Effects of oxygen scavengers (Sodium sulfite, sodium bisulfite, sodium dithionite, and sodium metabisulfite) on growth and accumulation of biomass in the green alga Asterarcys quadricellulare

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ABSTRACT

The present investigation aims to know the effects of oxygen scavengers such as sodium sulfite (Na₂SO₃), sodium bisulfite (NaHSO₃), sodium dithionite (Na₂S₂O₄), and sodium metabisulfite (Na₂S₂O₃) on growth and accumulation of biomass (chlorophylls) in green alga Asterarcys quadricellulare (A. quadricellulare). Alga is grown in tris-acetate-phosphate (TAP) medium, along with various concentrations of NaHSO₃, Na₂SO₃, Na₂S₂O₃, and Na₂S₂O₄ individually under in vitro conditions. To evaluate the effects of scavengers, in vitro grown algal cells were used to estimate the chlorophyll a and b, along with total chlorophylls. Augmented growth and total chlorophyll content (63.23 mg/L) was noticed in A. quadricellulare cultures grown in TAP with 3.2 mM of sodium sulfite medium. TAP with 0.4 mM sodium bisulfite medium enhanced the growth and total chlorophyll content (49.38 mg/L) in this alga. Similarly, improved growth and total chlorophyll content (44.46 mg/L) was observed in A. quadricellulare grown in TAP with 0.4 mM of sodium dithionite medium. TAP with 0.08 mM sodium metabisulfite medium augmented the growth and total chlorophyll content (41.01 mg/L) in this alga. Higher doses of scavengers inhibit the growth and reduced the total chlorophyll contents in A. quadricellulare. The present work will be helpful to standardize the oxygen scavenger resistance levels in various algal species for anaerobic experiments including in vitro production of hydrogen.

1. INTRODUCTION

Oxygen scavengers or absorbers are regularly used in product safety and are helpful in anaerobic experiments. In general, oxygen scavengers inhibit the growth of living organism, including photosynthetic organisms such as algae once if it crosses the critical level. However, it needs for certain experiments such as hydrogen production from algal cells through bypassing the Calvin cycle in photosynthetic process under in vitro conditions [1]. The inclusion of oxygen scavengers in the medium (generally tris-acetate-phosphate [TAP]) inhibits the photosynthetic O₂ in algal cultures, which, in turn, lead to hydrogen production through activation of hydrogenase [2]. It is a well-known fact that hydrogenase is oxygen-sensitive, which plays an important role in hydrogen production in algal species. Chemically, different types of oxygen scavengers are used for removing the oxygen such as sodium sulfite (Na₂SO₃), sodium bisulfite (NaHSO₃), sodium dithionite or sodium hydrosulfite (Na₂S₂O₇), sodium metabisulfite (Na₂S₂O₃), and ammonium sulfite [(NH₄)₂SO₃] [3]. Rapid screening of different photosynthetic algae using various oxygen scavengers is one of the best ways to find out the candidate species, biomass content, status of anaerobic condition, and level of hydrogen production [3,4]. Several works on algae were carried out using various oxygen scavengers for anaerobic condition in turn hydrogen production in Chlamydomonas reinhardtii, Chlorella vulgaris, Chlorococcum minutum, and Scenedesmus [3,5,6]. Among all the scavengers, sodium bisulfite was tested frequently with algal samples for growth and H₂ production [7].

Algae belong to primitive group of plants which divided into unicellular (microalgae) and multicellular (macroalgae) based on their size [8]. Algae, further, classified into terrestrial and aquatic species based on the habitat and aquatic (both marine and fresh water) species are available in bulk number. Furthermore, based on pigment accumulation, algae are divided into green, blue-green, red, and brown algae [9]. Microalgae own a simple life cycle and have short harvesting cycle which is an advantage to choose these species as biofuel resource [10]. Specifically green and blue-green algae are
popular for biofuel production due to their potential photosynthetic pathway [11-13]. Apart from energy sector, algae are a crucial staple for food, fodder, pharmaceutical, nutraceutical, and fertilizer industries. Some of the algae possess the significant quantities of vitamins, proteins, minerals, and lipids [14]. The nutrient content may vary from species to species which, in turn, depends on location, season, temperature, etc. However, certain countries neglecting algal species, and hence, they became orphans without use [15].

The quantity and quality of algal biomass indicates its growth and development. In general, unicellular microalgae are simple forms and chlorophyll is their primary photosynthetic pigment [16]. Hence, chlorophyll accumulation in algal species is an important step for their growth. In extent, research on enhancement of biomass production is a continuous process for algal species due to their enormous benefits [17,18]. In general, estimation of yield by means of biomass content is a regular process. Evaluation of the chlorophyll a, chlorophyll b, and total chlorophylls is one among the simplest ways to estimate the biomass of an organism [19]. Costache et al. [20] mentioned that chlorophylls are abundant in nature which provides green color to algae and have efficient light harvesting role in the process of photosynthesis. Considerable yield of any algal species is totally dependent on the growth and initial biomass accumulation. The biochemical processes involved in the conversion of biomass into biohydrogen were well established using model algal species such as C. reinhardtii and C. vulgaris in the recent past with the aid of oxygen scavengers [3,21]. Asterarcys quadricellulare is a fresh water green alga which belong to class Chlorophyceae [22].

In the current investigation, four oxygen scavengers, that is, sodium sulfite, sodium bisulfite, sodium dithionite, and sodium metabisulfite were tested in A. quadricellulare for its growth and accumulation of biomass which is useful for anaerobic and hydrogen production experiments even in other related algal species. To the best of our knowledge, this is the first report on the effects of oxygen scavengers in growth and biomass accumulation in A. quadricellulare.

2. MATERIALS AND METHODS

A. quadricellulare was a collection from the Department of Center for Advanced Study in Botany, Madras University, Chennai, India. Later, collected algal sample was stored as conventional glycerol stocks in −80°C and some of the cultures were maintained in agar plates and agar tubes for regular experiments. Before the preparation of various algal media for in vitro screening, all the washed glassware (Borosil, India) was desiccated in an oven (Kemi, K04.3, Ernakulam, India) for further usage. In the current work, TAP medium was used for algal cultures along with inclusion of oxygen scavengers, that is, Na$_2$SO$_3$ (0.0, 0.2, 0.8, 3.2, and 12.8 mM), NaHSO$_3$ (0.0, 0.4, 0.8, 1.6, and 3.2 mM), Na$_2$S$_2$O$_3$ (0.0, 0.16, 0.32, 0.40, and 0.80 mM), and Na$_2$S$_2$O$_4$ (0.0, 0.04, 0.08, 0.16, 0.20, and 0.40 mM) independently. Figure 1 illustrates the oxygen scavengers used for the current work. The pH of all the media was adjusted to 7.0 ± 0.1 with standard pH meter (Elico limited, India) and all the experiments were carried out in 30 mL serum vials which consist 5 mL of medium. It is mandatory to maintain empty space (minimum ⅓) in the serum vial to allow algal growth due to aeration facility [3]. Moreover, optimal growth of algae may take around 2–4 days depending on the genus or species or strain in the medium. All the 30 mL serum vials with medium were autoclaved using an autoclave (INLAB Equipment, Madras, India) at 121°C and 103421.36 Pa for 20 min. Further, autoclaved media were cooled down and later moved to laminar air flow chamber for algal inoculation (Hi-Tech products, No. 14, Chennai, India). All the inoculated vials were kept in the orbital shaker (RemiElektrotechnik Limited, Vasai, India) at 140 rpm under light conditions and cultures were grown at 25±1°C in continuous light (107.02 cd) conditions.

In the present work, estimation of chlorophyll content was carried out using modified Arnon’s [23] and Varaprasad et al. [24] methods with 2–3 days old cultures. One gram of fresh algal material of both controls and treated samples were ground with a mortar and pestle using 80% acetone. The homogenate was centrifuged at 5000 rpm for 5 min and the supernatant was saved and the residue was re-extracted with 80% acetone up to the formation of pale yellow residue. Using the supernatant absorbance, values were read at 645 and 663 nm in a ultraviolet (UV) visible Spectrophotometer (Shimadzu UV-1800, India). Total chlorophyll content was estimated using standard Arnon formula and expressed in mg/L fresh weight basis. Three replicates were used in each set and all the experiments were conducted thrice. All the statistical analysis was performed using excel program in personal computer.

3. RESULTS AND DISCUSSION

An effort has been made to understand the effects of Na$_2$SO$_3$, NaHSO$_3$, Na$_2$S$_2$O$_3$, and Na$_2$S$_2$O$_4$ on growth and biomass accumulation in green alga A. quadricellulare. The basic aim of the experiment is to estimate the hydrogen production rate in A. quadricellulare through activation of hydrogenase which, in turn, depends on anaerobic condition. Hence, in vitro screening was carried using four various oxygen scavengers and biomass was estimated using 2–3 days old cultures.

3.1. Effect of Na$_2$SO$_3$ on Growth and Chlorophyll Accumulation in A. quadricellulare

Cultures were initiated in TAP medium with different concentrations of sodium sulfite such as 0.2 mM, 0.8 mM, 3.2 mM, and 12.8 mM, along with control. The inclusion of sodium sulfite with various concentrations in TAP medium promotes the growth and chlorophyll content in A. quadricellulare at optimal concentrations. Optimal growth and high total chlorophyll content (63.23 mg/L) was observed in A. quadricellulare at 3.2 mM of sodium sulfite [Figure 2]. In contrast, high concentration of sodium sulfite reduces the growth and chlorophyll content. Overall A. quadricellulare cultures exhibited resistance at certain extent with this scavenger. Significantly, very few reports are available on the effect of sodium sulfite in algal culture and recently our laboratory established oxygen scavenging capacity of sodium sulfite using green alga C. minutum [3]. Sodium sulfite possesses high electron negativity and also sulfur and oxygen ratio was more when compared to other scavengers utilized in the current
investigation [25]. The severe oxygen removing capacity of sodium sulfite is one among the explanations for necrobiosis or death [26].

3.2. Effect of NaHSO₃ on Growth and Chlorophyll Accumulation in A. quadricellularae

Various concentrations of sodium bisulfite, that is, 0.0 mM, 0.4 mM, 0.8 mM, 1.6 mM, and 3.2 mM were used in TAP medium to know the growth pattern and chlorophyll accumulation in this alga [Figure 3]. Better growth and high total chlorophyll content (49.38 mg/L) was noticed in A. quadricellularae at 0.4 mM of sodium bisulfite. At higher doses of sodium bisulfite, reduction of growth was noticed in this algal species. Among oxygen scavengers, sodium bisulfite is a common reducing agent and it readily reacts with dissolved oxygen [2]. Sodium bisulfite reacts with O₂ and release sodium sulfate, SO₂, and water molecules. Algal cultures treated with high dose exhibited low levels of total chlorophyll accumulation in this species [Figure 3]. Similarly, moderate concentration of NaHSO₃ improved the growth in blue-green alga Anabaena, but high concentration inhibits the cell growth and photosynthesis and ultimately leads to death [2]. Probably, high concentration of scavengers may interact with other components in the TAP medium, and also, there is a chance of interruption of light penetration into the algal cells. Ma et al. [6] also proved that the addition of sodium bisulfite improves the growth and hydrogen production in C. reinhardtii at optimal dose.

3.3. Effect of Na₂S₂O₄ on Growth and Chlorophyll Accumulation in A. quadricellularae

In the current work, sodium dithionite was also used in various concentrations such as 0.0 mM, 0.16 mM, 0.32 mM, 0.40 mM, and 0.80 mM in the TAP medium to check the growth and biomass accumulation in A. quadricellularae [Figure 4]. Comfortable growth

Figure 3: Effect of NaHSO₃ on growth (a) and chlorophyll accumulation and (b) in Asterarcys quadricellularae (Chl a-chlorophyll a, Chl b-chlorophyll b, and T Chl-Total chlorophylls).

Figure 4: Effect of Na₂S₂O₄ on growth (a) and chlorophyll accumulation and (b) in Asterarcys quadricellularae (Chl a-chlorophyll a, Chl b-chlorophyll b, and T Chl-Total chlorophylls).
and high total chlorophyll content (44.46 mg/L) was noticed in this alga in TAP medium with 0.40 mM of sodium dithionite. In contrast, less growth was observed in \textit{A. quadricellulare} at 0.80 mM. Based on the observation, it is confirmed that \textit{A. quadricellulare} exhibited better resistance at optimal dose of sodium dithionite. Long back, Pow and Krasna [5] reported that the addition of sodium dithionite initiate low amount of photobiological hydrogen in green algae \textit{Scenedesmus obliquus} and \textit{C. vulgaris}. Kojima and Lin [27] also reported that the addition of sodium dithionite increases the hydrogen evolution (alternate pathway) in \textit{Chlorella pyrenoidosa} which indicates its role in enhancement of biomass is limited up to certain dose. In addition, sodium dithionite was also used to estimate the H$_2$ase activity in algal cultures [28].

### 3.4. Effect of Na$_2$S$_2$O$_3$ on Growth and Chlorophyll Accumulation in \textit{A. quadricellulare}

Different concentrations of sodium metabisulfite, that is, 0.0 mM, 0.04 mM, 0.08 mM, 0.16 mM, 0.20 mM, and 0.40 mM were used in TAP medium to check the growth pattern and chlorophyll accumulation in this alga [Figure 5]. Augmented growth and total chlorophyll content (41.01 mg/L) in \textit{A. quadricellulare} was observed in TAP with 0.08 mM sodium metabisulfite. At high concentration (0.40 mM), inhibition of algal growth was noticed. Till date, there was no report on oxygen scavenging capacity of sodium metabisulfite with algal cultures, and recently, our laboratory used it while working with \textit{C. minutum} [3]. However, Lee et al. [29] developed the sulfite-based oxygen scavengers for the preservation of plant-based food materials specifically using sodium metabisulfite. Overall current investigation standardized the optimal concentration of oxygen scavengers in this alga, which will be helpful for other algal species specifically for anaerobic experiments.

**Figure 5:** Effect of Na$_2$S$_2$O$_3$ on growth (a) and chlorophyll accumulation and (b) in \textit{Asterarcys quadricellulare} (Chl a-chlorophyll a, Chl b-chlorophyll b, and T Chl-Total chlorophylls).

### 4. CONCLUSIONS

Oxygen scavengers initiate the anaerobic conditions by removing oxygen in any culture. Hydrogen production in algal culture is the main aim of the current investigation, but it occurs through activation of hydrogenase which, in turn, is oxygen-sensitive. Enhanced growth and total chlorophyll content (63.23 mg/L) was noticed in \textit{A. quadricellulare} cultures grown in TAP with 3.2 mM of Na$_2$SO$_3$ medium. Similarly, augmented growth and total chlorophyll content (49.38 mg/L) was observed in \textit{A. quadricellulare} cultures grown in TAP with 0.4 mM NaHSO$_3$ medium. TAP with 0.4 mM of Na$_2$S$_2$O$_3$ medium improved the growth and total chlorophyll content (44.46 mg/L) in this alga. Augmented total chlorophyll content (41.01 mg/L) in \textit{A. quadricellulare} was observed in TAP with 0.08 mM Na$_2$S$_2$O$_3$ medium. Higher doses of scavengers reduced or inhibited the growth in this alga. Present standardization will be helpful for anaerobic experiments in algal cultures.

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All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agreed to be accountable for all aspects of the work. All the authors are eligible to be an author as per the International Committee of Medical Journal Editors (ICMJE) requirements/guidelines.

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