Chlorella Vulgaris Surface-Mount Photobioreactor with Vision-Based Growth Signature Prediction Optimized by Electromagnetism-Like Mechanism

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Abstract: Industrial waste disrupts the natural production of microalgae cultures. Cultivation of microalgae in a controlled environment highly results to biomass with lower contamination necessary as high-valued economic product. In response to the emerging challenges of sustainable energy production, the integration of computational intelligence and biosystems engineering is considered as an open research area. In this study, Chlorella vulgaris microalgae were cultivated in BG-11 growth medium on three customized surface-mount light bioreactors that are equipped with digital camera for growth monitoring in terms of accumulated biomass surface area and color reflectance intensity via IoT. Feature-based machine learning models predicted microalgae growth area in terms of water temperature, pH level and turbidity, and light intensity. Microalgae cultures were exposed to combinations of white artificial light source of 2000 ± 1000 lux and water temperature of 27 ± 5°C using Peltier plate to discriminate biomass growth within a 30-day cultivation period. A total of nine environmental conditions were employed to clearly discriminate the impacts of environmental stressors to microalgae growth. Combined neighborhood component analysis and ReliefF was used to select high impact color features of C, Ye, M, H, and S with biomass area. Electromagnetism-like mechanism optimized-RBNN bested RNN and generalized processing regression with R² of 0.985 and RMSE of 6.262. There is also considerable growth in biomass surface area for certain combinations of light intensity and water temperature (2125 ± 625 lux and 28.75 ± 3.25°C), and turbidity and water pH concentrations (3.85 ± 0.15 NTU and 8.025 ± 0.775). However, the photobioreactor with 27°C and 2000 lux exposure is considered having the exact optimum controlled environment condition in cultivating Chlorella vulgaris based on the generated growth in biomass surface area of 38.314%. This developed intelligent system is scalable for seamless microalgae production of any strains for renewable energy resource.

Keywords: bioreactor, computational intelligence, computer vision, microalgae

I. INTRODUCTION

The world is dependent on the use fuel to manufacture commodities, power large systems, and activate vehicle motion on the road [1]. Fossil fuel is the most consumed form of fuel resulting to a great contribution in the vulnerability of the atmosphere to the threat of global climate change due to perilous greenhouse gas (GHG). As the concentration of carbon dioxide (CO2) emission intensifies due to the production of first generation of biofuel which is drawn from food crops [2], the use of algal biomass has numerous environmentally safe production processes to offer given a land area constraint. Both seawater and freshwater resources are now easily damaged by industrial sector due to improper disposing of chemicals [3-4], with the added malpractices of household communities that are not connected to a local sewage system [5]. Algal biomass is the third generation of biofuel that includes both microalgae and macroalgae resources. This promising approach of utilizing microorganisms to yield high-value products with economic impact is one of the attractive sustaining solutions in maintaining ecological balance and speeding up industrial revolution [6-7]. Biofactories, on the average, produces enough microalgae that is a key indicator for domestication [8]. The challenge of cultivating microalgae is provided with developing an optimal controlled biosystem to have faster production and cleaner biomass. Using artificial intelligence and life cycle analysis, the environmental impact of cultivating microalgae and transforming it to biofuels have been predicted [1,9]. A planning tool for microalgae agriculturist and bio-economist had been developed that outlines the environmental impact and its corresponding profit based on the strain of microalgae to be cultivated and harvesting approach [10]. Microalgae are unicellular microscopic photosynthetic microorganisms or phototrophs that are classified to cyanobacteria or chloroxyxybacteria and eukaryotic which is similar to green algae or chlorophyta [11]. It can be transformed in energy forms of biodiesel [12], bioethanol [13], biohydrogen [14] and biogas [15]. Biodiesel extracted from microalgae requires the purity of lipids and fatty acids in order to refine good oil.
The Philippines comprises the geographical epicenter of tropical algae diversity in the world as there are abundant microalgae both in marine and freshwater natural habitats such as *Chlorella vulgaris* (*C. vulgaris*), *Spirulina platensis*, *Dunalieilllasalina*, *Nannochloropsisocceania*, and *Tetraselmis* sp. These strains of microalgae have been characterized to normally depend on environmental parameters such as dissolved oxygen (dO₂) and pH concentration [16-17]. Temperature effect was also analyzed based on the photosynthetic growth and biochemical reactions of *Nannochloropsisocceania* and it was proven that decrease of temperature to 18°C results to suboptimal production of lipids necessary for extracting biofuel [18].

Aside for being labor intensive, mass culturing of microalgae in outdoor environment is often results to algal crashes. The introduction of indoor mass culture of microalgae using controlled photobioreactors is the currently ideal tactic to generate clean biofuels. In this manner, the quality of biomass is maintained, monitored, and altered by adjusting the pre-harvest factors such as light intensity, temperature, carbon dioxide, pH and water nutrients [19]. A photobioreactor (PBR) is a closed-system bioreactor that houses photoautotrophs and exposes it to light source to generate its own biomass. Outdoor photobioreactors are comparably larger than the one being used in indoor type that is mostly used in mass production of microalgae in the industry scale. Indoor photobioreactors, on the other hand, are mostly used to produce laboratory-grade biomass. There is quite a number of fabricated photobioreactors that are made using transparent horizontal and vertical tubes and cylindrical vessels, raceway pond, flat plate [20], biocoil, and bubble column [21]. These geometrical shapes of photobioreactors have a fallback in analyzing the surface area of microalgae as growth indicator and high developmental cost due to customization of curves to allow proper pressure inside the chamber. The challenge of high cost in lighting system and directional illumination is still present. A centered-light photobioreactor was developed [22] imposing that pattern of illumination and its intensity are highly considerable factors to increase the biomass yield of microalgae. Light management involving triggering on and off of light source [23] and variation of illumination spectrum [24] had been materialized to verify its impact to biomass growth.

Biomass color is directly correlated with the light source spectrum. There is immediate increase in green chromaticity by cultivating *Dunalieillla tertiolecta* (*D. tertiolecta*) using red and blue lights after 10 days compared to using white light [25]. Conventionally, microscopic segmentation is the approach for microalgae growth detection [26]. However, the integration of computer vision with computational intelligence offers vast applications of speeding up monitoring, detection, classification, and prediction of microalgae growth signatures [27-28]. Microalgae growth has been monitored by using spectral analysis that defines the different light absorbance of microalgae biomass surface [29], and image processing which uses hue saturation value (HSV) and cyan-magenta-yellow (CMYe) color spaces [30].

Despite of the abovementioned scientific studies in cultivating microalgae for biofuel production, automation using artificial intelligence has not been comprehensively explored yet. The challenge of distinguishing the harvest maturity and growth signatures of microalgae using bare eyes of the agriculturist is a subjective approach and prone to misclassification. Hence, the integration of computer vision and computational intelligence is an open research in the field of algal technology. In this study, Chlorella vulgaris is in-vitro cultivated with combinations of varying artificial photosynthetic light intensity and water temperature to discriminate the best environmental configuration for higher yield. The growth signatures considered are based on the acquired images which are color and phytomorphological features. Surface-mount light photobioreactors equipped with RGB digital camera were developed with IoT-based wireless sensor network for seamless data collection. pH concentration and turbidity level are also monitored in relation to microalgae growth. Optimized feature-based machine learning models were developed to predict microalgae growth.

### II. MATERIALS AND METHODS

The developmental architecture for Chlorella vulgaris growth signature prediction based on environmental harvest factors using feature-based machine learning models is shown in Fig. 1. In order to mimic the natural environmental stressors, photobioreactors were developed in consideration with light and temperature variations only. Sensors were deployed to quantify environmental stressors. MATLAB R2020a is the only computational intelligence software used in this study. Minitab 19 was used in statistics and Python programming language was used for intelligent electronic system development.

![Development of microalgae photobioreactor](image1)

- Development of wireless sensor network
- Preparation of BG-11 growth medium and cultivation of microalgae
- Image acquisition
- Feature extraction and selection
- Development of feature-based machine learning models for prediction of microalgae growth
- Microalgae biomass surface area and color intensity in terms of environmental harvest factors

**Fig 1. Developmental architecture for chlorella vulgaris growth signature prediction based on environmental harvest factors using feature-based machine learning models**
A. Phenotype Description and Artificial Environment Conditions

*Chlorella vulgaris* is the strain of green microalgae that was cultivated in 30-day life cycle in a phytotron located in Las Piñas City, Philippines with coordinates of 14.4484° N and 120.9867° E. The microalgae culture was obtained from the Aquaculture Department (AQD) of the Southeast Asian Fisheries Development Center (SEAFDEC) in Binangonan, Rizal, Philippines. Photobioreactors were developed to contain Chlorella vulgaris culture with varying environmental conditions in terms of light intensity ranging from 1000 to 3000 lux and enclosed chamber water temperature of 22 to 32°C. Shown in Table 1 is the list of photobioreactor environmental test conditions with a total of 9 batches of microalgae cultivations. Each photobioreactor is configured based on the listed environmental conditions every 30 days. Only water temperature and light intensity were considered as controlled environmental stressors that will induce changes on growth rate of the microalgae culture contained on the vessel. The captured microalgae surface image has aspect ratio of 640 x 480 with horizontal and vertical resolutions of 96 dpi and bit depth of 24. There is a total of 864 images collected over a month of cultivation for all the photobioreactors.

| Test Chamber | Water Temperature (°C) | Light Intensity (lux) |
|--------------|------------------------|----------------------|
|              | Cond. 1 | Cond. 2 | Cond. 3 |
| PBR 1        | 22      | 27      | 32      |
| PBR 2        | 27      | 32      | 22      |
| PBR 3        | 32      | 22      | 27      |

B. Photobioreactor Development

Three photobioreactors were constructed to house Chlorella vulgaris. By definition, a photobioreactor utilizes light source to allow photosynthesis in the generation of biomass of phototropic microorganisms such as microalgae.

In this study, photobioreactors are constructed using white plastic sheets, with dimensions of 57 cm x 42 cm x 37 cm (Fig. 2). This vessel can handle 80 liters of liquid. Artificial photosynthetic lighting was employed using T8 white light spectrum LED that is placed at the chamber cover. It was carefully considered that the whole water surface area is benefited with this light source. As light intensity is directly proportional to its power, a metal oxide semiconductor field effect transistor (MOSFET) driver circuit enhanced by pulse width modulation (PWM) was constructed to calibrate the light source from 3 to 7 watts. To control the water temperature of the bioreactor, a tandem configuration of two Peltier device which is composed of bismuth telluride (Bi$_2$Te$_3$) and tellurium (Te), was placed on the sidewall of the photobioreactor. It exhibits thermoelectric principle by generating heat with the aid of external power source. This configuration increases the efficiency of Peltier plate to provide lower temperature despite of high ambient temperature. Each Peltier plate is rated 150 watts. Relay circuit is constructed for the controls of Peltier activation. Aerator was also placed to promote carbon dioxide dissolution on to the aqueous system.

One mote per photobioreactor is configured to monitor the environmental conditions. It is constructed by integrating TSL2561 light sensor, DS18B20 weather-proof temperature sensor, SE0198 turbidity sensor and Gravity pH sensor with Arduino Mega as the processing core in enabling the array of sensors (Fig. 3). The Arduino Mega microcontroller is then connected with Raspberry Pi 3 Model B+ for wireless transmission of acquired sensor data.

Fig 2. Design of (a) photobioreactor test platform and (b) the component placement of a single mote

Fig 3. System block diagram of a single mote placed on a photobioreactor to control light intensity and temperature configuration and monitor environmental stressors

To render data integrity on the acquisition module, each sensor was subjected to calibration. pH sensor underwent three-point calibration where each sensor probe is properly submerged to containers with pH 7, 4, and 10 calibration solution, resembling neutral, acidic, and basic concentrations, separately for two minutes.
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The sensor probe is rinsed with distilled water (dH2O) and dried with paper towel before contacting with another test solution. DS18B20 temperature sensor is corrected using two-point calibration technique by subjecting the sensor metallic probes to boiling and freezing point conditions [31]. TSL-2561 luminosity sensor is also corrected using two-point calibration technique by subjecting to laboratory light source with known luminosity and black box. Turbidity sensor is subjected to zero nephelometric test with agitations and signal-averaging. All sensor calibrations were done with room temperature ranging from 24 to 26 °C. The collected calibration data was used to generate the regression equations for each sensor and were embedded in Arduino Mega microcontroller. Overall, the developed photobioreactor is movable due to built-in wheels beneath the vessel itself and compact placements of electronic components. Single photobioreactor is considered an independent system, thus, can be placed to other place with no dependency with other photobioreactors because it has its own intelligent control in range the water temperature and light intensity needed for photosynthetic production.

C. Wireless Sensor Network Development

A self-configured data-centric wireless sensor network (WSN) composed of three client motes deployed in physical environment monitoring, one sink node, WiFi router and cloud connectivity and user machine is developed in this study (Fig. 4). It is used to monitor and record physical observations of environmental pre-harvest factors of Chlorella vulgaris microalgae. Motes were connected in star topology to the sink node. IP Logitech cameras were directly connected to the USB adapter of Raspberry Pi microcontroller and to the cloud via Internet capability. To ensure data storage, data duplication is manifested on the system by storing sensor data both in the Google cloud and individual Raspberry Pi local memory card. Time scheduling of hourly acquisition is employed. The collected images and data sensors were purged from the cloud storage using file transfer protocol (FTP) and secured control protocol (SCP) down to the local user machine in comma separated variable (.csv) file format. The data file is configured to record series of data array composed of 5 data elements with the following consecutive placement: [data acquisition timestamp, water temperature, light intensity, pH concentration, turbidity level]. No transmission delays were experienced due to close proximity of motes with sink node and WiFi router.

D. Growth Medium Preparation and Microalgae Cultivation

Water impurities due to industrial and household wastes adversely impacts the generation of biofuel from microalgae with good quality. With this very reason, there is a need to formulate artificial saltwater with no impurities. The preparation of BG-11 growth medium was done in a laboratory with analytical-grade chemicals. Shown in Table 2 is the composition of BG-11 growth medium and Table 3 presents the BG-11 stocks composition. BG-11 stock 3 is primarily composed of ferric ammonium citrate ([NH4]3[Fe(C6H5O7)3]) and partially of ethylenediaminetetraaetic acid disodium salt (EDTA 2Na). This medium results to no impurities yielding better quality of microalgae. In this study, 30 liters of distilled water and 6 liters of BG-11 growth medium were mixed and set as the primary solvent for each photobioreactor vessel.

BG-11 growth stocks 1, 2, 3 and 4 were first prepared before making the growth medium. These stocks were mixed with deionized water using laboratory glass bottles based on the measurement listed on Tables 2 and 3, and then mixed with other BG-11 growth medium compositions using glass rod that resulted to pure artificial saltwater.

E. Feature Extraction and Selection

Spectro-morphological microalgae growth signatures were extracted to differentiate daily growth with the impact of photosynthetic light intensity and water temperature level inside the photobioreactor. Color features were extracted from the following visible color spectrums: red, green, and blue (RGB), hue, saturation, and value (HSV), cyan, magenta and yellow (CMY), lumina, blue-difference, and red-difference (YCbCr), and lightness, green-to-red, blue-to-yellow (CIELab).

Fig. 5. Wireless sensor network architecture with sink node for IoT
Using 10,000 superpixels, which is based on simple linear iterative clustering (SLIC) algorithm and K-means clustering [4, 32], it was superimposed on to the raw microalgae surface images to segment non-vegetative pixels from vegetative which is microalgae and region of interest (ROI) in this application. The phytomorphological area of microalgae was computed using region properties of the annotated image.

There is a total of 16 image-based features considered as growth signatures for microalgae being cultivated where 15 of these are color elements. Using human bare eyes often results to subjective classification on the maturity of microalgae even with presence of artificial photosynthetic light, thus computer vision-aided color extraction is necessary. Hybrid neighborhood component analysis (NCA) and ReliefF algorithms were used for feature selection of abiotic environmental stressors of water temperature, pH, turbidity, and light intensity, and spectral signatures based on its variation impact to the extracted microalgae surface area (Fig. 5). NCA is configured with limited-memory Broyden-Fletcher-Goldfarb-Shanno algorithm as its solver, Hessian history size of 15 and line search method of weak Wolfe. By following the results of hybrid NCA-ReliefF, water temperature and pH concentration inside the photobioreactor have the greatest relevance to microalgae surface area among other abiotic environmental stressors. Conversely, color components of yellow, magenta, hue, cyan, and saturation resolves on providing highest correlation with microalgae surface area.

![Fig 5. Feature selection of (a) abiotic environmental stressors and (b) biomass spectral signatures based on its relevance to microalgae surface area using neighborhood component analysis and ReliefF](image)

**F. Development of Feature-Based Machine Learning Models for Prediction of Microalgae Growth**

The extracted features from the previous step were utilized in developing feature-based machine learning models for prediction of microalgae growth in terms of its surface area as captured by the camera. In this study, radial basis neural network (RBNN), recurrent neural network (RNN) and generalized processing regression (GPR) were configured to generate prediction of surface area based on environmental stressors and color signatures, separately. RBNN was optimized by using electromagnetism-like mechanism (EM) which is a physics-inspired metaheuristic algorithm materializing the principle of attraction and repulsion of charged particles in electromagnetic field [28]. In this study, it starts with initializing the 100 charged particles as uniformly distributed along the two-dimensional hypercube of electromagnetic field. Then, the electromagnetic activity of a sampled charged particle is tested, and its corresponding objective function value (y) is calculated with x as the spread factor value in constructing the RBNN architecture (1). The explored electromagnetic hypercube dimension ranges from 1 to 10 (Fig. 6a) and the maximum feasible step length ranges from 0 to 1 (Fig. 6b). New charged particle is labeled as best particle when the current charged particle exhibits lower objective function value. The EM optimization of RBNN terminates after 500 iterations with optimum convergence. It is exhibited with unchanging current best charged particle resembling the spread factor value of 14. It is noticeable that spread factor value dominantly follows the nature of the objective function value on a certain scale increase (Fig. 6c). RNN was optimized by constructing its architecture with 250-150-50 artificial neurons configuration on the hidden layer. On the other hand, GPR was optimized using Bayesian algorithm as solver, squared exponential for kernel function, constant basis function with beta of 24.8520 and sigma of 26.0293, exact predict method and fit m

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y = 0.0054 x^4 + 0.0254 x^2 - 0.047 x + 0.4327anumber{1}

**III. RESULTS AND DISCUSSIONS**

**A. Vision-Based Microalgae Photobioreactor**

Implementation of advanced technologies in biosystem setup for cultivating microalgae has fascinated both agriculturists, engineers, and scientists [1,3,6,9,10,18,21,22,24,25,29]. It involves the adjustment of light source, variations on nutrient injection rates, and innovations on the grow bed or bioreactor itself. The developed *Chlorella vulgaris* microalgae photobioreactor is equipped with artificial white light source and Peltier plates that adjust the light intensity and water temperature level in exploring what is the optimum combination of these environmental stressors will generate the significant biomass with consideration of cultivar life cycle (Fig. 7a).
It is noticeable that the developed automated photobioreactor is effective for moving its physical placement in a large phytotron as it has wheels beneath it. Moreover, the structure is compact and scalable for larger production of biomass. It employs adaptive bioreactor environment condition management by just choosing what type of microalgae will be cultivated. It means that the system automatically configures the suitable range for biomass production. With enough pre-harvest factors based on the requirements of photosynthesis, which are light energy, carbon dioxide, and water, the microalgae bioreactor generates comparable biomass. The greener colonies of eukaryotic species are the growing Chlorella vulgaris as expected because the bioreactor is subjected to minimum amount of light energy and temperature as part of the photosynthetic requirement (Fig. 7b). The overlaying of 10,000 superpixels over the raw captured image enables segmentation of biomass from non-algae pixels (Fig. 7c). Five color clusters were used to provide efficient segmentation which are white, red, green, blue, and black. Hence, the developed three photobioreactor acts independently one each other because of the embedded automations on separate systems which is comparable to locally dependent growth chambers using internally illuminated tubes [3] and eight series vertical columns [25]. The proposed surface-mount light photobioreactor was able to cultivate microalgae as the light is evenly spread over the water surface resulting to equal photosynthetic reaction among microalgae cultures throughout the cultivation period.

Fig 6. Particle charge response with varying electromagnetic hypercube dimension and maximum feasible step length, and its interaction with the spread factor of radial basis neural network

B. Feature-based Growth Signature Prediction of Chlorella Vulgaris

Based on hybrid neighborhood component analysis and ReliefF-selected spectral and abiotic environmental stressors, (2) and (3) are the corresponding regression models. \( A_{\text{biomass(c)}} \) denotes the biomass area that can be generated using the spectral signatures of microalgae with \( R^2 \) of 0.9788 where \( \theta, \varepsilon, \gamma, \psi \) and \( \mu \) are the spectral component values of hue, saturation, cyan, yellow, and magenta, respectively. On the other hand, \( A_{\text{biomass(e)}} \) denotes the biomass area that can be generated using the selected abiotic environmental stressors of temperature (\( \tau \)) and pH concentration (\( \rho \)) with \( R^2 \) of 0.93.

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A_{\text{biomass(c)}} = 53.1 - 22.9 \theta + 9.17 \varepsilon + 0.39 \gamma - 0.696 \psi + 0.229 \mu
\]  
(2)
Nine different combinations of controlled environmental stressors were separately configured on the customized photobioreactors. Matured microalgae biomass is visually recognizable with darker green color resembling the accumulation of photosynthetic pigment and thickening of algal strands (Fig. 7b). Captured microalgae surface images were quantitatively assessed by predicting its surface area based on environmental pre-harvest factors and spectral signatures using optimized general processing regression, recurrent neural network, and radial basis neural network. Machine learning modes were suffixed with numerical subscripts of 4 and 2 denoting the number of environmental stressors used as predictors in estimating biomass surface area such as GPR for general processing regression using the hybrid NCA and ReliefF-selected high impact features of water temperature and pH concentration. Equally, GPR\textsubscript{15} resembles GPR using the 15 original pre-selected spectral features of five color spaces, and RBNN\textsubscript{5} is for RBNN using yellow, magenta, hue, cyan, and saturation components as predictors. For predicting microalgae biomass surface area with environmental stressors as predictors, EM-RBNN\textsubscript{2} bested out GPR and RNN variants with RMSE of 6.262, R\textsuperscript{2} of 0.985 and MAE of 2.847 (Table 4). However, it is noticeable that RNN\textsubscript{2} and RNN\textsubscript{4} performed the shortest inference time of 4 seconds despite of its network looping in triple staged hidden layer. It is 87.8\% faster than the most sensitive RBNN\textsubscript{2}. For predicting microalgae biomass surface area with spectral signatures as predictors, EM-RBNN\textsubscript{5} performed the most accurate, sensitive, and responsive model with RMSE of 10.489, R\textsuperscript{2} of 0.944 and MAE of 5.160 (Table 4). RNN variants still turned out to have the fastest inference time of 81.42\% than RBNN\textsubscript{5}. It is evident that predictions using models requiring higher number of input features resulted to longer inference time. Among the included machine learning models, RBNN variants exhibited the most consistent performance from training, validation and testing phases depicting perfect sampling and no overfitting or underfitting is involved. Overall, RBNN\textsubscript{5} using water temperature and pH that is improved by EM resolved to have optimum performance in evaluating microalgae biomass surface area.

![Regression lines in predicting chlorella vulgaris surface area using (a) abiotic environmental stressors and (b) spectral signature of biomass](image)

**Fig. 8.** (a) Regression lines in predicting chlorella vulgaris surface area using (a) abiotic environmental stressors and (b) spectral signature of biomass

| Model | No. of Features | Training | Validation | Testing |
|-------|----------------|----------|------------|---------|
|       |                | RMS\textsubscript{E} | R\textsuperscript{2} | MAE | RMS\textsubscript{E} | R\textsuperscript{2} | MAE | RMS\textsubscript{E} | R\textsuperscript{2} | MAE | Inference Time (s) |
|       |                |          |            |        |          |            |      |          |            |      |                        |
| GPR   | 4              | 12.916   | 0.93 \textsubscript{1} | 11.36 \textsubscript{9} | 39.021 | 0.04 \textsubscript{4} | 30.04 \textsubscript{6} | 35.482 | 0.12 \textsubscript{6} | 25.40 \textsubscript{4} | 24.251 |
| GPR   | 2              | 6.284    | 0.20 \textsubscript{7} | 4.494 \textsubscript{2} | 6.549 | 0.97 \textsubscript{1} | 4.690 \textsubscript{6} | 10.695 | 0.93 \textsubscript{6} | 6.690 \textsubscript{4} | 12.744 |
| RNN   | 4              | 11.616   | 0.07 \textsubscript{2} | 9.131 \textsubscript{2} | 33.732 | 0.02 \textsubscript{4} | 26.41 \textsubscript{6} | 29.011 | 0.35 \textsubscript{9} | 22.28 \textsubscript{4} | 4.000 |
| RNN   | 2              | 12.389   | 0.10 \textsubscript{4} | 10.67 \textsubscript{5} | 29.309 | 0.23 \textsubscript{4} | 21.54 \textsubscript{8} | 39.070 | 0.16 \textsubscript{4} | 29.54 \textsubscript{6} | 4.000 |
| RBNN  | 4              | 0.893    | 0.97 \textsubscript{5} | 0.707 \textsubscript{4} | 10.086 | 0.90 \textsubscript{2} | 3.541 \textsubscript{5} | 13.535 | 0.90 \textsubscript{4} | 6.933 \textsubscript{6} | 36.116 |
| RBNN  | 2              | 0.893    | 0.98 \textsubscript{2} | 0.707 \textsubscript{2} | 10.086 | 0.93 \textsubscript{3} | 3.541 \textsubscript{5} | 6.262 | 0.98 \textsubscript{5} | 2.847 \textsubscript{5} | 32.784 |

|       |                |          |            |        |          |            |      |          |            |      |                        |
| GPR   | 15             | 10.835   | 0.00 \textsubscript{8} | 9.112 \textsubscript{5} | 42.930 | 0.19 \textsubscript{5} | 34.01 \textsubscript{3} | 28.578 | 0.29 \textsubscript{9} | 23.39 \textsubscript{7} | 48.148 |
Unlike a destructive testing of extracting biomass tissues from time to time in order to monitor its growth by weighting it [3, 25], this study is definite in predicting microalgal growth based on the intensity of color reflectance in various color spaces and abiotic environmental stresses as a nondestructive approach (Table 4).

C. Chlorella vulgaris cultivation and limnological interaction

The photobioreactor 1 (PBR1) with light intensity of 1,000 lux and water temperature of 22°C successfully cultivated a 26.422 cm² Chlorella vulgaris biomass surface area after a 30 days of cultivation period with 6.597 cm² growth from the first week baseline growth of 19.825 cm² (Fig. 9). Exposing Chlorella vulgaris and BG-11 growth medium to light intensity of 1,000 lux and increasing temperature up to 32°C substantially decreased the generated biomass surface by 51.707%. Photobioreactor 2 configured with condition 1 (PBR2-COND1) has total increase of 7.193 cm² of biomass surface after the cultivation period while PBR2-COND2 is characterized with an increase of 6.883 cm². PBR2-COND1 and PBR2-COND2 conditions primarily differ with the generated biomass surface area during first week with 20.166 cm² for PBR2-COND1 and 17.936 cm² for PBR2-COND2. Photobioreactor 3 configured with condition 1 (PBR3-COND1) having water temperature of 22°C and 3,000 lux generated an increase of 4.281 cm² over a period of one month where PBR3-COND3 having water temperature of 32°C and 3,000 lux is short of 0.04 cm².

Fig 9. Growth curve of chlorella vulgaris biomass surface area in three environmentally controlled photobioreactors

In terms of limnological parameters interaction, there is an increase of 14.075% in pH and 61.58% for the turbidity as water temperature is increased from 22°C to 32°C with constant 1,000 lux (Fig. 10). However, pH concentration is lightly weakened by 5.39% and turbidity is increased by 14.777% for the controlled environment emitting 2,000 lux. Like the PBR1, photobioreactor 3 that is equipped with 3,000 lux artificial lighting contained a noticeable increase of 31.579% for pH concentration and 2.473% in turbidity. With this, turbidity is highly reliable with light intensity and the growth of biomass on water surface as there is 59.107% deviation per 2,000 lux increase. Conversely, pH is highly affected by light intensity with 17.504% rise per 2,000 lux increase. There is also considerable growth in biomass surface area for certain combinations of light intensity and water temperature such as 1,500 to 2,750 lux and 25.5 to 32°C, and turbidity and water pH concentrations of 3.7 to 4 NTU and 7.25 to 8.8, respectively (Fig. 10). However, PBR2-COND2 with 27°C and 2,000 lux is considered as the exact optimum controlled environment condition in cultivating Chlorella vulgaris based on the generated growth in biomass surface area of 38.314%.

Fig 10. Impacts of limnological parameters inside the controlled photobioreactors to chlorella vulgaris biomass surface area
IV. CONCLUSIONS

This study demonstrated the development of a surface-mount light photobioreactor equipped with Pelletier plates, computer vision and wireless sensor network for the cultivation of Chlorella vulgaris, and the implementation of computational intelligence for its growth prediction. Three photobioreactors were constructed that worked independently with each other based on the configured automation in the adjustment of abiotic environmental stressors in terms of water temperature and light intensity that helps the photosynthetic process of microalgae tissue to grow inside this controlled environment chamber. Consumer-grade digital RGB camera visually monitors and captures images of the microalgae being cultivated and the images were used to extract biophysical signatures of microalgae. Array of pH, temperature, turbidity, and light sensors was placed inside the photobioreactors and connected to the IoT for seamless transmission of important environmental data. Artificial seawater was mixed using laboratory-grade chemicals to create pure nutrient medium for microalgae cultures. Cyan, yellow, magenta, hue, saturation, and biomass area are the highly selected spectro-morphological signatures for growth rate prediction of microalgae using hybrid neighborhood component analysis and ReliefF algorithms. Generalized processing regression, recurrent neural network and radial basis neural network optimized using electromagnetism-like mechanism were used in predicting the actual microalgae surface based on the pre-selected signatures. Accordingly, these are the following outcomes were obtained: EM-RBNN performed the best prediction accuracy and sensitivity with lower RMSE score, the best photobioreactor environment condition in cultivating Chlorella vulgaris exhibits 27°C water temperature and 2000 lux exposure, turbidity has direct relation with light intensity and accumulation of biomass tissue on water, and pH is highly affected by light concentration. Overall, there is significant benefit in cultivating microalgae in a closed environment biosystem as proper photosynthetic requirements will be met in all season. For future studies, it is recommended to employ techno-economic analysis to comprehensively assess the impact of materializing this biosystem in larger scale.

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