Extracellular Polysaccharide (EPS) in Microalga Porphyridium cruentum (Mutmainnah et al)

Growth Rate and Chemical Composition of Secondary Metabolite Extracellular Polysaccharide (EPS) in Microalga Porphyridium cruentum

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Abstract

Porphyridium cruentum microalgae have great potential to be developed as a source of active ingredients in various fields of health. It excretes secondary metabolites in the form of extracellular polysaccharides (EPS), potentially as antibacterial, antihyperglycemic, and immunostimulant components. This study aims to obtain the best culture density of P. cruentum, the active component of P. cruentum (EPS), and the chemical composition of EPS. The P. cruentum was cultivated on 15%, 20% and 25% culture stock, with an addition of silicate, Fe and vitamins in sterile seawater medium with salinity 35%, pH 8, temperature 25-27°C, with continuous aeration and 2500 lux continuous 24 hours, for 14 days of culture period. The results of this study indicate differences in growth rates and abundance of microalgae to each culture stock. The 15%, 20% and 25% culture stocks each showed stationary phase at days 10th, 12th and 14th, with EPS concentrations of 10 ppt, 12 ppt, and 15 ppt, respectively. Variation of P. cruentum culture showed different EPS results, with a density of 25% capable of producing the highest EPS extract of 15,000 mg.L-1. The EPS is known to contain glucose and carboxylic acid compounds that can be utilized in the health and industrial fields.

Keywords: culture, extracellular polysaccharides, Porphyridium cruentum.

INTRODUCTION

Porphyridium cruentum is one of the rare, single-celled, colon-free Rhodophyceae classes found in moist, freshwater, and dominant terrains found in marine waters [1,2]. The P. cruentum cell is about 4-9 μm in diameter, with cell structure consisting of nucleus, golgi body, mitochondria, chloroplasts, starch, mucus and vesicles. It contains red pigment phycoerythrin which is an additional pigment in the cell, dominant to cover the green color of chlorophyll. The pigment content is affected by the depth of the habitat, the deeper the habitats, phycoerythrin become more dominant, and it decrease in the shallower water. The P. cruentum contains chlorophyll a and has no chlorophyll b, but it has chlorophyll d. The red pigment P. cruentum masks the color of other photosynthetic pigments, so this red pigment becomes the main pigment that plays a role in the reception of light in the process of photosynthesis [1-4].

The P. cruentum cell is bound in the mucilage. Mucilage is a compound expressed by a cell constantly forming a capsule, and surrounds a cell, containing a sulfate polysaccharide or otherwise known as EPS. EPS is essentially a secondary metabolite of microalgae useful as a protective for microalgae cell in the conditions that do not support its life survival. As a source of nutritional reserves in nutritional deficiencies, microalgae cell coatings keep the cells protected from extreme temperature changes. EPS becomes one of the important components in P. cruentum function as antioxidant, antibacterial, antiviral, and anti hyperglycemic [2,5,6]. In addition P. cruentum contains carotenoids which are useful in improving the immune system [7,8].

Porphyridium cruentum has a carbohydrate biomass composition value of 32.1% (w/w), 34.1% crude protein. Mineral contents in 100 g of dried biomass are Na (1130 mg), Mg (629 mg), Ca (4960 mg), K (1190 mg), and Zn (373 mg). The fatty acid content is 1.6% for 16:0, 0.4% for 18:2ω6, 1.3% to 20:4ω6, and 1.3% for 20:5ω6. The biomass of P. cruentum contains pigment in the form of phycoerythrin characteristic of red color. Besides that, the biomass also contains vitamin K, tocopherol and carotene. These microalgae contain active components of phenol, sterols, terpenoids, flavonoids and polysaccharides that can be used as antiviral, antibacterial, antioxidant [2,9].

This microalgae is important to be explored because it has various benefits that can be applied in various fields, especially in medical field. Currently efforts to formulate alternative treat-
MATERIAL AND METHOD

The method used in this study is an experimental method, which is a deliberate method with certain controlled treatment in a place. This research was conducted at Fish Reproduction Laboratory and Parasitic and Fish Disease Laboratory, Faculty of Fisheries and Marine Sciences, University of Brawijaya, Malang.

Data Collection Cultivation of Porphyridium cruentum

Porphyridium cruentum microalgae seeds used in this research were obtained from Brackish Water Aquaculture Center (BBAP) Situbondo. The cultivation media using seawater that has been in sterilization by boiling. The tools used were sterilized using chlorine 15 g.L⁻¹ diluent, after diluted chlorine sprayed sufficiently into a glass jar container of 1.5 L capacity which has been filled with plain water and allowed to stand for 24 hours. Furthermore, the water is drained and the container is allowed to dry until it is completely dry before the tool is used. Cultivation with the addition of Fe and silicates and vitamins as much as 1 mL⁻¹. The lighting 2500 lux and aeration is continuously with 15%, 20% and 25% culture stock [10,11,12]. The microalgae cultivation is in laboratory scale that carried out for 7-14 days, and observed the density of microalgae growth daily using hemocytometer with the aid of 400x magnification light microscope [13].

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Extracellular Polysaccharide Extraction (EPS)
The EPS extraction process was conducted based on Raposo et al. [14], where fresh P. cruentum in culture medium was centrifuged at 10,000 rpm for 20-30 min at 4°C. The formed pellets are removed and only take the supernatant containing EPS. The supernatant is placed under a ventilated hotte to obtain a suspension containing EPS. After 3-4 days, the sample is placed in water bath at 80°C for 1 hour. Suspensions containing EPS are filtered using simple filter paper and the extraction is refined by precipitation with two variations of cold ethanol volume (as required). After the EPS obtained, previously treated with freeze dryer before and after the dialysis process. The process of dialysis is carried out by resuspending dry EPS in distilled water and the process of dialysis using cellulose membranes. This process is done several times.

EPS Chemical Composition Analysis

The chemical composition test of EPS by FTIR method is based on Mishra and Jha [15] research. Fourier Transform Infrared (FTIR) in general aims to detect functional groups of certain compounds, so that the compiler compound can be known in a more detail than just phytochemical test. The test was carried out by homogenizing 200 mg of Kbr as well as 2 mg of EPS powder in the mortar. Samples that have been homogeneous compacted and fed into tensor 37. The test sample is measured at a wavelength of 4000-500 cm⁻¹, resulting in FTIR spectra showing the absorption peaks of the sample wave number. Functional groups are determined by the absorption peaks of expressed wave numbers.

RESULT AND DISCUSSION

Biomass production of Porphyridium cruentum

The Porphyridium cruentum microalga growth with 15%, 20%, and 25% culture stock density reached peak on 10th day and decreased and stable on day 11 until day 14. Generally microalgae growth through several phases starting with phase lag (adjustment), logarithmic phase (growth), stationary phase (stop growing) and ending with death phase [13]. The abundance of P. cruentum which tends to be stable after passing through the stationary phase i.e. on the 8th day, can be caused by continuous aeration and lighting treatment, and the availability of nutrients needed by the microalgae to grow. Until at certain times, the growth will be completely stopped and die, i.e. in the conditions
where nutrients are no longer available in the growth medium of *P. cruentum*. Through this study, the time occurs at week 5 of culture, without the addition of nutrients, where the aeration and lighting process continues.

The growth of *P. cruentum* is influenced by internal factors such as the quality of microalgae itself. It is also strongly influenced by external factors such as availability of light (2500-3000 lux), temperature (25-27°C), media quality (sterile), and availability of nutrients (vitamins, Fe, silicate, phosphate and nitrate) [2,16].

There is a difference in the rate of growth in the variation of the microalgae culture stock, which is due to the microalgae cell density itself, as well as the uneven nutritional and lighting requirements that support the variation in growth rate. Prayitno explained that cell culture density affects the growth pattern of microalgae [17], which in high culture stock can shorten the phase of lag (phase of adjustment), and exponential phase increases faster. Denser culture stock conditions cause increasing competition between cells to obtain light and nutrients. Thus in conditions of lack of light, it can disrupt the process of cell division. In addition, if aeration forces are reduced, nutrient transport does not occur thoroughly, causing microalgae cells to not undergo good cell division that affects the time of growth rate.

In 200 mL culture stock, we obtained faster growth rate so that it reaching stationary phase faster when compared with both other culture stocks, that is 150 mL and 250 mL. This become indication that at amount of growth media 1 L, using 200 mL microalga density, is the best to obtain a fast growth rate. The growth of *P. cruentum* microalgae was observed daily for 14 days of culture, to obtain the amount of *P. cruentum* density presented by growth graph in Figure 1.

![Figure 1](image)

**Figure 1. Porphyridium cruentum** growth with 15%, 20% dan 25% culture stock density, with 2500 lux lighting and continuous aeration, for 14 days of culture.

### Extracellular Polysaccharide (EPS)

The concentration of EPS was obtained by centrifugation process followed by the extraction process, used 96% ethanol solvent with media and solvent ratio of 1:0.75 v/v. Based on microalgae culture results for 14 days of rainy season, at 15% culture density obtained EPS + 10.000 mg.L⁻¹ culture medium, 20% culture stock density obtained EPS + 12.000 mg.L⁻¹, and density stock culture 25% obtained EPS + 15.000 mg.L⁻¹ (Table 1).

A culture of *Porphyridium* sp. was performed in winter and extracellular polysaccharide values are obtained from 200 to 1000 mg.L⁻¹ [18]. In contrast to Setyaningsih et al. [13], where extracellular polysaccharide values were obtained from 12,500 - 21,500 mg.L⁻¹ for a culture time of 12 days. Differences in extracellular polysaccharide values can be influenced by differences in the culture medium used, which include differences in the nutrient composition of each medium, in addition to differences in temperature and illumination.

<table>
<thead>
<tr>
<th>P. cruentum Culture Stock (mL.L⁻¹)</th>
<th>EPS (mg.L⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>150</td>
<td>10,000</td>
</tr>
<tr>
<td>200</td>
<td>12,000</td>
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<tr>
<td>250</td>
<td>15,000</td>
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Biologically extracellular polysaccharides function to protect microalgae cells, play a role in ion exchange or reservoir, as a barrier wall that is difficult to penetrate by gas and water. It is also serves as a place of excretion of vitamins and hormones [1]. The number of EPS obtained shows good microalgae growth, and good cell metabolic processes, to produce continuous extracellular compounds. The resulting extracellular polysaccharides will be a source of nutrients for cells in the growth process. The more cell division occurs, the EPS excretion process will increase, and can extend the growth phase of microalgae caused by the availability of nutrient reserves of EPS.

The chemical composition of EPS is identified from the expression of the functional group via the FTIR test. The graph in Figure 2 shows the chemical compounds read through the functional groups such as wavelength absorption 3442.769 indicating the presence of hydroxylate bond (–OH), wavelength absorption at 2239.846 indicates the presence of carboxylic acid bond (–COOH), the CO bond is indicated by a long absorption wave 1625.614, while wavelength absorption 1438.650 and 1362.350 indicate the presence of phenol bonds, as well as at 1200-800 wavelength absorption indicates the presence of polysaccharide bonds.

Hasanah in 2016 performed FTIR test on P. cruentum EPS found that extracellular type of extracellular polysaccharide sugar was identified only maltoheptose composed of seven maltose disaccharides which is a component of glucose [10]. In contrast, Raposo et al. in 2014, found P. cruentum EPS was tested by GCMS (Gas Chromatografy Mass Spectrometry) method in which extracellular polysaccharides were identified to contain several major sugar components, including galactose (26.5-36.5 M%), glucose (22.5-24% M), and arabinose (16 M%). While other sugar components such as mannose, fucose, xylose, and rhamnose were also detected with low concentrations of 11, 9, 7 and 4 M% [14].

Figure 2. Extracellular polysaccharide functional group (EPS) Porphyridium cruentum, using FTIR test at 4000-500 cm$^{-1}$ wavelength.

It was reported that EPS in P. cruentum composition with several different fractionation treatments using TLC (Thin-layer Chromatography) and GC measurements, so it was known that EPS content included Xyl, Glc, and Gal sugar [19]. Through the method of m-hydroxybiphenyl assay, sodium rhodizionate assay and IR spectrophotography, it was found that EPS sugar content included uronic acid and sulfate compound, through IR spectrophotography on band 820 cm$^{-1}$ indicated hydroxyl primer sulfate group. This shows that the EPS extract contains sugar D- and L-Gal (Fig. 3).

Through a previous study of Geresh et al. in 1990 [20], it found α-d-glucopyranosyluronic acid-(1 → 3)-β-lactopyranose disaccharide from hydrolyzate acid Porphyridium sp. It indicates the configuration of each of GlcA and Gal are d- and -
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1. This disaccharide type is known as the polysaccharide constituent part for several types of red microalgae.

Differences in sugar composition in the EPS testing, allegedly caused by hydrolysis factors that have not been optimal, in this case may be the time of hydrolysis used has not been able to break the chemical bonds EPS Porphyridium cruentum so that no overall chemical composition EPS detected. The amount of galinisic acid hydrolyzed with 0.2 M H2SO4 at 80°C reached 2.5% for 28 hours and at 100°C reached 6% for 16 hours w/w [21]. It shows that hydrolysis values can be influenced by the time and temperature used. In addition Hasanah [10] confirmed the factors that can affect the hydrolysis level can be the time, temperature, type and concentration of the sample, and used solvent.

Figure 3. Unit Polysaccharide 1). Terminal Gal, 2). Xyl Terminal [19]

CONCLUSION

Variation of P. cruentum culture showed different EPS results, with a density of 25% capable of producing the highest EPS extract of 15,000 mg.L⁻¹. Increased chemical content of P. cruentum may be affected by culture modification by the addition of various fertilizers and vitamin supplements. The chemical composition of the dominant EPS is obtained by compounds of hydroxylate bonds including polysaccharides which can be utilized in the health and industrial fields.

REFERENCES


