High-performance Fuel Cell with Stretched Catalyst-Coated Membrane: One-step Formation of Cracked Electrode

Sang Moon Kim1,2,*, Chi-Yeong Ahn3,4,*, Yong-Hun Cho5, Sungjun Kim3,4, Wonchan Hwang3,4, Segeun Jang2,6, Sungsoo Shin2,6, Gunhee Lee2,6, Yung-Eun Sung3,4 & Mansoo Choi2,6

We have achieved performance enhancement of polymer electrolyte membrane fuel cell (PEMFC) though crack generation on its electrodes. It is the first attempt to enhance the performance of PEMFC by using cracks which are generally considered as defects. The pre-defined, cracked electrode was generated by stretching a catalyst-coated Nafion membrane. With the strain-stress property of the membrane that is unique in the aspect of plastic deformation, membrane electrolyte assembly (MEA) was successfully incorporated into the fuel cell. Cracked electrodes with the variation of strain were investigated and electrochemically evaluated. Remarkably, mechanical stretching of catalyst-coated Nafion membrane led to a decrease in membrane resistance and an improvement in mass transport, which resulted in enhanced device performance.

Polymer electrolyte membrane fuel cell (PEMFC) has been extensively studied as an eco-friendly alternative energy device in the aspect of high energy conversion efficiency and continuous production of the electricity without pollutant emissions1–4. Recently, there have been several technical issues for the commercialization of PEMFC including water management at the cathode in the membrane electrode assembly (MEA)5–8 and resistance reduction of the electrolyte membrane9,10. To improve transport of water generated from electrochemical reaction at the cathode, research has been conducted on inserting meso/macro pore structures such as platinum inverse opal structure and pore formers into the electrode, which methods need chemical post-treatments11–15. Moreover, there have been attempts to reduce the resistance of an electrolyte membrane by lowering the thickness of the membrane. This thinning method, however, has remained as a challenge due to the inferior mechanical properties of the thinned membrane16,17.

Herein, in order to address these issues in a simple and effective way, we deliberately generated cracks on the electrode composed of carbon supported Pt particles (Pt/C) by stretching out a membrane electrode assembly (MEA) and used the stretched MEA in order to enhance mass transport in the electrode and to reduce ohmic resistance simultaneously. Generally, cracks are generated from the elastic mismatch between two attached surfaces with different elastic modulus when the surfaces are deformed16,17. Cracks have commonly been considered as defects18–21 and many studies have focused on how to avoid them. However, recent studies have indicated that cracks can be used to detect the strain force as in an ultrasensitive sensor16, microfluidic channels17 and so on. It is noteworthy that the crack can be a useful tool rather than a defect, when it is properly generated with a purpose of achieving a specific goal.

1Department of Mechanical Engineering, Incheon National University, Incheon, 406-772, Korea. 2Global Frontier Center for Multiscale Energy Systems, Seoul National University, Seoul 151-744, Korea. 3Center for Nanoparticle Research, Institute for Basic Science (IBS), Seoul 151-742, Korea. 4School of Chemical and Biological Engineering, Seoul National University, Seoul 151-742, Korea. 5Department of Chemical Engineering, Kangwon National University, Samcheok 245-711, Korea. 6Division of WCU Multiscale Mechanical Design, Department of Mechanical and Aerospace Engineering, Seoul National University, Seoul, 151-742, Korea. *These authors contributed equally to this work. Correspondence and requests for materials should be addressed to Y.-E.S. (email: ysung@snu.ac.kr) or M.C. (email: mchoi@snu.ac.kr)
Results

Generation of cracks in the electrode by stretching out the catalysts-coated Nafion membrane. Figure 1 shows the schematic illustration of crack generation in the electrode. A Nafion membrane has a relatively lower elastic modulus than a porous electrode with Pt/C. From this difference in elastic modulus called elastic mismatch, when the two attached surfaces are stretched out, cracks are generated on the porous electrode. If the Nafion membrane is stretched over the elastic deformation region, then it would not recover its original shape, and this means that the membrane has become longer than its original length due to the applied strain. And then, the cracks on the electrode also maintain their deformed shape along with the deformed membrane.

Physical property of Nafion membrane. To investigate the stretching property of a Nafion membrane, we conducted a strain test on a Nafion 212 membrane with a thickness of ~50 μm. As shown in Fig. 2, the membrane showed the maximum tensile strength of ~31.11 MPa and elongation of ~300% till its break. (Table S1) This result shows that the membrane can endure the stress until reaching the maximum tensile strength before it tears off, and can be stretched out to ~3 times of its original length, which is an intriguing feature in terms of stretchability. Furthermore, the shape of the strain-stress curve in Fig. 2a presents that the region of elastic deformation completely which has recovered to its original length is under ~0.08 strain, and it shows that the plastic deformation which has incompletely recovered in the region is above ~0.08 strain. The plastic deformation region is much wider than the elastic region and we used the plastic deformation of the membrane to simultaneously reduce the thickness of the membrane and generate cracks in the electrodes composed of Pt/C particles. As shown in Fig. 2b, we measured the rate of changes of width, height and thickness of a Nafion membrane according to the variation of strains (0.5, 1.0, 1.5 and 2.0) after removing them. And, we ascertained that changes of the membrane thickness were spatially uniform after the removal of strains as shown in Fig. S1.

Morphological features of the generated cracks with variation of strains. To investigate the geometrical features of the cracks induced by applied strains, we observed the cracks with variation of strains by using a scanning electron microscopy (SEM) as shown in Fig. 3a. The catalyst-coated membranes were stretched out with variation of strains (S; 0.5, 1.0, 1.5, and 2.0) and areal fractions of the cracks with each strain in the corresponding SEM images were analyzed by using an image analysis program (ImageJ). As expected, the areal fractions of cracks increased as the value of strain increased. (~7.8% for S~0.5, ~13.4% for S~1.0, ~22.5% for S~1.5, and ~33.4% for S~2.0) Interestingly, we observed that the size of the cracks enlarged as the strain intensified. It implies that the cracks, or the macro-pores, in the catalyst layer can be controlled by the value of strain.

Improved performance with cracked membrane electrode assembly (MEA) in polymer electrolyte membrane fuel cell (PEMFC). To demonstrate the effect of cracks on the performance of PEMFC,
we incorporated a cracked MEA into a single cell. This single cell with the cracked MEA was operated in a fully humidified condition of H₂/O₂ (H₂/Air) and exhibited highly improved performance under all the conditions regardless of the strains compared with a conventional MEA, as shown in Fig. 3b,c. However, the MEA with a strain of ~2.0 applied exhibited more decreased performance than other samples under H₂/air condition. This resulted from the catalysts debonded by high strain. When looking into the surface morphology of the stretched MEA at a lower magnification of the SEM (Fig. S2), we observed that only the catalyst layer of the stretched MEA with a strain of ~2.0 was detached from the membrane (In some cases, we observed with the naked eye that large Pt/C aggregates in the catalyst layer were locally separated from the membrane.). The detached catalyst layer would induce degradation of performance due to the increase of interfacial resistance and loss of the Pt catalyst. And, the result indicates that there exists the upper limit of applied strain to the catalyst-coated membrane and the optimal strain for the high performance of the MEA. In our experimental sets, the stretched MEA with 1.5 strain applied exhibited the highest performance. It showed the maximum power density of ~0.72 W cm⁻² in the case of H₂/air conditions, which power density is higher than the conventional one (~0.59 W cm⁻²) by ~22%. (Table S2) These performance enhancements can be explained by the effects from a crack acting as a macro-pore and the thinned membrane. First, the water transport enhancement which was confirmed by the difference of the power density that increased as the current density increased contributed to the performance enhancement of the MEA. Second, the membrane thickness reduced by stretching out has lowered ohmic resistance that occurs during the PEMFC operation. To address the stability issue, we conducted accelerated durability tests (ADTs) by repeated polarization test for ~1000 cycles with fully humidified H₂/Air gases supplied to anode and cathode, respectively. Even after ADT tests, the maximum power density of the stretched MEA was higher than the conventional MEA by ~36.7% in the case of H₂/Air conditions under ambient pressure as shown in Fig. S3. Furthermore, the morphology of the cracks was maintained even after the durability test was carried out as shown in Fig. S4. There was no breakdown of the system even with the long term electrochemical stress. The results have come from the fact that pre-defined cracks from stretching out provide more available space that would distribute stress from the Nafion membrane when it swells and contracts. Hence, the stretched MEA with cracks have advantages in durability and long term stability than the conventional one.

**Electrochemical analysis of cracked MEA.** To quantitatively investigate the effect of water transport enhancement and reduced ohmic resistance, both electrochemical impedance spectroscopy (EIS) and cyclic
voltammetry (CV) were conducted (Fig. 4). As shown in Fig. 4a, comparable electrochemical active surface area (ECSA) was observed from the CV measurement, which indicates that the generated crack doesn’t affect the area of tri-phase boundary during the operation. When the EIS data were fitted and calculated by the equivalent circuit (Fig. 4b), we found relatively lower ohmic resistance in the case of the 1.5-strain-applied MEA than that of the conventional one by ~14%, which resulted from the thinned membrane in the stretching process. Moreover, the Warburg impedance of the 1.5-strain-applied MEA (0.0393 Ω cm²) was much smaller than that of the conventional one (0.0691 Ω cm²), although the current density of the 1.5-strain-applied MEA was higher than that of the reference at 0.6 V. (Table S3) Furthermore, results of EIS measurement at the same current density (~1.4 A cm⁻²) show less resistance in the case of stretched MEA (S~1.5) than that of the reference as shown in Fig. S5. It means that the water removal from the cathode catalyst layer has been improved due to the existence of cracks. The enhanced water transport also can be confirmed by calculating the oxygen gain. The data of the oxygen gain were obtained by calculating potential difference when oxygen and air are supplied in the current density range and they indicate the degree of mass transfer in the cathode catalyst layer. In Fig. S6, the calculated oxygen gain of the 1.5-strain-applied MEA showed much lower values than the conventional one, which means that the generated cracks in the cathode catalyst layer removed the produced water much easier and helped the MEA to supply fuel gas more effectively than the conventional one.

**Discussion**

To further investigate the mass transport effect while excluding the membrane thinning effect, we measured performances of the stretched MEA and the MEA with only the membrane stretched. The performance of the MEA with only the membrane stretched was slightly higher than that of the stretched MEA in the condition of H₂/O₂. However, in the condition of H₂/Air, the stretched MEA showed much higher performance than the MEA with only the membrane stretched as shown in Fig. S7. In view of the commonly known fact that the performance in the condition of H₂/Air is more affected by mass transport than that in the condition of H₂/O₂, the stretched MEA
with cracks displayed positive effect in improving mass transport. We also conducted EIS measurement of the stretched MEA and the MEA with only the membrane stretched. Fig. S8a shows that they exhibited comparable ohmic resistances. The radius of the circle of the stretched MEA was much smaller than that of the MEA with only the membrane stretched, which implies that the generated crack has improved mass transport. Moreover, oxygen gain calculation for the experiments also supports our results (Fig. S8b).

In this study, we have presented, for the first time, a novel strategy to improve water transport and reduce ohmic resistance by simply stretching out the Pt/C-coated Nafion membrane. With the use of the unique stretching property of the Nafion membrane which has the maximum elongation of ~300% until its break, we have incorporated electrode cracks with the variations of strain (0.5, 1.0, 1.5 and 2.0) into a single cell. The generated cracks effectively enhanced water transport in the cathode catalyst layer, which was confirmed by oxygen gain and EIS measurements. Moreover, the thinned membrane from stretching out was found to reduce the ohmic resistance. These combinational effects of improved water transport and reduced ohmic resistance of the 1.5-strain-applied MEA resulted in performance enhancement compared by ~22% of the performance of the conventional one. Our novel approach of using cracks that have been considered as defects surely possesses potential to be applied to other energy conversion and storage devices. This technique can also be helpful, with its thinned MEA, in reducing the total volume of fuel cell stacks for compact applications such as in vehicles.

**Methods**

**Preparation of membrane electrode assembly (MEA).** Catalyst ink was prepared by mixing water, 5 wt.% Nafion solution (DuPont) and isopropyl alcohol (IPA) (Aldrich) with the catalyst. 40 wt.% Pt/C (Johnson Matthey) was used for the cathode catalyst inks. The prepared catalyst ink was blended by ultrasonic treatment and sprayed onto the cathode side of bare Nafion 212 membrane to fabricate MEAs. The catalyst loadings were equally 0.2 mg cm\(^{-2}\) in the cathodes of the MEAs. These catalyst-coated membranes (CCMs) were dried at room

---

**Figure 4.** Electrochemical analysis. (a) Cyclic voltammogram (CV) of the cathode catalyst layers of a conventional MEA and the 1.5-strain applied MEA. (b) Electrochemical impedance spectroscopy (EIS) of a conventional MEA and the 1.5-strain applied MEA at 0.6 V compared with RHE. (c) Equivalent circuit of the PEMFC (\(L_W\) = inductance of the electric wire, \(R_{membrane}\) = internal membrane resistance, \(R_{cathode\ (anode)}\) = charge transfer resistance of the cathode (anode), \(CPE_{cathode\ (anode)}\) = constant phase element of the cathode (anode) and \(Z_W\) = Warburg impedance).
temperature for more than 12 hours. After the process of stretching out the CCMs, the prepared catalyst ink was sprayed onto the anode side with 0.2 mg cm\(^{-2}\) catalyst loading. Then, gas diffusion layers (GDLs, SGL 35 BC), Teflon type gaskets and one channel serpentine-type (Fig. S9) were put onto the anode and cathode without a hot-press process.

**Process of stretching out prepared CCMs.** As-prepared catalyst-coated membranes were stretched out by a stretcher machine (Intron Corp.), which can provide the strong binding force of the chuck and the uniform strain to the membrane. One side of the membrane was fixed and the other side of the membrane was pressed by two bite-blocks with the width of 5 cm. The initial length of the membrane was 2 cm and the membrane was stretched out with variation of strains (0.5, 1.0, 1.5, and 2.0). See detailed experimental process in Fig. S10. And the samples without the stretching process were also prepared as a reference.

**Physical analysis.** Field emission-scanning electron microscopy (FE-SEM) was conducted using a SUPRA 55VP microscope (Carl Zeiss) to measure the morphology of the various samples used in this paper. The samples were observed at SE mode without additional coating processes.

**Electrochemical measurements.** Prepared MEAs were assembled in a single cell (CNL PEM005-01, CNL Energy). For the single cell performance test at 70 °C, humidified H\(_2\) and O\(_2\) (air) gases were made to flow into the anode and cathode with the active geometric areas of 5.0 cm\(^2\), respectively. The stoichiometric coefficient of H\(_2\)/O\(_2\) (air) was 2.0/9.5 (2.0). Additionally, the relative humidity (RH) for the anode and cathode gases were 100%. To check repeatability, we tested five different samples which have the same applied strain. (Fig. S11). Electrochemical impedance spectroscopy (EIS) (Zennium, Zahner) of the single cells was measured at 0.6 V with an amplitude of 5 mV. The measurement was conducted in the frequency range from 0.1 Hz to 100 kHz. Other experimental conditions, such as temperature and gas humidification, were the same as the case for the single-cell operation at 70 °C with H\(_2\)/Air. The ZView program (Scribner Associates Inc.) was used to fit the EIS data, and a simple equivalent circuit was applied as shown in Fig. 4c. Cyclic voltamograms (CVs) were obtained at 100 mV s\(^{-1}\) between 0.05 and 1.20 V to measure the electrochemical active surface (EAS) of the prepared cathode catalyst layers at room temperature. Humidified H\(_2\) and N\(_2\) gases were supplied to the anode and cathode, respectively, and the RH was 100% during the CVs measurement. The anode with H\(_2\) gas flowing around was used as the reference and counter electrodes, and the cathode with N\(_2\) gas served as a working electrode.

**References**

1. Gasteiger, H. A. & Marković, N. M. Just a Dream—or Future Reality? *Science* **324**, 48–49 (2009).
2. O’Hayre, R. P., Cha, S.-W., Coella, W. & Prinz, F. B. Fuel cell fundamentals. 2nd ed. Ch. 1, 3–23 (John Wiley & Sons New York, 2006).
3. Barbir, F. PEM fuel cells. 2nd ed. Ch. 1, 1–16 (Springer, 2006).
4. Jang, J.-M. et al. The Analysis on the Activation Procedure of Polymer Electrolyte Fuel Cells. *Journal of Electrochemical Science and Technology* **2**, 131–135 (2011).
5. Kang, N. et al. High-performance hybrid catalyst with selectively functionalized carbon by temperature-directed switchable polymer. *Chemistry of Materials***25**, 1526–1532 (2013).
6. Sinha, P. K., Mukherjee, P. P. & Wang, C. Y. Impact of GDL structure and wettability on water management in polymer electrolyte fuel cells. *J. Mater. Chem.* **17**, 3089–3103 (2007).
7. Li, H. et al. A review of water flooding issues in the proton exchange membrane fuel cell. *Journal of Power Sources***178**, 103–117 (2008).
8. Tanuma, T. *Ex Situ Characterization Method for Flooding in Gas Diffusion Layers and Membrane Electrode Assemblies With a Hydrophilic Gas Diffusion Layer. Journal of Fuel Cell Science and Technology***12**, 061002 (2015).
9. Slade, S., Campbell, S., Ralph, T. & Walsh, F. Ionic conductivity of an extruded Nafion 1100 EW series of membranes. *Journal of the Electrochemical Society***149**, A1556–A1564 (2002).
10. Cho, H. et al. Multiplex lithography for multilevel multiscale architectures and its application to polymer electrolyte membrane fuel cell. *Nature communications***6**, 8484, doi:10.1038/ncomms9484 (2015).
11. Kim, O.-H. et al. Ordered macroporous platinum electrode and enhanced mass transfer in fuel cells using inverse opal structure. *Nature communications***4**, 2473, doi: 10.1038/ncomms3473 (2013).
12. Cho, Y.-H. et al. Improved mass transfer using a pore former in cathode catalyst layer in the direct methanol fuel cell. *international journal of hydrogen energy***37**, 11969–11974 (2012).
13. Beshetsenko, T. V., Kim, H.-T. & Kweon, H.-J. Cathode structure optimization for air-breathing DMFC by application of pore-forming agents. *Journal of Power Sources***171**, 433–440 (2007).
14. Zhao, J. et al. Addition of NH\(_4\)HCO\(_3\) as pore former in membrane electrode assembly for PEMFC. *International Journal of Hydrogen Energy***32**, 380–384 (2007).
15. Cheon, S.-Y. et al. Electrocatalytic Nickel Plating on Fibers for the Highly Porous Electrode. *Journal of Electrochemical Science and Technology***11**, 117–120 (2010).
16. Kang, D. et al. Ultrasensitive mechanical crack-based sensor inspired by the spider sensory system. *Nature***516**, 222–226 (2014).
17. Kim, H.-N., Lee, S.-H. & Suh, K.-Y. Controlled mechanical fracture for fabricating microchannels with various size gradients. *Lab on a Chip***11**, 717–722 (2011).
18. Nam, K. H., Park, J. H. & Ko, S. H. Patterning by controlled cracking. *Nature***485**, 221–224 (2012).
19. Livne, A., Bouchbinder, E., Svorclsky, J. & Fineberg, J. The near-tip fields of fast cracks. *Science***327**, 1359–1363 (2010).
20. Kundu, S., Fowler, M., Simon, L. & Grot, S. Morphological features (defects) in fuel cell membrane electrode assemblies. *Journal of Power Sources***157**, 650–656 (2006).
21. Guo, Q. & Qi, Z. Effect of freeze-thaw cycles on the properties and performance of membrane-electrode assemblies. *Journal of Power Sources***160**, 1269–1274 (2006).
22. Kim, S. & Mench, M. Physical degradation of membrane electrode assemblies undergoing freeze/thaw cycling: Micro-structure effects. *Journal of Power Sources***174**, 206–220 (2007).
23. Gouws, S. Voltammetric Characterization Methods for the PEM Evaluation of Catalysts. Ch. 5 (INTECH Open Access Publisher, 2012).
24. Yuan, X., Wang, H., Sun, J. C. & Zhang, J. AC impedance technique in PEM fuel cell diagnosis—A review. *International Journal of Hydrogen Energy***32**, 4365–4380 (2007).
Acknowledgements
This work was supported by the Global Frontier R&D Program of the Center for Multiscale Energy Systems funded by the National Research Foundation under the Ministry of Education, Science and Technology, Korea (2011-0031561) and the Institute for Basic Science (IBS) in Korea (IBS-R006-G1). Y.-H. C. acknowledges the financial support from the Basic Science Research Program (2013R1A1A2061636) through the National Research Foundation of Korea (NRF), which is funded by the Ministry of Education.

Author Contributions
S.M.K. and C.-Y.A. contributed equally to this work. M.C. and Y.-E.S. led this work. S.M.K., C.-Y.A., S.K., W.H., S.J., S.S. and G.L. performed experiments. All authors discussed the data and analysis. S.M.K., C.-Y.A., Y.-E.S. and M. C. wrote the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Kim, S. M. et al. High-performance Fuel Cell with Stretched Catalyst-Coated Membrane: One-step Formation of Cracked Electrode. Sci. Rep. 6, 26503; doi: 10.1038/srep26503 (2016).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/