Pilot scale photobioreactor system for land-based macroalgae cultivation

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Abstract

Marine macroalgae such as Ulva intestinalis have promising properties as feedstock for cosmetics and pharmaceuticals. However, since the quantity and quality of naturally grown algae vary widely, their exploitability is reduced – especially for producers in high-priced markets. Moreover, the expansion of marine or shore-based cultivation systems is unlikely in Europe, since promising sites either lie in fishing zones, recreational areas, or natural reserves. The aim was therefore to develop a closed photobioreactor system enabling full control of abiotic environmental parameters and an effective reconditioning of the cultivation medium in order to produce marine macroalgae at sites distant from the shore. To assess the feasibility and functionality of the chosen technological concept, a prototypal plant has been implemented in central Germany – a site distant from the sea. Using a newly developed, submersible LED light source, cultivation experiments with Ulva intestinalis led to growth rates of 7.72 ± 0.04 % day−1 in a cultivation cycle of 28 days. Based on the space demand of the production system, this results in fresh mass productivity of 3.0 kg m−2, respectively, of 1.1 kg m−2 per year. Also considering the ratio of biomass to energy input amounting to 2.76 g kWh−1, significant future improvements of the developed photobioreactor system should include the optimization of growth parameters, and the reduction of the system’s overall energy demand.

Keywords Macroalgae · Photobioreactor · Ulva intestinalis · LED light source · Production system · Closed system cultivation

Introduction

The green alga Ulva intestinalis is a typical representative of Chlorophyta and grows on rocky ground or as an epiphyte in the intertidal zones of nutrient-rich marine waters (Budd and Pizzola 2008). Appreciated for its high content of iron, magnesium, and selenium (Circuncisão et al. 2018), it is currently being cultivated for human consumption in Korea, Japan, India, and Indonesia (FAO 2018). Moreover, U. intestinalis appears highly promising as raw material for cosmetics and pharmaceuticals. Its antioxidant, fungicidal, antibiotic, antiviral, and anticarcinogenic properties have been assessed in numerous scientific studies (e.g., Farasat et al. 2014; Murphy et al. 2014; Kosanić et al. 2015). The biochemical substance responsible for the mentioned effects is called ulvan and consists of sulfated polysaccharides only present in the cell walls of the genus Ulva. However, with contents ranging from 2 to 60% in algal dry mass, the availability of these gelling carbohydrates largely depends on the quality of the algal material produced and thus on the production process itself (Kidgell et al. 2019). In this respect, a close look at the specifics of current production systems reveals a noteworthy potential for improvement.

Basically, macroalgae can be cultivated in three types of production systems: marine aquaculture, land-based tank or pond cultivation, and closed photobioreactor systems. Marine aquaculture is widely established but lacks the means to control cultivation parameters (Hafting et al. 2015): Algae growth rates are determined by oceanographic and meteorological parameters as well as the prevailing water quality in the production area (Buck and Grote 2018). Additionally, the crop quantity and quality are increasingly limited by the effects of climate change and ocean acidification (Kelly and Hofmann 2013) as well as proliferating pathogens, epiphytes, or herbivores (Hurd et al. 2014; Hafting et al. 2015; Kerrison et al. 2015; Fernand et al. 2017). These negative factors can be mitigated by cultivating algae in land-based...
tanks or ponds: Both the cultivation parameters and the presence of detrimental organisms can be monitored and partially controlled in these semi-closed systems. However, the key feature of such cultivation systems is their land-based but coastal location due to their constant need for fresh seawater (Pereira et al. 2012, 2013; Hafting et al. 2015). Consequently, both marine aquaculture and land-based tank cultivation compete with often more profitable uses of coastal regions, such as recreational activities, tourism, and fishery – a circumstance that is denominated critical for the development of macroalgae cultivation (Hafting et al. 2015). Overcoming these disadvantages, closed production systems can increase the degree of freedom in the choice of location, the biosecurity of the overall production process, and the value-added potential of the produced algae. Photobioreactor systems allow for the monitoring and fine-tuning of the growth environment and thereby lead to a production outcome predictable in quantity and quality. Albeit, the cultivation of macroalgae in closed photobioreactor systems is currently facing significant challenges preventing an implementation on an industrial scale: In the prototypes developed to date, insufficient temperature control and inadequate light supply limit the achievable growth rates. Also, the lack of an effective recycling strategy for the cultivation medium implies frequent medium changes and thus a seawater supply nearby (Chemodanov et al. 2017; Sebök et al. 2017, 2018; Mhatre et al. 2018).

To summarize, the current need for research revolves around a reliable provision of growth-promoting cultivation parameters for macroalgae production at sites distant from the shore. The aim is, therefore, to develop a closed photobioreactor system enabling full control of abiotic environmental parameters and an effective reconditioning of the cultivation medium. Special focus is laid on the supply of light in adequate intensity and spectrum for macroalgae production using energy-efficient light-emitting diodes (LED).

Materials and methods

A closed photobioreactor system was designed and implemented for the production of macroalgae in central Germany (Geebelsee, Thuringia). To assess the functionality of the overall system, cultivation experiments were undertaken with different algae species over the course of 2 years. The following sections cover the materials and methods used to develop, characterize, and evaluate the invented production system.

Land-based closed photobioreactor system

The designed photobioreactor system consists of numerous elements selected with regard to the functional requirements and energy efficiency of the macroalgae production process. To allow for a subsequent adaptation to local spatial circumstances or the variation of production quantities, particular attention has been paid to a modular system design. Implementing a basic setup, four breeding modules have been installed together with one conditioning and one system control module on the premises of the Geratal Agrar GmbH in Gebesee (D). A piping and instrumentation diagram in accordance with EN ISO 10628 is depicted in Fig. 1.
Breeding module

Each breeding module consists of an Intermediate Bulk Container (IBC) tank (100 × 120 × 116 cm) prepared for macroalgae cultivation as depicted in Fig. 2. To enable algae handling and harvesting, the inner bladder of the IBC was cut open and lined with an NR/SBR rubber band (500 × 30 × 0.1 cm) at its top edge. An aluminum plate (110 × 130 × 0.5 cm), reinforced with angle profiles on the upper and lower side, serves as tank cover and can be lifted using a rope (PP, ∅12 mm), and pulley block (Schöco 22B 6:1) attached to the ceiling. To provide for an uplifting flow within the breeding tanks (tumble culture), an aeration pump (AquaForte Air Pump V60, \( V = 3.6 \ m^3 \ h^{-1} \)) is connected to the aeration element attached to the underside of the tank lid. The dimensions of this aeration element (PVC pipes, ∅2 cm) have been varied in an iterative development process in order to achieve a uniform, gentle rotation of the algae in the breeding tanks. The implemented structure (80 × 60 × 80 cm) provides 4 outlet openings at a depth of 80 cm and achieves the desired, slowly rotating flow. Each tank has further been lined with a finely meshed net (100 × 100 × 120 cm, mesh size 1 mm), to eliminate risks of pipe clogging due to algae being sucked through the bottom tank valves. To test the influence of light supply on algae growth, different illumination setups have been implemented pairwise in the four breeding tanks: Two tanks (PBR 1 and PBR 2) are equipped with newly developed, submersible LED cylinders (396 LED emitting in 8 wavelength ranges, installed within a DURAN glass cylinder ∅20 cm × 1.2 m, custom made by Lucelab) allowing for a species-specific light supply in terms of intensity and spectrum. The remaining two tanks (PBR 3 and PBR 4) are fitted with four fluorescent tubes recommended as a light source for aquatic plants by their manufacturer (Dennerle, Trocal T5 Special Plant 39 W).

Conditioning module

The aim of the conditioning module is to ensure constant abiotic cultivation parameters as well as to minimize the fresh water and artificial sea salt demand of the production process. To this end, each breeding module is connected to the conditioning module through bottom tank valves and a collection pipe leading to a small sedimentation tank (PE, ∅50 × 115 cm). This tank is equipped with a perforated plate (∅ 5 mm) situated between the water inlets at the bottom level and the outlet positioned at a height of 50 cm. To further purify the cultivation medium from non-sedimenting particles, a cartridge filter cascade (1000, 500, 100, 50, 10, 1, 0.5 μm, Mahle) and a UV-C unit (Helix Max, Aqua Medic) have been integrated into the conditioning line. After these purification steps, the cultivation medium is being conveyed by a membrane pump (\( V = 140 \ L \ h^{-1} \), SHURflo 2088–573-534, Keller) to a cooling unit (Titan 4000, Aqua Medic), mixed with a nutrient solution added through a dosing pump (BL10 Black Stone, Hanna Instruments) and homogenized by a static mixer (PVC ∅25 mm, Kwerk). The conditioning line ends with a central overflow unit from which the cultivation medium is returned evenly and in equal proportions to the breeding tanks. The conditioning module has been designed to recycle the cultivation medium of all four connected breeding tanks on a daily basis. Thus, its recycling capacity sums up to 2800 L day\(^{-1}\) in total.

System control module

To improve user-friendliness and operational safety, the photobioreactor system has been automated as far as practicable. With all actuators (nutrient and light supply, cooling unit, and air and fluid pumps) connected to central programmable logic controllers (Controllino Maxi, Conelcom), the amount of work and time required for macroalgae cultivation can be sensibly reduced. In addition, a variety of sensors have been

Fig. 2 Breeding module: exterior (left) and interior view (mid, right) with net, aeration element, and LED light source
installed to improve general plant safety. Special attention was laid on the flow respectively the pressure in the pipes of the conditioning module. In addition to sensors monitoring the flow in the main pipe of the conditioning module, a process-integrated calculation of the four tank outflows \( V_{\text{out}} \) is implemented using the tank-specific inflow \( V_{\text{in}} \) (measured with FCH-midi-POM, BioTech) and the volume difference in the tanks \( \Delta V \) (via pressure difference \( \Delta p \), A-10, WIKA) in the considered time period \( \Delta t \) according to Eq. (1):

\[
V_{\text{out}} = V_{\text{in}} - \frac{\Delta V}{\Delta t} \quad \text{with} \quad \Delta V = -\frac{1}{\rho \cdot g \cdot A} \Delta p
\]  

(1)

\( \rho \) = medium density, \( g \) = gravity, \( A \) = sensor area.

An additional pressure sensor (WIKA, A-10) located in the collecting pipe between the sedimentation tank and the filter cascade provides information on the loading of the filter cartridges. The measured values are being transferred to a PC using custom-made software developed in order to monitor the production process. A VPN-secured internet connection enables remote monitoring of the system processes. Additionally, all pressure sensors are linked to an emergency stop switch which allows for an automatic system shut down in the event of increased (and damaging) pressure in the conditioning module. Altogether, the system control module is allowing for constant abiotic cultivation parameters and prevents major damage to the production system.

**Cultivation experiments**

All cultivation experiments were undertaken in the developed photobioreactor system presented in Sect. 2.1 with respect to the (known) physiological requirements of the chosen green macroalga *Ulva intestinalis*. Starting with a description of the origin of the algal biomass, the following subsections give an overview of both the overall experimental setup and the evaluation procedure used.

**Macralgal biomass**

The cultivation experiments are carried out with the alga *U. intestinalis* originating from the Baltic Sea (54°31’13.5”N 13°40’2.0”E, near Sassnitz, Germany). To avoid harming wild marine environments, algae are exclusively being collected in large populations and according to the wildlife-protecting indications summarized by Mac Monagail et al. (2017). On-site, the collected biomass is washed in fresh seawater, freed from sand, and examined for visible signs of grazers or epiphytes. Healthy phyllodes are placed in plastic bags (6 L), wrapped in cloths, and placed in polystyrene boxes (25 L) cooled with ice cubes. A sensor (Ebro, EBI 20) positioned between the algae bags is used to check the temperature in the cooling box. Temperature is kept below < 18 °C during the duration of the transport (~7 h) by gradually adding ice cubes.

**Experimental setup**

The cultivation experiments are carried out from July 6 to August 2, 2018 (28 days) in four breeding tanks holding 700 L of cultivation medium each. The operating schedule of the photobioreactor system provides for nutrient addition once daily, the continuous aeration of all tanks, and a photoperiod of 12:12. The fluid pump – and thus the entire treatment line – pauses four times a day for one hour to reduce pump deterioration. Cultivation conditions are monitored by continuously measuring the system parameters: pressure (WIKA, A-10), total flow of the cultivation medium (FCH-midi-POM, BioTech), inflow to the individual breeding tanks (FCH-midi-POM, BioTech), pH value (Jumo, tecLine 201,020) as well as the temperature of the cultivation medium (PT 100, SWAN) and air (Ebro, EBI 20). In addition, manual measurements of the turbidity (VisoTurb 700 IQ F, WTW) and salinity (areometer, Amarell) of the cultivation medium are carried out on a weekly basis.

Cultivation parameters are adjusted to the physiological requirements of the chosen algal species *U. intestinalis*. With a temperature of 20 °C and a salinity of 10, the cultivation medium made out of drinking water and an artificial sea salt mixture (Tropic Marin® CLASSIC) sticks closely to the natural seawater parameters in the collection zone – a marine environment with particularly low salinity. To enhance algal growth rates, initial stocking densities are kept low with 0.13 g L\(^{-1}\) (90 g per tank), and the nutrient supply is raised to 12 mg NH\(_4\) day\(^{-1}\) and 5.33 mg PO\(_4\) day\(^{-1}\) using a stock solution based on a universal mineral fertilizer (NovaTec, COMPO). The breeding tanks PBR 1 and PBR 2, equipped with submersible LED light cylinders, are provided with a light spectrum enhanced in the red and blue wavelength range (see Fig. 3) and a light intensity of 990 μmol photons m\(^{-2}\) s\(^{-1}\). The breeding modules PBR 3 and PBR 4 are equipped with standard aquarium fluorescent tubes providing a light intensity of 254 μmol photons m\(^{-2}\) s\(^{-1}\) at the water surface and a light spectrum with little intensity in the blue wavelength range, see Fig. 3. Thus, the number of replicates per light setting is \( n = 2 \).

**Calculation of biomass growth rate**

Fresh algal biomass is being weighed after using a standardized procedure to reduce adherent water. The algae are repeatedly centrifuged in a conventional salad spinner until the withdrawn water weighs < 1 g. Subsequently, biomass growth is evaluated using the biomass yield \( \Delta m \) during the cultivation
experiment defined in Eq. (2) and the specific daily growth rate \( \mu \) as defined in (Sterner and Elser 2002) with Eq. (3):

\[
\Delta m = m_{28} - m_0 \text{ in [g]}
\]

\[
\mu = \ln \left( \frac{m(t)}{m_0} \right) \cdot \frac{1}{t} \cdot 100 \text{ in [% day}^{-1}]\]

with the biomass \( m(t) \) at time \( t \), the initial biomass \( m_0 \) at time \( t=0 \) day, and the resulting biomass \( m_{28} \) at time \( t=28 \) days. To avoid damaging the delicate, filamentous \( U.\ intestinalis \), biomass weight is only assessed at the beginning (\( t=0 \) day) and the end (\( t=28 \) days) of the cultivation experiment.

### Energy consumption per production cycle

The energy consumption (EC) of the developed photobioreactor system has been calculated as shown in Eq. (4) using the energy demand (ED) of each system component as specified by its manufacturer and the daily operating hours (OH) during the cultivation process:

\[
EC = ED \cdot OH \text{ in [kWh day}^{-1}]\]

During the cultivation experiments with \( U.\ intestinalis \), additional measurements were undertaken with a power meter (Brennenstuhl, PM 231 E) for the feedback-controlled (temperature) cooling unit and the newly developed, prototypal LED light source.

### Results

The performance of the newly developed photobioreactor system is being ascertained by evaluating its overall functionality as well as the achieved results regarding the growth rates and energy consumption during the cultivation experiments with the green alga \( U.\ intestinalis \).

### Photobioreactor system

When developing the presented photobioreactor system, special focus was laid on the provision of optimal abiotic cultivation parameters for marine macroalgae, the reduction of freshwater demand to a minimum, and the realization of an energy-efficient production system with limited space demand. The following sections share the observations and findings made during the 24 months of operation of the prototypal plant.

### Species-specific cultivation parameters

The selected process technology allows for the adaptation of parameter values to the physiological needs of the specific macroalgae type being cultivated. The salinity of the cultivation medium can be adjusted manually by adding a defined amount of artificial sea salt (Tropic Marin CLASSIC) to freshwater; the temperature of the cultivation medium can be varied and automatically controlled through the use of a cooling unit; the nutrient supply is fully automated and adjustable using a fertilizer stock solution and a dosing pump. Similarly, the newly developed LED cylinders provide a light quality that can be adjusted between 0 and 1100 \( \mu \text{mol photons m}^{-2} \text{s}^{-1} \) in intensity and between 425 and 730 nm in the spectrum, see Fig. 4.

Altogether, the possibility of varying the cultivation parameters salinity and temperature of the cultivation medium, as well as the nutrient and light supply to the macroalgae, allows for the production of different species by simulating the conditions in their natural environments.
Moreover, by adjusting the named parameters closely to the physiological requirements of the cultivated algae, biomass yields and growth rates can – in theory – be maximized and optimized compared to the results achievable in marine aquaculture.

Recirculation of cultivation medium

As a prerequisite for a closed-loop production system, the recirculation of reconditioned cultivation medium is of crucial importance for the site independence of the developed cultivation plant. Accordingly, great attention was laid on the preservation and regeneration of the quality of the medium, especially in terms of salinity, nutrient content, and translucency. While the first two parameters can easily be readjusted whenever necessary, the latter requires a continuous and effective filtering treatment. Several filtering setups were tested during the 24 months of operation of the prototypical cultivation plant. Two major findings can be summarized as follows:

- The sedimentation tank substantially and reliably increases the functionality and operational safety of the conditioning module. In addition to the removal of particles, it allows for constant pressure levels in the collecting pipe leading to the fluid pump and thereby increases its longevity.
- Constant turbidity values ≤1.4 NFU can be guaranteed for at least 12 months of operation when a UV-C unit is being used in addition to a filtering cascade removing particles ≥0.5 μm, see Fig. 5.

Accordingly, production downtimes for maintenance can be reduced to four times a year and the intervals for total cultivation medium exchange were expanded to once a year.

However, it is important to notice that the water levels and salinity of the cultivation medium need to be adjusted once a month since losses occur through water evaporation and salt deposition on various system components.

Area and volume requirements

The space requirement of the developed photobioreactor system mainly depends on the number of breeding modules installed. With a basic tank size of 1.2 m² and an additional 0.5 m² of floor area needed for maintenance and algae handling, in total 1.7 m² are required for each breeding module. Considering a minimum height of 3 m to allow for cover lifting and algae handling, the volume to be envisaged per breeding module remains just under 6 m³. Thus, the space occupied by the four breeding modules installed at Geratal Agrar amounts to 6.8 m², respectively, 24 m³. In contrast, the conditioning and system control modules account for 4 m², respectively, 6 m³ together. Thereby, the total space demand for the basic setup of the photobioreactor system presented in this article sums up to 10.8 m², respectively, 30 m³.

Cultivation experiments

The developed photobioreactor system has been used for the cultivation of the macroalgae species *U. intestinalis*, *Fucus vesiculosus*, and *Palmaria palmata*. However, due to difficulties with the provision of vital algae as initial biomass and technical issues during the cultivation experiments, the experiments carried out with *F. vesiculosus* (temporarily defect light cylinders) and *P. palmata* (decreased vitality due to transport damage), did not lead to conclusive results. Hence, this article and the following subsections only present the results of the cultivation experiment achieved with the green alga *U. intestinalis*.

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**Fig. 5** Cultivation medium (f.l.t.r.): fresh medium; after 12 months when being reconditioned through filtering cascade including UV-C unit; after 4 weeks of operation when being reconditioned with filtering cascade excluding UV-C unit

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Growth rates of Ulva intestinalis

The cultivation experiments with *U. intestinalis* were started with an identical amount of initial fresh biomass (*m*₀) in each breeding tank: 90 g in PBR 1 to PBR 4. After 28 days, depending on the light source installed, the green macroalgae achieved biomass yields of 694.5 ± 8.5 g in breeding tanks equipped with submersed LED lights (PBR 1 and PBR 2) compared to 17 ± 6 g in those equipped with fluorescent tubes (PBR 3 and PBR 4). The resulting growth rates amount to 7.72 ± 0.04 % day⁻¹ in PBR 1 and PBR 2, respectively, to 0.61 ± 0.20 % day⁻¹ in PBR 3 and PBR 4. Interestingly, the algae also grew on the submersed light cylinders and thus increasingly reduced light supply over the course of time: As depicted in Fig. 6, both cylinders were fully grown over after 28 days. Table 1 summarizes the achieved results with *U. intestinalis* with reference to the light source and intensity provided in each breeding tank. Since all other parameters were kept constant and equal for all four tanks, the light parameters seem of primordial and decisive importance for the productivity of the grown algae.

Electricity consumption

The energy consumption of the developed photobioreactor system is highly dependant on the chosen cultivation parameter configuration, respectively, the physiological requirements of the macroalgae to be grown. When cultivating the green macroalgae *U. intestinalis*, the electricity consumption per breeding module amounts to 2.71 kWh day⁻¹ when using fluorescent tubes as light source, compared to 5.26 kWh day⁻¹ when using the newly developed LED cylinders. The conditioning module contributes with a daily electricity demand of 11.82 kWh day⁻¹ to the total energy demand when using the operating schedule described in Sect. 2.2.2. Adding up the energy demand for the sensory and control components, total consumption of 3.1 kWh day⁻¹ is to be allocated to the system control module. Thus, when cultivating *U. intestinalis*, the actual energy demand of the developed prototypical plant sums up to 30.86 kWh day⁻¹. However, the photobioreactor system would consume 25.76 kWh day⁻¹ in total if all four breeding tanks were equipped with fluorescent tubes as a light source. In contrast, providing an LED light cylinder for each breeding tank would result in increased total energy consumption of 35.96 kWh day⁻¹. Table 2 summarises the energy demand, operating hours, and electricity consumption of the system components.

Extrapolation of cultivation results

To merge the results of the cultivation experiments into key figures suited for extrapolation, the biomass yield achievable in a photobioreactor system either equipped with LED cylinders or fluorescent tubes is put into context with the energy, area, and volume required for its production. Table 3 summarizes the results and emphasizes the positive effects of the LED light source on biomass growth. Despite the higher energy demand compared to fluorescent light tubes, the ratio of biomass yield to energy consumption per growth
As a step toward this aim, a closed photobioreactor system was developed with the intention to provide optimal abiotic cultivation parameters for macroalgae growth while reducing the disadvantages of the cultivation systems known to date. The newly developed system allows for the production of macroalgae in central Germany – a site distant from natural seawater.

Compared to the photobioreactor systems for macroalgae cultivation constructed to date (Chemodanov et al. 2017; Sebök et al. 2017, 2018; Mhatre et al. 2018), the prototypical plant implemented in Gebesee allows for more comprehensive control of the light and temperature of the parameters, as well as an effective reconditioning and thus recirculation of the cultivation medium. In addition, the sensor-based monitoring and control of the process technology reliably prevents accidents and ensures a high level of user-friendliness. In cultivation experiments with the green algae *U. intestinalis*, the growth rates achieved with submersed LED cylinders just about lie within the range of 6.9–19.40% day$^{-1}$ documented in the literature for this species in 28-days, lab-scale experiments in natural seawater, and under natural sunlight (Fong et al. 2004; Ruangchuay et al. 2012). In contrast, the strikingly low growth rates achieved with fluorescent tubes lag behind literature values and thereby affirm the importance of adequate light parameters for macroalgae cultivation. Since the highest growth rates were achieved by Ruangchuay et al. (2012) using reduced stocking densities (0.05 g L$^{-1}$) and increased cultivation temperatures (25 °C), adjusting these abiotic parameters in further cultivation experiments seems promising. In addition, increasing the harvesting frequency in breeding tanks equipped with submersed LED light sources should lead to higher growth rates since the light supply was sensibly decreased from day 14 to 28 due to the light cylinders being grown over by the cultivated algae.

Nevertheless, the energy demand of the newly developed system is to be considered as too high: With 12 MWh per year for a macroalgae production output of 33 kg in total, the efficiency and profitability of the cultivation system need to be optimized. Looking more closely at the largest energy consumers, it has to be stated that almost 50% of the total energy consumption is caused by the newly developed LED cylinders. In contrast, the strikingly low growth rates achieved with fluorescent tubes lag behind literature values and thereby affirm the importance of adequate light parameters for macroalgae cultivation. Since the highest growth rates were achieved by Ruangchuay et al. (2012) using reduced stocking densities (0.05 g L$^{-1}$) and increased cultivation temperatures (25 °C), adjusting these abiotic parameters in further cultivation experiments seems promising. In addition, increasing the harvesting frequency in breeding tanks equipped with submersed LED light sources should lead to higher growth rates since the light supply was sensibly decreased from day 14 to 28 due to the light cylinders being grown over by the cultivated algae.

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Altogether, closed production systems such as the developed photobioreactor system can increase both the degree of freedom in the choice of location and the value of the produced macroalgae. However, to achieve the economic viability of these highly technical production systems, further
investigation should also include fundamental research on optimal cultivation parameters for the production of biochemically active substances such as ulvan as raw materials for pharmaceuticals and cosmetics.

## Conclusion

The developed photobioreactor system allows for comprehensive control of the cultivation conditions and effective treatment of the cultivation medium. Salinity, temperature, light, and nutrient supply can be adapted and maintained according to the physiological needs of different algae species, and the plant is independent of natural seawater. Through the sensor-based control and monitoring of the plant processes, damaging events can be prevented and a high degree of user-friendliness is being achieved. However, the energy demand of the plant is to be reduced and further cultivation experiments should be undertaken to optimize the cultivation parameters in order to maximize the growth rates and the production of biochemically active substances.

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## Author contribution

All authors contributed to the study’s conception and design. Material preparation, data collection, and analysis were performed by Tonia A. Schmitz. The first draft of the manuscript was written by Tonia A. Schmitz and revised by Eckhard Kraft. Funding for the presented work was acquired by Eckhard Kraft and Tonia A. Schmitz. All authors read and approved the final manuscript.

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## Data availability

The datasets generated and analyzed during the current study are available from the corresponding author on reasonable request.

## Declarations

### Conflict of interest

The authors declare no competing interest.

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