Modification of gas diffusion layers properties to improve water management

Martin Tomas¹ ∙ Indro S. Biswas² ∙ Pawel Gazdzicki² ∙ Lucie Kullova¹ ∙ Mathias Schulze²

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Abstract In this paper we report an approach to improve water management of commercial GDLs by introducing hydrophobicity patterns. Specifically, line and grid patterns have been created in the MPL side by laser radiation. For an in-depth investigation of these modified GDLs the current density distribution was monitored during fuel cell operation. Additionally, the physical properties of these materials were investigated by a number of ex situ methods such as Fourier transform infrared microscopy, electrochemical impedance spectroscopy and water vapor sorption. Furthermore, a comparison of the physical properties of the patterned GDLs with chemically modified GDLs (treated in H₂SO₄ and H₂O₂) is provided. Our results show a clearly improved homogeneity of current density distribution of the patterned GDLs compared to untreated GDLs. This observation is likely due to a reduced local hydrophobicity which facilitates water diffusion along the flow field of the fuel cell. However, performance of the fuel cell was not affected by the MPL irradiation.

Martin Tomas
mtomas@ntc.zcu.cz

1 University of West Bohemia, Univerzitní 8, 30614 Pilsen, Czech Republic
2 German Aerospace Center (DLR), Institute of Engineering Thermodynamics, Pfaffenwaldring 38-40, 70569 Stuttgart, Germany
Graphical Abstract

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Introduction

In a polymer electrolyte membrane fuel cell (PEMFC) the gas diffusion layer (GDL) is used for proper distribution of reactants to the catalyst layer and for the water management within the membrane electrode assembly (MEA) making the GDL a key component responsible for appropriate and continuous fuel cell operation [1, 2]. The GDL is typically divided into two parts—the macroscopically porous GDL backing and the microporous layer (MPL) [3]. Both parts are treated with poly(tetrafluoroethylene) (PTFE) to obtain specific hydrophobicity needed for fuel cell operation. Flooding of the active catalyst layer (CL), where the reactions take place [4] is avoided by transporting the water away from the reaction layer due to the hydrophobic behavior of GDL [5, 6]. Water produced inside the fuel cell can flood the pores of GDL and decrease the gas permeability of the layer promoting fuel starvation [7–9]. Moreover, excess water can drag the platinum particles from the CL, so that the electrochemical processes efficiency is lowered [10]. On the other hand, a drying out of the membrane has to be avoided to assure adequate ionic conductivity [11]. Furthermore, the MPL provides an effective current collection and a smooth, continuous interface between GDL and CL [12] reducing the ohmic losses in the fuel cell and enhancing chemical and mechanical stability of the CL and the membrane [1].

Within this paper we address the possibility of improving GDL properties by patterning it via laser radiation, i.e., by locally burning PTFE. The main idea of this modification was the fact that water management can be positively affected by varying the PTFE content in the GDL [11, 13]. In recent studies the non-homogeneous distribution of PTFE was investigated [11, 14, 15]. It was concluded that the appropriate ratio of hydrophobic and hydrophilic regions can improve the water management in a fuel cell [15].

We have also considered a treatment of the samples with sulfuric acid or hydrogen peroxide to affect the PTFE coating of carbon fibers [16] leading to a severe loss of hydrophobicity as reported by Borup et al. [17]. Then we have compared laser patterned GDLs with chemically treated samples to evaluate the differences between these two methods of the PTFE content modification.

Experimental

Sample preparation

To prepare a hydrophilic laser pattern, pristine GDLs (SGL Carbon Sigracet 24BC) were exposed to laser of 350 mW power (green laser, 532 nm). Simple patterns (parallel lines or grid) of burned channels were created by moving the laser with about 1 cm s\(^{-1}\) over the sample, see Fig. 1. Due to its more homogeneous structure in this study only the MPL side of the GDL was laser treated. For the assembly of the treated GDLs in a fuel cell the single channel flow field was oriented perpendicular to the pattern.

For chemical modification the GDL samples were exposed to 1 M hydrogen peroxide or 1 M sulfuric acid solutions for 24 h in 80 °C [16, 18]. These modified
samples were immersed in deionized (DI) water to remove the residues of solution. After that the samples were dried in a temperature chamber for 2 h at 80 °C.

**Fuel cell test conditions**

To test the pristine and patterned GDLs in fuel cell operation we used a 5 × 5 cm² cell with a single serpentine flow field. The cell was operated in galvanostatic mode with H₂ and air in co-flow configuration. The cell operation temperature was 80 °C. Above 0.3 A cm⁻² the H₂ stoichiometry and oxygen stoichiometry was set to 1.5 and 2.0, respectively. The relative humidity (RH) was controlled by the bubbler temperature and was set to 50%. The gas outlet absolute pressure was 1.5 bar.

A DLR home-made segmented board with 7 × 7 segments and 5 × 5 cm² was used for current density distribution monitoring [19, 20]. The board was installed at the anode side. In Fig. 1 the scheme of the flow field with the indicated segments is plotted.

The segmented board consists of current collector segments placed on the surface of an epoxy-glass resin matrix. For the local current measurement a resistor array is integrated in the printed circuit board. The gas supply is mounted from the back side of the measuring board. The back layer of the board is connected by conducting wires with the front (flow field) which is divided into segmented cells to avoid lateral conduction along the flow field. The current and voltage signals from the segmented board are measured with a multiplexer and a digital multimeter; the data acquisition system of the segmented cell is independent of the control system of the fuel cell test bench. The width of the channel and the land was 1.1 and 0.8 mm, respectively. The inlet is at segment G1, whereas segment A7 is the location of the gas outlet (see Fig. 1).

**Ex situ techniques**

For local Fourier transform infrared (FTIR) analysis of PTFE in the laser patterned MPL a Bruker Vertex 80v
FTIR spectrometer coupled with a Bruker Hyperion 3000 microscope was used. A typical FTIR microscopy image \[18\] consists of an array of numerous measuring points each measured in the attenuated total reflection (ATR) mode using a conical Ge crystal (diameter of the tip is 100 μm) that is pressed onto the sample for good mechanical contact, i.e., high signal to noise ratio. For high lateral resolution (1 μm) each measuring position \((32 \times 32 \, \mu m^2)\) was detected with a liquid nitrogen cooled mercury–cadmium–telluride focal plane array (FPA) detector consisting of 64 \times 64\ pixels. Typical spectra were recorded with \(2 \times 2\) binning \((32 \times 32\) superpixels) in the range of 600–4000 cm\(^{-1}\) with 4 cm\(^{-1}\) resolution and consist of typically 32 accumulated scans.

For detailed investigation of surface structure and morphology, the samples were scanned with a Jeol JCM-5000 electron microscope.

The electrical properties were studied with electrochemical impedance spectroscopy (EIS) using a Solartron 1287A/1260 facility. This method can be used to determine the dependency of electrical impedance on frequency of the applied electrical field. To ensure a proper contact between the sample and the electrode a special sample holder was developed and liquid electrodes (mercury) were used.

Through-plane conductivity of pristine, chemically modified and irradiated GDLs was studied in frequency range from 0.1 Hz to 20 MHz at ambient conditions. Excitation amplitude was set to 10 mV.

The thermal stability of the samples was investigated in the temperature range from 25 to 1000 °C with a thermogravimetry analyzer (MTGA, Q500) in a nitrogen atmosphere; the heating rate and the nitrogen flow were set to \(10 \, ^\circ C \, min^{-1}\) and 40 ml min\(^{-1}\), respectively. During the temperature increase, it is possible to detect the thermal decomposition of materials and the content of inert residues. Thermal stability measurements of GDLs provide information which can be relevant for MEA preparation processes such as hot pressing \[21\].

The absorption and desorption processes related to water uptake were studied with dynamic gravimetric water sorption analyzer (DVS ET, Surface Measurement Systems). During this measurement the mass of the sample is monitored and the humidity in the measuring cell is dynamically regulated. The difference in absorption and desorption processes is represented by sorption isotherm and its hysteresis.

**Results and discussion**

In this section first the characterization of the laser patterned GDLs by FTIR microscopy and scanning electron microscopy (SEM) is discussed. Secondly, the influence of the laser patterning on fuel cell performance is analyzed based on the local current density distribution. To better understand the behavior of the laser patterned layers the physical properties of these samples are compared with those of chemically modified GDLs.

By means of laser radiation simple patterns of straight lines (pattern 1) and grid (pattern 2) were created on the MPL surface (see Fig. 1). The distance between lines was approximately 2–3 mm and the line thickness was about 0.5 mm. The patterns were created by horizontal motion of the laser beam. The effect of the irradiation on the MPL with a laser beam is basically identical with a highly local heating of the material. This leads to burning, carbonization and/or evaporation of the surface components. It has been observed, that irradiation of the microporous layer media with 350 mW at 532 nm almost instantly burns holes or trenches into the material.

For local FTIR imaging of PTFE in the MPL the integrals of the C–F\(_2\) stretch vibrations were analyzed in the spectral range 1100–1300 cm\(^{-1}\), resembling the presence of PTFE \[18\]. Typical FTIR spectra of a non-lasered (high C–F intensity) and a lasered (low C–F intensity) region of the MPL are provided in Fig. 2a.

Figure 2b, c show, respectively, the FTIR microscopy image and a corresponding line scan of the MPL side (the signal corresponds to peak areas of the CF\(_2\) stretch vibrations in the range 1100–1300 cm\(^{-1}\)), which has been locally irradiated (two parallel lines corresponding to the blue color in the spectroscopic image) according to the procedure described above. Evidently within the burned lines the C–F\(_2\) vibrational intensity is virtually zero, while it is high outside the lines.

In this context it is noted that in some measurements an increased abundance of PTFE is observed between two lines compared to a pristine sample (Fig. 3). It can be speculated that parts of the evaporated or burned PTFE has been redeposited during the laser treatment.

The surface of the irradiated GDLs was scanned with the electron microscope (Fig. 2) whereby two imaging methods were used. In Fig. 2d the imaging mode based on the detection of secondary electrons is used. The burned lines are clearly visible as dark regions. However, the GDL backing is not recognizable: this fact suggests that only a small surface region of the MPL is affected. The same sample was scanned using the method based on backscattered electrons detection (Fig. 2e). This method provides additional information about the GDL surface. Obtained image shows the superficial deterioration of the MPL caused by laser irradiation.

For the performance tests the two patterns were prepared out of an MPL for 25 cm\(^2\) single cell and were mounted on the cathode side. The line patterned specimen was mounted in the test cell with the grafted trenches orthogonally to the
overall flow field channels; for the grid pattern, the mounting direction was irrelevant. To assemble the MEA a commercial catalyst coated membrane (Johnson Matthey) was used. The cell was equipped with a DLR home-made segmented board to monitor the current density distribution.

At the beginning of the tests, performance curves of MEAs containing either the pristine GDL or the modified GDLs were measured. As observed in Fig. 4b, the performance of the fuel cell is not significantly affected by laser irradiation of GDLs under the chosen experimental conditions.

However, the cells with the different GDLs exhibit substantial differences regarding current density distributions. In Fig. 4a current density distributions recorded at different loads using pristine and patterned MPLs are shown. In the provided test the humidification of gases was 50% RH. Evidently, in the case of the pristine MPL the maximum current density is located clearly in the gas inlet region. At the cell outlet the performance is worst. The

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Fig. 2  
(a) FTIR spectra of the C–F stretch of laser burned and pristine MPLs.  
(b) FTIR image of the MPL locally treated with laser radiation.  
The color scale corresponds to integrated C–F intensities.  
(c) FTIR line scan over the two laser burned lines from panel b.  
(d) SEM images of partially irradiated MPL recorded using secondary electrons.  
(e) SEM images of partially irradiated MPL recorded using backscattered electrons.
homogeneity of the distribution, represented by the value $a$ (percentage of values that are within ±20% of average), is around $a = 60–80\%$.

On the other hand, the current density distributions measured with the patterned MPLs show a maximum shifted to the centre of the cell with a significantly improved homogeneity with $a = 90–100\%$.

The improved homogeneity of the current density distribution along the flow field is clearly related to a decreased local hydrophobicity [11, 22]. Thereby, both tested patterns behave similarly suggesting that in the used co-flow configuration only a small change of MPL’s hydrophobicity is sufficient to obtain significantly more homogeneous current density distribution. In this context, it may be stated that a more homogeneous current density distributions would likely extend the lifetime of the fuel cell, as mentioned in [23, 24].

The electrical properties of the samples were measured using EIS. Thereby, an increase of electrical conductivity was observed probably due to the loss of the electrically insulating PTFE [27] during the immersion of GDLs in solutions. Experiments related to GDL resistivity were carried out using mercury electrodes to obtain renewable, uniform and smooth interface between the GDL and the electrode. The experimental setup allows the through-plane measurement under ambient pressure. The electrical properties of GDL depend on applied pressure and the resistivity can radically decrease during the compression in fuel cell, because the pores in the GDL without applied pressure are filled with air [28, 29]. Our method can be used to distinguish the samples with various content of PTFE (see Fig. 5a). Moreover, liquid electrode assures lower contact resistance compared to solid electrode. The electrical conductivity was calculated using the approximation (derived from [30]).

$$\sigma = \frac{h}{Rr^2};$$  

where $h$ is the thickness of the GDL, $R$ is the resistivity and $r$ is the radius of the mercury electrode (this formula is valid for $r \gg h$). As shown in Fig. 5a the electrical conductivity significantly increases in case of samples affected by sulfuric acid and hydrogen peroxide solutions. The surface of these samples is meant to be more influenced than the surface of irradiated samples. This fact is supported by other results obtained using FTIR and TGA methods.

The results of thermal stability measurement show significantly faster decomposition of chemically modified GDLs (Fig. 5b) compared to pristine and lasered samples. The steep decrease of sample mass between 500 and 600 °C is related to PTFE decomposition [31, 32]. Rapid mass loss above 600 °C can be caused by structural deterioration of GDL or change of functionalization of the carbon fiber surface (formation of the carboxyl groups). Solutions used for chemical modification may influence the carbon fibers and the diffusion layer becomes less stable during the temperature increase. Diluted solutions can oxidize carbon fibers [16] and this corrosion process results in formation of substances such as phenol and carboxyl groups [17].

The samples were also investigated with respect to water vapor sorption and desorption (see Fig. 5c). This measurement can be used for the determination of a volume density of
hydrophobic regions. Water vapor is stored in the hydrophilic regions and the increase of the sample mass can be detected. Thereby the amount of sorbed water is represented by the hysteresis. In case of pristine GDL the volume density of hydrophobic regions is larger than in case of chemically modified or irradiated GDLs leading to a narrow hysteresis. Similarly, the sample treated by sulfuric acid shows a small hysteresis suggesting virtually no impact of this treatment on the hydrophobicity. In contrast, hydrogen peroxide affects the PTFE coating leading to a severe volume density loss of the hydrophobic regions and a large hysteresis. A significant increase of sample mass was measured for irradiated samples pointing at their reduced hydrophobicity compared to other samples and supporting FTIR analysis where a loss of hydrophobic PTFE was clearly observed. Since the lasered area is larger in pattern 2 than in pattern 1, its hysteresis is larger accordingly (see Fig. 5c). The possible explanation is that the partly burned MPL strongly affected a water uptake of irradiated GDL and the water diffusion was facilitated [11, 33].

**Conclusion**

The GDL Sigracet 24BC was irradiated with laser beam and simple patterns of burned lines were created. The thermal, electrical and sorption properties were studied for pristine chemically modified and laser irradiated samples.
The performance test of MEA containing irradiated GDL was conducted and the current density distribution was investigated. Under the operation conditions used, the results show significantly more homogeneous current density distribution when patterned GDLs were used compared to pristine GDLs. This is likely due to an improvement of the water management changing local hydrophobicity leading to a facilitation of water diffusion and removal along the flow field. An influence of the laser irradiation on the performance of the fuel cell was not observed.

For detailed analysis of the physical properties of the patterned GDLs, further GDLs were modified by sulfuric acid and hydrogen peroxide for comparison. Electrical conductivity of chemically modified samples increased with respect to PTFE content decrease. Water sorption of the sample exposed to hydrogen peroxide increased due to the loss of the density of hydrophobic regions (the impact of sulfuric acid was weak). The thermal stability measurement of the chemically modified GDLs showed rapid decrease of sample mass during heating.

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