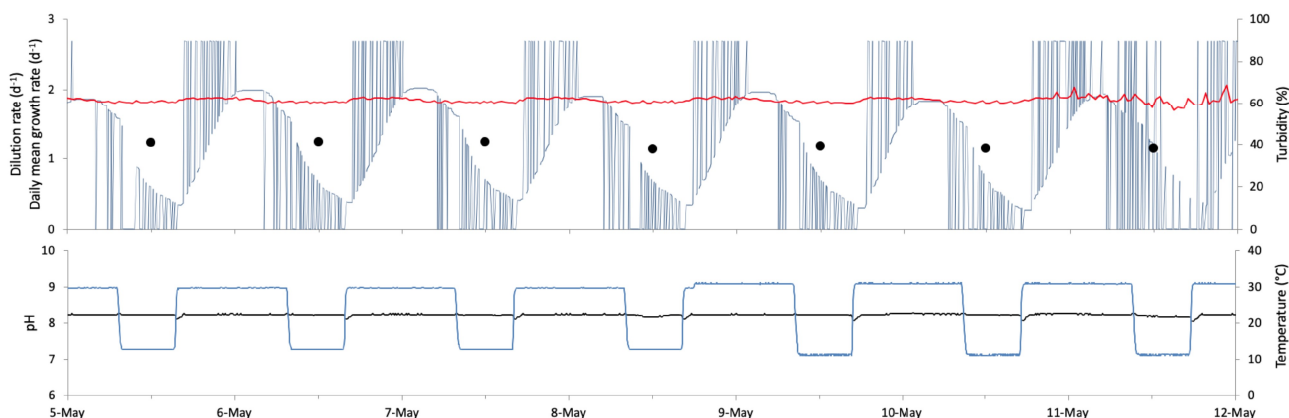


Fig. 3 Monitoring of selectiostat TL1 during 1 week. Upper graph, turbidity in red, dilution rate in blue, and mean daily growth rate in black dots. Lower graph, pH in black, and temperature in blue

Turbidity was well regulated by the automatic activation of the peristaltic pump, pH was stable, and temperature variations rapidly tracked the desired pattern between low and high temperatures, meaning that the temperature regulation through the cryostat was efficient. At the beginning of each high temperature (HT) period, the pump regularly reached its maximum speed to counterbalance growth and keep the turbidity stable by diluting the selectostat. The pump stabilized when half of the HT period was reached, which implies that culture growth slowed down. Finally, the pump activity reduced, with frequent stops, from  $\frac{3}{4}$  of the HT period and throughout the whole LT period. During that time, the growth of the culture is therefore very limited, as little dilution of the culture is then necessary to maintain a constant turbidity.

## Phenotype evolution in standardized cultivation conditions

### Growth rate

The recorded growth rates estimated in the standardized benchmark conditions ( $0.79 \pm 0.17$  /day) are consistent with values reported by Bonnefond et al. (2017), who obtained growth rates ranging between 0.5 and 0.6 /day at 20 °C, and up to 1.2 /day at the optimal growth temperature of the cultures. Growth rate (Fig. 4) was normalized according to the reference. The variations are slightly more important for culture TL1 than for TL2 (0.87–1.31 and 0.88–1.18, respectively). However, a T test comparing the growth rate of each culture with the control shows no significant difference (pvalues 0.66 for TL1 and 0.76 for TL2). This shows that the selection process did not affect the overall growth rate of the population throughout the experiment and contrasts therefore with previous results obtained on mono-strain experiments (Bonnefond et al. 2017; Fu et al. 2012). A possible explanation could be that different sub-populations are coexisting and alternatively growing depending on the conditions, thus reducing the growth rate variations.

### Nitrogen content

Nitrogen content in the cells tended to slightly decrease during the experiment while carbon content doubles and stabilizes from stress cycle S5 onwards for TL1 and S8 for TL2 (supplementary data, figure S2), leading to an overall increase in C/N ratio despite the strong variability between cycles. This increase is consistent with the cell volume increase in both cultures from stress cycle S6 onwards (Table 2). The slight evolution of cell nitrogen content, as the variability of the normalized C/N ratio (Fig. 4), could be linked to differential growth rates or result from different stresses when inoculating the bench-PSI Multi-Cultivator device. After 7 days, the initial delay may have induced spreading of the time when cells are N limited, and thus, different N limitations were reached in the control or in the cultures.

Table 2: Cell concentration, volume, and diameter in the Multi-Cultivator devices upon harvest for lipid analysis

Culture Stress cycle	Reference			Culture TL1			Culture TL2		
	cell/mL	Biovolume ( $\mu\text{m}^3$ )	Diameter ( $\mu\text{m}$ )	cell/mL	Biovolume ( $\mu\text{m}^3$ )	Diameter ( $\mu\text{m}$ )	cell/mL	Biovolume ( $\mu\text{m}^3$ )	Diameter ( $\mu\text{m}$ )
S1	1.20E+07	48	4.44	1.31E+07	44	4.31	1.33E+07	43	4.29
S2									
S3	1.93E+07	43	4.28	1.73E+07	42	4.26	1.87E+07	42	4.25
S4	1.08E+07	48	4.44	1.49E+07	41	4.22	1.45E+07	40	4.20
S5	1.63E+07	42	4.24	1.43E+07	49	4.43	1.57E+07	44	4.31
S6				6.79E+06	66	4.84	7.62E+06	53	4.51
S7									
S8	1.64E+07	43	4.30	1.42E+07	49	4.42	1.40E+07	51	4.46
S9	1.58E+07	41	4.22	1.00E+07	47	4.33	1.11E+07	49	4.40
S10	1.46E+07	46	4.39	2.19E+07	54	4.55	2.43E+07	48	4.39
S11	1.22E+07	42	4.26	8.19E+06	62	4.71	9.26E+06	54	4.51



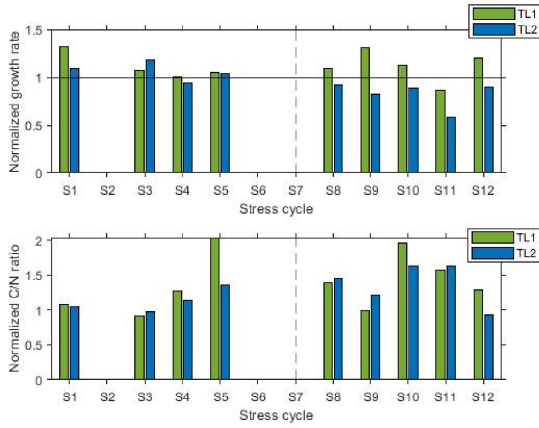


Fig. 4 Normalized growth rates\* and normalized C/N ratio\* of cultures in standardized conditions (PSI Multi-Cultivator device) throughout the experiment, triplicates. No data for stress cycles S2 and S6; cultures collapsed during stress S7. \*Calculated as fold change in growth rate or C/N ratio normalized to the reference

### Total lipid content and lipid classes

The mass of total lipid per cell gradually increased along the selection process. From stress cycle S1 to S12, a pro-

gressive enrichment of total lipid content per cell was observed. The gain between two successive stress cycles from S3 to S11 ranged from 8 to 20%. This led in fine to a cellular lipid content increased by 2.1 for TL1 and 1.8 for TL2, whereas the reference culture stayed constant at an average value of  $6.16 \pm 0.62$  pg/cell (Fig. 5). The cell lipid contents of both cultures are not significantly different ( $p$  value of 0.32). The lipid productivity showed the same increasing trend. At the end of the experiment, total and neutral lipid contents were similar to those measured by Da costa et al. (2016, 2017) on their lipid-rich *T. lutea* strain (11.2 pg/cell and 4.3–5.1 pg/cell, respectively) in similar growing conditions (continuous  $200 \mu\text{mol photons/m}^2/\text{s}$  white light, 20–23 °C, Conway medium).

Cellular lipid content for each lipid class (5.7–7.8 pg/cell for total lipids, 3.1–7.5 pg/cell for neutral lipids, 1–3.1 pg/cell for phospholipids, and 1–2.4 pg/cell for glycolipids) was slightly higher than those measured (via HPTLC) by Carrier et al. (2018) (3–6 pg/cell for total lipids, 0.4–3.1 pg/cell for triacylglycerol, 0.9–1.3 pg/cell for phospholipids, and 1.4–1.7 pg/cell for glycolipids). The quantitative measurements of each major lipid class after fractionation (Fig. 5) show a cellular enrichment of every class from stress cycle S5 onwards.

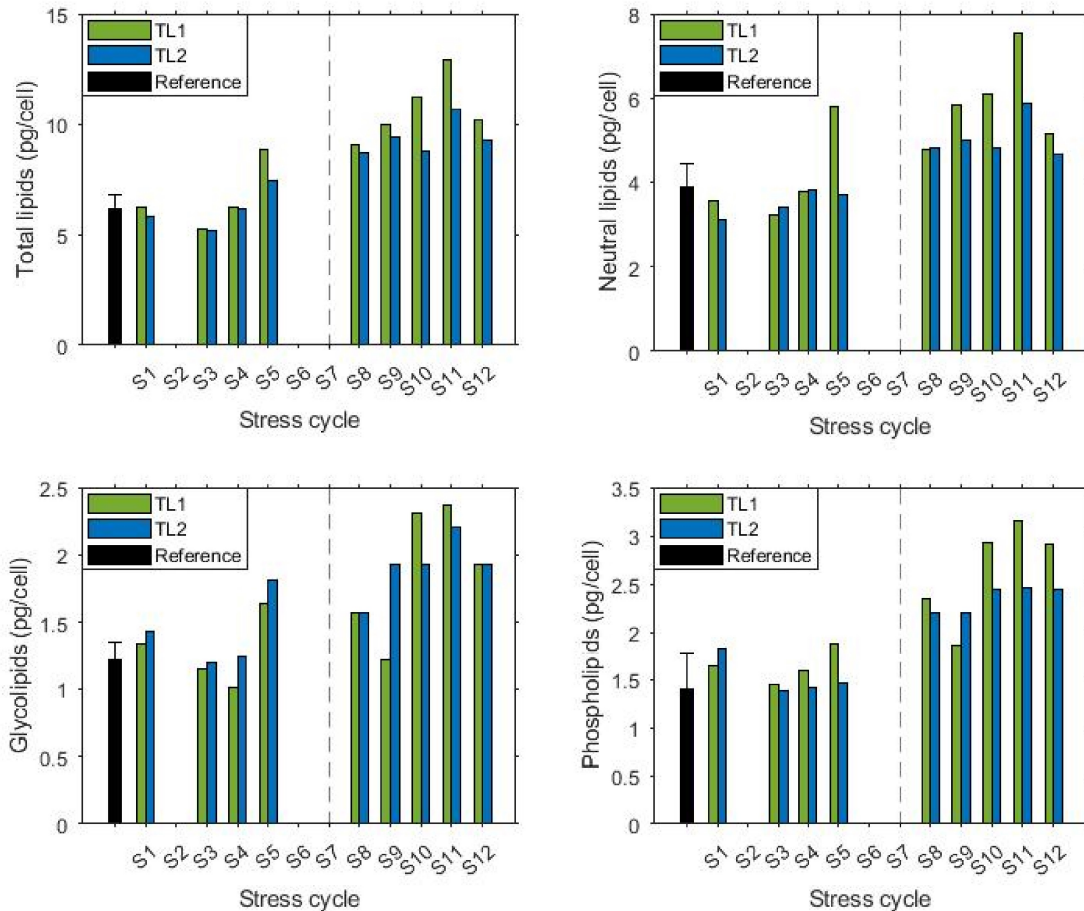


Fig. 5 Evolution of cellular total lipid and lipid class contents (pg/cell) throughout the stress cycles; cultures in standardized cultivation conditions in the Multi-Cultivator. No data for stress cycles S2 and S6; cultures collapsed during stress S7. Reference is an average of 9 measurements on independent cultures

## Fatty acids composition

The main fatty acids partitioned to phospholipids were 14:0, 16:0, 18:1 n-9, 18:3 n-3, 18:4 n-3, and 22:6 n-3 in the 2 stressed and reference cultures. The same fatty acids were incorporated into glycolipids together with 18:2 n-6. Overall, fatty acid profiles of polar lipids (GL + PL) (Table 3) were similar to those reported by Marchetti et al. (2018), and closely similar to total fatty acid profile (16% for 14:0, 14.5% for 16:0, 4.2% for 16:1 n-7, 20.1% for 18:1 n-9, 3.6% for 18:3 n-3, 17.4% for 18:4 n-3, 2.5% for 18:5 n-3, and 8.5% for 22:6 n-3) obtained by Volkman et al. (1989). However, they differed from the profiles obtained

by Renaud et al. (1995) who cultivated *T. lutea* at 15, 20, 25, and 30 °C. They obtained significantly higher amounts of 14:0 (except at 15 °C), 18:4 n-3 (>20%, > 22%, respectively) and 18:5 n-3 but only at the lowest temperature (15 °C). On the other hand, they observed lower amounts of 16:0, 18:1 n-9, 18:2 n-6, 22:5 n-6, and DHA except at 15 °C (< 9.6%, < 8.7%, < 4.8%, < 1%, and < 9.7%, respectively). However, Renaud et al. (1995) and Volkman et al. (1989) studied total lipids, not polar lipids; the observed variations can therefore be caused by the specific fatty acid composition of polar lipids, which can greatly differ from the overall fatty acid composition.

Fatty acid (% TFA)	Reference	TL1-S1	TL1-S12	TL2-S1	TL2-S12
<i>14:0</i>	<i>13.99 ± 3.23</i>	<i>15.21</i>	<i>11.75</i>	<i>16.94</i>	<i>8.01</i>
ISO15:0	0.22 ± 0.10	0.22	0.09	0.12	0.04
15:0	0.55 ± 0.43	0.70	0.16	0.29	0.13
ISO17:0	0.19 ± 0.16	0.30	0.22	0.00	0.39
<i>16:0</i>	<i>15.26 ± 2.34</i>	<i>13.58</i>	<i>14.91</i>	<i>16.39</i>	<i>9.61</i>
17:0	0.25 ± 0.21	0.35	0.06	0.25	0.06
18:0	0.54 ± 0.18	0.52	0.31	0.35	0.30
<b>Total SFAs</b>	<b>31.25 ± 5.14</b>	<b>31.02</b>	<b>27.69</b>	<b>37.75</b>	<b>18.67</b>
14:1 n-7	0.13 ± 0.13	0.36	0.16	0.16	0.00
14:1 n-5	0.23 ± 0.09	0.35	0.21	0.50	0.22
<i>16:1 n-7</i>	<i>3.71 ± 0.25</i>	<i>3.27</i>	<i>3.72</i>	<i>3.62</i>	<i>4.01</i>
16:1 n-5	0.15 ± 0.10	0.06	0.05	0.21	0.00
17:1 n-10	0.08 ± 0.07	0.13	0.03	0.14	0.08
<i>18:1 n-9</i>	<i>16.22 ± 3.71</i>	<i>10.63</i>	<i>20.75</i>	<i>12.81</i>	<i>22.35</i>
18:1 n-7	0.35 ± 0.22	0.40	1.17	0.35	0.00
18:1 n-5	0.12 ± 0.11	0.00	0.05	0.00	0.00
22:1 n-13 + 11	0.09 ± 0.13	0.13	0.00	0.15	0.17
<b>Total MUFAs</b>	<b>21.32 ± 3.62</b>	<b>15.42</b>	<b>26.18</b>	<b>18.07</b>	<b>26.87</b>
16:2 n-6	0.20 ± 0.17	0.22	0.03	0.25	0.00
16:2 n-4	1.05 ± 0.12	0.77	0.76	1.18	0.82
<i>18:2 n-6</i>	<i>4.48 ± 0.94</i>	<i>4.37</i>	<i>4.94</i>	<i>3.90</i>	<i>6.10</i>
20:2 n-6	0.14 ± 0.04	0.13	0.08	0.15	0.09
16:3 n-4	0.07 ± 0.02	0.11	0.05	0.13	0.07
16:3 n-3	0.26 ± 0.26	0.29	0.09	0.16	0.13
18:3 n-6	0.16 ± 0.08	0.06	0.14	0.08	0.22
<i>18:3 n-3</i>	<i>6.47 ± 0.56</i>	<i>6.97</i>	<i>7.90</i>	<i>6.75</i>	<i>7.94</i>
20:3 n-3	0.16 ± 0.06	0.04	0.00	0.05	0.11
16:4 n-3	0.06 ± 0.03	0.03	0.05	0.09	0.02
16:4 n-1	0.16 ± 0.08	0.18	0.10	0.08	0.11
<i>18:4 n-3</i>	<i>14.68 ± 2.99</i>	<i>16.40</i>	<i>15.79</i>	<i>14.82</i>	<i>23.64</i>
20:4 n-6	0.10 ± 0.07	0.12	0.05	0.06	0.04
22:4 n-6	0.09 ± 0.06	0.05	0.00	0.18	0.05
<i>18:5 n-3</i>	<i>3.77 ± 0.69</i>	<i>5.57</i>	<i>3.25</i>	<i>3.35</i>	<i>5.50</i>
20:5 n-3	0.26 ± 0.10	0.45	0.24	0.35	0.18
21:5 n-3	0.64 ± 0.26	0.78	0.55	0.50	0.58
<i>22:5 n-6</i>	<i>2.30 ± 0.44</i>	<i>2.39</i>	<i>1.87</i>	<i>2.27</i>	<i>1.04</i>
22:5 n-3	0.13 ± 0.07	0.14	0.06	0.23	0.07
<i>22:6 n-3</i>	<i>11.86 ± 1.92</i>	<i>14.08</i>	<i>10.12</i>	<i>11.94</i>	<i>7.12</i>
<b>Total PUFAs</b>	<b>47.43 ± 6.63</b>	<b>53.55</b>	<b>46.13</b>	<b>44.18</b>	<b>54.47</b>
<b>Others</b>	0.89	0.61	0.28	1.21	0.81
<b>Total n-3 PUFAs</b>	<b>39.29 ± 6.95</b>	<b>44.77</b>	<b>38.05</b>	<b>39.23</b>	<b>45.29</b>
<b>Total n-6 PUFAs</b>	<b>7.46 ± 1.80</b>	<b>7.356309</b>	<b>7.11</b>	<b>4.89</b>	<b>7.54</b>
<b>n-3/n-6 ratio</b>	<b>5.13</b>	<b>6.09</b>	<b>5.35</b>	<b>5.55</b>	<b>6.00</b>

Table 3 Proportions of fatty acids in the polar lipids representing more than 0.1% TFA at the beginning and the end of the experiment. Cultures in standardized conditions (PSI Multi-Cultivator device). Reference is an average of 8 measurements on independent cultures. Major FAME (> 2% TFA) of the fatty acids profile are emphasised in italics

The fatty acid profiles at S1 were similar to the reference for both cultures, except 18:1 n-9 which is slightly lower compared to the reference culture (10.63% and 12.81% for TL1 and TL2, respectively,  $16.22 \pm 3.71\%$  for the reference). Similarly, the total saturated fatty acid (SFAs) and PUFA proportions were similar; the lower total monounsaturated fatty acid (MUFAs) in the cultures is linked to their lower 18:1 n-9 content. At S12, the selection process seemed to have impacted the fatty acid composition, especially of polar lipids. The two major SFAs (14:0 and/or 16:0) diminished while the proportion of C18-MUFAs (18:1 n-9) and most C18-PUFAs (18:2 n-6, 18:3 n-3 and 18:4 n-3) increased. Changes in PUFAs (in particular DHA) fluctuate between the two photobioreactors.

The cellular content of DHA partitioned to polar lipids nearly tripled between the beginning and the end of the ALE experiment. It was mainly due to the important increase of the total lipid cell content. DHA increase is significant from stress cycle S8 onwards (Fig. 6). Indeed, in the first half of the experiment, before cycle S7, the DHA content partitioned to polar lipids varied between 50 and 100 fg/cell. In the second half of the experiment, it reached 200 fg/cell during stress cycles S8 and S9, then increased of  $13 \pm 3\%$  between stress cycles S9 to S12 for TL1, and 20 to 23% for TL2. This difference could result from the intense stress during cycle S7 which led to the culture crash. It may have caused a stronger selection, and possibly the emergence of a sub-population with a higher DHA content.

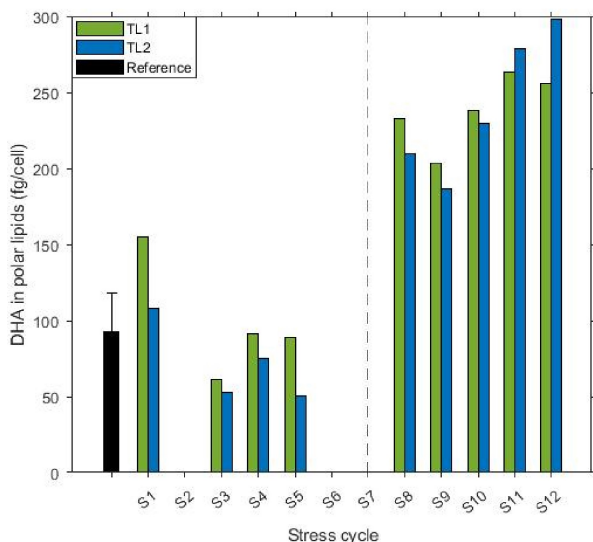


Fig. 6 Evolution of cellular DHA content partitioned to polar lipids. No data for stress cycles S2 and S6; cultures collapsed during stress S7. Reference is an average of 9 measurements on independent cultures

## Genomic analysis

Samples for genomic analysis were taken and analyzed at the end of stress cycle S1, S8, and S12. Through the SNP search, mutations corresponding to a total change of genotype (disappearance of the initial genotype) were detected

in both cultures and for each sequenced sample. It demonstrates an evolution of the initial population rather than a simple selection of some initial characteristics.

The overall presence indexes were calculated from crosscomparison of the global SNPs with the initial ones. Presence indexes inferior to 0.15 mean that the corresponding strain is likely to be absent from the population. They ranged between 0.08 and 0.16 except for strains CCMP 463 (0.40) and RCC 1344 (0.50) (Table 4). This demonstrates that these two strains dominate the population, since the beginning of the experiment. Both are originated from low latitudinal coastal areas from north hemisphere: CCMP 463 was isolated in the Caribbean archipelago (Turks and Caicos Islands) and RCC 1344 along the Spanish Atlantic coast (Bendif et al. 2013). This data suggests that, out of the initial twelve-strain assemblage, only CCMP 463 and RCC 1344 compose the population undergoing the stressed cultivation and the final population of *T. lutea*. The 10 other strains were cleaned up during the build-up phase (batch growth and turbidostat stabilization) and during first stress cycle (S1) of the experiment.

Total mutation and selection events detected at stress cycle S1, S8, and S12 stayed quite stable for both cultures throughout the experiment (5874, 4743, and 4783 for TL1 and 4870, 3725, and 5232 for TL2, respectively). This is surprising considering the culture crash at stress cycle S7, with the loss of a large fraction of the cells and the observed change in the lipid phenotype, suggesting that this event had little impact on the genomic diversity of the population.

Few SNPs led to a complete change of the genotype, with the disappearance of the initial one: 49 for culture TL1 and 17 for culture TL2. Further analyses on those specific mutations were carried out to determine whether they could be partially responsible for the phenotype changes observed on the final populations. For culture TL2, 2 out of the 17 SNP mutations were localized in coding regions, and 9 out of 49 for culture TL1. Putative-associated functionality search on these genes remains on the prospective side, as very little is known on the function of genes in *T. lutea*. Researches of homologous genes in Uniprot database did not lead to associated functionality for 7 of the genes; 2 could be GAG-POL transposons. The last 4, all detected in culture TL1, could be impacting a ClpC protease, a dihydrolipoamide acyltransferase part of the pyruvate dehydrogenase complex, a trehalose-phosphate synthase, and a ribonucleotide-diphosphate reductase. Extensive studies of these candidate genes must be conducted to confirm their potential roles. Especially, the ClpC protease could act as a chaperone during heat shocks by targeting denatured proteins and addressing them for degradation (Uniprot database, identified function in *Staphylococcus aureus*). If that role was confirmed in *T. lutea*, it would be very interesting to investigate how it impacts the operation of the protein and potentially link it to the dynamic stress protocol.



## Discussion

In the selectiostat devices, the cell growth rate presented daily oscillations (Fig. 3), with constant mean daily growth rate. This behavior was different from observations on singlestrain continuous cultures by Bonnefond et al. (2017). It might be due to the coexistence of several sub-populations in the selectiostats, which grow alternatively and thus reduce the apparent variations in the growth rate. A closer look at daily growth rate pattern shows a clear phasing with temperature oscillations. This is probably due to the synchronization of cell division with temperature as already observed in *D. salina* (Bonnefond et al. 2016). Most of the cells divide at the same time during the first half of each HT period, generating an increase in turbidity.

The selection process occurring in the selectiostats seemed to favorably impact lipid metabolism with higher lipid content in all lipid classes; however, polar lipids doubled while neutral lipids increased only by 1.5-fold. This is also visible in the calculated proportions (supplementary data, figure S1), which show a slight decrease for neutral lipids and concomitantly a slight increase in phospholipids while glycolipids remain stable throughout the experiment. These data, and notably the polar lipids changes, are consistent with the measured increase of the cell volume during the experiment (Table 2). Indeed, this means that more cell membranes are produced, thus increasing the quantity of phospholipids, but not necessarily more chloroplasts since glycolipids compose thylakoidal membranes and not cell membranes. The accumulation of a pool of MUFAs and PUFAs partitioned to polar lipids inside the cell is also likely to allow quicker membrane acclimation to temperature fluctuations. Lipid accumulation (neutral and polar) might also contribute to the cell metabolic response for keeping cellular redox homeostasis. Indeed, fatty acids are highly reduced molecules, which can be used as an alternative sink for electrons produced by the photosynthetic chains, especially at colder temperature, avoiding an over-reduction of the NADP pool (Zhang et al. 2016).

When growth temperature of *T. lutea* was increased from 10 to 20 °C, Thompson et al. (1992) measured a strong metabolic shift from 16:1 n-7 to 18:1 n-9 and greater production of FA 18:5 n-3. This acclimation strategy is coherent with the lipid adaptation obtained in the ALE experiment. Since 18:1 n-9 is the first unsaturated fatty acid in the PUFA biosynthesis pathway, this could be a metabolic shift of the cells to prioritize the PUFA biosynthesis pathway over the one of SFAs. The rationale behind the 18:5 n-3 shifts is unclear as it diminishes of 42% in TL1 and increases of 64% in TL2 between stress cycle S1 and S12. However, most C18-PUFAs (18:2 n-6, 18:3 n-3, and 18:4 n-3) increased from cycles S1 to S12. These modifications in the FA profile, and more specifically the accumulation of C18 fatty acids, along with the increase in cell bio-

volume, suggest a strategy to maintain membrane fluidity for adapting to fluctuating temperatures. It is consistent with previous observations and conclusions when studying both cell adaptation with a similar approach (Bonnefond et al. 2017) and cell acclimation (Renaud et al. 1995).

Our new population of *T. lutea* fits the major criteria to be used as feed for bivalve larvae (Rico-Villa et al. 2006), i.e., a rapid growth and an easy cultivation (these parameters were not impacted by the evolution process of the experiment). The larger cell diameter (around 5  $\mu\text{m}$  with an increase detected from stress cycle S6 onwards, Table 2) is likely to present an improved nutrition value compared to *T. lutea* wild-type. Indeed, the observations made by Da Costa et al. (2016) on the growth and survival of *C. gigas* larvae fed with *T. lutea* suggest that a high level of NL within total lipids impedes the larvae's survival. The new mixed-population, with higher polar lipids' content compared to NL, is likely to provide a more balanced lipid source as the fatty acid profile of polar lipids (see below) has hardly been affected. Moreover, the importance of PUFAs for the development of *C. gigas* larvae has been extensively studied (Chu and Webb 1984; Rico-Villa et al. 2006; Da Costa et al. 2016). It was shown that EPA and DHA contents in the range of 7–17% and 7–14% of TFA, respectively (Rico-Villa et al. 2006), were necessary to ensure a good larval development. *Tisochrysis lutea* produces only traces of EPA but its DHA content is within the necessary proportion range. However, the cells of our new population are richer in total lipids compared to the reference; therefore, it could be a better source of DHA for oysters.

The presence indexes calculated from the SNPs are representative of the probability to exist in the evolving population. It therefore does not mean that the two remaining strains were equally represented in the population. Moreover, as the first sampling was done at the end of S1, it is not possible to know which of the 10 strains disappeared during the build-up phase (because of a slower growth), and which disappeared during the first stress cycle (because they could not acclimate). Strain CCMP463 presented the same pattern in both culture during the experiment: the presence index increased between S1 and S8, then stabilized between S8 and S12. On the other hand, strain RCC 1344 behaved slightly differently in both cultures. Its presence index increased from 0.11 to 0.43 in culture TL1, but varied between 0.42 and 0.45 in culture TL2. This deviation could be a possible explanation for the differences measured on the fatty acid profiles in both cultures.

Finally, Costas et al. (2014) suggested that *I. galbana* wild strain had a “greater genetic variability in fitness,” with a high adaptive potential to thermal stress compared to the other species they studied. Our data tends to confirm this result for *T. lutea*, as the final population was able to grow outside its initial thermal niche.

Table 4: Presence index of the different strain in the cultivated population, calculated from the SNPs, at stress cycles S1, S8, and S12. Major presence indexes (> 0.10) are emphasised in italics

Strain	TL1-S1	TL1-S8	TL1-S12	TL2-S1	TL2-S8	TL2-S12
CCAP 927/14	0.008	0.037	0.028	0.031	0.054	0.028
<i>CCMP 463</i>	<i>0.174</i>	<i>0.315</i>	<i>0.353</i>	<i>0.190</i>	<i>0.370</i>	<i>0.375</i>
RCC 179	0.012	0.036	0.036	0.036	0.060	0.060
<i>RCC 1344</i>	<i>0.112</i>	<i>0.427</i>	<i>0.420</i>	<i>0.448</i>	<i>0.420</i>	<i>0.413</i>
RCC 3691	0.013	0.033	0.013	0.033	0.033	0.007
RCC 3692	0.023	0.068	0.008	0.030	0.015	0.008
RCC 3693	0.056	0.028	0.065	0.046	0.019	0.028
RCC 3699	0.039	0.072	0.02	0.033	0.039	0.013
Argenton 1998	0.030	0.060	0.045	0.015	0.015	0.045
IFMG	0.028	0.037	0.009	0.019	0.019	0.009
NIVA 4-91	0.026	0.017	0.009	0.017	0.017	0.026
S5-1%	0.036	0.018	0.018	0.00	0.018	0.009

This study demonstrated the efficiency of ALE with an enhanced population of *T. lutea* enriched in PUFA-rich polar lipids. Two strains out of the 12 composing the initial population turn out to survive the adaptation experiment. The observed mutations suggest that evolution took place upon selection. The lipid phenotype drastically changed during the second half of the experiment, with a twofold increase in total lipid content and a threefold increase in the DHA partitioned to polar lipids. As growth rate was not affected, the gain in lipid and DHA is directly a gain in productivity.

There is currently a lack of data to assess the stability of the metabolic changes after months of cultivation conditions back to normal. A possibility to keep them for longer periods of time, while continuing to improve their performance, is to maintain the selection pressure, but this strategy is expensive, with high manpower costs. Screening the population in order to isolate the clones presenting the desired phenotype will prevent the strain to be outcompeted by individuals in minority. On top of this, cryopreserving the strain will definitely guaranty constant properties when inoculating a new reactor.

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**Data availability** Not applicable

## Compliance with ethical standards

**Conflict of interest** The authors declare that they have no conflicts of interest.

**Code availability** Not applicable

**Ethical approval** This article does not contain any studies with human participants or animals, performed by any of the authors.

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