Characterization of *Chlorella sorokiniana* and *Chlorella vulgaris* fatty acid components under a wide range of light intensity and growth temperature for their use as biological resources

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**ABSTRACT**

This study aims to characterize the potential of three strains of microalgal species (*Chlorella sorokiniana* KNUA114 and KNUA122; *C. vulgaris* KNUA104) for use as feedstock, based on their fatty acid compositions. Each strain was molecularly identified using four marker genes (*ITS*, SSU, *rbcL*, and *tufA*) and phylogenetically characterized. *C. sorokiniana* and *C. vulgaris* collected from Ulleung Island, South Korea, were homologous with other known species groups. Samples’ fatty acid components were measured using GC/MS analysis in growth temperatures of 10°C, 25°C, and 35°C. The growth rate of *C. sorokiniana* strains was higher than that of *C. vulgaris* under high-temperature conditions, confirming the potential industrial applicability of the former as feedstock material. Additionally, saturated fatty acid contents and productivities increased as biological resources of the *C. sorokiniana* strains were higher than those of *C. vulgaris* under high light intensity and temperature conditions. These results suggest that the fatty acid components of *C. sorokiniana* strains may potentially be used as biological resources (e.g., feedstock).

1. Introduction

Microalgae are important primary producers in the process of photosynthesis. Microalgal species have distinct characteristics and are adapted to a wide range of environments (Andersen and Hessen, 1991; Dufossé et al., 2005; Ralas et al., 2019). Previous research confirmed that microalgae could be used as biological resources (Dufossé et al., 2005; Becker, 2007; Vanthoor-Koopmans et al., 2013). Microalgal nutritional compounds, such as vitamins, minerals, fibers, fatty acids, and amino acids, have been investigated to determine the appropriate use for different species (Caporgno and Mathys, 2018). Microalgae do not require supplementation with organic carbon sources during cultivation as they carry out photosynthesis. The higher photosynthetic efficiency and growth rate of microalgae compared to other photosynthetic organisms explain its relatively faster and higher biomass accumulation (Chisti, 2007, 2010; Otelé and Pire, 2001; Herrero et al., 2006; Spolaore et al., 2006; Randhir et al., 2020). Microalgal biomass contains large amounts of unsaturated fatty acids, lutein, food particles, and other useful materials.

Mass cultivation of microalgae will be required to meet the potential demand for biomass in various industries, leading to the development of the open pond type raceway system (Ryu et al., 2019; Banu et al., 2020). This system can effectively produce large amounts of microalgal biomass at a low cost but has a risk of exposure to stress from external environments, such as light and extreme temperatures (Lee et al., 2018; Akizuki et al., 2020; Ma et al., 2020). Microalgae are photosynthetic organisms, but the light intensity and temperature required for optimal growth differ between species (Lee et al., 2018; Ma et al., 2020). Indeed, the range of light intensity and temperature under which microalgal growth is inhibited varies between species (Isdepsky and Borowitzka, 2019; Rayen et al., 2019). Therefore, it is important to select a suitable species for biomass production of microalgae using the open pond type raceway system (Isdepsky and Borowitzka, 2019; Rayen et al., 2019) and one which can adapt to conditions of high light intensity and temperature (Lee et al., 2018; Rayen et al., 2019; Akizuki et al., 2020). Such environmental conditions have been...
shown to increase the lipid and saturated fatty acid content of microalgae (Han et al., 2016; Hong et al., 2019; Yaisamlee and Sirikhachornkit, 2020), further suggesting the importance of adaptation to these conditions in the microalgal species we aim to select.

Microalgae fatty acid components account for 10%-20% of the organism’s dry weight. These compounds are used as biological resources in biofuel and dietary supplements as they contain a varying number of carbon atoms, such as medium-chain (C<sub>10</sub>-C<sub>14</sub>), long-chain (C<sub>16</sub>-C<sub>18</sub>), and very-long-chain (C<sub>20</sub> or more) molecules and derivatives (Oles and Pire, 2001; Hu et al., 2008). C<sub>16</sub> and C<sub>18</sub> fatty acids, in particular, are the main biological resources for biofuel and dietary supplement feeding (Herrero et al., 2006; Lin and Lin, 2011; Zhou et al., 2011). Methyl ester is the base unit used to produce these fatty acids. Several methods, such as blending, pyrolysis, transesterification, and catalytic digestion, have been employed for the possible application of the abovementioned fatty acid components as biological resources (Ma and Hanna, 1999; Spolaore et al., 2006; Zhou et al., 2011). A previous study has reported that an increasing interest of consumers in biological resources has led to the rise of demand for high-quality fatty acid ingredients (Oles and Pire, 2001; Kumar et al., 2019). This demand in fatty acids for biological resources has led to the rise of conditioning of nutrients, such as nitrogen and phosphorus, to enhance the fatty acid content of microalgae (Anto et al., 2019; Gupta et al., 2019; Huang et al., 2020). Therefore, nutrient deprivation-induced stress affects the production of microalgae biomass and may impair lipid productivity (Yang et al., 2018). Furthermore, the consumption rate of nitrate and phosphorus in the culture medium corresponds to the growth of microalgae conditioning (De-Bashan et al., 2005; Xin et al., 2010; Nigam et al., 2011; Wan et al., 2013). However, few studies have investigated the changes in microbial production under conditions of high light intensity and temperature, or the impact of these growth conditions on fatty acid content and saturated fatty acids and corresponding biomass and lipid productivity. The production of biological resources normally involves two purification processes: supercritical fluid extraction and subcritical water extraction, which are used to extract natural products from various raw materials, including plants, algae, and microalgae (Oles and Pire, 2001; Hidayontz et al., 2004; Herrero et al., 2006). Microbes are employed in these processes to produce linoleic acid (C<sub>18:2</sub>ω6) and α-linolenic acid (C<sub>18:3</sub>ω3) from oleic acid (C<sub>18:1</sub>ω9). Furthermore, microalgae can produce hexadecadienoic (C<sub>16:2</sub>) and hexadecatrienoic (C<sub>16:3</sub>) acids from palmitic acid (C<sub>16:0</sub>) (Liperoti et al., 2009; Zhou et al., 2011). Based on these results, the extraction of fatty acids from microalgae is considerably important for their application as biological resources.

In this study, Korean *Chlorella sorokiniana* KNUA114 and KNUA122, and *Chlorella vulgaris* KNUA104 were aseptically isolated from Ulleung Island. Previous observations of these strains suggest that Chlorellaceae is one of the important families of organisms that can be used as biological resources worldwide (Morimura and Tamiya, 1954; Taub and Dollar, 1968; Ray and Barton, 1991; Park et al., 2020). The phylogenetic differences between the isolated *C. sorokiniana* and *C. vulgaris* strains were confirmed through morphological and molecular characterization. Furthermore, the value of *C. sorokiniana* and *C. vulgaris* strains as a potential biological resource was determined by analyzing microalgal growth and fatty acid composition. Based on our findings, the values of *C. sorokiniana* and *C. vulgaris* were compared; *C. sorokiniana* strains were found to be more suitable as fatty acid sources than *C. vulgaris* due to comparatively faster growth and greater productivity of the former under high light intensity and temperature conditions.

2. Materials and Methods

2.1. Sampling and isolation of microalgae

Microalgal bloom samples were collected from on Ulleung Island, South Korea, between April and October 2017. The two collection sites were the Okcheon stream (for *C. sorokiniana* KNUA114 and KNUA122; 37°28′24.0″N, 130°53′06.8″E; Figure 1A) and the Namseo stream (for *C. vulgaris* KNUA104; 37°28′01.5″N, 130°50′12.4″E; Figure 1B) (Abou-Shanab et al., 2011). Samples were then transferred to the laboratory and 10-mL aliquots were inoculated into 100 mL of BG-11 medium in a 250-mL flask (Rippka et al., 1979; Bolch and Blackburn, 1996). The flasks were placed in an orbital shaker (VS-202D, Vision Scientific, Bucheon, South Korea) rotating at 160 rpm and incubated at 25 °C with illumination by a fluorescent lamp (400 μmol photons) under light-dark cycle of 16:8 h until algal growth could be observed. Grown algal cultures (100 μL) were streaked onto BG-11 agar plates. Subsequently, single colonies were aseptically transferred onto fresh BG-11 agar plates; this step was repeated until a pure algal culture was obtained (Stanier et al., 1971).

2.2. Morphological characterisation of microalgal species

Before the morphological analysis (Safi et al., 2014; Krienitz et al., 2004), the isolated algal strains were grown in BG-11 medium for 10 days. A pure culture of algal cells was harvested by centrifuging at 3,000 rpm for 5 min (Centrifuge 5810R, Eppendorf AG, Hamburg, Germany) and cells were washed with distilled water and inspected using a Nikon Eclipse E100 biological microscope (Tokyo, Japan) at 1000× magnification.

2.3. Molecular identification

Before genomic DNA extraction, algal cells were harvested by centrifugation at 3,000 rpm for 5 min (Centrifuge 5810R, Eppendorf AG, Hamburg, Germany) and washed with distilled water. Cells were then mixed with 400 μL DNA extraction buffer (100 mM Tris HCl, 1 M KCl, and 10 mM EDTA), 8 μL RNase A (50 mg/mL, Elips-Biotech, Daejeon, South Korea), and sterile glass beads in a 2-mL microfuge tube. The tube was vortexed for 5 min and incubated at 65 °C for 30 min. Cellular debris was removed by centrifuging for 10 min at 13,000 rpm and 4 °C. Subsequently, 400 μL of the supernatant was transferred into a new tube and DNA extraction was performed using a DNA purification kit (Wizard DNA Clean-Up System, Promega, Madison, WI, USA). Marker genes were amplified by PCR using internal transcribed spacer (ITS) region ITS1-F and ITS54-R (Polle et al., 2008; Koetschan et al., 2010); small subunit rRNA gene (SSU) NS1-F and NS8-R (White et al., 1990; Friedli, 1995); large unit ribulose biphosphate carboxylase (rbcL); rbcL-M379-F, rbcLFP-R, GhrbC-L, and GrhB-R (Hayuda et al., 2000; Vieira et al., 2016); and elongation factor tu gene (tufA) tuF-A and tuf-R (Vieira et al., 2016). The PCR primers used in this study are listed in Table S1. Amplified DNA fragments were purified using a PCR purification kit (Spin type-200) (Elips-Biotech, Daejeon, South Korea), then cloned using the pGEM®-T Easy Vector kit (Promega, Madison, WI, USA) (Weisburg et al., 1991). Cloned DNA samples were sequenced at the Genotech facility in Daejeon, South Korea. Finally, the phylogenetic analysis was performed using ITS-, SSU-, RbcL-, and tufA-specific primers generated according to methods described in Friedli (1995) and Buchheim et al. (2011) and MEGA version 7.0 (Kumar et al., 2008, 2016). The best-fit nucleotide substitution models for ITS (Kimura 2-parameter), SSU (Tamura-Nei), rbcL (General Time Reversible), and tuAF (General Time Reversible) were selected using MEGA 7.0, based on the Bayesian information criterion. These models were used to build maximum likelihood phylogenetic trees with 1000 bootstrap replications (Felsenstein, 1985).

2.4. Identification of the optimal temperature and light intensity

The optimal temperature and light intensity conditions for the culture of each microalgal strain were investigated using a PhotobioBox, a high-throughput system used to analyze photosynthesis (Heo et al., 2015). Each strain was grown from aseptically-transferred, isolated colonies (1.0 ± 10<sup>6</sup> cells/mL, 200 μL), inoculated to 96-well plates containing BG-11.
medium and incubated for three days at temperatures ranging from 6–28 °C and light intensities ranging from 300–1,000 μmol photons m⁻² s⁻¹.

2.5. Laboratory cultivation

A 10-day-old culture of each strain originating from aseptically-transferred, isolated colonies (15 mL) (diluted to OD₆₈₀ ~0.3, corresponding to the following cell densities: KNUA104, 391.43 ± 16.05 cells x 10⁶ ml⁻¹; KNUA114, 486.32 ± 29.51 cells x 10⁶ ml⁻¹; and KNUA122, 523.41 ± 21.73 cells x 10⁶ ml⁻¹) was inoculated into flasks containing 150 mL of BG-11 medium (performed in triplicates). Flasks were then incubated for 14 days on orbital shakers (VS-202D, Vision Scientific) at 160 rpm in rooms illuminated by fluorescent lamps (300 μmol photons m⁻² s⁻¹) under controlled temperature (10 °C, 25 °C, or 35 °C) and a light:dark cycle of 16:8 h. The growth of each strain was subsequently measured as previously described (Hong et al., 2016). Algal density was estimated by measuring the OD₆₈₀ values with an Optimizer 2120 UV spectrophotometer (Mecasys, South Korea) (Griffiths et al., 2011). The above procedure was scaled up and further performed under the same conditions, and algal cells were inoculated into 2 L of BG-11 medium in a flask (maximum volume of flask; 5 L). The algal density of the scale up culture was estimated by measuring OD₆₈₀ values.

2.6. Productivity analysis

The dry weights of the microalgal cultures were measured to analyze biomass productivity at the stationary phase. Dry weight was calculated by filtering 10 mL of culture through a pre-weighed GF/C Whatman filter (47 mm, nominal pore size 0.7 μm), which was subsequently dried at 105 °C for 24 h. The dried filters were cooled to room temperature and reweighed (Zhu and Lee, 1997).

2.7. Lipid extraction and gas chromatography/mass spectrometry (GC/MS) analysis

Each microalgal strain was grown for 10 days and the dilutions of all samples (OD₆₈₀ = 1.0) were freeze-dried. Thirty milligrams of pulverized, freeze-dried algae were subjected to lipid extraction using a mixture of chloroform and methanol (1:1) as previously described (Hong et al., 2016). Chloroform was evaporated from the extracted mixture using a rotary evaporator. The crude lipid was treated with methanol and a potassium hydroxide solution to facilitate transesterification. To isolate fatty acid methyl esters (FAMEs), hexane was added to the mixture and the reaction was heated at 30 °C for 10 h. Subsequently, the hexane layer was isolated and analyzed using GLC-90 (SUPELCO, Bellefonte, PA, USA) and used as an external standard to determine FAME contents. The fatty acid components present in FAMEs derived from microalgae were analyzed using a 6890N gas chromatograph (Agilent Technology Inc., Santa Clara, CA) equipped with a 5973N mass selective detector (Agilent Technology) and a HP-5MS capillary column [30 m x 0.25 mm (internal diameter) x 0.25 μm (film thickness); Agilent Technology Inc.]. The gas chromatography (GC) oven temperature was maintained at 120 °C for 2 min and subsequently increased to 300 °C at a rate of 5 °C/min and maintained at 300 °C for another 22 min. The injection volume was set to 1 μL, with a split ratio of 20:1. Helium was used as the carrier gas at a

Figure 1. The collection sites on Ulleung Island, South Korea. (A) Okcheon Stream, Ulleung-eup, Ulleung-gun, Gyeongsangbuk-do (C. vulgaris KNUA104 and C. sorokiniana KNUA122; 37°28‘24.0’’N, 130°53‘06.8’’E) (B) Namseo Stream, Seo-myeon, Ulleung-gun, Gyeongsangbuk-do (C. sorokiniana KNUA114; 37°28‘01.5’’N, 130°50‘12.4’’E). (C) Light microscope images of C. vulgaris KNUA104, (D) C. sorokiniana KNUA114, and (E) C. sorokiniana KNUA122 (Scale bar = 5 μm). (F) A 250-mL flask containing C. vulgaris KNUA104, (G) C. sorokiniana KNUA114, and (H) C. sorokiniana KNUA122.
Figure 2. Molecular phylogenetic analysis according to the Maximum Likelihood (ML) tree. Numbers at the nodes indicate the bootstrap probabilities (>50%) of the ML analyses (1000 replicates). (A) Internal transcribed spacer (ITS) region, (B) small subunit rRNA gene (SSU), (C) large unit ribulose bisphosphate carboxylase (rbcL), and (D) elongation factor tu gene (tufA). The green, blue, and red boxes indicate C. vulgaris KNUA104, C. sorokiniana KNUA114, and C. sorokiniana KNUA122, respectively.
constant flow rate of 1 mL/min. The injector and MS source temperatures were set to 250 and 230 °C, respectively. In the electron impact mode, an acceleration voltage of 70 eV was used for sample ionization and the scanning range was increased from 50 to 550 m/z. The Wiley/NBS libraries were used as reference databases (Furuhashi and Weckwerth, 2013).

2.8. Statistical analysis

We compared individual data points using Student's t-test, using a P-value of 0.05 as the statistical significance level. All experiments were performed at least in triplicates and all the results were expressed as mean ± SD.

Figure 3. Growth pattern analysis using a PhotoBiobox and measurement of optical density. Growth rate matrix for (A) C. vulgaris KNUA104, (B) C. sorokiniana KNUA114, and (C) C. sorokiniana KNUA122 at temperatures ranging from 6 °C–30 °C and light intensities from 300–1,000 μmol photons m⁻² s⁻¹.
3. Results and discussion

3.1. Molecular identification and phylogenetic analysis of isolated C. sorokiniana and C. vulgaris strains

The aseptic isolation of microalgae from a mixed culture with bacteria was performed by spreading or streaking on a BG-11 culture medium (Chang et al., 2013). The C. sorokiniana strains KNUA114 and KNUA122, as well as the C. vulgaris strain KNUA104, all lacked a flagellum, a common feature of species belonging to the genus Chlorellaceae (Figure 1C-E). In previous reports, Chlorellaceae species were described as round (1–4 μm diameter), with green-colored, cup-shaped chloroplasts (Darienko et al., 2010; Safi et al., 2014; Krienitz et al., 2004). Our morphological observations of the strains used in this study are similar to those of other Chlorellaceae members (Figures 1 and 2).

The ClustalW analysis by using MEGA 7.0 revealed that the gene sequences of our strains aligned with those of Chlorellaceae and other algal species described in previous studies (Hoshina et al., 2010; Luo et al., 2010). The sequences of four marker genes (ITS, SSU, rbcL, and tufA) were compared between the isolated species and those available in the NCBI database, the results of which are summarized in Table S2. According to the BLAST results, KNUA114 and KNUA122 are homologous and are closely related to the C. sorokiniana strains, while the KNUA104 strain is closely related to other C. vulgaris strains (Figure 2; Table S2).

3.2. Analysis of the growth patterns under high light intensity and temperature conditions

In previous research, Chlorellaceae have been shown to grow optimally under high light intensity rather than low-light (Mandal and Mallick, 2012; An et al., 2016). Our PhotoBiobox analysis revealed that C. sorokiniana KNUA114 and KNUA122 strains were red-colored under high light intensity, unlike C. vulgaris KNUA104. This suggests that the former strains had a higher photosynthetic efficiency than C. vulgaris KNUA104 (Figure 3). Moreover, C. sorokiniana KNUA114 and KNUA122 showed better growth patterns than C. vulgaris KNUA104 under various temperature conditions, particularly at high temperatures (Figure 4), reaching the stationary phase within 10 days of inoculation (Figure 4B).

Furthermore, C. sorokiniana KNUA114 and KNUA122 strains grew better at high temperatures than at low temperatures, whereas C. vulgaris KNUA104 did not grow well at high temperatures (Figure 4C). Our results revealed that the C. sorokiniana KNUA114 and KNUA122 strains had higher photosynthetic efficiencies and growth rates than C. vulgaris KNUA104 strain under high light intensity and high-temperature conditions.

Light intensity and temperature are known to influence microalgal growth and, indirectly, biomass productivity (Renaud et al., 2002; Ras et al., 2013; Xu et al., 2019). Furthermore, each microalgal species has a unique optimum for light intensity and temperature, hence selecting a microalgal species suitable for the biomass production environment is one of the strategies to increase biomass productivity (Yoo et al., 2010; Alvarez-Díaz et al., 2017; Aketo et al., 2020). Growth results showed the characteristics of C. sorokiniana (KNUA114 and KNUA122 strains) and C. vulgaris (KNUA104) for light intensity and temperature. Under light intensity conditions, the C. sorokiniana KNUA114 and KNUA122 strains showed a wide range of tolerance throughout the tested light intensity. Moreover, the growth of C. sorokiniana KNUA114 and KNUA122 strains was not inhibited under 600 μmol photons m⁻² s⁻¹, while that of C. vulgaris KNUA104 was significantly inhibited.

High light intensity inhibits microalgal growth, but the wide range of growth investigated (300–1000 μmol photons m⁻² s⁻¹) suggests high applicability for biomass production of C. sorokiniana KNUA114 and KNUA122 strains. Furthermore, the C. sorokiniana KNUA114 and KNUA122 strains showed higher growth rates and productivity than C. vulgaris KNUA104 at 25 and 35 °C. Therefore, we conclude that the C. sorokiniana KNUA114 and KNUA122 strains, which showed tolerance to a wide range of light intensity, high growth rate, and high lipid productivity among the tested microalgal strains, are more suitable biological sources than C. vulgaris KNUA104.

Figure 4. Growth curve of C. sorokiniana KNUA114, C. sorokiniana KNUA122, and C. vulgaris KNUA104 at (A) 10 °C, (B) 25 °C, and (C) 35 °C under a light intensity of 300 μmol photons m⁻² s⁻¹.
3.3. Fatty acid components of C. sorokiniana KNUA114 and KNUA122, and C. vulgaris KNUA104

Strains were harvested during the stationary phase and analyzed for fatty acid productivity and chain composition using an SPV reaction and GC/MS, and we found that C. sorokiniana KNUA114 and KNUA122 strains had higher fatty acid contents under high-temperature conditions. Furthermore, the fatty acid components of C. sorokiniana KNUA114 and KNUA122 strains were incomparable with those of the C. vulgaris KNUA104 strain, as the growth of the latter was inhibited at high temperatures (Figure 4C, Table 1). Previous research has described saturated fatty acids (SFAs) as biological resources for biofuel production (Chang et al., 2013; Wei et al., 2018). These fatty acid components are commonly found in microalgae (Hong et al., 2016), which may be used as a potential source for these valuable compounds (Oles and Pire, 2001; Alberts, 2003; Herrero et al., 2006; Spolaore et al., 2006; Knothe, 2008; Mohammady, 2011). The fatty acid composition of microalgae grown in this study varied with the temperature conditions used; under higher temperatures, SFA concentration was higher. In particular, 40%–50% of fatty acids of C. sorokiniana KNUA114 and KNUA122 were composed of SFAs when grown at 35 °C (46.78 and 44.40%, respectively). Based on these results, the positive influence of temperature on fatty acid composition in microalgal biomass was confirmed, as well as an increase in lipid productivity and SFAs concentration. Similar trends have also been reported in previous research (Pasquet et al., 2014; Wei et al., 2015), highlighting the potential use of C. sorokiniana KNUA114 and KNUA122 as a biological resource which can adapt to extreme temperature stress.

Our findings suggest that C. sorokiniana KNUA114 and KNUA122 produce fatty acid components, such as palmitic (C16:0), hexadecenoic (C16:1), hexadecadienoic (C16:2), hexadecatrienoic (C16:3), stearic (C18:0), oleic (C18:1 ω9), linoleic (C18:2 ω6), and α-linolenic acid (C18:3 ω3) more efficiently than C. vulgaris KNUA104, particularly at high light intensity and temperature conditions. Taken together, our findings suggest that C. sorokiniana KNUA114 and KNUA122 strains can be used as a potential source of fatty acids, due to their autotrophic ability and versatility in growing under various environmental stress conditions, as opposed to the C. vulgaris KNUA104 strain.

4. Conclusion

In this study, we investigated three strains of two species belonging to the Chlorella genus, collected on Ulleung Island, South Korea. We demonstrated that the C. sorokiniana KNUA114 and KNUA122 strains could be cultured more efficiently than the C. vulgaris KNUA104 strain with a wide range of light intensity and temperature conditions. Our findings demonstrate that C. sorokiniana KNUA114 and KNUA122 strains generated higher yields of fatty acid components compared to C. vulgaris KNUA104 under high light intensity and temperature conditions, suggesting their potential use as biological raw materials. Future research should focus on screening for the genes encoding the various fatty acids in C. sorokiniana KNUA114 and KNUA122 strains to further evaluate the usefulness of these strains as biological sources of fatty acids. We assume that C. sorokiniana KNUA114 and KNUA122 strains were investigated as a potential feedstock for fatty acid components compared with C. vulgaris KNUA104 strain and the ability of these strains to autotropically produce fatty acids that can be used as biological resources was demonstrated in this study.

Declarations

Author contribution statement

Hyun-Sik Yun: Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data. Young-Saeng Kim: Conceived and designed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data. Ho-Sung Yoon: Conceived and designed the experiments; Wrote the paper.

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Table 1. Fatty acid productivity and composition of C. sorokiniana KNUA114 and KNUA122, and C. vulgaris KNUA104, under different cultivation temperatures.

| Strain | KNUA104 | KNUA114 | KNUA122 |
|--------|---------|---------|---------|
| Temperature (°C) |         |         |         |
| 10     | 25      | 10      | 25      |
| 35     | 10      | 25      |         |
| SFAa   | 21.24   | 26.29   | 20.70   |
| MUFAc  | 9.02    | 10.07   | 2.44    |
| UFAb   | 78.76   | 73.71   | 79.30   |
| Productivity (g/L) |         |         |         |
| Biomass | 0.48    | 0.71    | 0.52    |
| Lipid   | 0.08    | 0.14    | 0.11    |

a Percentage of saturated fatty acids.

b Percentage of unsaturated fatty acids.
c Percentage of monounsaturated fatty acids.
d Percentage of polyunsaturated fatty acids.
