Data Article

Data on growth, productivity, pigments and proximate composition of indigenous marine microalgae isolated from Cox’s Bazar Coast

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A B S T R A C T

Data on growth, productivity, pigments and proximate composition of the four different indigenous marine microalgae (isolated from Cox’s Bazar Coast) were collected to compare the growth performance, pigments and nutritional composition. Chlorella sp., Nannochloropsis sp., Tetraselmis sp. and Chaetoceros sp. are the four different marine microalgae. Growth curve was determined as the prerequisite to identify the stationary phase for each of the isolated microalgae. Data on growth curves were collected in terms of cell density and optical density to observe the growth rates and division per day. Isolated species were mass cultured in commercial culture medium. When the culture reached at stationary phase, microalgae were extracted to determine productivity, pigments, and proximate composition. The data of productivity (volumetric, areal and lipid productivity), pigments (Chlorophyll a, b, c, carotenoids, and phycobiliproteins), and proximate composition (protein, lipid, and carbohydrate) were significantly (p < 0.05) different among the four different microalgae. Therefore, this data will contribute to the selection

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of potential microalgae species through proper characterization for vast industrializations.

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Specifications Table

| Subject | Food Science, Aquatic Science |
|---------|------------------------------|
| More specific subject area | Microalgae growth, productivity, pigments and proximate composition |
| Type of data | Table and Chart |
| How data were acquired | Microscopic and spectrophotometric analysis for growth; biochemical and spectrophotometric analyses for productivity and pigments; biochemical analysis for proximate compositions. |
| Data format | Raw and analyzed |
| Parameters for data collection | Conway culture medium was used for algal growth. Four different microalgae were cultured with three replicates. Mass culture was sustained until stationary phase to harvest dried biomass. |
| Description of data collection | For growth curve: cell density and optical density. For productivity: volumetric productivity, areal productivity and lipid productivity. For pigments: chlorophyll a, b, c, carotenoids, total phycobiliproteins, allophycocyanin, phycocyanin and phycoerythrin. For proximate composition: protein, lipid and carbohydrate. |
| Data source location | Microalgae lab, Department of Aquaculture, Faculty of Fisheries, Chattogram Veterinary and Animal Sciences University, Khulshi-4225, Chattogram, Bangladesh |
| Data accessibility | Data are available with this article and also at http://dx.doi.org/10.17632/znns9msdx6.1 |

Value of the Data

- This data will contribute to the selection of potential microalgae species through proper characterization on the basis of biomass production, lipid production and pigment accumulation for vast industrializations. This data provides a basis in understanding the profile of four different indigenous marine microalgae species of Bangladesh.
- This data will be useful for the mass production of selected microalgae for specific use. Aquaculture industry can use the information to formulate the feed as well as to enhance the coloration of fishes.
- Growth and productivity data will be useful to plan proper culture method; pigments data will be useful to select species for pigment production. In addition, proximate data will be useful to select potentially commercial species, especially for animal feed and biofuels production.

1. Data Description

Growth curves of the four microalgae species, cultured in Conway medium, are shown in this data (Fig. 1). Onset of death phase (7-10 days) varied among the four species.

The volumetric, areal, and lipid productivity of the four microalgae species, cultured in Conway medium, are shown in this data (Table 1). All the data of productivity significantly (p < 0.05) varied among the four different species.

This data shows chlorophyll a, b, c, carotenoids, and phycobiliproteins (phycocyanin, allophycocyanin and phycoerythrin) production of the four microalgae species, cultured in Conway medium (Table 2, Fig. 2, 3). Table 2 shows that chlorophyll a, chlorophyll b, chlorophyll c, and
Fig. 1. Growth curve in terms of cell density (cells/ml × 10^6) and optical density (Absorbance) of marine microalgae Chlorella sp. (A), Chaetoceros sp. (B), Nannochloropsis sp. (C), Tetraselmis sp. (D). Wavelengths for measuring optical density were 780 nm, 750 nm, 780 nm and 480 nm for A, B, C and D, respectively. Values are means with standard deviation. CD and OD represents cell density, and optical density, respectively.

Table 1
Volumetric, areal, and lipid productivity of marine microalgae, cultured in Conway medium, and isolated from Cox’s Bazar Coast. Values with different letters within each column are significantly different (p < 0.05).

| Species           | Volumetric productivity (mg/L/day) | Areal productivity (mg/cm²/day) | Lipid productivity (mg/L/day) |
|-------------------|------------------------------------|---------------------------------|--------------------------------|
| Chlorella sp.     | 0.39 ± 0.03^c                      | 0.76 ± 0.06^c                   | 0.047 ± 0.003^b                |
| Chaetoceros sp.   | 0.61 ± 0.08^a                      | 1.21 ± 0.17^a                   | 0.109 ± 0.003^a                |
| Nannochloropsis sp.| 0.45 ± 0.04^bc                     | 0.89 ± 0.07^bc                  | 0.108 ± 0.004^a                |
| Tetraselmis sp.   | 0.57 ± 0.06^ab                     | 1.12 ± 0.07^ab                  | 0.051 ± 0.014^b                |

Values are means ± SD of triplicate measurements.

Table 2
Chlorophyll a, b, c, and carotenoids content of marine microalgae, cultured in Conway medium, and isolated from Cox’s Bazar Coast. Values with different letters within each column are significantly different (p < 0.05).

| Species           | Chlorophyll a (µg/L) | Chlorophyll b (µg/L) | Chlorophyll c (µg/L) | Carotenoids (µg/mL) |
|-------------------|----------------------|----------------------|----------------------|---------------------|
| Chlorella sp.     | 0.48 ± 0.05^c        | 0.19 ± 0.05^b        | 0.06 ± 0.05^c        | 0.56 ± 0.03^b       |
| Chaetoceros sp.   | 1.30 ± 0.09^b        | 0.04 ± 0.02^c        | 0.29 ± 0.01^a        | 1.36 ± 0.22^a       |
| Nannochloropsis sp.| 0.48 ± 0.04^c        | 0.05 ± 0.003^c       | 0.01 ± 0.01^c        | 1.68 ± 0.05^a       |
| Tetraselmis sp.   | 2.68 ± 0.04^a        | 1.23 ± 0.02^a        | 0.10 ± 0.01^b        | 1.51 ± 0.14^a       |

Values are means ± SD of triplicate measurements.
carotenoids varied significantly \((p < 0.05)\) among the different species. Fig. 2 shows that total phycobiliproteins \((0.0137 \pm 0.0013 - 0.0253 \pm 0.0015 \text{ mg/g})\) varied significantly \((p < 0.05)\) among the four microalgae; similarly, Fig. 3 shows that the three different phycobiliproteins, phycocyanin \((0.0017 \pm 0.0005 - 0.0027 \pm 0.0007 \text{ mg/g})\), allophycocyanin \((0.0100 \pm 0.0006 - 0.0197 \pm 0.0011 \text{ mg/g})\), and phycoerythrin \((0.0018 \pm 0.0002 - 0.0029 \pm 0.0003 \text{ mg/g})\) varied significantly \((p < 0.05)\) among the four different species.

Finally, this data shows protein, lipid and carbohydrate content \(% \text{ dry weight}\) of the four different indigenous marine microalgae, culture in Conway medium (Fig. 4). Protein \((42.7 \pm 4.0 - 56.7 \pm 0.9 \% \text{ dry weight})\), lipid \((12.1 \pm 0.4 - 25.2 \pm 2.6 \% \text{ dry weight})\) and carbohydrate \((17.1 \pm 1.4 - 23.2 \pm 2.3 \% \text{ dry weight})\) data varied significantly \((p < 0.05)\) among the four different species.
2. Materials and Method

2.1. Media and Method

2.1.1. Filtration, sterilization and preservation of seawater

Seawater was collected from Saint Martin’s coast of Bay of Bengal, Bangladesh. After collection, the water was preserved for the settlement of solid particles. Then water was filtered, using 45 μm glass microfiber filters (GF/C). After that, the filtered water was autoclaved at 121°C temperature and 15 lbs./inch² pressure for 15 minutes. The sterilized water was stored at 20–21°C temperature following Reda et al. [1].

2.1.2. Conway medium preparation

Conway medium includes micronutrients, trace metal solution, and vitamin [2]. Pure Conway medium was used for Chlorella sp., Nannochloropsis sp., and Tetraselmis sp. culture; however, Conway medium + silicate solution was used for Chaetoceros sp. culture. Table 3 shows the amount of different constituents. To prepare 1L Conway media, 1 mL of solution A, 0.5 mL of solution B and 0.1 mL of solution C, were added with 28–30 g/L autoclaved seawater (Table 3).

2.1.3. Collection of microalgae, culture and maintenance

The four different types of indigenous marine microalgae (Chlorella sp., Nannochloropsis sp., Tetraselmis sp., and Chaetoceros sp.) were collected from previously preserved samples of the Microalgae laboratory of Department of Aquaculture, Faculty of Fisheries, Chattogram Veterinary and Animal Sciences University, Bangladesh. The pure samples were cultured in Conway culture medium. Stock was scaled up, and then sub cultured for growth curve determination.

2.2. Determination of growth curve

Data of the growth curves were determined prior to the start of the microalgae culture to determine the data of productivity, pigments, and proximate composition. A total of 300 mL of culture volume was maintained in a sterile 500 mL borosilicate Erlenmeyer flask for each species with three replicates. Out of the 300 mL, 270 mL was culture medium and 30 mL was stock culture. The culture was continued up to the death phase. Depending on the species, culture duration was varies which were 9 (Chlorella sp.), 11 (Nannochloropsis sp.), 9 (Chaetoceros sp.) and 8 (Tetraselmis sp.) days respectively. Water salinity was maintained 30.1 ± 0.11 g/L, pH was maintained 7.72 ± 0.17, and gentle aeration was provided 24 hrs which offered 4.53 ± 0.53 mg/L
Table 3
Constituents of microalgae culture medium.

(A) Main Mineral Solution

| Names of Chemicals          | Quantity          |
|-----------------------------|-------------------|
| NaNO3/KNO3                  | 100.00g/116.00g   |
| Disodium EDTA (C_{10}H_{16}N_{2}O_{8}) | 45.00g           |
| H_{3}BO_{3}                 | 33.60g            |
| Na_{2}PO_{4}.4H_{2}O        | 20.00g            |
| FeCl_{3}.6H_{2}O            | 1.30g             |
| MnCl_{2}.4H_{2}O            | 0.36g             |
| Trace metal solution        | 1.00mL            |

Dissolving in deionized/distilled water and make the volume 1 L.

(B) Trace Metal Solution

| Names of Chemicals          | Quantity |
|-----------------------------|----------|
| ZnCl_{2}                    | 2.10g    |
| CoCl_{2}.6H_{2}O            | 2.00g    |
| (NH_{4})_{3}MO_{2}.4H_{2}O  | 0.90g    |
| CuSO_{4}.5H_{2}O            | 2.00g    |

Dissolving in deionized/distilled water and make the volume 1 L.

(C) Vitamin

| Names of Chemicals          | Quantity |
|-----------------------------|----------|
| Thiamine, B1                | 0.20g    |
| Cyanocobalamin, B12         | 0.01g    |

Dissolved in deionized/distilled water and make the volume 100 mL.

(D) Silicate Solution

| Names of Chemicals          | Quantity |
|-----------------------------|----------|
| Sodium silicate (Na_{2}SiO_{3}) | 20.00g   |

Dissolving in deionized/distilled water and make the volume 1 L.

dissolved oxygen. Growth curve was determined on basis of cell density \((\text{cell/mL})\), and optical density \((\text{absorbance at 780nm} \text{ for Chlorella sp.}, \text{ at 780nm for Nannochloropsis sp.}, \text{ at 480nm for Tetraselmis sp.}, \text{ and at 750nm for Chaetoceros sp.})\).

2.2.1. Cell density

Microalgal cells were counted using hemacytometer [3] every day during the data collection of growth curve. The hemacytometer and its cover slip (Bright-line improved Neubauer hemacytometer, 0.0025 mm², 0.1 mm deep chambers, Assistent, Germany) were cleaned using Milli-Q water (Millipore Corp.) prior to the fill up of the chambers with culture samples. Evenness of cell distribution was checked under low power magnification \((4 \times \text{ and } 10 \times)\) of the microscope (Nikon E600). Cells were counted for both chambers of the hemacytometer under magnification of \(20 \times\); Lugol’s iodine was added to culture aliquots for fixation and staining to facilitate counting. The formulae to calculate the cells are as the following:

\[
\text{Cell count calculation (cell/ml) for 5 squares} = \frac{\text{Total number of cells counted}}{10 \times 4} \times 10^6
\]

Where 10 represented the 10 squares of the 2 hemacytometer chambers and \(4 \times 10^{-6}\) represented the volume of samples over the small square areas, that were equivalent to 0.004 mm³ (0.2 mm × 0.2 mm × 0.1 mm), expressed in cm³ (ml).

\[
\text{Cell count calculation (cell/ml) for 25 squares} = \frac{\text{Total number of cells counted}}{50 \times 4} \times 10^6
\]
Where 50 represented the 50 squares of the 2 hemacytometer chambers and $4 \times 10^{-6}$ represented the volume of samples over the small square areas, that were equivalent to 0.004 mm$^3$ (0.2 mm $\times$ 0.2 mm $\times$ 0.1 mm), expressed in cm$^3$ (mL).

2.2.2. Optical density

Optical density of culture aliquots were measured using a spectrophotometer (UV-VIS Double beam, Model-T80, HANNA), every day during the data collection of growth curve. The culture medium for the species was used as the blanks. The absorbance were measured at the wavelength 780nm for Chlorella sp., 780nm for Nannochloropsis sp., 480nm for Tetraselmis sp., and 750 for Chaetoceros sp. [4].

2.3. Design of the cultures for productivity, pigment, and proximate data

Microalgae were cultured up to stationary phase, in large sterile borosilicate Erlenmeyer flasks with three replicates of each species, thus to collect the data of productivity, pigments and proximate composition. 1.5 L pure Conway medium was taken into each flasks for Chlorella sp., Nannochloropsis sp., and Tetraselmis sp.; while, 1.5 L Conway medium with silicate solution was taken into each flasks for Chaetoceros sp. Then 5% pure culture stock was added to each flask. Each species was cultured separately to inhibit contamination for maintaining pure culture. The culture were conducted at 25.2 $\pm$ 0.7°C temperature with 24 hrs light at 150 $\mu$E m$^{-2}$ s$^{-1}$ intensity. Water salinity was maintained 30.1 $\pm$ 0.11 g/L, pH was maintained 7.72 $\pm$ 0.17, and gentle aeration was provided 24 hrs which offered 4.53 $\pm$ 0.53 mg/L dissolved oxygen. Culture was kept in environmental chamber to control the temperature and light throughout the experiment. Culture was maintained until the stationary phase. Culture volume data was recorded every day, while, biomass was determined in every alternate days to determine productivity. For chlorophyll data, 10 mL of each culture was filtered using glass microfiber filter papers (47 mm Ø Whatman® GF/C) at their stationary phase. For Carotenoids data, 1 mL aliquot solution of each culture was collected in 15 mL centrifuge tube. Finally, all the cultures were harvested at their stationary phase based on the growth curve experiment. The cultures were centrifuged to harvest the microalgae (Hitachi* High-speed Refrigerated Centrifuge, himac CR 21g-II). The harvested microalgae was dried at 60°C temperature for 3 days to inhibit contamination, using hot air oven, and subsequently, preserved at normal refrigerator (4°C) for pigments and proximate analysis.

2.4. Determination of productivity

Volumetric [5], areal [6], and lipid [7] data were determined to estimate productivity. Biomass data is the prerequisite of estimating the productivity data. All the data of productivity was calculated at the stationary phase of different microalgae cultures.

2.4.1. Biomass (Dry Weight Basis)

Biomass was estimated every alternate day using 1 mL microalgae samples from each cultures, filtered through pre-weighed (Rinsed with 1 mL distilled water, and oven dried at 60°C for 4 hrs followed by 1 hr desiccation) glass microfiber filter paper. Then the filter paper with biomass was oven dried again at 60°C for 4 hrs followed by 1 hr desiccation. After that, the dry biomass concentration was calculated by dividing the difference between the weights of the dried filter paper (pre and post filtration) by the filtered volume [3].

2.4.2. Volumetric productivity

Volumetric productivity (VP) indicates the average daily productivity of a culture based on dry weight. Following equation was used to calculate the volumetric productivity:

$$
VP \text{ (mg L}^{-1} \text{ day}^{-1}) = \frac{(X_n - X_0)}{N}
$$

Where, $X_n$ = Final biomass, $X_0$ = Initial Biomass and $N$ = Culture days
2.4.3. Areal productivity

Areal productivity \((AP)\) is the productivity of an area occupied by the microalgae. Following equation was used to calculate areal productivity:

\[
AP \text{ (mg cm}^{-1} \text{ day}^{-1}) = (VP \times V)/A
\]

Where, \(VP\) = Volumetric Productivity, \(V\) = Total Volume of the culture, \(A\) = surface area occupied ground.

2.4.4. Lipid productivity

Lipid productivity \((LP)\) is the amount of lipids produced by microalgae in a day during stationary phase. The lipid productivity was calculated using lipid content and volumetric productivity at the stationary phase. Following equation was used to calculate the lipid productivity:

\[
LP \text{ (mg L}^{-1} \text{ day}^{-1}) = VP \times (% \text{ lipid}/100)
\]

Where, \(VP\) = volumetric productivity of the PBR and, \(\% \text{ lipid}\) = lipid content.

2.5. Determination of pigments

2.5.1. Extraction of microalgae for chlorophyll determination

For extraction, 10 mL of each sample was filtered (47 mm Ø Whatman® GF/C glass microfiber filter papers). Filter papers with samples were put into airtight plastic bags, and stored frozen (−20°C) for 3 weeks. After 3 weeks, each filter paper with sample was kept submerged in a centrifuge tube with 2–3 mL 90% aqueous acetone solution (Mixing of 90 parts of Acetone with 10 parts of MgCO₃ Solution), and macerated at 500rpm for 1 min. Then the sample volume was adjusted up to 10 mL with 90% aqueous acetone solution. Subsequently, the samples were steeped for 2 hrs at 4°C temperature. After 2 hrs, the samples were clarified by centrifuging in closed tubes for 20 mins at 500g. Then the clean extract was separated into new tubes.

2.5.2. Determination of chlorophyll

Chlorophyll was determined according to Aminot et al. [8]. The clean extract was transferred to a 1 cm cuvette, and measured optical density \((OD)\) at 750, 664, 647 and 630nm wavelength. OD at 664, 647, and 630nm were used for chlorophyll determination, where OD at 750nm was used as turbidity correction factor. Values of OD at 750nm was subtracted from each of the pigment OD values of the other wavelengths before using them in the equations below:

\[
C_a(\mu\text{g/L}) = 11.85(OD664) - 1.54(OD647) - 0.08(OD630)
\]
\[
C_b(\mu\text{g/L}) = 21.03(OD647) - 5.43(OD664) - 2.66(OD630)
\]
\[
C_c(\mu\text{g/L}) = 24.52(OD630) - 7.60(OD647) - 1.67(OD664)
\]

Where, \(C_a, C_b,\) and \(C_c\) = concentrations of chlorophyll \(a, b,\) and \(c,\) respectively, and OD664, OD647, and OD630 = turbidity corrected optical densities (with a 1-cm light path) at the respective wavelengths.

After determining the concentration of pigment in the extract, following calculation was applied to determine the amount of pigment per unit volume:

\[
\text{Chlorophyll (\mu\text{g/L})} = \frac{\text{Chlorophyll a} \times \text{Extract Volume in mL}}{\text{Volume of sample in L}}
\]

2.5.3. Determination of carotenoids

1 mL aliquot of the algal suspension of each culture were taken at their stationary phase, and centrifuged at 1000g for 5 mins to obtain pellet. Afterwards, the pellet was extracted with 3 mL 2:1 of ethanol:hexane \((v/v)\).Then the pellet with the solvent was shaken vigorously, and centrifuged again at 1000g for 5 mins. Thus, the hexane layer was separated, and its absorbance was determined spectrophotometrically at the wavelength of 450nm. The amount of extracted carotenoids from the samples in micrograms was determined by multiplying the absorbance \((A_{450})\) with 25.2 [9].
2.5.4. Extraction of phycobiliproteins

The cultures were centrifuged at 6,000 rpm at room temperature for 15 mins to harvest the pellet. The cell pellets were rinsed 2–3 times with distilled water. The harvested biomass was dried in oven at 40°C overnight. Dried powder (40 mg) was then soaked in 10 mL phosphate buffer (pH 7.0; 0.1 M), mixed well using vortex mixture, and then stored at 4°C for 24 hrs. Phycobiliproteins were extracted by centrifuging at 6000 rpm for 10 mins. Finally, the supernatant was collected and absorbance was measured spectrophotometrically (UV-VIS Double beam, Model-T80, HANNA) at the wavelength 562, 615, and 652nm; phosphate buffer was used as blank.

2.5.5. Spectrophotometric estimation of phycobiliproteins

The amount of phycocyanin (PC), phycoerythrin (PE) and allophycocyanin (APC) in the sample was calculated from the absorbance using the following equations and the extinction coefficients from Siegelman and Kycia [10]:

\[
\text{PC (mg/mL)} = \frac{A_{615} - (0.474 \times A_{652})}{5.34}
\]

\[
\text{APC (mg/mL)} = \frac{A_{652} - (0.208 \times A_{615})}{5.09}
\]

\[
\text{PE (mg/mL)} = \frac{A_{562} - (2.41 \times \text{PC}) - (0.849 \times \text{APC})}{9.62}
\]

Total phycocyanin, phycoerythrin, and allophycocyanin (mg/g) were calculated according to Silveira et al. [11] as follows:

\[
P = \frac{\text{Pigment concentration V}}{\text{DB}}
\]

Where, V= solvent volume, DB= Dried biomass

Total phycobiliproteins (mg/g) were further calculated from the sum of the phycocyanin, phycoerythrin, and allophycocyanin contents in dried microalgae biomass.

2.6. Determination of proximate compositions

2.6.1. Carbohydrate determination

Carbohydrate was determined according to Dubois et al. [12]. For each sample, 5 mg freeze dried biomass was taken to prepare a 25 mL well mixed (tissue homogenizer) solution using distilled water. Afterwards, 1 mL from 25 mL solution was taken for each samples, and then 1 mL of 5% phenol solution and 5 mL of sulfuric acid were added into it. Then, the samples were kept in a cold water bath. When cooled, absorbance of the solution was taken at 488nm wavelength using spectrophotometer to estimate carbohydrate. To produce a calibration graph, 1000 μg/L of standard (glucose) stock solution was prepared, and subsequently, a series of standards at various dilution (20 μg /L, 40 μg /L, 60 μg /L, 100 μg /L and 140 μg /L) were also prepared from the stock solution. The same procedures as described for carbohydrate analysis were applied for the standard series; a standard graph was plotted according to the standard results obtained from the absorbance, and the carbohydrate composition for every sample was determined accordingly.

2.6.2. Protein determination

Protein was determined according to Lowry et al. [13]. For each sample, 5 mg of freeze dried microalgal biomass was taken to prepare a 25 mL well mixed (tissue homogenizer) solution using distilled water. 0.5 mL from 25 mL solution was taken for each samples for protein analysis. 50 mL of Reactive 2 (2g of Na₂CO₃ in 100 mL of 0.1 NaOH) and 1 mL of Reactive 1 (1% NP tartrate) were mixed. After that, 0.5 mL sample was added with 0.5 mL of 1N NaOH, and it was kept in a hot water bath at 100°C for 5 mins. Subsequently, the samples were cooled in a cold water bath, and 2.5 mL of the prepared mixed reagent was added 10 mins after cooling. After
that, 0.5 mL of Falin reagent was added to the mixed reagent, and then kept in a dark places for 30 mins. The absorbance of the mixed solution was measured using spectrophotometer at 750 nm wavelength. To develop a calibration graph, 2000 μg/L of standard (albumin) stock solution was prepared, and a series of standards were prepared (20 μg/L, 40 μg/L, 80 μg/L, 100 μg/L and 200 μg/L) from the stock solution. The same procedures as described for protein analysis were applied for the standard series; a calibration line was plotted according to the absorbance, and the protein composition for each sample was determined accordingly.

2.6.3. Lipid determination

Lipid was determined according to Bligh and Dyer [14], and Folch et al. [15]. For each sample, an aluminum dishes were labeled and weighted as initial weight. Then 50 mg of each sample was taken in a centrifuge tube, and diluted into 5x volume using distilled water. Then, 3 mL 2:1 of methanol:chloroform (v/v) was mixed with the sample homogenously using tissue homogenizer. After that, all the tubes were centrifuged for 4 mins at 1000 rpm at 4°C; the supernatants were transferred into clean tubes by Pasteur pipette, and placed them in ice. Again, 3 mL 2:1 of methanol:chloroform (v/v) was mixed with the sample homogenously. After that, the tubes were centrifuged at same conditions again, and supernatants were transferred to the previous tubes of supernatants. In this combined supernatant, 1.5 mL of 0.9% NaCl was mixed using vortex mixture. Then the tubes were kept in the refrigerator for 1 hr at 4°C temperature. After 1 hr, the tubes were centrifuged for 10 mins at 1000 rpm at 4°C temperature. The upper layer of methanol and chloroform was discarded, while, the lower layer was transferred in previously made aluminum dish. The solvent was then evaporated at 60°C by hot air oven. Afterwards, the aluminum dishes were weighed to get the final weight. Finally, initial weight was subtracted from the final weight to get the lipid weight in the samples.

2.7. Statistical analysis

Mean and standard deviation of mean were calculated using MS excel. When assumptions were met, ANOVA was applied to test the significance of the difference of productivity, pigments, and proximate composition among the four different microalgae. IBM SPSS (v. 26.0) was used for ANOVA.

Ethical Statement

These data were collected complying ARRIVE guidelines. Ethical approval is not a prerequisite of starting the data collection procedure for microalgae.

Declaration of Competing Interest

The authors disclose that they have no known conflict of interests that may have influenced either the data collection or the presentation of the data.

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