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Biohydrogen production using green microalgae as an approach to operate a small Proton Exchange Membrane Fuel Cell

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Abstract

In this paper the wild-type of Chlorella sorokiniana, a green microalga isolated from Algerian Sahara soil, is tested for its ability to produce hydrogen in a 500 ml photobioreactor coupled to a small Proton Exchange Membrane Fuel Cell (PEMFC). The strain grown in heterotrophically conditions (Tris-Acetate-Phosphate medium) under continuous light is transferred, in medium without sulfur to make anaerobiosis and activate the reversible hydrogenase, inducing the hydrogen production process. The evolution of hydrogen and oxygen concentrations in the photobioreactor during the sulfur-deprived conditions is measured. The difference of total carbohydrates amount and cell morphology at the beginning and the end of experience is also studied. The results of the
coupled system show that the produced biohydrogen can be used to operate a PEM Fuel Cell with good performances under standard conditions.

Keywords: Biohydrogen production; Chlorella sorokiniana strain Ce; PEM Fuel Cell (PEMFC).

1. Introduction

The growing preoccupation for environment and global warming in conjunction with the adherence of signatory countries to Kyoto Protocol including Algeria lead to use clean and renewable energy. Currently, several technologies developed energy systems based on the use of hydrogen as an optimal energy vector for the future. Except its biological production, all the other forms of generation (steam reforming, thermochemical cracking, gasification of coal ...) are consuming fossil energy, electricity and heat. However, biological production occurs under ambient temperature and pressure conditions does not require energy input. This production way is an environment-friendly process and it permits the valorisation of some natural resources, too often neglected.

The hydrogen gas photobiologically produced (i.e. biohydrogen) is considered as a secondary metabolite obtained by microorganism metabolism released under particular conditions. According to microorganism species and biochemical process, different biological ways of hydrogen production are reported in the literature [1]. Biological hydrogen generation can be classified into four categories: (i) direct biophotolysis using green microalgae [2-4] and some cyanobacteria [5], (ii) indirect biophotolysis with other cyanobacteria species and certain nitrogen-fixing bacteria [6,7], (iii) photofermentation of waste and effluents [8] and (iv) darkfermentation of rich sugar wastes [9,10].

Up to now the large utilization of hydrogen produced by biological way in industrial processes is difficult due to its low conversion rate [11]. However, the generation of electricity via small fuel cells using biohydrogen as fuel seems to be a promising application.
Unfortunately, few studies report the practicability of coupling the biohydrogen production to the operation of fuel cell. Electricity generated by the use of biohydrogen in fuel cell varies depending on the microorganism involved, carbon source used, experimental and physiological conditions. Recently, Wünschiers and Lindblad [12] used green algae Scenedesmus obliquus to produce hydrogen gas and developed a software interface to read several parameters such as power generation by the bioreactor connected to Proton Exchange Membrane Fuel Cell (PEMFC). Dante [13] investigated the possibility to exploit directly in a fuel cells stack, the hydrogen produced by Chlamydomonas reinhardtii. He and al. [14] showed that several strains of Rhodobacter capsulatus generate biogas containing hydrogen which is successfully used as feed for a small PEM Fuel Cell system. More recently, García-Peña and al. [15] established a semi-continuous biological system to produce hydrogen and generate electricity by coupling the bioreactor to a fuel cell using a seed sludge as inoculum.

In the same context the present study attempts to investigate the feasibility of electricity generation from green microalgae and to assess the effectiveness of a photobioreactor coupling to a fuel cell. The main purpose of this paper is to estimate and measure the response of a small PEM Fuel Cell by injection of unpurified hydrogen gas produced by Chlorella sorokiniana strain Ce locally isolated from the soil in the Algerian Sahara and incubated under sulphur-deprivation conditions. The next section is dedicated to the description of the experimental setup, materials and methods. The photobioreactor coupled with a small PEM Fuel Cell is also illustrated in this section. The last section is devoted to the presentation of some obtained results and associated discussions. The comparison of the generated electricity by the proposed system with those given in the literature will be done.
2. Experimental methodology

In this section, the growth of the microalgae used and biohydrogen production process are described with the main analyse methods employed to evaluate the production rate of biohydrogen and the degradation rate of carbohydrate during sulfur-deprived conditions. The physiological cell behaviour and the system of photobioreactor coupled with small PEM fuel cell are also detailed.

*Chlorella sorokiniana* strain Ce (wild-type) isolated from microflora sampled from soil of Algerian Sahara was used (Fig. 1). It firstly grew in a Tris-Acetate-Phosphate (TAP) solid medium (pH 7.20) under continuous cool white fluorescence lamps ($\approx 100$ photon·m$^{-2}$.s$^{-1}$). Secondly, to produce hydrogen, the culture with an initial cells concentration of $5 \times 10^6$ cell·ml$^{-1}$ has been transferred in TAP liquid medium without Sulfur at incubation temperature of $23 \pm 2^\circ$C with stirring under the same light intensity as indicated by Chader et al. [4] and Melis et al. [16,17].

![Sampling area](image)

**Figure 1.** Photography of a sampling area in Algerian Sahara.

Hydrogen and oxygen concentrations were measured in the gas phase by a single polarographic probe. This probe operated at a voltage, which is sequentially inverted by means of an electronic switch ($+0.7$ V for H$_2$, $-0.7$ V for O$_2$). The amount of total
carbohydrates was determined by the anthrone-sulfuric acid method suitably adapted for microalgal biomass calculated by means of a calibration curve using D+ glucose dissolved in distilled water [4,18].

Cell morphology of *Chlorella sorokiniana* strain Ce used in this study as a function of sulfur-deprived conditions associated to biohydrogen production was observed at the beginning and the end of experiments under light microscope (Zeiss, Germany) operated at a magnification of 200×.

Experimental setup consists on a photobioreactor coupled with a small PEM Fuel Cell is illustrated in Fig. 2. A glass bioreactor of 500 ml volume, containing the liquid culture described above, is used for hydrogen production, surmounted by syringe to sample the gas from this photobioreactor and to inject it directly through a pipe system to a Proton Exchange Membrane Fuel Cell (H-TEC GmbH. Lübeck, Germany). It’s characterised by an electrode surface of 16 cm² with a size of 105×200×130 mm³ (height×width×depth) producing a power of 1.2 W and maximum operating voltage of 0.9 V. A resistance of 10 Ω is used as the load of the PEM Fuel Cell. Voltage and current values are measured with a multimeter TUV 3275.

**Figure 2.** A system of photobioreactor coupled with a small PEM Fuel Cell.
3. Results and discussion

3.1 Hydrogen production

Under sulfur-deprived conditions sealed Chlorella sorokiniana culture exposed to a continuous light, becomes rabidly anaerobic medium after 12 to 50 h due to the significantly increased of respiration illustrated by oxygen consumption as shown in Fig. 3. After a short time of about 24 h, the hydrogen production begins due to the induction of hydrogenase activity. Indeed, compared to others green microalgae [16,19], the study strain starts earlier the hydrogen accumulation and can sustain production for high O₂ partial pressures.

The obtained results show that Chlorella sorokiniana strain Ce accumulates hydrogen gas under anaerobic conditions such as reported in the literature for several green microalgae [2,4,16]. Indeed, anaerobiosis (under continuous light) is necessary and sufficient for induction of the reversible hydrogenase and for photobiological hydrogen production [20,21]. As indicated by Melis et al. [16], the photobiological hydrogen production is based on the principle to circumvent the severe oxygen sensitivity of the reversible hydrogenase by separating photosynthetic oxygen evolution and carbon accumulation (photosynthesis and growth phase) of the consumption of cellular metabolites (oxidative respiration and hydrogen production phase).

![Graph showing hydrogen production and oxygen consumption](image)

**Figure 3.** Hydrogen production and oxygen consumption by Chlorella sorokiniana cells under sulfur deprivation.

The total gas volume as well as the hydrogen one increase until to reach maximum values respectively 116 ml and 73 ml after 168 h of sulfur deprivation. At this time the concentration
of the hydrogen in the mixture containing N₂, O₂, CO₂ and H₂ [4] is about 63%. The total gaseous mixture produced is divided into five volumes which will be separately injected in a PEM Fuel Cell in order to generate electricity as will be detailed in the third subsection below.

3.2 Physiological cell behaviour

*Chlorella sorokiniana* in the log-linear of growth has small ellipsoid form or slightly ovate with parietal chloroplast and distinctive cell wall. This cellular morphology change severely under sulfur-deprived conditions (Fig. 4). As shown in this figure, cell volume increased and its form becomes spherical due to a large metabolism and degradation of endogenous reserves and carbohydrates (i.e. oxidative respiration) associated to water molecules production. Indeed, before sulfur deprivation this strain accumulates substantial starch in the cells (growth period) degraded during anaerobic period (hydrogen production period). The carbohydrates content decreased down to 54% of the highest starch accumulated level. Similar results concerning cell morphology changes were observed by Zhang et al. [19].

![Morphology of Chlorella sorokiniana cells under sulfur deprivation.](image)

**Figure 4.** Morphology of *Chlorella sorokiniana* cells under sulfur deprivation.
3.3 Electricity generation

In order to know if the biohydrogen produced by *Chlorella sorokiniana* strain Ce is able to generate electricity and to estimate the response of the Proton Exchange Membrane Fuel Cell, the unpurified gas produced is injected into five volumes: 8 ml, 10 ml, 15 ml, 25 ml and 43 ml. A typical voltage response of fuel cell to this gas injection (8 ml containing 5.04 ml of hydrogen) is illustrated in Fig. 5 below.

![Voltage versus time response of PEM Fuel Cell obtained with 8 ml volume of total gas (5.04 ml of hydrogen) produced by *Chlorella sorokiniana* culture.](image)

**Figure 5.** Voltage versus time response of PEM Fuel Cell obtained with 8 ml volume of total gas (5.04 ml of hydrogen) produced by *Chlorella sorokiniana* culture.

Once the gas is injected, the voltage increased rapidly due to the hydrogen conversion into the proton exchange membrane and tends to level off to reach a steady state. Indeed, the ionic equilibrium at the surface of the membrane [22] is obtained after about two minutes giving a maximum voltage of 20.15 mV during 19 seconds. Those values depend on the injected gas volume and the discontinuous injection mode used in this study. After the steady state phase, the voltage decreases slowly until a value of 0.1 mV at the end of experiment. The voltage
curve described above is similar to voltage distribution in PEMFC operated with pure hydrogen under standard conditions [23,24].

Furthermore, concerning the performance of the PEM Fuel Cell for different volumes of hydrogen, the results presented in Table 1 show that these hydrogen amounts generate maximum voltages between 20.1 and 89.1 mV. This interval of efficiencies conversion of biohydrogen is better compared to maximum voltages obtained by the same fuel cell coupled continuously with culture of *Scenedesmus obliquus* obtained by Wünschiers and Lindblad [12]. *Chlorella sorokiniana* generates a maximum current of 8.9 mA with 27.09 ml of hydrogen injected. Using a close hydrogen amount, this maximum current value is higher than the one reported by García-Peña et al. [15] with a dark fermentation process.

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The results of this study show that the hydrogen produced by *Chlorella sorokiniana* strain Ce and injected through a PEMFC was efficiently converted into electricity that has generated measurable voltage and current in agreement with the characteristics of the fuel cell used. We plan to investigate in a future work the permanent coupling of hydrogen bioreactor and a FEM Fuel Cell.
4. Conclusion

*Chlorella sorokiniana* strain Ce culture produced a high amount of biohydrogen under sulfur-deprived conditions. The hydrogen production starts early despite the high oxygen partial pressure in photobioreator. This photobiological process is accompanied by some morphological and physiological changes of cells. Hydrogen content in the mixture gas injected in Proton Exchange Membrane Fuel Cell has been converted in electricity. This fuel cell is successfully operated using the produced biohydrogen and has generate similar response that obtained with pure hydrogen under standard conditions.

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