How platinum oxide affects the degradation analysis of PEM fuel cell cathodes

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HIGHLIGHTS

• New diagnostic routine to determine platinum oxide related voltage losses explicitly

• Oxide related voltages losses depend only on half potential voltage.

• Voltage loss break down of end of test voltage losses into 6 different contributors.

• Unassigned voltage loss is severely underestimated if oxide formation is neglected

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ABSTRACT

In this work, proton exchange membrane fuel cell cathodes are degraded with accelerated stress tests. These PtCo containing cathodes are analyzed at begin of life and end of test with a dedicated diagnostic procedure. For every individual load point, the oxygen trans port resistance and voltage losses due to the formation of platinum oxides were obtained in addition to commonly measured electrochemical surface area, high frequency resistance, as well as cathode ionomer resistance. These data were used to break down the voltage losses into six different contributors. With this break down, performance gains and per formance losses were determined at end of test. At low current densities, it was found that voltage losses due to degradation are dominated by the loss of specific activity and catalyst surface area in line with the state of the art knowledge. But by quantifying the losses from platinum oxide formation explicitly, we show that end of test an unassigned voltage loss is not only present at highest current densities, but already at low current density. More precisely, the unassigned voltage loss shows a linear increase with decreasing half

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Introduction

The proton exchange membrane fuel cell (PEM FC) is a promising technology for automotive drivetrain applica tions [1]. State of the art cathode materials - like the PtCo catalyst on a carbon support - are prone to lifetime limiting degradation under a wide range of relevant oper ation conditions [1,2]. Most importantly, degradation arises from cathode degradation. Cathode degradation itself, today, arises majorly from half cell voltage cycles. These lead on the one hand to the loss of catalyst surface area and on the other hand to support corrosion. Degradation from freezing, over temperatures and contaminates are less important [3-5]. Accordingly, detailed understanding of this degradation under relevant driving conditions must be gained. Only with this insight, hybridization strategies can be developed in order to reach the necessary fuel cell lifetime targets while using the minimum amount of precious metals.

Conventionally, activity losses are believed to be the root cause for performance losses of PEM FCs, especially at high cathode loadings [6–11]. For cathodes containing pure plat inum as catalyst material, these voltage losses are related to the loss of electrochemical surface area and can be well esti mated from the Tafel relation. The electrochemical surface area loss originates from platinum dissolution, Ostwald ripening, particle agglomeration and also from corrosion of the carbon support material. These mechanisms are dis cussed in a vast number of publications, including some comprehensive reviews [10,12–15]. However, cathodes con taining low amounts of Pt alloy catalyst can suffer from four additional degradation phenomena.

The first phenomenon is a reduction of the specific activity, caused by a change of the catalyst morphology due to deal loying [16–18]. For example, Pt based core shell structures were found to loose specific activity in response to an increasing outer shell thickness arising from metal leaching [19–21].

Besides the impact on activity, de alloying causes also the second phenomenon, namely increased Ohmic losses. These increased Ohmic losses originate from cation contamination of the ionomer, blocking its sulfonic head groups. For example, Cai et al. [22] used Co^{2+} doped membranes (sulfonic site blockage up to 46%) to investigate the potential influence of leached out cations on the proton conductivity in the membrane and the cathode catalyst layer. They found increasing high frequency resistance and proton transport resistance with increased cation concentration especially under dry operating conditions. Accordingly, a cation impact on the proton conductivity in membrane electrode assemblies without ex situ membrane doping, especially at low

humidity, is not unexpected [23–25]. However, so far no in crease of high frequency resistance or proton transport resistance in alloy containing membrane electrode assem blies was found if cathode cation leaching is the only reason for ionomer contamination [17].¹

The third phenomenon causing voltage losses in Pt alloy containing PEM-FC-cathodes with low catalyst loadings is an increased oxygen mass transport resistance. This in crease can be induced by changes of the catalyst support (carbon corrosion) as well as the loss of electrochemical surface area (ECSA) [8,17]. Carbon corrosion, on the one hand, can reduce the hydrophobicity of the carbon support, which increases the amount of flooding in the catalyst layer under wet operation conditions. Correspondingly, oxygen transport related voltage losses increase in the presence of liquid water in the cathode electrode [26,27]. On the other hand, carbon corrosion can also cause a collapse of the support structure, reducing the porosity. This densification of the support material leads to an increase of the transport related voltage losses even in the absence of liquid water in the cathode [8,28,29]. The ECSA based increase of the oxygen transport resistance arises from the ionomer thin film. The thin film overcoats the active sites and provides a resistance to oxygen transfer based on the oxygen flux per active cata lyst surface area. Therefore, reducing the surface area is, at a given current density, equivalent to an increase of the local oxygen flux. This results in a larger oxygen mass transport resistance (R₀₂) [30-33].

The fourth phenomenon is the limitation of the cell per formance of membrane electrode assemblies (MEAs) at high current densities. This limitation of yet unclear origin occurs already at begin of life, especially when the cathodes have low catalyst loadings [30,34-36]. The related losses increase drastically during cathode aging [8,11,35]. So far, different authors already explored this phenomenon, trying to trace it back either to a changing reaction mechanism at sufficiently low half cell voltage [37,38] or to an increased oxygen trans port resistance [34,39,40]. As there is no explicit explanation, the origin of these losses is still under debate. Doubts about the reaction mechanism hypothesis mainly arise from the harsh increase of the Tafel slope up to several hundreds of mV at high current densities observed in Refs. [8,36]. These slopes are much steeper than the slopes normally reported in liter ature for a changed reaction mechanism (up to 140 mV/dec.) [25,38]. Oppositely, the experimentally observed R_{02} increase was never strictly linked to these unassigned losses, as an increase would be required that seems far too large to explain

¹ Nevertheless, Papadias et al. cannot sufficiently disprove the hypothetical relevance of cation leaching for increased Ohmic losses, as only polarization data under 100% relative humidity are reported [17].

the unassigned voltage loss.² However, due to the technical inability to measure oxygen transport resistances under the conditions of the polarization curve,³ it was not possible to disregard the hypothetical R_{O2} increase, yet.

In this work, unassigned voltage losses of unknown origin were obtained as well. However, these losses occur even at low current densities. Linked with these unassigned voltage losses is a shortcoming of all voltages loss analyses so far: the omission of oxide related performance losses. Although it is clear that PtOx reduces the activity of the catalyst by "poisoning" part of the active sites [17,38,41], it also seems plausible that additional electronic resistances arise in the catalyst particles. Fantauzzi et al. and Kirchhoff et al. simu lated the cell voltage dependent oxidation of Platinum nano particles [42,43]. Accordingly, surface oxides are already formed at cell voltages as low as 750 mV.⁴ If the cell voltage decreases the surface oxides are altered to subsurface/bulk oxides and the electronic resistances increase. Direct evidence for Ohmic losses from oxides is given by Neff et al. as they show the increase of resistivity with increasing oxygen con tent in PtOx films [46]. Zhu et al. show how the Ohmic in plane resistance of a polycristalline Pt film jumps up (and slowly grows afterwards) once the atmosphere is changed from H₂ to air [47]. Ignoring the PtOx related voltage loss results in the following problem: By comparing polarization curves at the begin of life (BoL) and end of test (EoT) and disregarding the PtOx formation based voltage losses, it is inherently assumed that losses from PtOx remain constant during degradation. But this assumption is not reasonable due to the strong de pendency of PtOx formation on the half cell voltage [42,43,48–50], which is reduced as consequence of degrada tion. Accordingly, the cell voltage at a given current decreases over lifetime due to degradation, the losses from PtOx should also be expected to decrease [48-51].

 2 According to Ref. [11] the oxygen mass transport related voltage loss and the R_{O2} can be calculated with the following Eqs.

$$\eta_{\text{RO2}} \quad \frac{\mathbf{R} \cdot \mathbf{T}}{\mathbf{F}} \cdot \left(\frac{1}{4} + \frac{\gamma}{\alpha}\right) \cdot \ln\left(\frac{p_{\text{O2,ch}} \quad \frac{\mathbf{R} \cdot \mathbf{T}}{4\mathbf{F}} \cdot \mathbf{i} \cdot \mathbf{R}_{\text{O2}}}{p_{\text{O2,ch}}}\right)$$

$$R_{O2} = \frac{F}{R \cdot T} \cdot \frac{p_{O2,ch}}{i_{lim}}$$

where i_{lim} represents the limiting current of the used MEA. With these equations voltage losses up to several hundreds of mV can only be explained by two fold increase of the R₀₂. This is only possible if the limiting current decreases drastically to unreasonably low values.

³ The simple limiting current based method requires low cell voltages, at which flooding or overheating of the used MEA can occur, leading to changed conditions compared to those during polarization curve measurements.

⁴ The effect of platinum oxide on the degradation of the PEM FC cathode itself cannot be suppressed arbitrarily [44], as the typical operation window of the cells in PEM FC vehicles lies between 0.6 and 0.85 V. However, the voltage losses from Plat inum oxide can be reduced, e.g. by implying a dynamic operation strategy that contains frequent phases with cell voltages lower than 750 mV to reduce the formatted oxides [45].

The aim of these work is to identify, whether the intro duced degradation phenomena are influencing the perfor mance of the analyzed MEAs and how the quantification of PtOx related losses affect the interpretation of the degradation analysis of sate of the PEM FC cathodes. In the following, the new diagnostic procedure, presented in Section Testing, is implemented to assign voltage losses in polarization curves to PtOx, high frequency resistance (HFR), oxygen mass transport resistance (R_{O2}) , electrochemical active surface area (ECSA), specific activity (SA) and proton transport resistance (R_{H+}) . Consequently, this procedure is used in Section Results and discussion to study voltage losses during cathode degrada tion, whereby the focus in this contribution lies on analyzing the PtOx losses and their influence on the unassigned voltage losses occurring at high current densities. It will be shown that R₀₂ remains constant over life time, while PtOx related losses decrease. Finally, it becomes apparent, that the unas signed voltage loss increases over life time, impacting high and intermediate current densities.

Experimental

Materials and equipment

Experimental data were gathered using a 5 cm² MEA design based on the studies of Baker and Caulk [52]. A picture of the used flow field and the MEA is provided in Appendix A. A cathode loading of 0.25 mg_{Pt}/cm² PtCo catalyst with a Pt: Co molar ratio of 70:30 in the raw powder, an un alloyed Pt anode with a loading of 0.05 mg_{Pt}/cm² an ionomer with a low equiv alent weight and a persulfonic acid based membrane with a thickness of 18 μ m were used. For gas diffusion layers Sigracet 25BCE were used on both sides, compressed with a pneumatic hardware to a compression in the active area of 1 MPa. All electrochemical data and operation conditions were gathered with a fully automatized single cell test bench from Horiba Fuel Con and an additional external potentiostat (Zahner Zennium).

Testing

The basic test sequence consists of four parts. It begins with a conditioning procedure, consisting of several step changes in current and voltage, to activate the MEAs. Afterwards the MEAs are characterized at BoL with the developed diagnostic procedure, described in detail below. After the diagnostic procedure, the used MEAs were degraded with accelerated stress tests (AST) [15,53,54]. The ASTs were ended after 31,500 cycles and the diagnostic procedure was repeated to charac terize the MEAs at EoT.

Fig. 1 shows the developed diagnostic procedure. It starts with determining the ECSA using cyclic voltammetry (CV) (Fig. 1a). The CVs were recorded with a *Zahner Zennium* between 0.06 and 0.7 V (vs. H_2 anode) under N_2 atmosphere with 20 mV/s and were analyzed to derive ECSA and the H_2 crossover [8,11]. The electrochemical surface area was obtained by averaging the *H* desorption and *H* adsorption charge (using a specific charge of 210 C/cm²_{Pt}). After the CV, the diagnostic procedure shown in Fig. 1b–d is run with a cell temperature of 80 °C, H_2 /air flows of 5 Nl/min, an inlet pressure on both sides of 2.0 bar and

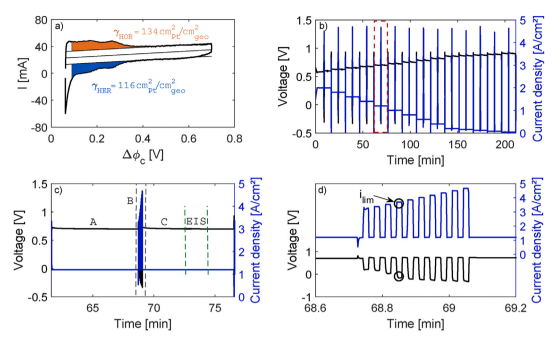


Fig. 1 – The developed diagnostic procedure: a) Measured CV under N₂ operation for ECSA determination; b) Overview of the developed diagnostic procedure containing the 13 measurement sequences between 2.0 and 0.0 A/cm² c) Exemplary sequence showing the three different phases of the procedure at 1.2 A/cm² d) Zoom into the 10 current changes conducted in phase B.

a relative humidity of 100% at the cell inlet. With the high stoichiometries (>10 at 1.8 A/cm²) and the specific flow field design, it is ensured, that the 5 cm² active area is exposed to differential conditions – gradients along the channel in pres sure, relative humidity, temperature and/or oxygen partial pressure are negligible. The diagnostic procedure consists of 13 different measurement sequences of 15 min, each belonging to a specific current density between 0 and 2.0 A/cm². In each sequence different "diagnostic tools" contained in three phases are used to extract the desired cell properties, as depicted in Fig. 1c.

In phase A, the cell is conditioned for 7.5 min at the desired current density to set up the liquid water profiles (in plane and through plane) in the MEA (Fig. 1c). In phase B, one potentiostatic and nine galvanostatic steps are carried out. Thereby, during the first step the voltage of 0.2 V is set for 1 s and the average current density is measured. Afterwards the steady state current density of phase A is set again for 1 s, followed by nine galvanostatic steps. In these steps the cur rent density measured in the potentiostatic step is increased step wise to obtain even lower voltages. In each of these steps the high current densities are maintained for 1 s, followed by a 1 s hold at the steady state current density of phase A. The aim of this sequence is two fold. On the one hand, at these low voltages PtOx is completely reduced. On the other hand, from this sequence the R_{02} for the load point and its corresponding liquid water profile is estimated. For this estimation, the cur rent density at the reversal point shown in Fig. 1d is used. For more details about the procedure and its legitimization, the reader is referred to Ref. [55].

In phase C the current density of phase A is held for another 7.5 min. Thereby, a voltage decay is seen that is attributed to the re formation of PtOx. The maximum voltage (~1 s after the start of phase C) is considered to be "PtOx free".⁵ The average voltage of the last 2 min of phase C is referenced as equilibrium voltage. This voltage is used to extract the so called equilibrium polarization curves from the 13 test sequences shown in Fig. 1b. From the PtOx free and equilibrium voltage, the losses due to PtOx formation can be determined. The strict linkage of the voltage decay in phase C to the formation of PtOx is reasonable as all other potential causes can be ruled out [55]: Firstly, the contribu tion from double layer charging within in observed voltge decay is insignificant as the capacitive current is in the range of several μ A/cm², assuming 100 mF/cm² as double layer capacity. Secondly, as phase B is too short to change the temperature of the whole cell setup, the temperature in the catalyst layer and the GDL adapts within 200 ms after the last limiting current measurement and the temperature in phase C is constant. Thirdly if liquid water is absent the rH profile is also settled within ms after phase B. Consequently, no changes of the Ohmic losses in phase C are expected. Fourthly, the resettling of the oxygen profile after phase C is finished after 25 ms and we assume that the liquid water profile and the mass transport related losses within in phase C are constant, too.

Fifthly, if liquid water is present in the GDL and/or cathode electrode, we argue that there is not enough time to evaporate a lot of water and the mass transport related losses are also unchanged in phase C. This rationalization of the unchanged liquid water profile over phase C is further supported by our later discussed findings (Fig. 5), that the PtOx losses at BoL are

 $^{^5\,}$ The term "PtOx free" used in this context is somewhat loosely as in the first second of the hold time some surface adsorbates will have formed.

Table 1 – Operation conditions during the AST including variations of the holdtime at the lower and upper potential limit.

| General Operation Conditions | | |
|------------------------------|-----------------------------------|-------------|
| | Potential shape | Square Wave |
| | Lower Potential Limit (LPL) | 0.6 V |
| | Upper Potential Limit (UPL) | OCV |
| | Temperature | 90 °C |
| | Relative Humidity (rH) | 100% |
| | Inlet Pressure (p _{in}) | 2.0 bar |
| | Cycle number | 31,500 |
| AST 1 | Hold time | 2.5 s |
| AST 2 | Hold time | 10 s |
| AST 3 | Hold time | 20 s |

almost independent on current density a misinterpretation of the mass transport losses due to varying liquid profiles would cause a notable dependence instead.

In addition to the PtOx loss determination, EIS spectra between 50 Hz and 30 kHz are recorded from minute 3 to minute 4.5 of phase C (Fig. 1c). These spectra are then fitted with the help of a transmission line model to derive HFR and R_{H+} according to Ref. [56]. A detailed picture of the HFR and R_{H+} determination is given in Appendix B. In order to estimate the specific catalyst activity (SA), the current density at 0.9 V is estimated by extrapolating the measured PtOx free polariza tion curve between 0.04 and 0.12 A/cm². This current is divided by the derived ECSA.

After the diagnostic procedure, the used MEAs were degraded with accelerated stress tests (AST) as summarized in Table 1.

The AST consisted of 31,500 cycles, with each consisting of potentiostatic steps between open circuit voltage and 0.6 V, with different holding times at each potential. After 31,500 AST cycles the diagnostic procedure is carried out again to obtain the desired EoT MEA properties. The complete process containing conditioning, BoL diagnostic, AST and EoT diag nostic was run with two identical cells for AST 1 and 3. In Fig. 2, exemplary BoL and EoT polarization curves obtained with the diagnostic procedure are shown for one cell. The solid lines represent the so called equilibrium polarization curves, while the dashed lines are the PtOx free polarization curves, the influence of PtOx on the cell performance can easily be derived.

Voltage loss break down

In this section, it is shown how the performance change of the degraded MEA can be broken down into single voltage loss contributions, which are quantified with the gathered data. In Fig. 2 the solid black and blue lines represent the measured equilibrium polarization curve at BoL and EoT ($U_{Cell,equi}^{BoL/EoT}$), respectively. They contain the reversible cell voltage U_{rev} and the various overvoltages:

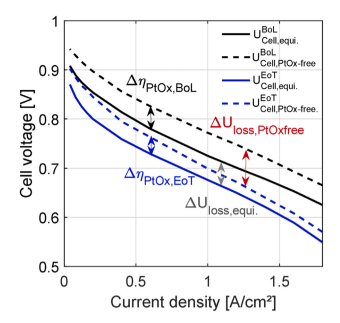


Fig. 2 – Exemplary polarization data BoL and EoT obtained with the developed diagnostic routine.

$$U_{Cell,equi.}^{BoL/EoT} = U_{rev} \quad \eta_{HFR}^{BoL/EoT} \quad \eta_{a}^{BoL/EoT} \quad \eta_{act}^{BoL/EoT} \quad \eta_{H+}^{BoL/EoT} \quad \eta_{RO2}^{BoL/EoT}$$

$$\eta_{PtOx}^{BoL/EoT} \quad \eta_{?}^{BoL/EoT}$$
(1)

These overpotentials are Ohmic losses, η_{HFR} , related to the proton transport through the membrane and the Ohmic re sistances of the electrically conductive components of the cell, activity losses of the anode, η_a , (neglected in the rest of this work due to high anode activity) and losses related to cathode. These cathode losses are the activation over potential, η_{act} , losses related to the transport of protons through the cathode η_{H+} , losses attributed to the oxygen transport from the gas channels to the catalyst η_{RO2} and losses related to the formation of surface oxides η_{PtOx} . Furthermore, possible unassigned overpotentials are referred as $\eta_{?}$. The PtOx free polarization curves is described by setting η_{PtOx} to zero:

$$U_{Cell,PtOx free}^{BoL/EoT} = U_{rev} \eta_{HFR}^{BoL/EoT} \eta_a^{BoL/EoT} \eta_{act}^{BoL/EoT} \eta_{H+}^{BoL/EoT} \eta_{RO2}^{BoL/EoT}$$
$$\eta_{?}^{BoL/EoT}$$
(2)

As $U_{Gell,equi}^{BoL/EoT}$ and $U_{Gell,PtOx}^{BoL/EoT}$ are measured (as described in Section Testing), the PtOx losses are derived from their difference:

$$\eta_{\text{PtOx}}^{\text{BoL/EoT}} \quad U_{\text{Cell,PtOx} free}^{\text{BoL/EoT}} \quad U_{\text{Cell,equi}}^{\text{BoL/EoT}}$$
(3)

To quantify the degradation caused performance losses designated as $\Delta U_{loss,equi}$ and $\Delta U_{loss,PtOx}$ free in Fig. 2, the differ ence between the BoL and EoT polarization is determined as:

$$\Delta U_{\text{loss,equi}} \quad U_{\text{Cell,equi}}^{\text{BoL}} \quad U_{\text{Cell,equi}}^{\text{EoT}} \tag{4}$$

$$\Delta U_{\text{loss,PtOx free}} \quad U_{\text{Cell,PtOx free}}^{\text{BoL}} \quad U_{\text{Cell,PtOx free}}^{\text{EoT}} \tag{5}$$

The obtained differences of the BoL and EoT overpotentials are described in Eqs. (6) and (7):

$$\Delta U_{\text{loss,equi.}} \quad \Delta \eta_{\text{ECSA}} + \Delta \eta_{\text{SA}} + \Delta \eta_{\text{HFR}} + \Delta \eta_{\text{RH}+} + \Delta \eta_{\text{RO2}} + \Delta \eta_{\text{PtOx}} + \Delta \eta_{\gamma}$$
(6)

$$\Delta U_{\rm loss,PtOx free} \quad \Delta \eta_{\rm ECSA} + \Delta \eta_{\rm SA} + \Delta \eta_{\rm HFR} + \Delta \eta_{\rm RH+} + \Delta \eta_{\rm RO2} + \Delta \eta_{\rm ?} \tag{7}$$

The first and the second term of Eq. (6) account for losses which are related to a reduced ECSA and a changed catalyst activity. For the estimation of these losses Eqs. (8)-(11) are used:

$$\Delta \eta_{act} \quad \eta_{act}^{\text{BoL}} \quad \eta_{act}^{\text{EoT}} \quad b \cdot \log_{10} \frac{i_{0.9V}^{\text{BoL}}}{i_{0.9V}^{\text{EoT}}} \right)$$
(8)

$$\Delta \eta_{\rm ECSA} = b \cdot \log_{10} \left(\frac{\rm ECSA^{\rm BoL}}{\rm ECSA^{\rm EoT}} \right)$$
(9)

The current density $i_0^{Bol/EoT}$ is obtained from extrapolation (or interpolation) of the PtOx free polarization curve. The ECSA^{BoL/EoT} used in Eq. (9) stems from CV measurements (see Section Testing). *b* represents the Tafel slope:

$$b = 2.303 \cdot \frac{\mathbf{R} \cdot \mathbf{T}}{\alpha \cdot \mathbf{F}} \tag{10}$$

With R being the gas constant, T being the cell temperature in Kelvin, α being the cathodic transfer coefficient (α 1 ac cording to Ref. [57]) and F being the Faraday constant. Accordingly, the Tafel slope is approximately 70 mV/dec [8,11,38]. The losses related to a changed specific activity are calculated with Eq. (11):

$$\Delta \eta_{\rm SA} \quad \Delta \eta_{\rm act} \quad \Delta \eta_{\rm ECSA} \tag{11}$$

The third part of the performance loss is linked to changes of the HFR extracted from the impedance spectroscopy described in Section Testing. This performance loss is described in Eq. (12):

$$\Delta U_{HFR} \quad i \cdot A \cdot (HFR^{BoL} \quad HFR^{EoT}) \tag{12}$$

where i describes the current density and A the active area of the used MEA (5 cm²). The fourth contributor to the perfor mance losses is the change of proton conductivity in the cathode. This voltage loss is estimated with Eq. (13) [56]:

$$\Delta \eta_{H+} = \frac{1}{3} \cdot \mathbf{i} \cdot \mathbf{A} \cdot (\mathbf{R}_{H+}^{BoL} - \mathbf{R}_{H+}^{EoT})$$
(13)

where R_{H+}^{BoLEOT} is extracted from the impedance spectra (see Appendix Fig. A3). The fifth term of Eq. (6) is related to the comparison of the mass transport behavior of the fresh and aged MEA. An equation derived in the work of Zhirul et al. [11] is used to calculate the mass transport related voltage loss at end of test:

$$\Delta \eta_{\text{RO2}} = \frac{R \cdot T}{F} \cdot \left(\frac{1}{4} + \frac{\gamma}{\alpha}\right) \cdot \ln \left(\frac{p_{\text{O2,ch}}}{p_{\text{O2,ch}}} - \frac{R \cdot T}{4F} \cdot \mathbf{i} \cdot \mathbf{R}_{\text{O2}}^{\text{BoL}}}\right)$$
(14)

where $R_{O2}^{Bol,EoT}$ is derived from the limiting current measure ments described in Section Testing, α represents the cathodic transfer coefficient and γ is the ORR reaction order with respect to oxygen partial pressure (γ 0.54 according to Ref. [57]). The oxygen partial pressure in the channels of the flow field is described with $p_{O2,ch}$. Eq. (15) describes the influ ence of the sixth term of Eq. (6) on the cell performance,

$$\Delta \eta_{\rm PtOx} \quad \eta_{\rm PtOx}^{\rm EoL} \quad \eta_{\rm PtOx}^{\rm BoL} \tag{15}$$

The last part of Eq. (6) is attributed to unassigned voltage losses introduced in Section Introduction and is addressed by : $\Delta \eta_{?} \quad \eta_{?}^{\text{EoL}} \quad \eta_{?}^{\text{BoL}}$ (16)

whereby $\eta_{?}^{\text{Bol/EoT}}$ is obtained from Eq. (6), as all other contributors are calculated with Eqs. (8)–(15) and the different measured BoL and EoT values are specified. The Eqs. introduced above are valid under the assumption that the cell temperature is constant over the active area and that the difference between the oxygen and hydrogen concentration at the cell in and outlet is insignificant.

Results and discussion

This section is divided in four parts. In the first three parts the BoL and EoT data gathered with the diagnostic procedure are discussed. In Section Break down of degradation caused voltage losses finally, the break down of the EoT perfor mance loss into the different contributors is analyzed.

High frequency resistance, cathode proton transport resistance and oxygen mass-transport resistance

Fig. 3 shows the measured HFR, R_{H+} and R_{O2} at low, medium and high current densities. Fig. 3a–c represent the high fre quency resistances derived from the EIS measurements. The differences between the BoL and EoT values are insignificant. Consequently, it can be concluded, that the proton conduc tivity of the membrane is not significantly reduced, e.g., by leached out Co^{2+} from the cathode or other membrane degradation effects.

Fig. 3d–f shows the obtained R_{H+} . Even though there is a considerable scattering of the data (e.g., compare the BOL values), the proton transport resistance generally decreases at EOT. The reduction of R_{H+} is possibly a result of reduced cathode thickness following carbon corrosion, which is known to become relevant at voltages as high as 0.95 V [35,58–60]. In addition to the reduced proton transport pathway, the reduced cathode layer thickness leads to an increasing amount of ionomer per volume, resulting in lower values for cathode proton resistance, as discussed, e.g., in Refs. [61,62]. The influence of the changed cathode layer thickness on the Ohmic behavior of the analyzed MEAs is insignificant, as the HFR is mainly determined by the proton conductivity of the membrane.

In Fig. 3g l the extracted mass transport resistances are summarized. For this cathode property the difference be tween BoL and EoT is also insignificant. This result is in contradiction to the expectation derived from literature,

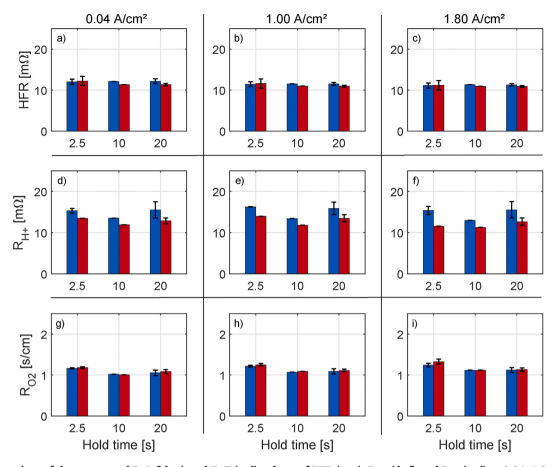


Fig. 3 – Overview of the measured BoL (blue) and EoT (red) values of HFR (a–c), R_{H+} (d–f) and R_{O2} (g–i) at 0.04, 1.0 and 1.8 A/ cm² for all three accelerated stress tests (variation of hold times). (For interpretation of the references to color/colour in this figure legend, the reader is referred to the Web version of this article.)

where carbon corrosion leads to an increase of the oxygen mass transport resistance [28,29,35]. But it is in line with findings recently published by Harzer et al. [8], where carbon corrosion was evident from SEM images although no signifi cant increase in R_{O2} was detected. Generally, we observed no difference in impact of the different degradation procedures on electron, ionic and oxygen mass transport.

Catalyst activity

Fig. 4 shows the electrochemical surface area (ECSA), current densities at 0.9 V, and the derived specific activity (SA) at BoL and EoT normalized to the BoL values. The loss of current densities at 0.9 V increases with increasing hold time between 2.5 and 10 s but no significant difference between 10 and 20 s is found (Fig. 4a). These losses can be divided in losses of ECSA and SA. The ECSA losses are shown in Fig. 4b. Comparable results can be found in literature [60,63], where higher ECSA losses were found with increasing hold times at the upper potential limit and lower potential limit.

The loss of SA (Fig. 4c) is a well known degradation phe nomenon for alloy catalyst [16,17,64,65]. For pure Pt catalyst or catalyst with a very low alloy content, the specific activity shows only a slight decreases at EoT [17,66,67]. Consequently the loss of ECSA determines the voltage losses related to a changed activation overpotential [11,35]. The loss of SA for PtCo catalyst used in this work can be related to dealloying effects. The loss of the non precious metal due to degradation results in a loss of the activity enhancing effect due to altered electronic structure of the catalyst material [17,65].

Platinum oxide related overvoltages

In Fig. 5a, the PtOx related overvoltages at BoL and EoT are shown as a function of cell voltage. But as the PtOx formation is driven by the potential on the cathode side it is necessary to consider the PtOx related overvoltages at BoL and EoT as function of half cell potential (Fig. 5b) to avoid a misinterpre tation of development of the PtOx related losses over time. The half cell potential is obtained by correcting the measured cell voltage $U_{equi}^{BoL/EoT}$ with $\eta_{HFR}^{BoL/EoT}$, $\eta_{H+}^{BoL/EoT}$ and $\eta_{RO2}^{BoL/EoT}$. In general, Fig. 5b shows that the PtOx related overvoltages already exist at half cell potentials higher than 720 mV. This is in line with the recently published work of Kirchhoff et al. who show that oxide formation on platinum nanoparticles can already begin at 700 mV [43]. The PtOx related overvoltages increase with

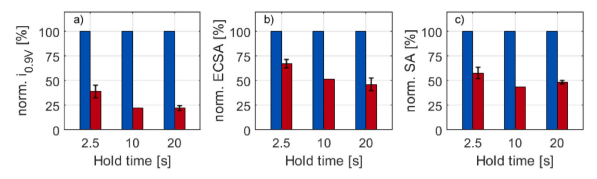


Fig. 4 – Overview of the normalized current densities at 0.9 V (a), electrochemical surface area (b) and specific activity (c) values at BoL (blue) and EoT (red) for all three accelerated stress tests (variation of hold times). (For interpretation of the references to color/colour in this figure legend, the reader is referred to the Web version of this article.)

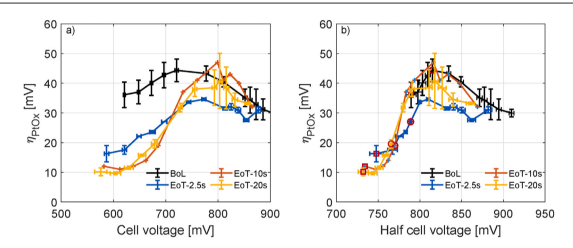


Fig. 5 – a) η_{PtOx} as function of the cell voltage BoL and EoT; b) η_{PtOx} as function of the half cell potential BoL and EoT (red circles: $\eta_{PtOx}^{BoL/EoT}$ @ 1.0 A/cm², red squares: $\eta_{PtOx}^{BoL/EoT}$ @ 1.8 A/cm²). (For interpretation of the references to color/colour in this figure legend, the reader is referred to the Web version of this article.)

increasing half cell potential (decreasing current density) be tween 750 and approximately 820 mV. After this maximum, the overvoltage decreases as the current density is further reduced.

To interpret this behavior, first, it needs to be said that to the best of the authors knowledge no comparable data in the literature exist, i.e. data that explicitly represents the half cell potential dependency of voltage losses attributed to PtOx formation. Commonly, in papers investigating PtOx, the accu mulated charge measured in the PtOx reduction peak of a CV is used for oxide quantification [48-50,68]. In all these works, the amount of PtOx increases with increasing half cell potential. This suggests that the PtOx overpotential would increase with half cell potential, too and contradicts the observed maximum in Fig. 5. The authors believe that the increase of the PtOx overvoltage between 750 mV and 820 mV half cell potential is caused by higher kinetic loss due to an increasing oxide coverage of the catalyst particles [17,69] and an increased loss from electron transport through (subsurface) oxides. Accord ingly, the increase of oxide overvoltage with increasing half cell potential, arises not uniquely from kinetic oxide losses, but at least partially from Ohmic oxide losses, which also depend on current density. Consequently, the decrease of the PtOx losses at voltages higher than 820 mV represents lower kinetic and electronic losses from PtOx that arise from lower current den sities (even though increasing Oxide loadings might further increase, as shown in Refs. [48–50]).

At half cell potentials lower than 750 mV η_{PtOx} seems to level off to a finite value. We speculate that this 10 mV offset is related to the adsorption of anions from the ionomer on the catalyst surface. As Jinnouchi et al. showed by DFT simulations for Pt (111), adsorption of sulfate containing anions is thermo dynamically favored between half cell potentials of 400 and 780 mV. These adsorbates suppress the oxide formation, poi son the catalyst and result in an inhibition of the ORR [44,69]. Accordingly, the authors believe that the "PtOx free" polariza tion curve has not only reduced losses from PtOx, but is also not impacted (or at least less impacted) from anion adsorption. This hypothesis is potentially in line with the findings of [70] about the regeneration of sulfonate adsorbates.

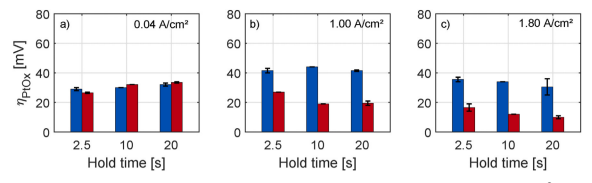


Fig. 6 – Overview of the averaged BoL (blue) and EoT (red) PtOx overvoltages at 0.04 (a), 1.0 (b) and 1.8 A/cm² (c). (For interpretation of the references to color/colour in this figure legend, the reader is referred to the Web version of this article.)

Fig. 6 shows the measured PtOx related overvoltages at 0.04, 1.0 and 1.8 A/cm² at BoL and EoT extracted from Fig. 5b. At 0.04 A/cm² (Fig. 6a) it can be seen that almost the same PtOx related overvoltages are obtained at BoL and EoT in dependent of the chosen hold time. In Fig. 6b and c, the overvoltages at 1.0 A/cm² respectively at 1.8 A/cm² are shown. At these current densities η_{PtOx} is reduced significantly at EoT due to decreased half cell potential at higher current densities. And as the activity decreases with increasing hold time (see Fig. 4) lower half cell potentials are obtained at EoT. Consequently, the reduction of η_{PtOx} is more severe if the hold time is increased from 2.5 to 10 s. But no further reduction of η_{PtOx} is obtained for a hold time of 20 s, as comparable activity losses are measured for hold times of 10 and 20 s (see Fig. 4).

Break down of degradation-caused voltage losses

In this section the difference between the different over voltages defined in Section Voltage loss break down at BoL and at EoT obtained for the three ASTs are compared to each other. The data described in the sections before are used to break down these losses into the different contributors of Section Voltage loss break down and the origin of the unassigned voltage loss is discussed.

Fig. 7a represents the equilibrium polarization curves at BoL and EoT for differently aged samples. In Fig. 7b–d, the break down of the measured degradation related voltage losses at 0.04, 1.0 and 1.8 A/cm² following Eqs. (8)–(16) is depicted. In general one can see the following: As the HFR did not show significant changes over life (Fig. 3a–c), additional HFR related voltage losses (according to Eq. (12)) are absent. The same is true for oxygen transport related losses, as the oxygen transport resistance was stable, too (Fig. 3g–i). Oppositely, the cathode proton resistance did show some improvement for all variants of the AST (Fig. 3d–f). This improvement is converted with the help of Eq. (13) and leads to negative voltage loss changes. PtOx losses were seen to be reduced at EoT in Fig. 6 for all but the smallest currents. The work of Zago et al. also shows, that the catalyst activity is less influenced by PtOx for aged samples [44].

Accordingly, in Fig. 7c and d the difference is also counted as gain (bars heading negative). Increases in the voltage losses arise majorly from a reduction in activity, whereby ECSA and SA contribute almost equally. The related bars are heading up, starting below zero at the value of total gain. The gap between the upper end of the ECSA related voltage losses and the black tick marks the additional unassigned voltage losses at EoT. They will be discussed in the end of this sec tion, when the impact of current density and hold time was rationalized. According to Eqs. (9) and (11) that ECSA and SA related voltage losses are independent of current density and therefore constant in Fig. 7b-d. Increasing gains (with increasing current density) arise on the one hand from the trend in cathode proton losses with hold time. But no clear trend can be identified given the error bar of the cathode proton resistance (Fig. 3d-f). On the other hand, increasing gains (with increasing current density) arise from reduced oxide related voltage losses. The root cause is simply a reduced oxide formation with decreasing half cell potential EoT, as shown in Fig. 5b. Thereby, the PtOx gain is almost identical for 1.0 and 1.8 A/cm², simply due to the curvature of the oxide voltage losses with half cell potential (Fig. 5b compare highlighted marks at 1.0 and 1.8 A/cm² BoL and EoT). On the other hand, the gain from a reduced cathode proton resistance also increases due its direct dependence on current density (Eq. (13)).

Discussing the impact of hold times, it is evident, that the total losses (black ticks) increase from 2.5 to 10 s. The outcome of the AST with 20 s hold time is then very close to the AST with 10 s hold hardly larger than the error bar (~ 5 mV). This result is in line with earlier studies [11,63], which found that ASTs with the same upper potential limit tend to the same EoT voltage losses. As expected, ECSA and SA losses increase from 2.5 to 10 s, but not further when the hold time is 20 s. Regarding the trend in the PtOx related voltage losses, one can see the lowest gain for the AST with 2.5 s hold time due to the higher EoT half cell potential. It seems that there is a little less gain in the case of 20 s hold time. However, this potentially unexpected difference between the ASTs with 10 and 20 s hold time is in the order of 5 mV and therefore small. Nonetheless, it might point to slightly different EoT states of the samples evidently particle size and shape have a tremendous impact on PtOx build up [42,43].

In the end, the increase in unassigned voltage loss shall be discussed. Evidently, the loss is negligible at 0.04 A/cm^2 and increases with current density up to 35 mV. This insight is

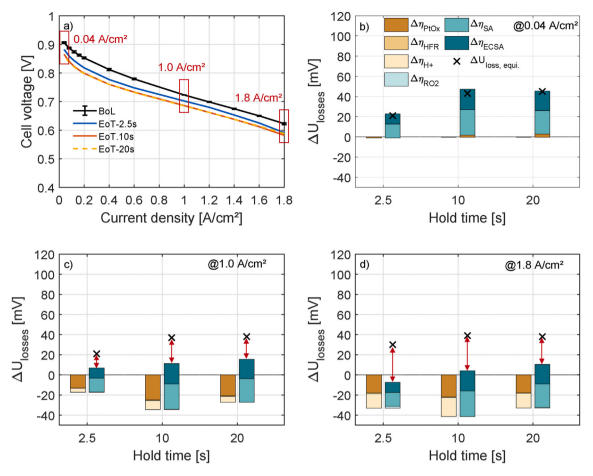


Fig. 7 – a) Averaged begin-of-life polarization curve and the obtained end-of-test polarization curves for the differently degraded samples; b-d) Break down of the additional voltage losses at end-of-test at 0.04 (b), 1.0 (c) and 1.8 A/cm² (d) into the different contributors introduced in Section Voltage loss break down.

surprising, as the almost perfect vertical shift in the polari zation curves EoT (Fig. 7a) suggests that degradation trivially causes activity losses only. However, if the PtOx related voltage losses are taken into account, unassigned losses remain or rather turn out larger than without considering PtOx.

Earlier, others have identified an unassigned voltage loss, too [30,34–36]. However, in the current work the discussion of

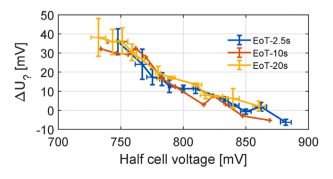


Fig. 8 – $\Delta \eta_{?}$ as function of the half cell potential EoT.

the unassigned voltage loss is based on the direct determi nation of the oxygen transport resistance and PtOx related voltage losses. Accordingly, with this additional diagnostic information, the often argued "increased oxygen transport resistance EoT" can be excluded as no change in oxygen transport resistance over the course of AST testing is seen (Fig. 3g–i). A similar argument is derived in Ref. [8] derived from the comparison of stationary limiting currents at BoL and EoT. Two hypotheses seem more plausible to explain the unassigned voltage loss. The first hypothesis bases on an increased Tafel slope over the course of AST testing. Support of this idea arises from Fig. 8, which displays the unassigned voltage losses as a function of half cell potential at EoT.

Indeed, a linear increase with half cell potential is seen. Therefore, the unassigned losses could be caused by a dis regarded increase of the Tafel slope at EoT. This increase will lead to a steeper slope of the polarization curve and conse quently the unassigned voltage losses would be reduced. However, assuming an increasing Tafel slope is counter intuitive as PtCo is known to exhibit a larger Tafel slope compared to Pt [71–73]. But likely the catalyst is more Pt like at EoT due to leaching and accordingly should have lower Tafel slope at EoT compared to BoL not larger.

The second hypothesis that might be plausible bases on catalyst poisoning due to sulfonate adsorption on the catalyst surface [74,75]. Indeed, Kodama et al. found an increasing voltage loss from Nafion at Pt with decreasing half cell po tential (Fig. 6 in Ref. [76]). Accordingly, at EoT more sulfonate adsorption losses could be expected at a given current. This expectation can be roughly quantified: Reading from Fig. 6 in Ref. [76], the losses from Nafion at Pt increase from ~ 40 mV at a half cell potential of ~900 mV to ~60 mV at a half cell po tential of ~700 mV. This means at least a voltage loss of ~10 mV can be attributed to the interaction between Nafion and the catalyst if there is a reduction of the half cell potential of 100 mV. Fig. 5b in this work shows, that the half cell po tential at 1.8 A/cm² decreases from ~800 mV to ~700 mV over the cause of AST testing (for all ASTs). This decrease of half cell potential yields 35 mV of unassigned loss (Fig. 8), which is indeed more than the 10 mV expected from Ref. [76]. Never theless, one should also be aware that [76] is a study of Nafion on Pt (111) in an RDE setup and therefore a considerably different system.

Conclusion

In this work, PEM FC cathodes containing PtCo catalyst, degraded with square wave accelerated stress tests of vary ing hold time, were analyzed with a dedicated diagnostic procedure. For every individual load point, the oxygen transport resistance and the voltage losses due to the for mation of PtOx were obtained in addition to commonly measured electrochemical active surface area, high fre quency as well as proton transport resistance. These data were used to break down the end of test voltage losses into six different contributors.

The high frequency resistance and the oxygen mass transport resistances shows no significant changes end of test. Consequently, no performance loss was attributed to an increased mass transport or to an increased Ohmic resistance end of test. In contrast, the proton resistance decreased for the analyzed samples, leading to performance gains after the accelerated stress tests. This behavior can be explained by a loss of carbon, leading to an increased volumetric ionomer content. As the electrochemical active surface area and the specific activity were reduced due to degradation, activity related performance losses are obtained. The PtOx related losses are decreased at end of test, except at the smallest current densities. Most noticeably, it was shown that PtOx related voltage losses show almost the same dependency on half cell voltage for all begin and end of test samples. Taking all these different contributors into account, additional unassigned voltage losses remain.

Interestingly, the unassigned voltage loss was not only present at highest current densities as found elsewhere [30,34,36], but showed a steady increase with current density. When plotting it against half cell voltage, it was found to even be independent from the accelerated stress tests, or in other words: The unassigned voltage loss at end of test showed the same linear increase with cathode overvoltage for all accel erated stress tests. We speculate that the origin of the unas signed loss might arise from ionomer adsorption that, based on an earlier publication, depends on half cell voltage [76]. In the end it can not be in scope of membrane electrode as sembly testing to unambiguously identify the root cause of the unassigned voltage loss, but with this new diagnostic pro cedure we believe to have made a good attempt in revealing its true impact.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A

For the experiments a 5 cm² MEA design based on the studies of Baker and Caulk [52] with a gas flow field of 50 cm² shown in Fig. A1 and an active MEA area of 5 cm² (Fig. A2: MEA arranged on the flow field) are used.

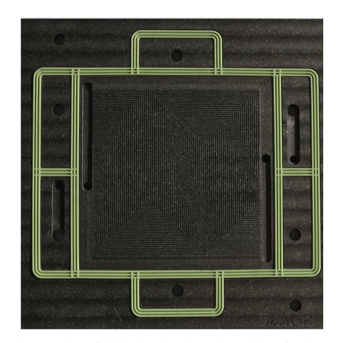


Fig. A1 – Flow-field-design based on the studies of Baker and Caulk [52].

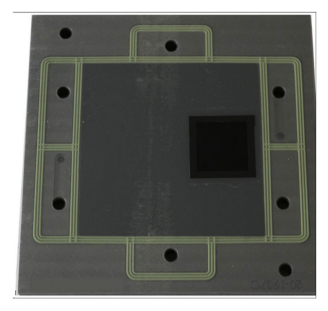


Fig. A2 – Flow-field with used GDL and 5 cm^2 MEA.

Appendix B

The high frequency resistance and the cathode ionomer resistance are determined by analyzing the recorded imped ance spectra between 50 Hz and 30 kHz. Therefore a trans mission line model derived from Makharia et al. [56] is used to fit the measured data. With this fit the HFR can be extracted by determing the high frequency intercept on the real impedance axis and the R_{H+} is extracted from the 45° branch of the fit (Fig. A3).

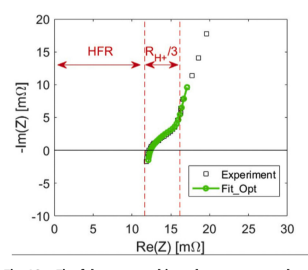


Fig. A3 – Fit of the measured impedance spectra and determination of the high frequency resistance and cathode ionomer resistance according to Ref. [56].

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