Testing fuel cell catalysts under more realistic reaction conditions: accelerated stress tests in a gas diffusion electrode setup

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Abstract

Gas diffusion electrodes (GDE) setups have very recently received increasing attention...
as a fast and straight-forward tool for testing the oxygen reduction reaction (ORR) activity of surface area proton exchange membrane fuel cell (PEMFC) catalysts under more realistic reaction conditions. In the work presented here, we demonstrate that our recently introduced GDE setup is suitable for benchmarking the stability of PEMFC catalysts as well. Based on the obtained results, it is argued that the GDE setup offers inherent advantages for accelerated degradation tests (ADT) over classical three electrode setups using liquid electrolyte. Instead of the solid-liquid electrolyte interface in classical electrochemical cells, in the GDE setup a realistic three-phase boundary of (humidified) reactant gas, proton exchange polymer (e.g. Nafion) and the electrocatalyst is formed. Therefore, the GDE setup not only allows accurate potential control but also independent control over the reactant atmosphere, humidity and temperature. In addition, the identical location transmission electron microscopy (IL-TEM) technique can easily be adopted into the setup enabling a combination of benchmarking with mechanistic studies.

**Keywords**

Accelerated stress tests; Gas Diffusion Electrode; Platinum; Carbon Supported High Surface Area Catalysts; Fuel Cells;
1. Introduction

Benchmarking activity and stability of proton exchange membrane fuel cell (PEMFC) catalysts using classical three-electrode setups and aqueous electrolyte solution is a popular approach [1, 2]. The electrode potential can be controlled independent of the reaction conditions allowing not only to compare the performance of different PEMFC catalysts, but also to investigate how potential excursions affect the catalyst stability and the degradation mechanism. For the latter, accelerated degradation tests (ADTs) are employed in combination with techniques such as identical location transmission and scanning electron microscopy (IL-TEM and IL-SEM) [3-12] and scanning flow cells (SFC) coupled to inductively coupled plasma - mass spectrometry (ICP-MS) [13, 14]. Thus, degradation mechanisms such as metal dissolution and particle detachment can be related to certain excursions in the electrode potential as well as catalyst properties such as particle size, carbon support, etc. The aim of ADTs thereby is to apply realistic conditions and at the same time to reduce the time in which a performance loss is observed significantly [15-17]. Although this is a contradictory goal, certain ADT protocols are commonly accepted as a compromise, in particular the one of the Fuel Cell Commercialization Conference of Japan (FCCJ) [18, 19] simulating load-cycle and start-up/shutdown conditions of a proton exchange membrane fuel cell (PEMFC).
Despite the achievements made, the current approach also has disadvantages. For example, it is not clear if and how an aqueous electrolyte environment as compared to a realistic three-phase boundary of humidified reactant gas (proton exchange polymer, e.g. Nafion) influences the degradation of the active component and the carbon support, respectively. In aqueous electrolyte environment it is found that for certain Pt/C fuel cell catalysts and simulated start-up/shutdown conditions the degradation mechanism is restricted to particle loss, most likely due to carbon corrosion [3]. However, at simulated load cycle conditions the main observed degradation mechanism changes to particle migration and coalescence [20]. By comparison, in membrane electrode assembly (MEA) tests the reported degradation mechanism is in general platinum dissolution [21], and in some cases extensive carbon corrosion. More subtle mechanisms such as particle migration and coalescence cannot be identified.

Only few efforts exist to combine a simple setup with a more realistic environment [22-29]. Here we demonstrate how our recently introduced gas diffusion electrode (GDE) setup [30-32] can be used for benchmarking the stability of high surface PEMFC catalysts under realistic reaction conditions. Gas diffusion electrode setups have very recently received increasing attention allowing to combine the ease of use known from electrochemical three-electrode setups and liquid aqueous environment with reaction
conditions closer to applications [33]. In the work presented, it is shown that in the GDE setup parameters such as the reactant gas atmosphere and its humidity can be easily controlled and their influence on the catalyst stability be studied. Thus, the setup offers inherent advantages over the conventional three-electrode setups for ADTs. In addition, the IL-TEM technique can easily be adopted into the setup enabling a combination of stability benchmarking with mechanistic studies.

2. Experimental

2.1. Chemicals, materials, and gases.

Ultrapure water (resistivity > 18.2 M\(\Omega\) cm, total organic carbon (TOC) < 5 ppb) from a Milli-Q system (Millipore) was used for acid/base dilutions, catalyst ink formulation, and the GDE cell cleaning. The following chemicals were used in ink formulation and electrolyte preparation: Isopropanol (IPA, 99.7+ %, Alfa Aesar), 70 % perchloric acid (HClO4, suprapure, Merck), potassium hydroxide hydrate (KOH·H2O, Suprapur, Merck), commercial Pt/C catalysts (46.0 wt. % TEC10E50E and 50.6 wt. % TEC10E50E-HT, Tanaka kikinzoku kogyo as well as HISPEC 3000, Johnson Matthey), and Nafion dispersion (D1021, 10 wt. %, EW 1100, Fuel Cell Store). A Nafion membrane (Nafion 117, 183 \(\mu\)m thick, Fuel Cell Store) and a gas diffusion layer (GDL) with a microporous layer (MPL) (Sigracet 39BC, 325 \(\mu\)m thick, Fuel Cell Store) were
employed in the GDE cell measurements. Before use, the Nafion membrane was prepared and activated as follows: after cutting several circles with a diameter of 2 cm from a sheet of Nafion membrane, the membranes were treated for 30 minutes at 80 °C in 5 wt.% H₂O₂, followed by rinsing with Milli-Q water. Then, the membranes were treated for 30 minutes at 80 °C in Milli-Q water followed by rinsing with Milli-Q water. Finally, the membrane was treated for 30 minutes at 80 °C in 8 wt.% H₂SO₄, again followed by a rinsing with Milli-Q water. All membranes were kept in a glass vial filled with Milli-Q water. The following gases from Air Liquide were used in electrochemical measurements: Ar (99.999 %), O₂ (99.999 %), and CO (99.97 %). TEM characterization of the respective Pt/C catalysts can be found in the supporting information (Figure S1).

2.2. Gas Diffusion Electrode cell setup.

An in-house developed GDE cell setup was employed in all electrochemical measurements that was initially designed for measurements in hot phosphoric acid [24]. The design used in the present study has been described before [31]. In short, it was optimized to low temperature PEMFC conditions (< 100 °C) by placing a Nafion membrane between catalyst layer and liquid electrolyte; no liquid electrolyte is in direct contact with the catalyst [31]. A photo of the parts of the improved GDE setup is shown in Figure 1.
The cell body above the Nafion membrane is made of polytetrafluoroethylene (PTFE). A platinum mesh and a reversible hydrogen electrode (RHE) were used as a counter electrode (CE) and a reference electrode (RE), respectively. The counter electrode placed inside of a glass capillary tube with a glass frit on the bottom, which avoids the trapping of gas bubbles in the hole of the Teflon cell and consequently helps increasing the reproducibility of the measurement. All potentials in this study are referred to the RHE potential. For initial cleaning, the Teflon upper part was soaked in mixed acid (H$_2$SO$_4$:HNO$_3$ = 1:1, v:v) overnight. Subsequently it was rinsed thoroughly by ultrapure water, and boiled in ultrapure water twice. Between the measurements the Teflon upper part was boiled in ultrapure water twice.

**Figure 1.** Photo of the individual components of the GDE cell employed in the present study (left) and different applied ADT protocols (right). The Nafion membrane and the GDL (including the catalyst layer fabricated on it) are sandwiched between the upper and lower cell body. The upper cell body is pressed to the lower cell body by a metal.
ring and a clamp. A gas humidifier (gas bubbler) is connected to the reactant gas. The counter electrode is inserted into a glass capillary to avoid gas capture at the Teflon cell. Optionally, instead of applying a catalyst layer on the GDL an Au-TEM grid with a thin catalyst layer can be sandwiched between GDL and Nafion membrane (not shown). The ADT protocols are simulating (i) load cycles, (ii) start-up/shutdown conditions, and (iii) a combination of both.

2.3. **Catalyst ink formulation and catalyst layer application.**

2.0 mg of the respective commercial 46.5 wt. % Pt/C catalyst powder and 10.09 µL (or 9.36 µL) of the 10 wt. % Nafion dispersion were mixed with 4.74 mL (or 5.15 mL) of IPA:ultrapure water (1:3, v:v) mixed solvent. The glass vial containing the mixture was placed in the ultrasonic bath and sonicated for 15 minutes. Subsequently the mixture was dispersed using a horn sonicator (Q500, QSonica) for 1 minute. Before the application of the catalyst ink on the GDL, circular pieces (φ = 20 mm) of the GDL were punched from a larger sheet. The catalyst ink was sprayed onto the GDL (MPL coated side) by using a pump (Harvard apparatus 11 plus) with a flow rate of 0.5 mL/min and an ultrasonic spray nozzle (Sonozap narrow spray atomizer). During the spraying, the catalyst ink was constantly sonicated by an ultra sonicator (Sonozap ultrasonic atomizer) with a power of 1.5 W and frequency of 130 KHz and the GDL was heated on a heating plate (140 °C) and covered with an iron mask so that a circular catalyst layer (φ = 3 mm) was formed at the center of the GDL.
2.4. Electrochemical measurements.

The electrochemical measurements were performed using a computer controlled potentiostat (ECi 200, Nordic Electrochemistry). The measurements were performed with 4 M HClO₄ aqueous solution in the upper Teflon compartment of the GDE setup applying different temperatures as reported previously [31]. Prior to the measurements, the electrode was purged from the backside (through the gas diffusion layer) with Ar gas and the catalyst was cleaned by potential cycles between 0.06 and 1.10 V<sub>RHE</sub> at a scan rate of 500 mV s⁻¹ until a stable cyclic voltammogram could be observed (ca. 50 cycles). The resistance between the working and reference electrode (~10 Ω) was compensated to effective value of around 1 Ω by using analog positive feedback scheme of the potentiostat. The resistance was determined online using an AC signal (5 kHz, 5 mV) [34].

To determine the stability of the Pt/C electrocatalysts, ADTs inspired by the Fuel Cell Commercialization Conference of Japan [18] were performed. Three different ADT protocols were tested: (i) a protocol simulating load-cycles, where the electrode potential was modulated with a square wave and stepped between 0.6 and 1.0 V<sub>RHE</sub> with a holding time of 3 s at each voltage for a total of 9000 potential cycles, (ii) a protocol simulating start-up/shutdown conditions [35], where the electrode potential was cycled
with a scan rate of 0.5 V s\(^{-1}\) between 1.0 and 1.5 V\(\right\)RHE for a total of 9000 potential cycles and (iii) a mixed protocol combining (i) and (ii), which consisted of 500 potential cycles of the load-cycling protocol followed by 10 potential cycles of the start-up/shutdown protocol, repeated overall 18 times; in the following we refer to this protocol as mixed protocol. In order to monitor the H\(_{\text{upd}}\) area and the change of the quinone/hydroquinone (Q/HQ) redox peak [36, 37] after each 1000 potential cycles (500 cycles for procedure (iii)) six cyclic voltammograms were recorded in Ar atmosphere at a scan rate of 0.5 V s\(^{-1}\). The electrochemically active surface area (ECSA) of the catalyst was determined by conducting CO stripping voltammetry before and after the ADT. We did not record further CO stripping during the ADT to minimize the potential negative impact of CO stripping on the surface area [38-40]. For the CO stripping measurements, the working electrode was held at 0.05 V while purging first CO through the GDL for 5 minutes and thereafter Ar for an additional 5 minutes. The ECSA was determined from the CO (Q\(_{\text{CO}}\)) oxidation charge recorded at a scan rate of 50 mV s\(^{-1}\). Unless otherwise stated, all measurements were performed at room temperature.

2.5. TEM grid preparation.

For IL-TEM investigations, a gold finder grid (400 mesh; Plano, Germany) was used. The catalyst ink was diluted by a factor of 1:10. Afterwards, 10 \(\mu\)L of diluted catalyst
ink were pipetted onto the gold finder grid. In order to avoid overlapping of the catalyst particles, after approximately 10 s the droplet of the catalyst was carefully absorbed off from the grid with a filter paper. Afterwards the grid was dried, and the catalyst investigated in a Technai Spirit (FEI) with an accelerating voltage of 80 kV before and after accelerating degradation test. The TEM grid was placed between a Nafion membrane and a GDL (without a catalyst layer on it) during the ADT. The ADT for the IL-TEM experiment was conducted by applying the load-cycling protocol (1200 cycles) at 60 °C, 100 % relative humidity (RH).

3. Results and discussion

3.1. Comparison of different ADT protocols.

The aim of the present study was to employ our recently developed GDE setup mimicking PEMFC environment to ADT studies. At first, we compared different ADT protocols for their suitability to monitor the catalyst degradation in the GDE setup. The different ADT protocols are designed to apply as realistic conditions as possible and at the same time to significantly reduce the time in which a performance loss is observed. For PEMFCs a variety of different ADT protocols can be found in literature, here we tested the popular approach suggested by the Fuel Cell Commercialization Conference
of Japan [18, 19] simulating load-cycle and start-up/shutdown conditions, respectively. Load-cycle conditions are simulated by stepping the applied electrode potential between 0.6 and 1.0 V_{RHE} and holding the potential for three seconds each. 0.6 V_{RHE} is a typical cell potential under full load, whereas 1.0 V_{RHE} is close to the open circuit potential of Pt and represents the cell voltage at idle conditions. Start-up/shutdown conditions are simulated by scanning the applied electrode potential between 1.0 and 1.5 V_{RHE} with a scan rate of 0.5 V s^{-1}, which simulates an increase in cathode potential when introducing hydrogen in an air filled flowfield or replacing the hydrogen by air during the start-up and shut-down period, respectively [35]. For a better understanding of the ADT protocols the reader is referred to ref. [18]. The total duration of the load-cycle treatment lasted 9000 cycles, which is a total of 16 and 19 hours in Ar and O_2 atmosphere, respectively. The degradation is monitored by cyclic voltammograms recorded in Ar atmosphere after each 1000 cycles as well as CO stripping experiments before and after of the ADT protocol (therefore, the total time varies as for the ADT protocol measured in Ar and O_2, respectively, as the O_2 needed to be replaced from the flow field before measuring the cyclic voltammograms). The start-up/shutdown treatment lasted 9000 cycles as well, which is a total of 6 and 9 hours in Ar and O_2 atmosphere, respectively.
In Figure 2 the effect of both ADT protocols employed in the GDE setup is demonstrated on a commercial Pt/C catalyst with an average particle size of 2-3 nm, see FigureS1. On the left-hand side, the cyclic voltammograms (CVs) recorded every 1000 cycles in Ar atmosphere are plotted, whereas on the right-hand side the CO stripping curves recorded before and after the ADT treatment are shown. For Pt/C, the ECSA loss can be monitored based on the changes in $H_{\text{upd}}$ area in the CVs recorded in Ar atmosphere as well as changes in the area under the CO stripping curves. Further information concerning the degradation of the catalyst can be derived from the change in the “oxide region” ($\sim 0.75 \, V_{\text{RHE}}$) and the formation of a Q/HQ redox peak ($\sim 0.6 \, V_{\text{RHE}}$). As seen in Figure 2 a, b), the behavior of the Pt/C catalyst upon the load cycling treatment (i) is qualitatively the same as reported previously for ADT measurements performed in half-cells with liquid electrolyte [37]. The Pt/C catalyst loses continuously ECSA and the reduction peak in the “oxide region” decreases, whereas its peak potential slightly shifts to higher potentials. After 9000 cycles a total loss in ECSA of $41.5 \pm 2.3 \%$ is determined. Interestingly this is roughly 10 % more loss than what is observed for the same catalyst in a conventional cell with liquid electrolyte ($32.0 \pm 1.6$), see Table 1 and FigureS2, 3 for a summary of the ECSA losses observed for different catalysts.
Figure 2. Influence of different ADT protocols performed in Ar atmosphere on the electrochemical behavior of a commercial Pt/C catalyst (TEC10E50E). On the left-hand
side, the cyclic voltammograms (sweep rate 0.5 V s\(^{-1}\)) recorded every 1000 cycles in Ar atmosphere are shown, whereas on the right-hand side the initial and final CO stripping curves are shown (solid line) together with the subsequent cyclic voltammogram (solid line); both recorded at a sweep rate of 0.05 V s\(^{-1}\). (a,b) Step protocol simulating load cycles, (c,d) CV protocol simulating start-up/shutdown conditions, (e,f) mixed protocol combining the two prior protocols. The measurements were performed at room temperature.

In contrast to protocol (i), the effect of applying the start-up/shutdown protocol (ii) in the GDE setup is more difficult to compare with previous measurements using conventional half-cells and liquid electrolyte [20, 37, 41]. When applying protocol (ii) in the GDE setup, see Figure 2 c, d), the CVs recorded in Ar atmosphere quickly lose the features of Pt/C and become significantly tilted with time. This indicates massive carbon corrosion and an increase in cell resistance with time, which is confirmed by the online superposition of an AC signal (5 kHz, 5 mV). The same trend is seen in the CO stripping curve recorded after the ADT procedure. Due to the significant distortion of the CVs and CO stripping measurements a proper analysis of the surface area loss from determining the \(H_{\text{upd}}\) or the CO stripping charge is highly questionable. Most likely the prolonged exposition to high electrode potentials leads to the massive oxidation not only of the catalyst layer, but also the MPL and possibly even the GDL. Note that in our measurements the amount of carbon in the catalyst layer is significantly less than the carbon in the MPL.
A more representative degradation of the catalyst is achieved by the presented mixed ADT protocol (iii), which combines load cycles with start-up/shutdown cycles in a sequence of load cycles and start-up/shutdown events, Figure 2e, f). Although only a limited number of high potential excursions are applied, the evolving features in the CV recorded every 1000 cycles are distinctively different than in the load-cycle protocol (i): i.e., the CVs exhibit a strong increase in double layer capacitance and the formation of a pronounced Q/HQ redox peak is observed. Although also for this ADT, it cannot be excluded that the observed features are related to the degradation of the catalyst layer as well as the MPL and the GDL, the loss in CO stripping area is solely related to the loss in ECSA of the catalyst. In contrast to the commonly applied start-up/shutdown protocol (ii), the measurements from the mixed ADT protocol can be analyzed with confidence and thus we suggest using such a mixed procedure instead of the protocol (ii) proposed by the FCCJ for simulating start-up/shutdown conditions. Despite of the fact that the catalyst experiences only 180 high potential excursions in total, it is seen that in the mixed protocol (iii) the ECSA loss (54.3 ± 0.1 %) is more than 10 % higher than upon applying the load-cycle protocol (i) (41.5 ± 2.3 %), see Table 1.

3.2. The influence of the particle size and heat treatment.
We performed the same measurements (ADT protocol (i) and protocol (iii)), on an additional commercial catalyst (TEC10E50E-HT) with larger particle size, i.e. ca. 4-5 nm instead of 2-3 nm, see FigureS1. As expected and shown previously in ADT using conventional electrochemical cells [42], the results obtained in the GDE setup demonstrate a significantly higher loss in ECSA for the catalyst with smaller particle size (2-3 nm, TEC10E50E) than for the one with larger particle size (4-5 nm, TEC10E50E-HT). Upon applying the load cycling protocol (i) using the Pt/C catalyst with 4-5 nm particle size, the ECSA loss is 19.2 ± 0.5 %, i.e. it is reduced to less than half as compared to the Pt/C with 2-3 nm particle size (ECSA loss of 41.5 ± 0.1 %). Applying the mixed protocol (iii) the stability improvement is even more significant, i.e. an ECSA loss of 20.7 ± 0.4 % is detected as compared to 54.3 ± 0.1 %. An obvious explanation for the observed stability improvement is the larger particle size of the active Pt phase. It should be mentioned however, that the particle size might not the only contributing factor to the enhanced stability. As these are commercial samples, the exact synthesis routes are not publicly available, the catalyst notation however, suggests that the larger particle size results from a heat treatment of the Pt/C catalyst with 2-3 nm particle size. Such treatments result not only in increased particle sizes due to agglomeration and sintering, but also influence the carbon support properties [43]. For a
more detailed study of the influence of the particle size on the catalyst stability, which is
not the scope of this work, therefore, the use of in-house synthesized catalysts using the
toolbox approach [44] is suggested.

3.3. The influence of the reactant gas conditions.

Above discussed ADT measurements were performed under Ar atmosphere. Performing
the same measurements in O$_2$ atmosphere using the Pt/C catalyst with 2-3 nm particle
size, leads to small but distinctive differences, see Figure 3. The startup/shutdown
protocol (ii) faces the same limitations as when performed in Ar atmosphere and is
therefore not discussed further. However, in the CVs of the load-cycle protocol (i), it
can be seen that (as compared to the ADT protocol (i) performed in Ar atmosphere) the
formation of the Q/HQ redox peak is more pronounced indicating an impact of the O$_2$
atmosphere on the carbon support oxidation. This is confirmed in the CO stripping
measurements, where a small, but distinctive difference in ECSA loss is seen whether
the ADT protocol is performed in Ar (41.5 ± 2.3 %) or O$_2$ (48.1 ± 1.6 %) atmosphere.
Interestingly the difference in ECSA loss between measurements performed in Ar and
O$_2$ atmosphere is less pronounced applying the mixed ADT protocol (iii). This finding
might be related to the fact that during the high potential excursions the catalyst
experiences enhanced oxidation, independent of the reacting atmosphere. However,
applying the same procedures (i) and (iii) with the Pt/C catalyst with ca. 4-5 nm particle
size (TEC10E50E-HT), no influence of the reaction gas atmosphere on the recorded
ECSA loss is discernable, i.e. 19.2 ± 0.5 % in Ar and 18.2 ± 1.2 % in O₂ gas atmosphere.
The results therefore confirm previous reports stating that the gas atmosphere affects the
surface chemistry of the carbon supports [28, 45] but they also show that the influence
of the gas atmosphere on the ECSA losses is individually dependent on the Pt/C
catalyst.
Figure 3. Influence of different ADT protocols performed in O\textsubscript{2} atmosphere on the electrochemical behavior of a commercial Pt/C catalyst (TEC10E50E). On the left-hand side, the cyclic voltammograms (sweep rate 0.5 V s\textsuperscript{-1}) recorded every 1000 cycles in Ar atmosphere are shown, whereas on the right-hand side the initial and final CO stripping curves (solid line) are shown together with the subsequent cyclic voltammogram (solid line); both recorded at a sweep rate of 0.05 V s\textsuperscript{-1}. (a,b) Step protocol simulating
load-cycles, (c,d) CV protocol simulating start-up/shutdown conditions, (e,f) Mixed protocol combining the two prior protocols. The measurements were performed at room temperature.

Table 1. The total loss in ECSA in % during the different ADT protocols as determined from the CO stripping measurements. The error is the standard deviation of three independent measurements.

| Catalyst            | Cell type  | ADT protocol         | Gas Atmosphere |
|---------------------|------------|----------------------|----------------|
|                     |            |                      | Ar             | O₂              |
| TEC10E50E            | Conventional cell | load-cycle protocol (i) | 32.0 ± 1.6    | 37.2 ± 2.5    |
| TEC10E50E            | GDE cell   | load-cycle protocol (i) | 41.5 ± 2.3    | 48.1 ± 1.6    |
|                     |            | mixed protocol (iii)  | 54.3 ± 0.1    | 57.4 ± 1.8    |
| TEC10E50E-HT         | GDE cell   | load cycle protocol (i) | 19.2 ± 0.5    | 18.2 ± 1.2    |
|                     |            | mixed protocol (iii)  | 20.7 ± 0.4    | 24.0 ± 1.5    |

3.4. Influence of reactant gas humidity.

Another interesting factor for the degradation of PEMFC catalysts is the effect of the relative gas humidity. This factor is not accessible in standard electrochemical three-electrode setups with liquid aqueous electrolyte but can be easily studied in the GDE setup. To demonstrate the effect of the relative gas humidity, we performed a load cycle ADT in dry oxygen gas (protocol (i)). To be able to monitor the real loss in ECSA, before and after the ADT, CO stripping measurements were performed in humidified gas. The results are summarized in Figure 4. It is seen that in dry Argon gas, the $H_{upd}$ region in the CVs – recorded as part of the load cycle protocol (i) – decreases indicating
a significant reduction in active Pt surface area. This sudden loss in active Pt surface area can be correlated to a significant deactivation in dry gas conditions. However, the final CV recorded under re-humidified conditions demonstrates that the active Pt surface area can to a large extend be recovered. That is, after applying the load cycle protocol (i) on the TEC10E50E Pt/C catalyst with a particle size of 2-3 nm, the ECSA loss in dry O2 gas condition was ca. 50 %, whereas switching back to humidified O2 gas, the overall ECSA loss is only ca. 28 %, which is less than applying the load cycle protocol (i) under fully humidified conditions.

The results can be interpreted in the following way. During the load cycle protocol (i), the active Pt surface area is significantly reduced due to a loss in proton conduction from the dry Nafion in the catalyst layer. Thus, only part of the catalyst is actually subjected to degradation while the other part remains unaltered. This results in an overall lower degradation when switching back to fully humidified conditions. Such behavior, however, is not expected in real PEMFC devices where the power output is decisive. In contrast to the potentiostatic ADT measurements, under amperometric conditions inaccessible Pt surface area due to dry gas conditions are expected to lead to increased degradation in the accessible part of the catalyst layer. To demonstrate this, we designed an amperometric ADT protocol that switches 600 times from load (i.e. 100 mA
cm$^2_{\text{geo}}$) to open circuit potential (OCP) and back, holding the respective condition for 3 s each. Comparing the same TEC10E50E Pt/C catalyst under dry and humidified conditions, respectively, a significantly higher ECSA loss was observed for the dry O$_2$ gas (ca. 65 % loss) as compared to the humidified O$_2$ gas (ca. 10 % loss) conditions. In contrast to the potentiodynamic load cycle protocol (i), re-humidifying the catalyst after applying the amperometric ADT protocol, could only recover a small part of the ECSA (total loss of ca. 57 % instead of ca. 65 % loss)

**Figure 4.** Influence of humidification of O$_2$ gas on the ECSA loss of a commercial Pt/C
catalyst (TEC10E50E) upon applying an amperometric ADT consisting of 600 steps between load conditions (100 mA cm$^{-2}_{geo}$) and OCP, maintaining each condition for 3 s each. (a,b) Cyclic voltammograms and CO stripping (sweep rate 0.05 V s$^{-1}$) before and after ADT protocol applied under humidified O$_2$ gas. (c,d) Cyclic voltammograms and CO stripping (sweep rate 0.05 V s$^{-1}$) before and after ADT protocol applied under dry O$_2$ gas. The measurements before applying the ADT protocol were recorded under humidified conditions. In addition, the effect of re-humidifying the O$_2$ gas after applying the ADT protocol is shown. The measurements were performed at room temperature.

3.5. IL-TEM measurements in GDE setup

An advantage of half-cells with liquid electrolyte - as compared to MEA test - is the possibility to perform IL-TEM measurements to analyze the degradation mechanism leading to the loss in active surface area. Here, we demonstrate that the same is feasible in the GDE setup and even elevated temperatures can be used, see Figure 5. By placing the TEM grid between the membrane electrolyte and GDL, the IL-TEM method can be applied straightforward. For the demonstration, a catalyst with lower Pt loading (20 wt. %) was used to facilitate the ability to follow the change of individual particles. The typical degradation phenomena like migration and coalescence (yellow circles) and particle detachment (red circle) can be clearly seen to occur as consequence of the load-cycle treatment.
Figure 5. IL-TEM micrographs of a commercial 20 wt. % Pt/C catalyst (HiSPEC 3000) before and after load-cycle ADT treatment for 1200 cycles at 60 °C.

Conclusion

We demonstrate the application of our recently introduced gas diffusion electrode (GDE) setup for benchmarking the stability of PEMFC catalysts under realistic reaction conditions. It is shown that the popular ADT protocol proposed by the FCCJ to simulate load-cycle conditions leads to suitable results for describing the degradation behavior of the catalyst, whereas the protocol proposed for simulating start-up/shutdown conditions leads to massive corrosion of the carbon in the catalyst layer as well as MPL and GDL, thereby obstructing the ability to track the surface area loss of the catalyst. Therefore, we propose a combination of the two protocols limiting the number of high potential excursion as feasible alternative. Such treatment would also be closer to a realistic
condition, where not exclusively load-cycle or start-up/shutdown conditions are applied, but both cases occur. Using these ADT protocols a significant influence of the particle size on the ECSA loss is observed, although it is acknowledged that the (assumed) heat treatment of the catalyst also changes the carbon support. Furthermore, it is shown that the gas atmosphere (reactant or inert gas as well as dry or humidified gas) influences the degradation behavior. In contrast to previous reports, it is demonstrated that the influence of the presence of oxygen on the degradation depends on the ADT protocol and the catalyst. No systematic difference between liquid and polymer environment was seen, however, it is found that when applying high potential excursions, the influence of reactant gas atmosphere gets less pronounced (for those cases where a clear influence is observed).

More intriguing is the observed influence of the reactant humidity, which cannot be studied in conventional cells using liquid electrolyte. It is found that under dry conditions only parts of the catalyst layer participate in the reaction. In a potentiostatic ADT protocol, almost all of the initial surface area can be re-established by switching back to humidified conditions, i.e. the apparent loss in ECSA is reversible. This is, however, not the case in a more realistic amperometric degradation protocol. This finding highlights that to investigate such effects, additional ADT protocols are needed.
to ensure that the entire catalyst layer is indeed participating the reaction.

Last but not least, it is demonstrated that the IL-TEM technique can easily integrated in the measurements allowing a combination of surface area loss determination and mechanistic insights. This is even possible at 60 °C, i.e. close to the actual operation temperature of a PEMFC. To sum up, it is demonstrated that the introduced GDE setup offers substantial advantages over standard degradation measurements in electrochemical cells employing liquid electrolyte.

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