THE EFFECT OF SHIFTS IN MEDIUM TYPES ON THE GROWTH AND MORPHOLOGY OF SPIRULINA PLATENSIS (ARTHROSPIRA PLATENSIS)

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Key words: Arthrospira, Spirulina platensis, Provasoli, Ultrastructure, Zarrouk.

ABSTRACT

In order to reduce the cultural cost and increase the biomass of Spirulina platensis cultures, two basic culture media were used in the present study, the Zarrouk (Z9) and Provasoli’s enriched seawater (PES) media. They were re-modified into five media by adding varied concentrations of NaCl, NaHCO₃ and seawater (salinity). S. platensis could grow well in all of these media. The Z9 medium provided the best growth rate at the end of culture and the PES medium provided the best initial growth curve slope. After the media were shifted from PES to Z9, the long algal filaments fragmented to form many short filaments within 5 days of culture. Those short filaments then straightened into less spiral forms due to the ingrowths of active cell division after 7 days of culture. This media shifted batch culture is simple to operate with the lowest cost and the highest biomass in laboratory scale culture.

I. INTRODUCTION

It is widely recognized that microalgae can be used as animal feed and human food [3]. The nutritional value of microalgae has been substantiated by numerous studies and compared well with other conventional food products. The food potential of the genus Spirulina seems particularly promising. Spirulina filaments, may be easily separated from their medium, have high digestibility, a mild flavor, and contain up to 70% excellent quality protein [16]. Spirulina is commercially exploited for human food supplements, fodder, and pharmaceuticals such as anti-tumor drugs [6] and anti-HIV-1 [1]. Spirulina species are usually grown in open systems, such as raceway ponds and in closed photobioreactors [15] for mass cultures. Despite the plethora of mass cultured Spirulina, little attention has focused on the impact of the media shift used in the production of Spirulina. Spirulina production as well as its photosynthetic activity and growth-physiology is greatly restricted by the culture media. There have been few studies that reported on the morphological changes involved with inducing and accelerating cell division when Spirulina has been cultivated in various media.

The aim of the present study was to increase the Spirulina biomass in batch cultures by shifting the cultures to different media. This attempt may reduce the cost of Spirulina platensis cultures, especially for ultrahigh-density cultures. Special attention was given to the morphological alterations of the alga, more surprising, was the response of Spirulina to the shift of media.

II. MATERIALS AND METHODS

1. Algal Cultures and Media Preparations

Stock cultures of Spirulina platensis, preserved in the laboratory of the Department of Aquaculture, National Taiwan Ocean University, Keelung, Taiwan, were used as the test materials.

Two modified media, Provasoli’s enriched seawater (PES) medium [10] (Table 1) and Zarrouk (Z9) medium [19] (Table 2) were re-modified to five media as follows:

1. PES medium (as Table 1) in which the salinity of the seawater was reduced to 20‰. The pH value was 8.5.
2. PEF medium in which the nutrients were the same as the PES medium, but freshwater was used instead of seawater with 16.8 g/L NaHCO₃ added for a pH buffer and osmoregulant. The pH value was 9.7.
3. Z9 medium (as Table 2), pH value was 9.7.
4. Z9-1 medium, the same as Z9, but the concentration of NaHCO₃ was reduced to half, and 8.4 g/L NaCl was added for osmotic balance. The pH value was 9.7.
5. Z9-2 medium, the same as Z9, but NaCl (16.8 g/L) was added instead of NaHCO₃. The pH value was set to 8.5 by adding NaOH.
Table 1. The nutrient elements of Provasoli’s enriched seawater (PES) medium (g/L of H2O).

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaNO₃</td>
<td>75</td>
</tr>
<tr>
<td>NaH₂PO₄ • 2H₂O</td>
<td>5.6</td>
</tr>
<tr>
<td>NH₄Cl</td>
<td>26.8</td>
</tr>
<tr>
<td>Na₂EDTA</td>
<td>4.36</td>
</tr>
<tr>
<td>FeCl₃ • 6H₂O</td>
<td>3.15</td>
</tr>
<tr>
<td>Biotin</td>
<td>0.001</td>
</tr>
<tr>
<td>H₂BO₃</td>
<td>2.24</td>
</tr>
<tr>
<td>Thiamin HCl</td>
<td>0.2</td>
</tr>
<tr>
<td>(Vitamin B₁)</td>
<td></td>
</tr>
</tbody>
</table>

Add 1 mL of this stock solution to 1 L of seawater as PES medium.

Table 2. The nutrient elements of Zarrouk (Z9) medium.

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaHCO₃</td>
<td>16.8 g/L of water</td>
</tr>
<tr>
<td>KNO₃</td>
<td>3.0</td>
</tr>
<tr>
<td>K₂SO₄</td>
<td>1.0</td>
</tr>
<tr>
<td>NaCl</td>
<td>1.0</td>
</tr>
<tr>
<td>H₂PO₄</td>
<td>0.25 mL/L of water</td>
</tr>
</tbody>
</table>

Plus 1 mL each of solutions A and B for each liter of the above.

<table>
<thead>
<tr>
<th>Solution A (g/L of water)</th>
<th>Solution B (g/L of water)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H₂BO₃</td>
<td>NH₄VO₃</td>
</tr>
<tr>
<td>MnCl₂ • 4H₂O</td>
<td>CaCl₂</td>
</tr>
<tr>
<td>ZnSO₄ • 7H₂O</td>
<td>FeSO₄</td>
</tr>
<tr>
<td>CuSO₄ • 5H₂O</td>
<td>EDTA</td>
</tr>
</tbody>
</table>

The pH value was measured using a digital pH meter (SUNTEX, SP-701, Taiwan).

The _Spirulina platensis_ stock cultures were transferred into two 3-L flasks with cotton plugs, with 2.5 L of Z9 medium (as Z9 algal stock solution) and 2.5 L of PES medium (as PES algal stock solution) respectively. Samples were cultured in a plant incubator (Firstek, Taiwan). The irradiance was 170 μmol photons • m⁻² • s⁻¹ [18] and 12:12, L:D photoperiod, at 35°C [9].

Because the algal bodies (length and cell number in trichome) of _Spirulina platensis_ vary, a spectrophotometer (Metertek SP-830, Taiwan) was used to estimate the absorbance of the algal cultures. The optical densities of the algal cultures were determined at 650 nm wavelength. Subsequently, the absorbance values were transferred to biomass of day weight which was determined in 1 L of algal culture solution. The cells were washed 3 times with distilled water and dried at 80°C for 24 hrs. The experiments in this study were performed three times, and the average counts were recorded.

2. Shift of Culture Media

When the absorbance of Z9 and PES algal stock solution reached 2.0, ca. 60 mL of Z9 algal stock solution was transferred into five 3-L flasks with 2.5 L of the five media, Z9, Z9-1 Z9-2, PEF and PES, (Z9 shifted to Z9, Z9-1, Z9-2, PEF and PES), respectively. In addition, ca. 65 mL of the PES algal stock solution was transferred (shifted) to a 3-L flask with 2.5 L of Z9 medium (PES shifted to Z9). The initial culture absorbance was set to 0.08 by adding the algal stock solutions. Cultures were incubated in the same plant incubator with the irradiance at 170 μmol photons • m⁻² • s⁻¹ and a 12:12, L:D photoperiod at 35°C.

The culture materials were observed and photographed under light microscope (Zeiss, Axioscope) and electron microscope.

In this culture test, a Z9 culture group was remained the same therefore serving as the control group.

3. Preparation for Electron Microscopy Study

The algal cultures were collected in 15-mL centrifuge tubes centrifuged at 1200 × g for 20 min (HERMLE Z360K) followed by separate fixation in 0.1 M sucrose solution containing 4% glutaraldehyde and 0.1 M sodium cacodylate buffer (pH 7.0) at 4°C for 2 hrs. They were then rinsed twice with a 0.1 M sodium cacodylate buffer containing 10 mM CaCl₂, and sucrose concentration successively reduced to 0.05 M. This treatment was followed by two rinses by a pure (sucrose-free) 0.1 M sodium cacodylate buffer containing 10 mM CaCl₂.

Post-fixation was performed with 2% OsO₄ in 0.1 M sodium cacodylate buffer containing 10 mM CaCl₂ for 1 hr at 4°C.

Thereafter, all materials were rinsed four times with a sodium cacodylate buffer containing 10 mM CaCl₂, three times
Table 3. The initial slopes (from day 1 to day 23) of the growth curves and the growth rates of all the culture groups. The formula of growth rate is: $\mu = \ln x_2 - \ln x_1 / d_2 - d_1$, where $x_1$ and $x_2$ are the absorbance of day intervals $d_1$ and $d_2$.

<table>
<thead>
<tr>
<th>Culture groups</th>
<th>Z9</th>
<th>Z9-1</th>
<th>Z9-2</th>
<th>Z9-PES</th>
<th>Z9-PEF</th>
<th>PES-Z9</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial slopes</td>
<td>0.08</td>
<td>0.089</td>
<td>0.095</td>
<td>0.139</td>
<td>0.121</td>
<td>0.152</td>
</tr>
<tr>
<td>Growth rates</td>
<td>1.25</td>
<td>1.16</td>
<td>0.89</td>
<td>0.75</td>
<td>0.7</td>
<td>1.31</td>
</tr>
</tbody>
</table>

Table 4. The average biomass of all groups at day 40 of culture.

<table>
<thead>
<tr>
<th>Culture groups</th>
<th>Z9</th>
<th>Z9-1</th>
<th>Z9-2</th>
<th>Z9-PES</th>
<th>Z9-PEF</th>
<th>PES-Z9</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gram of dried weight/L</td>
<td>(±0.26)</td>
<td>(±0.24)</td>
<td>(±0.22)</td>
<td>(±0.22)</td>
<td>(±0.25)</td>
<td>(±0.28)</td>
</tr>
</tbody>
</table>

with aqueous ethanol (50%) and gradually dehydrated in ethanol (50, 70, 85, 95, 100%). Dehydrated materials were prepared for transmission electron microscope (TEM, Hitachi HF100) and scanned by electron microscope (SEM, Hitachi S-800).

For TEM, some dehydrated materials were rinsed in propylene oxide (three times, 30 mins each), followed by infiltration in propylene oxide-Spurr’s resin in a decreasing ratio from 2:1 (2 parts propylene oxide: 1 part Spurr’s resin) to 1:1, each for 4 hrs. Samples were then suspended in pure Spurr’s resin for two days at 4°C in darkness before embedding in Spurr’s resin [14]. The thin-sections were stained with uranyl acetate and lead citrate according to Smith and Croft [13].

For SEM, some dehydrated materials were dropped onto specimen holders and then dried with a critical-point-drying machine (Hitachi-HCP-1). Finally, they were coated onto an ion coater (Joel, JCF-1100E) for 220s.

III. RESULTS

As shown in Fig. 1, after 40 days of culture, the absorbance of the Z9, Z9-1 and PES-Z9 culture groups were above 3.0. The PES-Z9 culture group exhibited the best growth condition. It took only 13 days for the PES-Z9 group to reach 3.0 of absorbance, and reached as high as 3.5 (the biomass was 10.29 g dry weight/L) at day 40 of culture, and still showed a trend toward increase. The Z9 culture group showed the slowest growth increase (the initial slope) until day 17. The Z9 group absorbance reached 3.0 at the end of culture (The biomass was 7.13 g dry weight/L as shown in Table 4). The Z9-PES, Z9-PEF and Z9-2 culture groups never reached 3.0 and showed a decrease in growth from day 24 for Z9-PEF and Z9-PES, and from day 25 for Z9-2. As shown in Table 4, the biomass of those groups were under 6.0 g dry weight/L.

As shown in Fig. 1 and Table 3, Z9 to PES (the Z9-PES culture group) and PES to Z9 (the PES-Z9 culture group) exhibited better initial slopes than the other culture groups. Over 23 days of culture the initial slope of the PES-Z9 culture was the best.

The initial slopes for the Z9-PES and Z9-PEF culture groups were better than Z9, Z9-1 and Z9-2. Alga cultured in Provasoli modified media could accelerate cell division, which resulted in an initial phase with the best initial growth curves slopes. In contrast, the growth rates of the Z9, Z9-1 and Z9-2 culture groups were better than Z9-PES and Z9-PEF at the end of culture. Zarrouk modified media were the most suitable ones for steady growth, and produced the better biomass at the end of culture. The PES-Z9 culture group combined the two best factors, the best initial growth slope and the best growth rate, and therefore produced the best biomass.

The final algal filaments from all culture groups were long (ca. 433 µm in length) with ten or more spirals (Fig. 2). The trichome, a chain of cells within the thin filament sheath (Fig. 3) were ca. 6.5 µm in diameter. In the media shift tests, the PES-Z9 culture group, fragmented sections, the hormogonia (Fig. 4) appeared on the long algal filaments within 5 days of culture after the media were shifted. All of the long filaments then became many short filaments (Fig. 5). However, the media shift between the same incubated media, such as from PES to PEF and from Z9 to Z9-1or Z9-2 rarely induced the formation of hormogonia in this alga.

The media shift also resulted in morphological changes in the newly formed short algal filaments after 7 days of culture. Scanning electron microscope (SEM) observation found that the newly formed short filaments straightened and became less spiral (Fig. 6) in contrast to the spiral filament (Fig. 7) nor...
mally found on *Spirulina platensis* cultured in Z9 medium. These filaments were also observed under transmission electron microscope. It was found that the ingrowths (cell division) (Fig. 8) frequently appeared in the newly formed filament in contrast to the more spiral filament (Fig. 9).

**IV. DISCUSSION**

The filamentous alga *Spirulina* is common in lakes with a high soda content and high pH. *Spirulina platensis* can therefore be clearly defined as an obligate alkaliphile [2]. Cultures of this alga are more exceptional than other microalgal cultures. In the present study, it was clearly found that the bicarbonate, NaHCO$_3$, was important for sustained *Spirulina* cultures. This may be because NaHCO$_3$ maintains the high pH environment and osmoticum required by the *Spirulina*. In contrast to the Z9 (Zarrouk medium) culture group, the NaHCO$_3$ concentration in the Z9-1 culture group was reduced to half and the Z9-2 culture group was without NaHCO$_3$. Both groups did not exhibit enhanced initial growth slopes and their growth rates were less than the Z9 culture group at the end of culture (as shown in Table 3). The PEF culture group, with added high NaHCO$_3$ (16.8 g/L) content, showed a small increase in the initial slope better growth rate than the Z9-1 and Z9-2 culture groups at the end of culture. Schlesinger *et al.* [12] also reported that *S. platensis* was alkaliphile and required high sodium content to survive in alkaline environments. Diluted seawater was frequently used instead of NaHCO$_3$ in some commercial mass cultures for reducing costs in Taiwan. However, the results from the present study indicate that *Spirulina* cultures using NaHCO$_3$ for media produced a better
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Fig. 6. The straight short filament with slight spiral after the media were shifted within 5 days of culture.

Fig. 7. Ten-day-old normal spiral filament without the media shift treatment cultured in Zarrouk media.

Fig. 8. Ingrowths (arrow heads) frequently appeared in the short slight spiral filament.

Fig. 9. The normal spiral filament shown without the ingrowths.

Biomass than cultures based on diluted seawater. This result was consistent with the study by Schlesinger et al. [12]. They reported that sodium deprivation caused rapid death in *S. platensis*.

High NaHCO$_3$ content has the additional function of excluding medium contamination by other organisms. Vonshak et al. [17] reported that the contaminant, *Chlorella*, declined rapidly when 16 g/L of bicarbonate was added to the medium.

Although *Spirulina platensis* favors alkaline environments, even at high external pH values, it is capable of maintaining an appreciable pH gradient across its cytoplasmic membrane [5]. The dependency on active sodium-proton antiporters for maintenance of a relatively acidic internal pH [8] that is energetically depleted may delay the logarithmic phase of the growth, causing a lower initial growth slope as shown by the results from algae that were cultured in the alkaline Zarrouk modified media in the present study.

The PES-Z9 culture group biomass was the highest in the present study. This was because the media shift accelerated the algal cell division and formed hormogonia that fragment.
had therefore increased the rate of the initial growth slopes. Although the cultures were first cultivated in PES media, they were later cultivated in Z9 media. The alga growth increased in its’ new favorable environment. The highest produced biomass (i.e. 10.29 g dry weight/L) of the present study was consistent with the results of the complex photobioreactor cultures by Qiang et al. [11]. The present study used normal batch cultures, which were the simplest Spirulina cultures with the lowest cost. It is important to point out the cost of high-tech algal cultures determines the commercial viability of a production crop.

The media shift in the present study could alter the morphologies of the algal filaments after they were shifted from Provasoli medium to Zarrouk within 5 days of culture. This was evidenced in the SEM and TEM observations, in which the short filaments produced many ingrowths that straightened the spiral filament into a less spiral form. In contrast, when the growth of *S. platensis* was inhibited by ethyl methane-sulphonate, their filaments became varied lengths with more spars [4]. Mühling et al. [7] also reported the helix orientation altered in *Arthrospira* when the environmental factor changed, such as temperatures.

A combination of Zarrouk and Provasoli media produced a new medium for ultrahigh-density culture of *Spirulina platensis*. This combination complicated the concentrations of the nutrient elements considerably, resulting in unstable growth with many unknown problems that require further study. The macro-elements of these two media are similar, but the micro-elements of the Zarrouk medium are more complex than those in the Provasoli medium. Provasoli medium includes thiamin HCl, biotin and vitamin 12. I supposed that the vitamins made the alga obtain active cell division when cultured in Provasoli medium. The high NaHCO₃ content in the Zarrouk medium restrained the vitamin effects when they had been added.

In conclusion, the *Spirulina* media shift cultures were easy to operate and ready to use with the lowest cost and high biomass in comparison with other culture methods (i.e. raceway, closed photobioreactors etc.). This finding could facilitate the small scale culture of *Spirulina platensis* in laboratory for the materials for further scientific study, and may apply to out door mass culture for commercial production.

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