Development of Bipolar Plate Stack Type Microbial Fuel Cells

Seung-Hun Shin, Youngjin Choi, Sun-Hee Na, Seunho Jung, and Sunggyun Kim

Department of Chemistry and Department of Advanced Technology Fusion, Bio/Molecular Informatics Center, Konkuk University, Seoul 143-701, Korea. E-mail: skim100@konkuk.ac.kr

Received October 22, 2005

Microbial fuel cells (MFCs) have gained much attention as they can use various forms of biomass to generate electricity. Unlike enzyme fuel cells in which enzymes are used as biocatalysts, MFCs utilize living microorganisms to directly convert chemical energy to electricity. Among many advantages of MFCs are environmental friendliness, a large selection of microorganisms, and abundance of fuels.

Although many different types of MFCs have appeared, the current design has not been changed much from a prototype since the basic concept of MFCs was introduced in the 60's. In this design, a single electrode plate is functioning either as an anode or as a cathode. This type of monopolar electrodes does not cause any problem for the purpose of demonstration or development of the MFCs. However, when it comes to the real application where the higher voltage is required to drive external electric devices, monopolar electrode systems may not be realized. Since a single MFC generates about 0.5 V, several single cells should be connected in series to achieve higher voltage. A simple connection by conducting wire between an anode in one single cell and a cathode in the other single cell can cause unwanted problems such as i) non-uniform potential distribution on the electrode surface due to the point contact between an electrode plate and a connecting wire, ii) lower output voltage due to the potential drop occurring between an anode and a cathode, and iii) a large size of the whole cell.

In this paper, we have tried to overcome these problems by designing bipolar electrode plates. In this design, both anode and cathode exist in one plate. Since there are no connecting parts between anode and cathode, problems associated with non-uniform current distribution or voltage drop could be minimized. Also overall cell size can be made small simply by stacking the bipolar plates in series. Although bipolar plates electrodes have already been used in proton exchange membrane fuel cells where the bipolar plates mostly function as a reactant-supplying path and an electrical connection, actual electrochemical reactions can take place in our design. It is first time, to our knowledge, to apply this concept to MFCs. Here we have shown the basic concept of a bipolar stack type MFC and some preliminary results.

Experimental Section

Preparation of microorganisms. P. vulgaris (ATCC 6059) was bought from KCTC (Korean Collection for Type Cultures) and maintained on a nutrient agar plate at 4 °C. The experimental culture was aerobically grown in a nutrient badge containing 10 g·L⁻¹ of NaCl, 10 g·L⁻¹ of tryptone, and 1.8 g·L⁻¹ of glucose at 37 °C. This culture was daily sub-cultured with 5% inoculums for three days, and the cells were harvested by centrifuging at 3000 g and then washed three times with 0.05 M phosphate buffer of pH 7.0. This procedure was carried out at 4 °C. The washed microorganisms were suspended in the phosphate buffer and the weight was adjusted to 10 mg·mL⁻¹ by dry mass. The cell concentration was determined by UV-visible spectrophotometry and a cell colony counting method.

Chemicals. We employed an MFC system operated under the electron mediator which carried electrons from the microorganism to the anode. Thionin (Aldrich, USA) was used as the mediator since it gave the best performance among molecules tested. Glucose (Aldrich) was used as a substrate although other carbohydrates could be utilized. These compounds were used without further purification. Other chemicals were of reagent grade or better. Solutions were prepared using deionized water of 18 MW·cm obtained...
from the ion exchange filter system (Younglin, Korea).

**Bipolar stack fuel cell systems.** Two types of bipolar stack MFCs have been constructed: One type used ferri-cyanide reduction as a cathodic reaction and the other, oxygen reduction. However, the anodic compartment in both designs comprised of *P. vulgaris*, substrate, and a mediator.

Figure 1 shows the single cell-stacked MFC using bipolar plates. Each bipolar plate was made of carbon block and machined into an H-beam shape. It has two compartments at each side that can be used as an anodic and cathodic compartment. Reticulated vitreous carbons (45 pores per inch, E-Tech, USA) were used as electrode material and firmly attached to the carbon plate for electrical connection. Between each bipolar plate was set a semi-permeable membrane. In this figure, five single cells are connected in series and yet only 6 plates are required including end plates. All the plates were strongly tightened by four long screws. Since anode and cathodes are in one bipolar plate, Ohmic drop and non-uniform current distribution could be minimized. An anodic reaction utilized electrons produced by glucose oxidation in *P. vulgaris*, which was transported to the anode by thionin. In the meantime, ferri-cyanide reduction was utilized at the cathode.

Another design in which the oxygen reduction reaction was employed at the cathode is shown in Figure 2. One face of a bipolar plate was grooved for the oxygen passage. Oxygen was catalytically reduced to water by the catalytic layer coated on the Nafion surface. The catalyst made of Pt-embedded carbon nano powder (De Nora North America, USA) was dispersed in 5% Nafion solution (Dupont, USA) and sprayed unto the Nafion membrane before subject to the hot pressing at 120 °C. Oxygen gas was preheated by passing through the hot water before feeding to the cathode. This time two single cells were connected in series.

**Electrical measurements.** The discharge curve was recorded only after the open circuit voltage was stabilized with nitrogen gas flowing through the cell. Discharging was done by connecting an external resistor of various values between the anode and the cathode to obtain a polarization curve. The cell voltage with time was then recorded with a battery tester (Model WBCS-3000, Wonatech, Korea) connected to a personal computer. Current was simply calculated using the Ohm’s law, \( I = \frac{V_{\text{cell}}}{R_{\text{load}}} \). The power was calculated by multiplying voltage and current, \( P = V_{\text{cell}} \cdot I \).

**Results and Discussion**

**Electrochemical behaviors of thionin and ferri-cyanide.** Figure 3 shows voltammograms of thionin (panel A) and Fe(CN)\(_{6}^{3-}\) (panel B) in an fuel cell experimental condition. Thionin shows a reversible voltammetric behavior characteristic of adsorbed species. This certainly makes thionin a very promising electron mediator that can shuttle between microorganisms and the electrode. Note that due to the adsorption effect, current is not proportional to the concentration. Panel B shows that Fe(CN)\(_{6}^{3-}\) reduction is diffusion-controlled and easily reduced to Fe(CN)\(_{6}^{4-}\), indicating that this species is an ideal cathodic fuel. Determined from the peak potential separation, the cell voltage of ca. 0.5 V is expected.

**Fuel cell performance test.** Panel A in Figure 4 is the voltage output as a function of different external loads for the five single cells connected in series, in which ferri-cyanide reduction reaction was utilized at a cathode. The open circuit voltage of 2.5 V indicates that the single fuel cell generates 0.5 V which in turn confirms our bipolar
stack type MFCs are working as expected. As the higher load was applied, the less output voltage was obtained. A new load was connected after the cell voltage was restored to the original value. However, once the higher load (smaller resistance) was applied, a full recovery was not achieved. The fact that physiological conditions of a microorganism are not the same as those in initial stage and the Nafion membrane function is slowly degraded by the thionin adsorption could be some of possible explanations.

Panel B is the plot of power density vs cell obtained from Panel A. Higher voltage resulted in lower the current. Fuel cell assessment can be made from the power by multiplying voltage and current. Maximum power of 0.13 mW·cm$^{-2}$ in this case is regarded very low compared with those of inorganic fuel cell using oxygen and hydrogen as fuels. This number may not be taken seriously because the output power quite depends on the healthiness of microorganisms and the initial culture conditions.

Although the power density from the MFCs is quite low, MFCs can still find many applications in a real world. For the electric appliances consuming small power such as a small motor and electronic calculator, a single MFC can be used. In the case that the large power is needed, electricity from the MFCs can be stored in capacitors for the later use. We used a 1 F supercapacitor of 5.5 V voltage limit to store electrical energy. Figure 5A shows the output voltage variation as a function of time. After the induction period of ca. 200 s, voltage slowly increased to 2.3 V. Thus stored electricity could be used to lighten the electric bulb (Figure 5B).

Another example is the MFC using oxygen reduction reaction at a cathode. Using oxygen gas is advantageous over ferricyanide because oxygen can be supplied from air. However, since the incomplete reduction of oxygen to hydrogen peroxide may damage the cell and poison the cathode, the complete reduction to water is very important. We used Pt-dispersed carbon powder as a cathode material.

Panel A in Figure 6 shows the discharging curve as a function of the external load. In this case two single cells were connected in series. The open circuit voltage of ca. 1.2
V corresponds to the sum of two single fuel cell voltages. The discharging pattern is similar to that of a MFC consisted of five single cells. Panel B is the plot of power density vs cell obtained from Panel A. Maximum power density of 0.023 mW·cm⁻² was found to be lower than that in Figure 4. This is because we did not use RVC as an anode but rather

**Figure 5.** MFC output voltage change as a function of time (panel A) and the picture of powering of an electric bulb (panel B) from electricity stored in a 1 F supercapacitor.

**Figure 6.** Voltage output as a function of different external loads for the two single cells stacked in series. Oxygen reduction reaction was utilized at a cathode. Anodic volume is 1.50 mL. Each anodic compartment contains 6 mL of *P. vulgaris* suspended solution, 24 mL of 1 mM thionin, 6 mL of 1 mM glucose, and 114 mL of buffer solution.

**Figure 7.** MFC output voltage change as a function of time (panel A) and the picture of powering of an electric bulb (panel B) from electricity stored in a 1 F supercapacitor.
used a graphite cell body as an electrode. A supercapacitor was also used to store energy generated from this MFC. Figure 7A shows the output voltage change as a function of time. After the induction period of ca. 500 s, voltage slowly reached to 2.3 V. Thus stored electricity was used to lighten the electric bulb (Figure 7B).

Conclusions

In this paper, we have shown the concept and some results of bipolar stack type MFCs. Contrary to the wire-connected MFC, we used bipolar plates functioning both as an anode and as a cathode to construct MFCs in which each single cell was connected in series. The overall voltage was the mathematical sum of individual fuel cells and no decrease in performance was found. We constructed two MFCs: one with ferricyanide reduction and the other with oxygen reduction reaction at a cathode, but both with the same glucose oxidation reaction catalyzed by \textit{P. vulgaris} at an anode. Electrical energy generated from both MFCs was stored in a supercapacitor to drive electrical devices such as a motor and an electric bulb. We hope that this type of MFCs can find applications in a real world.

Acknowledgement. This work was supported by Korea Sanhak Foundation for the financial support in the program year of 2004.

References