A Cyanobacteria-Based Biofilm System for Advanced Brewery Wastewater Treatment

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Featured Application: Post-treatment of brewery wastewater through an attached cyanobacterial growth system with simultaneous production of bioethanol.

Abstract: Algal/cyanobacterial biofilm photobioreactors provide an alternative technology to conventional photosynthetic systems for wastewater treatment based on high biomass production and easy biomass harvesting at low cost. This study introduces a novel cyanobacteria-based biofilm photobioreactor and assesses its performance in post-treatment of brewery wastewater and biomass production. Two different supporting materials (glass/polyurethane) were tested to investigate the effect of surface hydrophobicity on biomass attachment and overall reactor performance. The reactor exhibited high removal efficiency (over 65%) of the wastewater's pollutants (chemical oxygen demand, nitrate, nitrite, ammonium, orthophosphate, and total Kjeldahl nitrogen), while biomass per reactor surface reached 13.1 and 12.8 g m^{-2} corresponding to 406 and 392 mg L^{-1} for glass and polyurethane, respectively, after 15 days of cultivation. The hydrophilic glass surface favored initial biomass adhesion, although eventually both materials yielded complete biomass attachment, highlighting that cell-to-cell interactions are the dominant adhesion mechanism in mature biofilms. It was also found that the biofilm accumulated up to 61% of its dry weight in carbohydrates at the end of cultivation, thus making the produced biomass a suitable feedstock for bioethanol production.

Keywords: Leptolyngbya sp.; attached growth; brewery wastewater; carbohydrates; bioremediation; supporting materials; hydrophobicity; attachment efficiency; biofilm photobioreactor

1. Introduction

The cultivation of photosynthetic microorganisms (microalgae and cyanobacteria) to produce a wide range of high value products such as pigments, proteins, polysaccharides, vitamins, and fatty acids [1], as well as electricity in bioelectrochemical systems [2], has been investigated extensively. Wastewater treatment systems utilizing photosynthetic microorganisms have recently emerged as a promising alternative to conventional biological processes such as activated sludge [3]. However, the treatment of agro-industrial, industrial and domestic wastewaters with algae/cyanobacteria has gathered significant research interest over the last decade and produced promising results in both organic and inorganic pollutant removal [4–9]. This method offers several advantages, such as reduced operational cost as photosynthesis replaces mechanical aeration requirements, decreased CO₂ emissions [10], and the high cost of using chemical fertilizers for microalgal growth is avoided since microorganisms use the nutrients contained in the wastewaters [11]. Additionally, the produced biomass can be utilized as a platform for liquid (i.e., biodiesel and bioethanol) and gaseous (i.e., methane and biohythane) biofuels [1,11,12]. Brewery
wastewaters are produced in high quantities (3–10 L per L of beer produced) from beer brewing processing industries worldwide [13] and in 2019 the total global beer production amounted to about 1.91 billion hectoliters [14]. Brewery wastewaters are characterized by significant organic (2000–6000 mg·L⁻¹ of chemical oxygen demand, COD), nitrogen and phosphorous loadings (25–80 and 10–15 mg·L⁻¹, respectively) and they have been traditionally treated aerobically [13,15]. Until today, a limited number of studies on brewery wastewater treatment using axenic microalgal cultures [6,16] have been conducted in suspended cultivation systems. Recently, brewery wastewater was treated by our research group using a cyanobacterial-bacterial consortium dominated by the filamentous cyanobacterium *Leptolyngbya* sp. either without pre-treatment [4,5] or after pre-treatment with electrocoagulation [4,5].

Biomass harvesting/processing is often a challenging task. Photosynthetic culture cell densities are typically low (as low as 0.5 g·L⁻¹ in conventional open pond photobioreactors [17]) and separation methods such as chemical flocculation, gravitational sedimentation, filtration, and centrifugation can be complex, time-consuming, or expensive [3,18]. Techno-economic analyses indicate that the cost of algal/cyanobacterial biomass harvesting and dewatering can amount to 20–30% of the total cultivation expenses in high-rate algal ponds [17,19].

The introduction of attached cultivation (biofilm) systems could be a step towards addressing these issues. Biofilms are microbial communities living in a self-produced matrix of extracellular polymeric substances (EPSs) that provide structural integrity and mediate their adhesion to surfaces [20]. Under nutrient limitation and sufficient irradiation/carbon, cyanobacteria tend to over exceed their typical carbohydrate content in the form of EPSs, instead of channeling energy towards cell synthesis ("carbon overflow" metabolism) [20,21]. Along with lower quantities of secreted proteins and nucleic acids, EPSs allow high cell density colonization of abiotic and biotic surfaces through hydrophobic or hydrophilic cell–cell and cell–surface interactions [20]. These properties enable efficient biomass harvesting by scraping [22], as biofilms have a similar consistency to centrifuged biomass [23].

Algal/cyanobacterial biofilms also show high potential in wastewater treatment applications. The biofilm matrix is typically dominated by synergistic micro-consortia and protects microbial populations from exogenous environmental stresses (dehydration, antibiotics, chemicals, heavy metals, UV radiation, protozoan predators, etc.) [20,24]. Therefore, the microenvironment formed is less vulnerable to toxicity and variations in wastewater composition. Due to these attributes, several photosynthetic biofilm systems have been reported to increase pollutant removal rates and biomass production performance compared to suspended cultivation systems [25,26]. Moreover, the structural integrity and viscoelasticity [27] of biofilms minimizes the amount of residual biomass that ends up in the cultivation system outlet [28], thus reducing secondary pollution and disinfection costs [28]. However, the complexity of photosynthetic biofilms entails standardization and the scalability of this technology remains a considerable challenge. It is crucial that further research is carried out on the physicochemical/biological mechanisms that take place and establishing reactor design guidelines (geometry, growth surface materials, etc.).

This study introduces for the first time in the literature a simple cyanobacterial biofilm system designed for the post-treatment of biologically-treated, via the activated sludge process, brewery wastewater. This technology provides a viable alternative to conventional brewery wastewater treatment setups. The flat-plate, semi-immersed, photobioreactor was designed for the attachment of a microbial consortium dominated by the filamentous cyanobacterium *Leptolyngbya* sp. and the effectiveness of the biofilm system was compared in terms of pollutant removal and biomass production to suspended culture systems using the same *Leptolyngbya*-based microbial consortium [2] to treat brewery wastewater. The attached microbial mass was also characterized in terms of carbohydrates, proteins, and lipids to evaluate its potential in future applications. Moreover, two different biofilm reactor surfaces (glass and polyurethane) were tested to investigate the effect of
substratum hydrophobicity on microbial attachment and growth, a subject of debate in the literature [29].

2. Materials and Methods

2.1. Wastewater Characterization

The brewery wastewater used in this study was collected from a local brewery (Patras industrial zone, Greece). It was treated aerobically in the brewery’s activated sludge wastewater treatment facility, before being discharged into the central wastewater treatment unit of the industrial zone. After sampling from the outlet of the unit’s secondary settling tank, the wastewaters were filtered through 0.45 µm pore membrane filters to remove suspended solids and then stored at −20 ºC until use. The filtered wastewaters were analyzed for dissolved chemical oxygen demand (d-COD), nitrate (NO\textsubscript{3}−-N), nitrite (NO\textsubscript{2}−-N), ammonium (NH\textsubscript{4}+-N), orthophosphate (PO\textsubscript{4}^{3−}-P), total Kjeldahl nitrogen (TKN) and pH, according to Standard Methods [30]. The pH values were determined with a HANNA HI 5521 multiparameter instrument (Table 1).

| Parameter | Concentration | Units |
|-----------|---------------|-------|
| d-COD     | 190 ± 10      | mg L\textsuperscript{−1} |
| NO\textsubscript{3}−-N | 16 ± 0.5 | mg L\textsuperscript{−1} |
| NH\textsubscript{4}+-N | 5 ± 0.7 | mg L\textsuperscript{−1} |
| TKN       | 15 ± 0.9      | mg L\textsuperscript{−1} |
| PO\textsubscript{4}^{3−}-P | 5 ± 0.3 | mg L\textsuperscript{−1} |
| pH        | 8.5           | -     |

Table 1. Physicochemical characterization of the treated brewery wastewater.

2.2. Culture Identification and Inoculum Preparation

Microbial samples were collected from the secondary settling tank of the wastewater treatment plant of Patras University’s and were cultivated autotrophically in a nutrient medium (containing (in mg L\textsuperscript{−1}) KNO\textsubscript{3}, 200; MgSO\textsubscript{4}·7H\textsubscript{2}O, 100; CaCl\textsubscript{2}·2H\textsubscript{2}O, 50; K\textsubscript{2}HPO\textsubscript{4}, 108; KH\textsubscript{2}PO\textsubscript{4}, 56) suitable for algae/cyanobacteria enrichment [31]. The samples were cultivated for approximately 30 days under constant illumination of 2000 lux (or 27 µmol m\textsuperscript{−2} s\textsuperscript{−1}) provided by two cool white lamps (Phillips, 120 cm, 36 W), without any mechanical air supply. Temperature ranged between 24 and 26 ºC and pH gradually rose from 7 to 9.

After the initial cultivation period, a microbial consortium consisting of mixotrophic aggregates was obtained. Samples of the microbial culture were placed in sedimentation chambers and, using phase contrast and epifluorescence microscopy (Nikon Eclipse SE 2000 microscope), two cyanobacterial species were identified. Leptolyngbya sp. filaments in aggregate form dominated the culture (relative biomass fraction: 85%), while a second Chroococcus-like population (relative biomass fraction: 5%) was also identified. The culture also contained a significant number of heterotrophic bacteria and protozoa (relative biomass fraction: 10%). Microbial population percentages are expressed as fractions of the total biomass of the microbial community, including cyanobacteria filaments and large-sized heterotrophic bacteria (1–2 µm) [4,5]. The culture was then acclimatized in the brewery wastewater (Section 2.1), under agitation and the conditions described above, without any significant alterations in microbial population composition. This culture was used as an inoculum for all experiments when in the mid-exponential growth phase (biomass concentration of 300 mg L\textsuperscript{−1}).

2.3. Experimental Apparatus

The experiments were conducted in a flat-plate photobioreactor with constant wastewater recirculation to establish the formation of a liquid film that semi-immersed the biomass within the wastewater medium. A scheme of the biofilm photobioreactor setup is presented in Figure 1. It consisted of a 0.5 L magnetically stirred glass buffer vessel and a
The experiments were conducted in a flat-plate photobioreactor with constant irradiance of 41 \( \mu \text{mol m}^{-2} \text{s}^{-1} \) (3000 lux) generated by fluorescent lamps (Phillips, 120 cm, 36 W) placed above the apparatus. Temperature ranged between 24 and 26 °C, while pH was not adjusted as the wastewater’s initial value of 8.5 was considered favorable for cyanobacterial cultivation. Changes in pH and temperature conditions were monitored using a HANNA HI 5521 multiparameter instrument.

Figure 1. Schematic representation of the experimental setup.

Initially, the buffer vessel contained 0.5 L of culture (90% v/v brewery wastewater medium and 10% v/v inoculum prepared as described in Section 2.2). At the start of each experiment, the peristaltic pump was activated, generating a recirculating flow of 100 mL min\(^{-1}\) from the buffer vessel towards the top of the tilting structure. The culture medium flowed through three nozzles towards the bottom end of the structure, forming a thin liquid film over the tilted surfaces (Figure 1) and then drained back into the buffer vessel. Water losses due to evaporation were compensated for by the addition of distilled water into the recirculation system.

Both experimental setups were under constant average irradiance of 41 \( \mu \text{mol m}^{-2} \text{s}^{-1} \) (3000 lux) generated by fluorescent lamps (Phillips, 120 cm, 36 W) placed above the apparatus. Temperature ranged between 24 and 26 °C, while pH was not adjusted as the wastewater’s initial value of 8.5 was considered favorable for cyanobacterial cultivation. Changes in pH and temperature conditions were monitored using a HANNA HI 5521 multiparameter instrument.

2.4. Determination of Total Biomass & Nutrient Concentrations

To determine pollutant and suspended biomass concentrations, samples were taken from the stirred buffer vessel. Total suspended biomass concentration (expressed as total suspended solids, TSS) was determined by filtration through 0.45 μm pore membrane filters [30]. The filtrate was analyzed for dissolved chemical oxygen demand (d-COD), nitrate \((\text{NO}_3^-\text{-N})\), nitrite \((\text{NO}_2^-\text{-N})\), ammonium \((\text{NH}_4^+\text{-N})\), orthophosphate \((\text{PO}_4^{3-}\text{-P})\), total Kjeldahl nitrogen (TN), and pH according to Standard Methods [30].

To estimate the amount of biomass that adhered to the biofilm reactor surface, two 1 cm-wide strips of attached biomass were carefully scraped using a hand scraper. The obtained biomass was then re-suspended in 10 mL of distilled water, stirred, and subsequently filtered through 0.45 μm pore membrane filters for total biomass determination [30]. The obtained concentration was converted to biomass per biofilm area. The total amount of biomass produced in the biofilm setup was calculated by summing the suspended
and attached fractions of biomass per reactor working volume. Similarly, percentages of attached biomass (attachment efficiency) were calculated by dividing the attached biomass by the total amount of biomass.

2.5. Biomass Harvesting and Determination of Intracellular Compounds

After the end of each experiment, the treated wastewater was drained into the buffer vessel and biomass was harvested from the growth surface by scraping. After re-suspension in 50 mL of distilled water, it was centrifuged, washed twice to remove nutrient residues and other impurities, and dried overnight at 105 °C. Previous studies utilizing the same Leptolyngbya-based microbial consortium have shown that it accumulates up to 40% of its dry biomass weight in carbohydrates [4,5], and for this reason carbohydrates were chosen as the target biomass compound of the present study. Intracellular sugar content (expressed as glucose equivalents) was estimated photometrically according to Dubois et al. [32]. Firstly, 1 mg of dried biomass reduced to powder was mixed with 5 mL of distilled water. Then, 1 mL of this suspension was then mixed with 1 mL of 5% (w/v) phenol solution and 5 mL sulfuric acid solution and measured at 490 nm after 30 min [4,5,32]. To determine protein content, 10 mg of dried harvested biomass were suspended into 5 mL of phosphate buffer solution (pH = 7.3). The resulting suspension was sonicated for 40 min in a sonication bath (Julabo USR 3) for cell lysis [4,5,33]. After filtering through a 0.45 µm pore membrane filter, the protein content was determined following Bradford [34]. To determine lipid content, 20 mg of dried biomass underwent extraction with 2:1 (v/v) mixture of chloroform and methanol, washing with 0.88% (w/v) KCl solution, and drying over anhydrous Na2SO4. The solvent was evaporated, and the extracted oil was quantified gravimetrically [4,5,35].

2.6. Statistical Analysis

All analyses were carried out in duplicate with a relative standard deviation not exceeding 5%. Results were expressed as mean ± standard deviation. Statistically significant differences between data were evaluated using the t-student confidence interval for 95% probability. In this statistical analysis, the data were assumed to follow the t-student distribution and the confidence interval for the difference of a pair of mean values was calculated.

3. Results & Discussion

3.1. Attachment Efficiency

After activating the wastewater recirculation system, the culture started flowing over the inclined biofilm surface initiating adhesion and biofilm formation. A different initial adhesion rate was observed for the two surface materials tested (Figure 2). In the case of glass, which is a mildly hydrophilic material [36], biomass attachment was rapid and efficient, surpassing 90% after two days of cultivation. On the contrary, adhesion on the polyurethane surface did not occur at an equally high rate, as after two days of cultivation 42.6% of the total biomass remained in suspension. Following this initial adhesion phase, the fraction of biomass that remained attached exceeded 98.3% and 97.4% for glass and polyurethane, respectively, nine (9) days into the experiments. At the end of cultivation, both experiments exhibited complete biomass attachment (>99%), while statistically significant differences on biomass attachment were not observed between the two surface materials.

There have been conflicting reports on whether hydrophobic or hydrophilic surfaces enhance adhesion [28,29,36]. Hydrophobic surfaces are generally considered more susceptible to biofilm formation; however, factors such as surface roughness, wettability, hydrodynamics, and the nature of EPSs produced by the microorganisms play equally important roles [29,36,37]. In this study, the more hydrophilic surface (glass) favored initial adhesion; thus, it is recommended for attached cultivation of Leptolyngbya sp. This finding could indicate that the Leptolyngbya-based consortium utilized in this study produces EPSs of a mostly hydrophilic nature. Various species of cyanobacteria have been reported to se-
crete hydrocolloidal EPSs which draw extensive research interest due to their applications in the bio-materials field (hydrogels, etc.) [38].

The predominant role of EPSs and cell-to-cell interactions in biofilm formation is further indicated by the high attachment efficiency after the initial adhesion phase. The microbial community appeared to create an EPS/cell “coating” on the substratum surface, negating its impact on adhesion efficiency. Similar amounts of biomass remained attached on both materials after this point as the biofilm matured (irreversible adhesion). Mazumder et al. reported that microbial accumulation on surfaces can significantly decrease their hydrophobicity and binding characteristics [39].

3.2. Pollutant Removal

Although primary aerobic wastewater treatment is generally effective at removing organic pollutants, it often does not reduce inorganic pollutant concentrations (especially phosphorus) to a level that allows safe disposal or reuse and, therefore, requires secondary physicochemical treatment [40]. The post-treatment biofilm reactor introduced in this study aims to effectively remove residual pollutant concentrations for safe effluent disposal. Figure 3a presents the results of NO$_3^-$-N, NH$_4^+$-N, and PO$_4^{3-}$-P removal over the course of the 15-day cultivation period. Similar inorganic pollutant removal rates and no statistically significant differences were observed for both surface materials used. NH$_4^+$-N is usually the preferred nitrogen substrate for cyanobacterial cultivation as its assimilation is less energy demanding and, in this study, was rapidly consumed after two days of cultivation [41]. However, it should be noted that under the alkaline pH conditions of these experiments (pH rose from 8.5 to 9.45) a large fraction of the initial ammonium concentration evaporated in the form of ammonia gas [41]. NO$_3^-$-N gradually reduced by 67.0% (glass) and 67.6% (polyurethane) within the first ten days of cultivation, while TKN concentration followed a similar trend as, in both cases, its concentration dropped below 5 mg L$^{-1}$. It is worth noting that NO$_2^-$-N concentrations did not exceed 0.05 mg L$^{-1}$ at any point during the experiments. The microbial consortium was also effective at removing phosphorus compounds. PO$_4^{3-}$-P concentrations reduced by 76.2% and 75.5% for the glass and polyurethane growth surfaces, respectively. After post-treatment, all pollutant concentrations were found to comply with European regulations for effluent disposal (Articles 4 and 5 of the Urban Wastewater Directive, 91/271/EEC) [42].
In absence of an external mechanical CO2 feed, the microbial consortium utilized organic pollutants as its primary carbon source. In the case of the glass growth surface, 71.6% of the initial d-COD was removed, while for polyurethane the corresponding removal percentage was 76.2% (Figure 3b). Note that statistically significant differences in d-COD removal rates were not observed between the two surface materials. *Leptolyngbya* sp. is known to be a mixotrophic cyanobacterium and for this reason it has been utilized for treatment of wastewaters containing high concentrations of organic pollutants in past research [4,5,43,44]. Tsolcha et al. investigated the performance of a *Leptolyngbya*-based mixotrophic culture in the treatment of various agro-industrial wastewaters (cheese whey, poplar sawdust, and grass hydrolysates) containing d-COD concentrations of up to 3660 mg·L\(^{-1}\) and recorded removal percentages of up to 93% without the provision of an additional inorganic carbon source [44].

Table 2 presents pollutant removal percentages of the present study compared to those obtained by Papadopoulos et al., who investigated post-treatment of the same wastewater using a suspended *Leptolyngbya*-based culture under identical pH, light, and temperature conditions [4]. The biofilm photobioreactor manages to equal or exceed all the removal percentages recorded in suspended cultivation. The greatest deviation between the two cultivation systems is in d-COD removal. Specifically, in attached cultivation 70–83% of the initial d-COD was removed within nine (9) days, in comparison to 24% in suspended cultivation over the same period. This behavior can be interpreted if the secretion of EPSs is considered. Under nitrogen limitation, which occurred throughout the experiments, cyanobacteria produce exopolysaccharides and other slowly biodegradable macromolecules [20,21] that add to the existing organic loading of the wastewater, thus reducing observed d-COD removal rates as cultivation progresses. Namely, Papadopoulos et al. documented an extracellular polysaccharide concentration as high as 80 mg·L\(^{-1}\) in nitrogen limitation conditions [4]. On the contrary, in the case of the biofilm reactor, it is hypothesized that a large fraction of these EPSs remained bound within the biofilm matrix resulting in a higher COD removal rate.
Similar growth rates were observed for both growth surface materials tested, although the use of a glass growth substratum appears to slightly enhance biomass growth in comparison to polyurethane. The corresponding maximum growth rates were 66 mg·L\(^{-1}\) (glass) and 59 mg·L\(^{-1}\) (polyurethane). After an exponential growth phase during the first 6–10 days of cultivation, total biomass concentration stabilized at around 406 mg·L\(^{-1}\) for glass and 392 mg·L\(^{-1}\) for polyurethane. In terms of attached biomass per biofilm surface area, the respective calculated values were 13.1 and 12.8 g·m\(^{-2}\).

### Table 2. Pollutant removal percentages after nine (9) days of attached cultivation (glass and polyurethane growth surfaces) and suspended cultivation

| Substance    | Attached Cultivation | Suspended Cultivation ¹ |
|--------------|----------------------|-------------------------|
|              | Glass | Polyurethane | Glass | Polyurethane | ¹ |
| NO\(_3^+\)-N | 67.2  | 76.5 | 50.0 |
| NH\(_4^+\)-N | 100   | 100 | 100 |
| PO\(_4^{3-}\)-P | 66.0 | 65.4 | 57.0 |
| d-COD       | 83.2  | 70.7 | 24.0 |

¹ Adapted by Papadopoulos et al. (2020) [4].

#### 3.3. Biomass Production

The sum of suspended and attached biomass per reactor working volume and reactor surface growth surface over the course of the 15-day cultivation period is presented in Figure 4. Similar growth rates were observed for both growth surface materials tested, although the use of a glass growth substratum appears to slightly enhance biomass growth in comparison to polyurethane. The corresponding maximum growth rates were 66 mg·L\(^{-1}\)·d\(^{-1}\) (glass) and 59 mg·L\(^{-1}\)·d\(^{-1}\) (polyurethane). After an exponential growth phase during the first 6–10 days of cultivation, total biomass concentration stabilized at around 406 mg·L\(^{-1}\) for glass and 392 mg·L\(^{-1}\) for polyurethane. In terms of attached biomass per biofilm surface area, the respective calculated values were 13.1 and 12.8 g·m\(^{-2}\).

![Figure 4. Total biomass concentration per working reactor volume (left y-axis) and reactor surface area (right y-axis) over cultivation time for glass and polyurethane growth surface materials.](image)

The work of Papadopoulos et al. [4] provides a reference point to assess the performance of the biofilm reactor introduced in the present study. Both biofilm setups (glass/polyurethane surfaces) significantly enhanced biomass growth rates compared to suspended cultivation. Specifically, on the ninth day of cultivation the suspended culture produced 360 mg·L\(^{-1}\), i.e., 11% less than the biofilm cultivation experiments. Biofilms have been reported to enhance biomass growth in many studies [28]. For example, Gao et al. studied synthetic secondary effluent treatment using CO\(_2\)-aerated attached and suspended growth membrane bioreactors and reported a significant increase in biomass productivity and nutrient removal in the case of the biofilm reactor [26]. Similarly, Huang et al. compared the performance of a membrane biofilm photobioreactor to a suspended growth
setup utilizing the same BG-11 cultivation medium and reported a 30% increase in biomass productivity when using biofilm reactors [45]. In this study, it is suspected that the thin water film flowing above the biofilm surface enables more efficient CO2 transfer between the gas (atmosphere) and liquid phases thus providing an additional inorganic carbon source. On the contrary, in cultures that are fully immersed in the liquid medium CO2 diffusion can be a slow process and limit photosynthetic growth [18]. Cyanobacteria mixotrophy is known to enhance growth rates and photosynthesis efficiency compared to strictly autotrophic or heterotrophic conditions [46]. Moreover, the thinness of the liquid film allows more efficient light transmission. The brewery wastewater treated in this study has a brown/yellow color and subsequently absorbs part of the available light before it radiates the cyanobacterial cells. Radiation absorptivity is proportional to the distance light travels in a material (Beer–Lambert Law) [47]. Thus, reducing the light path offers a significant advantage to the biofilm reactor in comparison to suspended systems where light travels several centimeters through the liquid phase.

3.4. Biomass Composition

After cultivation, the attached biomass was scraped off the growth surface and analyzed in terms of carbohydrate, protein, and lipid contents (Figure 5). While inoculum carbohydrate content was 38.4%, the respective content of the harvested biofilm biomass was over 60% for both surface materials tested. The significantly increased carbohydrate content can be attributed to the extensive nitrogen limitation that occurred in both experiments. Nitrogen is an essential nutrient for protein and nucleic acid synthesis and in cases of shortage cyanobacteria tend to store photosynthetic energy and carbon in the form of glycogen as carbohydrate [48] instead of initiating new cell synthesis. As cultivation progressed protein and lipid contents slightly decreased from 27% and 4% to around 22–23% and 3–3.5%, respectively.

![Carbohydrate, protein, and lipid contents of dry biomass weight (DBW) taken from the culture inoculum and the biofilm harvested from glass/polyurethane (PU) growth surfaces.](image)

Papadopoulos et al. reported carbohydrate content of 43.65% when utilizing the same microbial consortium in suspended cultivation mode for wastewater treatment, despite operating under similar conditions [4]. As discussed in Section 3.2, the cyanobacterial population applied in these experiments secretes large quantities of EPSs (up to 80 mg·L·1) under conditions of stress/nutrient limitation. Biomass attachment ensures that a large fraction of these molecules remain within the biofilm matrix instead of dissolving into the wastewater, thus raising the carbohydrate content of the harvested biofilm as cyanobacterial
EPSs comprise mostly of exopolysaccharides [21]. Therefore, biofilm cultivation could be an effective strategy for carbohydrate-enriched biomass production.

3.5. Practical Applications and Research Perspectives

The photosynthetic biofilm approach described in this study could complement conventional brewery wastewater treatment systems such as activated sludge. Life cycle analyses indicate that photosynthetic systems provide a less energy-demanding option (~47% operating cost reduction) with a positive environmental impact [49]. The use of biofilms can further enhance the financial viability of photosynthesis-based wastewater treatment facilities as it entails efficient, low-cost, biomass harvesting [23]. The biomass composition of this study’s microbial consortium adds a new dimension to the sustainability of the process. Due to its high carbohydrate content, the biofilm biomass can be used as feedstock for bioethanol, a product of significant commercial value [50]. To realize the system’s full biotechnological potential more research should be carried out on analyzing the biofilm’s EPSs profile, as certain cyanobacterial EPSs could be sustainable source materials for the production of hydrocolloids, bio-sourced materials, and biological agents [21,38].

4. Conclusions

A novel biofilm photobioreactor was introduced for brewery wastewater post-treatment. By semi-immersing the biomass within the liquid medium, the flat-plate bioreactor design enabled efficient gas transfer and light transmission that enhanced pollutant removal and biomass production. Both growth surface materials tested (glass/polyurethane) managed to achieve complete biomass attachment (>99%); however, the hydrophilic material (glass) performed better at the initial stages of microbial adhesion. This finding suggests that for mature cyanobacterial biofilms, the effect of the EPSs produced prevails over the supporting surface hydrophobicity. Biomass harvesting and dewatering are considered the bottleneck that hinders wastewater treatment using algae/cyanobacteria in full-scale applications. The reactor design suggested by the present study successfully addresses these issues without sacrificing water treatment performance or produced biomass quality. Namely, d-COD removal exceeded 70% for both materials tested, a significant increase compared to suspended cyanobacteria-based systems for brewery wastewater post-treatment. Moreover, due to extensive nitrogen limitation, EPSs retention and efficient irradiation, the harvested biomass accumulated a remarkable carbohydrate content of 61%. The carbohydrate-enriched biofilm could be utilized as a substrate for high yield bioethanol production as well as EPSs extraction. Overall, the results suggest that a flat-plate photobioreactor design using a hydrophilic supporting material could be a simplistic yet efficient approach for sustainable, low-cost, brewery wastewater post-treatment.

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Abbreviations

Dissolved Chemical Oxygen Demand  d-COD
Total Kjeldahl Nitrogen  TKN
Nitrate Nitrogen  NO₃⁻-N
Nitrite Nitrogen  NO₂⁻-N
Ammonium Nitrogen  NH₄⁺-N
Orthophosphate Phosphorus  PO₄³⁻-P
Extracellular Polymeric Substances  EPSs
Polyurethane  PU

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