A one-compartment hydrogen peroxide semi-fuel cell was fabricated using a metallic anode (Mg or Al) and Prussian blue as the cathode to improve the power density of the device.
COMMUNICATION

Membraneless Hydrogen Peroxide Micro Semi-Fuel Cell for Portable Applications†

Seyed Ali Mousavi Shaegh, a,*c Seyed Mohsen Mousavi Ehteshami, b,c* Siew Hwa Chan, b,c* Nam-Trung Nguyen d and Swee Ngin Tan e

Abstract

This communication describes the first one-compartment hydrogen peroxide semi-fuel cell as a micro-power source for miniaturized applications. This power source is inspired by taking advantage of the high energy content of metals and design flexibility, light weight and ease of packaging. Benefiting from oxidation of metals, including magnesium (Mg) or aluminium (Al), at the anode and reduction of hydrogen peroxide (H₂O₂) at the cathode, theoretical specific energies of 8600 Wh kg⁻¹ and 17000 Wh kg⁻¹ are achievable. Fabricating a cell with Mg as the anode and Prussian blue (PB) as the cathode, an open-circuit potential of 2.3 V with the net maximum power density of 7.5 mW cm⁻² were achieved. These results are the highest ever reported for one-compartment H₂O₂-based power source. This cell design can provide a platform for fabrication of a new generation of power sources for portable and miniaturized applications.

Portable microelectronic devices including cyborg insects, insect-scale robots and medical point-of-care diagnostic sensors, require micropower sources for autonomous missions and operations. Practically, such micropower sources should have a light weight and a high energy density.

Micro fuel cells have been investigated as micropower sources for portable applications. 4.5 But few issues still remain to be addressed. First, two reagents representing fuel and oxidant must be introduced to anode and cathode compartments, separately. This generally imposes the need of ancillary components adding complexity for the design. Second, a membrane should be sandwiched between electrodes to provide charge transport between them. Such a membrane-electrode-assembly (MEA) increases the total weight of a micro fuel cell and adds complexity to miniaturization and packaging.

In order to address the above mentioned issues, membraneless micro fuel cells, in various configurations, have been investigated. Membraneless microfluidic fuel cells 6.7 usually exploit co-laminar flows containing liquid fuel and oxidant reagents in a micro channel. Fuel-oxidant interface represents a virtual membrane to keep reagents separated. Power densities of up to tens of mW cm⁻² can be reached. But, generally, microfluidic fuel cells need continuous pumping of two streams into the microchannel to provide a confined liquid-liquid interface to avoid fuel crossover. This characteristic results in a reduced net power generation and prevents the deployment of the fuel cell in practical applications.

Ideally, membraneless fuel cells with selective electrodes should provide a prospect to implement anode and cathode in a single compartment. Such a configuration may facilitate delivery of reactants to the electrodes with less required power. Tominaka et al. 8.10 introduced a one-compartment membraneless microfuel cell with a selective electrocatalyst for oxygen reduction in an air-breathing cathode. Each fuel cell was able to use methanol, ethanol or 2-propanol containing a supporting electrolyte in a reservoir fabricated in silicon or bendable polymeric substrates. Using full passive delivery of fuel and oxidant, a net power of 10 µW was achieved.

Furthermore, an ideal design of a membraneless microfuel cell should exploit only one reactant as both fuel (electron donor) and oxidant (electron acceptor) in a single-compartment design. This feature facilitates fabrication of very compact fuel cells. Hydrogen peroxide has a unique characteristic, usually considered as a carbon-free energy carrier, 11, which can be exploited as both fuel and oxidant in a membraneless one-compartment fuel cell. Hydrogen peroxide benefits from easy transport and handling in aqueous phase. 12

Yamazaki et al. 13 used hydrogen peroxide as both electron donor and electron acceptor in an alkaline one-compartment fuel cell. A maximum OCP of 0.13 V was achieved using Au, Ni, Pt and Pd wires as a selective anode and an Ag wire as a selective cathode.

Running a direct hydrogen peroxide fuel cell in acidic conditions is more desirable for hydrogen peroxide-based energy production and storage. 14 Hydrogen peroxide can be formed under acidic environment via the two-electron reduction of oxygen using solar
power. In addition, $\text{H}_2\text{O}_2$ is more stable in an acidic environment with a lower decomposition kinetics which increases the utilization efficiency of $\text{H}_2\text{O}_2$.\cite{13}

Yamada et al.\cite{16} fabricated a direct acidic membraneless fuel cell running only on hydrogen peroxide. An OCP of 0.5 V with a maximum power density of 10 $\mu$W cm$^{-2}$ were achieved using $[\text{Fe}^-\text{tpc}]^2$ as a cathode material and a Ni mesh as anode. Both electrodes were immersed in an acetate buffer with 0.3 M $\text{H}_2\text{O}_2$ at a pH of 3. Recently, we improved the OCP and maximum power density to 0.6 V and 1.55 mW cm$^{-2}$ using, ferric ferrocyanide ($\text{Fe}_4^{11-}[\text{Fe}^{11-}\text{CN}_6]^3$), coated on a carbon-fibre-paper as a cathode material and a Ni mesh as anode. A 0.5 M $\text{H}_2\text{O}_2$ solution containing 0.1 M HCl was used as both fuel and oxidant.\cite{14} More recently, Yamada et al. used other cyanide complexes coated on carbon cloth as cathode materials and a Ni mesh as anode, and achieved an OCP of 0.78 V and a maximum power density of 1.2 mW cm$^{-2}$.\cite{17}

It is very appealing to improve the performance of one-compartment fuel cell for microelectronic portable applications. In this communication, we report a small scale single-compartment semi-fuel cell using $\text{H}_2\text{O}_2$ in acidic condition with improved power output compared to other membraneless hydrogen peroxide fuel cells ever reported. The cathode was comprised of Prussian blue coated on carbon-fibre-based paper.\cite{14} The anode was made of Al or Mg plates. Both anode and cathode were immersed in a reservoir containing 0.5 M $\text{H}_2\text{O}_2$ with a supporting electrolyte of 0.1 M HCl, shown in Fig 1. Procedure of PB synthesis was adopted from elsewhere.\cite{18} PB was coated on carbon-fiber-paper by spraying a solution of PB containing Naflon.\cite{14} Both Mg and Al plates had a thickness of 0.25 mm purchased from Alfa-Aesar and Good Fellow, respectively. A gap between the electrodes was maintained by a silicon rubber slab as an insulating spacer with a thickness of ca. 1 mm.

The overall redox reactions happening in Mg-PB and Al-PB cells are $\text{Mg} + \text{H}_2\text{O}_2 + 2\text{H}^+ \rightarrow \text{Mg}^{2+} + 2\text{H}_2\text{O}$; $E^0 = 4.15$ V vs. SHE (1) $\text{Al} + \text{H}_2\text{O}_2 + 2\text{H}^+ \rightarrow \text{Al}^{3+} + 2\text{H}_2\text{O} + e^-; E^0 = 3.43$ V vs. SHE (2) Hydrogen peroxide has a specific energy of 478 Wh kg$^{-1}$\cite{16} while Mg-$\text{H}_2\text{O}_2$ and Al-$\text{H}_2\text{O}_2$ cells have specific energies of 8600 Wh kg$^{-1}$ and 17,000 Wh kg$^{-2}$. For comparison, lithium-ion batteries and lithium-air batteries have theoretical specific energy of less than 1000 Wh kg$^{-1}$ and 12,000 Wh kg$^{-1}$.\cite{20}

OCP of two cells was monitored for 600 seconds before running other experiments. Mg-PB cell produced an OCP of ca. 2.3 V while OCP of Al-PB cell was ca. 1.25 V (see Fig S1 at ESI†). Both values are higher than the theoretical OCP of a membraneless hydrogen peroxide fuel cell (1.09 V). The actual cell open-circuit potentials are lower compared to the theoretical ones. This is attributed to the occurrence of parasitic reactions in the cell causing a significant drop in the cell potential.

Linear potential sweep voltammetry experiment was carried out to study the potential–current characteristics of the device using only 2 ml solution in a 13 mm-diameter container. As shown in Fig 2, a maximum power density of ca. 4.9 mW cm$^{-2}$ at potential of 1.1 V was achieved using Mg plate as the anode and PB as the cathode. The device using Al as anode material with PB cathode produced a maximum power density of 3.3 mW cm$^{-2}$ at ca. 0.58 V. These power densities are three and two times higher, respectively, than that produced by a membraneless hydrogen peroxide fuel cell using Nickel mesh anode and PB-based cathode immersed in a 50 ml solution (1.55 mW cm$^{-2}$)\cite{14}. For Mg anode with PB cathode immersed in a beaker including 50 ml solution, a maximum power density of up to 7.5 mW cm$^{-2}$ was achieved. Higher power output is mainly because of improved mass transport to the electrodes.

In order to investigate the long term performance and stability of the device, chronoamperometry test was carried out for both cells using Mg and Al anodes. As shown in Fig 3, cell with Mg anode showed a sharp drop in current density compared with the cell with Al anode. The overall decrease of current density for both cells might be attributed to $\text{H}_2\text{O}_2$ consumption, degradation of electrode electrochemical activity and losses due to mass transfer. In the case

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{fig1.png}
\caption{Schematic of the 2 mL one-compartment semi-fuel cell running on $\text{H}_2\text{O}_2$ with a supporting electrolyte of 0.1 M HCl. The anode consists of either a Mg or a Al plate. The cathode is made of Prussian blue coated on carbon paper.}
\end{figure}
of the Mg anode, it was observed that a large area of the electrode was covered by bubbles, limiting H₂O₂ transport to electro-catalytic active sites. The source of the bubbles could be either hydrogen or oxygen molecules produced from metal corrosion or hydrogen peroxide decomposition (see ESI for corresponding chemical consumption).

To investigate the prevailing mechanism responsible for the degradation of performance of the cell with Al as anode, the solution in the reservoir was changed with a fresh solution, after the first experiment. It was observed that the subsequent current generation followed the same trend of the first operation (see ESI† for experimental results). This implies that the cell performance degradation is mainly attributed to mass transfer which is the result of a combination of bubble formation around anode and H₂O₂ consumption.

Conclusion

Miniaturized one-compartment membraneless H₂O₂ semi-fuel cells using magnesium and aluminium as anode materials and Prussian blue as cathode material with high performance were realized. Open-circuit potential and maximum power density were improved considerably compared to the previous one-compartment membraneless H₂O₂ fuel cells reported in the literature. Since the membrane is removed from the structure of the device, there is no requirement for a tight membrane-electrode assembly with passive delivery of reagents to electrodes. In addition, this design benefits from an ease of operation with potential of fuel recirculation to improve performance. These features result in reduced mass of inactive materials and a light-weight assembly and packaging structure, ideal for portable applications. High specific energies with high theoretical OCPs, make H₂O₂ semi-fuel cells very appealing for further developments.

The performance of the fabricated device was mainly restricted by a combination of electrochemical kinetics and mass transport. More investigations should be focused on cell chemistry to determine parasitic reactions deviating the cell potential from theoretical values (see ESI†). Synthesised nano-electrocatalysts with three-dimensional structure can be explored in preference to plain metallic thin plates in order to enhance electrocatalytic reactions. In addition, microfabrication techniques can be adopted for making the miniaturized electrodes, reservoir and assembly.

Notes and references

†Electronic Supplementary Information (ESI) available: [details of any supplementary information available should be included here]. See DOI: 10.1039/c000000x/
