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Lipid production of Chlorella vulgaris and Pseudokirchneriella subcapitata

Ana L Gonçalves, José CM Pires^{*} and Manuel Simões

Abstract

Background: The depletion of fossil fuel reserves has stimulated the search for sustainable sources of energy that are carbon-neutral or renewable. In this context, microalgae are a promising energetic resource. They are photosynthetic microorganisms that use CO_2 as carbon source, with high specific growth rates. Furthermore, some species present high lipid content that can be easily converted into biodiesel. Accordingly, this study aims to analyze the effect of light supply (one of the most important culture parameters) on lipid production of selected microalgae, *Chlorella vulgaris* and *Pseudokirchneriella subcapitata*.

Methods: Both microalgal species were cultured under different light irradiance values (36, 72, 96, and 126 μ E m⁻² s⁻¹) and for each light irradiance value, three light/dark ratios (10:14, 14:10, and 24:0) were tested. Lipid contents of both microalgae were then determined using a recently developed colorimetric method.

Results/Conclusions: *P. subcapitata* presented higher lipid productivity than *C. vulgaris*. High lipid concentration was achieved in microalgal cultures with higher light irradiance values. However, for 96 and 126 μ E m⁻² s⁻¹, lipid productions of both microalgae were not significantly higher than with 72 μ E m⁻² s⁻¹, which means that microalgal light saturation point may be achieved.

Keywords: Chlorella vulgaris, Pseudokirchneriella subcapitata, Lipid production, Light effect, Microalgae

Background

Economic development and increased standards of living have resulted in an increased demand for energetic sources. In the last decades, fossil fuels have been extensively used as a source of energy. Consequently, there are now concerns about the low availability of this non-renewable energy resource and about the environmental impacts associated to burning of fossil fuels [1-3]. Attentions are now focused on the production of renewable and carbon-neutral energetic sources, such as biodiesel. Biodiesel consists of a mixture of esters with long-chain fatty acids, such as lauric, palmitic, stearic, and oleic [2,4]. This renewable energy source can be produced from vegetable oils, animal fats, microorganisms' oils, or waste cooking oil [2,4-7].

Microalgae are known for their high lipid contents (20% to 50% of dry weight) and high growth rates, being a promising oil source for biodiesel production [8,9]. However, producing biodiesel from microalgal lipids is not economically viable, requiring several optimizations to reduce

* Correspondence: jcpires@fe.up.pt

LEPAE, Departamento de Engenharia Química, Faculdade de Engenharia, Universidade do Porto, Rua Dr. Roberto Frias, Porto 4200-465, Portugal production costs and to achieve current demand for biofuels [10]. Different studies have reported that lipid contents differ according to the microalgal strain and their cultivation conditions [11,12]. To analyze the lipid content of microalgae, the most applied method is organic solvent extraction followed by chemical characterization of lipid profiles using gravimetric analysis [13], thin-layer chromatography, high-performance liquid chromatography, or gas chromatography [14,15]. However, this procedure is costly, timeconsuming, and requires large amounts of toxic and harmful solvents and biomass [5,9,10,16]. Additionally, extraction efficiencies of these methods strongly depend on water content present in microalgal biomass, requiring a previous drying process [9].

Microscopic and colorimetric methods showed to be simple and rapid procedures for screening lipid content of microalgae. Microscopic methods require the use of dyes, such as triptan blue or Nile red, which are selectively incorporated in cells and bind to the target compounds, while colorimetric ones, also known as spectrophotometric methods, are based on the color development of the analyzed solution after reaction of the molecules of interest



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with specific ions [17-19]. Application of these methods requires little amounts of biomass and reagents [20,21]. Nile red (a lipid-soluble fluorescent dye) has been frequently used in the determination of neutral lipid contents in animal cells [22], bacteria [23], and microalgae [20,21,24]. This method has received much attention due to the rapid and extraction-free determination of neutral lipids and also due to its sensitivity [21,25]. However, this method is not universal as the applied concentrations of Nile red should be defined for each organism. The establishment of a specific amount of Nile red for each organism is required because this dye acts by penetrating the cells and binding to cytosolic neutral lipids, which means that different affinities of cell membranes to uptake Nile red will result in false determination of lipid contents [18,21]. Another methodology used to easily determine microalgal lipids is to quantify free fatty acids (FFA). Lipids can be accumulated in free or bound forms; in microalgae, they are mainly stored as bound fatty acids (FA). Thus, previous saponification (hydrolysis of FA) will allow the quantification of FFA as they can react with copper or cobalt salts, forming colored complexes that can be detected by colorimetric/spectrophotometric methods [26-28]. The method applied in this study is based on the fact that FFA can react with copper or cobalt salts with the formation of colored complexes that can be extracted by chloroform in the presence of triethanolamine (TEA). The amount of copper in the chloroform phase is then colorimetrically determined by adding diethyldithiocarbamate to develop a yellow-colored product that can be detected by measuring absorbance at 440 nm. This method was derived from the methods proposed by Wawrik and Harriman [19] and Chen and Vaidyanathan [18].

The aim of this study was to evaluate the influence of different light supplies on microalgal lipid productivities. The colorimetric method derived from those proposed by Wawrik and Harriman [19] and Chen and Vaidyanathan [18] was applied to cultures of *Chlorella vulgaris* and *Pseudokirchneriella subcapitata* grown under different light supplies: irradiance values of 36, 72, 96, and 126 μ E m⁻² s⁻¹ and photoperiods of 10:14, 14:10, and 24:0.

Methods

Microorganisms cultivation and harvesting

C. vulgaris and *P. subcapitata* (obtained from the American Type Culture Collection) were cultured in modified Organisation for Economic Co-operation and Development test medium [29], with the following composition (per liter): 24 mg MgCl₂·6H₂O, 36 mg CaCl₂·2H₂O, 30 mg MgSO₄·7H₂O, 3.2 mg KH₂PO₄, 0.16 mg FeCl₃·6H₂O, 0.2 mg Na₂EDTA·2H₂O, 0.37 mg H₃BO₃, 0.83 mg MnCl₂·4H₂O, 6 µg ZnCl₂, 3 µg CoCl₂·6H₂O, 0.02 µg CuCl₂·2H₂O, 14 µg Na₂MoO₄·2H₂O, and 100 mg NaHCO₃. Nitrogen was supplied in the form of NaNO₃ for *C. vulgaris* and in the form of NH₄Cl for *P. subcapitata*

(30 mg L⁻¹). Experiments were performed in 500-mL flasks operating in batch with a working volume of 400 mL. The cells were cultivated for 12 days using the growth medium described above. The experimental conditions were the following: (1) initial cell concentration of 0.05 to 0.08 g L⁻¹, (2) room temperature (22.2 \pm 1.2°C), and (3) continuous aeration with the injection of atmospheric air at the bottom of the flasks. The assays were carried out under different irradiance values: 36, 72, 96, and 126 µE m⁻² s⁻¹. For each value, different photoperiods were evaluated: 10:14, 14:10, and 24:0 (light/dark). All the experiments were conducted in triplicates.

Growth monitoring

Duplicate samples were collected at 24-h intervals, and biomass concentration was determined by measuring optical density at 683 nm, OD_{683} [30], using a V-1200 spectrophotometer (VWR International China Co., Ltd, Shanghai, China). The relationship between OD_{683} (*y*) and cell dry weight (*x*) of *C. vulgaris* and *P. subcapitata* was established by linear regression: *y* = 1.841*x*(R^2 = 0.997) *and y* = 2.732*x*(R^2 = 0.993), respectively.

Standard curves

Sodium salts of different fatty acids (decanoic acid, lauric acid, myristic acid, sarkosyl acid, and palmitic acid) were purchased from Sigma-Aldrich, St. Louis, MO, USA (\geq 98% purity). Stock solutions (5 mM) of each salt were prepared in a solvent mixture of chloroform and methanol (2:1), according to the method proposed by Chen and Vaidyanathan [18]. Serial dilutions were prepared, and 300 µL of each dilution was used for lipid quantification, as described below.

Saponification of cellular lipids

For each experiment performed, 1.5 mL was collected on the first and last day of culturing. These algal suspensions were centrifuged at 12,000 rpm for 10 min in an Eppendorf 5424 centrifuge (Eppendorf, Hamburg, Germany), and cell pellets were stored at -20°C until lipid extraction. The cell pellets were thawed and resuspended in 20 μ L of Tris-HCl (1 M, pH 8.0). Then, 480 μ L of saponification reagent (25% methanol in 1 M NaOH) and approximately 50 mg of 0.1-mm glass beads (Sigma-Aldrich) were added. Cell disruption was promoted by vigorous vortexing for 3 min. Then, another 500 μ L of saponification reagent was added, and the mixture was saponified at 90°C for 30 min and vortexed every 5 min to hydrolyze ester bonds of membrane lipids and triglycerides [18,19].

Lipid extraction and colorimetric detection

After saponification, the samples were cooled down to room temperature, and 600 μ L of the saponified mixture was added to an Eppendorf tube (Eppendorf) containing 900 μ L of a solvent mixture of chloroform and methanol

(2:1). After vortexing for 2 min, the samples were centrifuged at 5,000 rpm for 2 min. A volume of 500 µL of the organic phase was then transferred to a new Eppendorf tube containing copper reagent (9 vol. aq. 1 M TEA, 1 vol. 1 M acetic acid, 10 vol. 6.45% w/v Cu $(NO_3)_2 \cdot 3H_2O$). The procedure applied to standards was similar. To 300 µL of standard solutions, 300 µL of saponification reagent and 300 µL of chloroform and methanol mixture (2:1) were added. After vortexing and centrifugation in the same conditions, 500 µL of the organic phase was added to a new Eppendorf tube, containing the same volume of copper reagent. Samples and standards were vortexed and centrifuged again. After this step, 200 µL of the organic phase was carefully transferred to a quartz cuvette, where 200 μ L of color-developing reagent (1% w/vsodium diethyldithiocarbamate in 2-butanol) was added. In this step, it is important that none of the aqueous phase is transferred since it contains copper ions that will produce false signal during colorimetric detection. Absorbance at 440 nm was then measured using a V-1200 spectrophotometer (VWR International China Co., Ltd). A solution containing 200 µL of chloroform and methanol (2:1) mixture supplied with 200 µL of color-developing reagent was used as a blank [18,19].

Statistical analysis

Lipid quantification was performed in triplicate, and data were analyzed using paired samples *t* test from the statistical software SPSS 17.0 (Statistical Package for the Social Sciences Inc., Chicago, IL, USA). Significance level was set at p < 0.05.

Results and discussion Microalgal growth

Figure 1 shows the growth curves of C. vulgaris and P. subcapitata corresponding to different light supplies at room temperature and aerated with CO_2 at atmospheric concentration. For almost all cultures, the lack of an adaptation phase or a very short one was observed: the exponential phase started before completing the first day of culture. However, for photoperiod of 10:14 and low irradiance values, both microalgal species showed longer adaptation phases. Generally, almost all cultures presented an exponential growth phase with the same duration, reaching the stationary phase on the seventh day of culture. Similar behavior was observed by Jacob-Lopes et al. [31] when the effect of light cycles on the cultures of the cyanobacterium Aphanothece microscopica Nägeli was analyzed. However, for cultures where light supply was higher (longer light periods and higher irradiance values), stationary phase was not reached at all before the end of the experiment. This results indicate that nutrients can be a growth-limiting factor. Two possible limiting nutrients are nitrogen and phosphorus. According to the study performed by Bhola et al. [32], *C. vulgaris* reached its maximum concentration for nitrogen and phosphorus supplied at 5 and 0.04 g L⁻¹, respectively. In this study, nitrogen and phosphorus were supplied at 0.03 and 0.0032 g L⁻¹, respectively. As these values are lower than those used in the study performed by Bhola et al. [32], the possibility of nutrient limitation is very high.

Regarding the maximum biomass concentration achieved in each culture, the data presented in Figure 1 show that this value was higher for the microalga *P. subcapitata* for the photoperiod of 10:14 and light irradiance of 96 μ E m⁻² s⁻¹ (0.808 ± 0.021 g L⁻¹, representing ten times more than the initial concentration of the culture). This kinetic parameter did not present statistical differences varying the light irradiance (*p* > 0.05), but it was statistically different for other photoperiods (*p* < 0.05). The achievement of the highest value in a discontinuous light supply may be related with possible photooxidation [33,34]. The oxygen generated by photosynthesis may accumulate in culture medium, reaching values that in combination with intense light can damage microalgal cells.

Lipid analysis

The method described in this study enables the simple and rapid quantification of lipids from small amounts of microalgal biomass during cell growth experiments. Long-chain fatty acids used in this study showed linear responses for concentrations between 0 and 190.5 mg L⁻¹, as shown in Figure 2. One-sample *t* test showed that the extinction coefficients, in liters per milligram per centimeter, for all fatty acids were not significantly different (p > 0.05), indicating a similar behavior between fatty acids of different chain lengths. For this reason, lipid quantification was assessed by using an average of extinction coefficients for all fatty acids (Figure 3): $y = 1.327 \times 10^{-2} x + 2.911 \times 10^{-2} (R^2 = 0.999)$, where *y* is the absorbance measured at 440 nm and *x* is the fatty acid concentration in milligrams per liter.

Figure 4 shows the lipid yields of C. vulgaris and P. subcapitata determined for different light supplies. Lipid yields achieved range from 4.611 \pm 0.939 to 28.234 \pm 0.581 mg g^{-1} cell dry weight (CDW) for *C. vulgaris* and from 8.562 ± 1.204 to 39.287 ± 1.879 mg g⁻¹ CDW for P. subcapitata. Similar values were reported by different authors [35,36]. Ranjan et al. [36] achieved lipid yields between 6.0 and 60 mg g^{-1} CDW when applying organic solvent extraction to recover lipids from the microalga Scenedesmus sp. Using the same extraction procedure, Lee et al. [35] achieved lipid yields of 49.0 mg g^{-1} CDW for *C. vulgaris*. The effect of light supply on lipid yields was similar for both microalgal strains (p > 0.05). Lipid yields were significantly lower (p < 0.05) for the lowest irradiance (36 μ E m⁻² s⁻¹). Concerning the other light supplies, lipid yields were not statistically different (p > 0.05). Similar results were obtained by Solovchenko et al. [37] who showed that the

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accumulation of total fatty acids by the microalga *Parietochloris incisa* increased with irradiance (35, 200, and 400 μ E m⁻² s⁻¹). Renaud et al. [38] also demonstrated that lipid yields from *Isochrysis* sp. grown under light irradiance values ranging from 107 to 140 μ E m⁻² s⁻¹ increased from 7.9 to 10.5 pg cell⁻¹. The same study has revealed that for higher irradiance values, lipid yields did not significantly



increase [38]. Weldy and Huesemann [12] also showed that lipid concentration in Dunaliella salina cells increased with higher irradiance value. Different photoperiods did not influence lipid yields achieved for both microalgae; however, for 96 μ E m⁻² s⁻¹, continuous light supply led to a significant increase in lipid yields. To our knowledge, this is the first study demonstrating the effects of different photoperiods on lipid yields from microalgae. Higher lipid yields observed when irradiance value is increased may be due to the storage of excessive light energy into chemical energy in the form of FA, thus avoiding photooxidative cell damage [39]. Non-increasing lipid yields under the highest irradiance values studied, 96 and 126 μ E m⁻² s⁻¹, and for longer light periods are a consequence of the photooxidative mechanism: when exposed to high irradiance values, chlorophyll molecules are excited forming a very unstable form, which, in turn, reacts with dissolved oxygen and transfers energy to it. The excited oxygen then reacts with fatty acids to form lipid peroxides, resulting in the reduction of FA concentrations [40]. Another possible reason is that cells can store excessive light energy into chemical energy in the form of other compounds, such as polysaccharides [41].

Conclusions

The use of colorimetric/spectrophotometric methods for rapid screening of microalgal lipids constitutes a promising and accurate technique to determine lipid productivities from microalgae. *P. subcapitata* presented higher lipid productivity than *C. vulgaris*. High light supplies (irradiance values greater than 72 μ E m⁻² s⁻¹) led to the achievement of higher lipid accumulation for both microalgae. However, no statistically significant differences in lipid accumulation were observed for the studied photoperiods. Under the highest light irradiance value with continuous light supply, a slight decrease in lipid productivity was observed that can be associated with photooxidation. Another possible reason is the storage of excessive light energy into chemical energy in the form of other compounds, such as polysaccharides.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

ALG carried out the experimental assays and statistical analysis and drafted the document. JCMP participated in the design of the study and coordination. MS conceived the study and participated in its coordination. All authors read and approved the final manuscript.

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