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# Self-supported Pt-CoO networks for oxygen reduction: electrocatalysts combine high specific activity with high surface area

Gustav W. Sievers<sup>a,b</sup>, Anders W. Jensen<sup>a</sup>, Jonathan Quinson<sup>a</sup>, Alessandro Zana<sup>a,c</sup>, Francesco Bizzotto<sup>c</sup>, Mehtap Oezaslan<sup>d,e</sup>, Alexandra Dworzak<sup>d,e</sup>, Jacob J. K. Kirkensgaard<sup>f,g</sup>, Thomas E. L. Smitshuysen<sup>h</sup>, Shima Kadkhodazadeh<sup>h</sup>, Mikkel Juelsholt<sup>a</sup>, Kirsten M. Ø. Jensen<sup>a</sup>, Kirsten Anklam<sup>b</sup>, Hao Wan<sup>a</sup>, Jan Schäfer<sup>b</sup>, Klára Čépe<sup>i</sup>, María Escudero-Escribano<sup>a</sup>, Jan Rossmeisl<sup>a</sup>, Antje Quade<sup>b</sup>, Volker Brüser<sup>b</sup>, Matthias Arenz<sup>a,c</sup>

<sup>a</sup> Department of Chemistry, University of Copenhagen, Universitetsparken 5, DK-2100 Copenhagen Ø Denmark

<sup>b</sup> Leibniz Institute for Plasma Science and Technology, Felix-Hausdorff-Strasse 2, 17489 Greifswald, Germany

<sup>c</sup> Department of Chemistry and Biochemistry, University of Bern, Freiestrasse 3, CH-3012 Bern, Switzerland

<sup>d</sup> Department of Chemistry, School of Mathematics and Science, Carl von Ossietzky University of Oldenburg, 26111 Oldenburg, Germany

<sup>e</sup> Institute of Technical Chemistry, Technische Universität Braunschweig, D-38106 Braunschweig

<sup>f</sup> Niels Bohr Institute, University of Copenhagen, Universitetsparken 5, DK-2100 Copenhagen, Denmark

<sup>g</sup> Department of Food Science, University of Copenhagen, Rolighedsvej 30, DK-1958 Frederiksberg C, Denmark

<sup>h</sup> Technical University of Denmark, Anker Engelunds Vej 1, 2800 Lyngby

<sup>i</sup>Regional Centre of Advanced Technologies and Materials, Šlechtitelů 27, 783 71 Olomouc, Czech Republic Corresponding authors: <u>matthias.arenz@dcb.unibe.ch; sievers@inp-greifswald.de</u>

Substantial progress has been achieved in the development of Pt-based catalysts for the oxygen reduction reaction (ORR) – the process taking place at the cathode of fuel cells. In the last years, several concepts based on Pt-alloy catalysts have been presented that significantly exceed the Department of Energy (DOE) targets for Pt related ORR mass activity (MA). With few exceptions, most concepts achieve their high ORR activity by increasing Pt specific activity (SA) at the expense of a lower electrochemically active surface area (ECSA). In the potential region controlled by kinetics, which is used in the DOE targets for fuel-cell performance, such lower surface area is more than counterbalanced by the high SA. At higher overpotentials, which are often applied in real systems, however, a low ECSA leads to limitations in the reaction rate not by kinetics, but mass transport in form of a local O<sub>2</sub> resistance. Herein we report on self-supported, nanoporous platinum-cobalt-oxide (Pt-CoO) networks that combine high SA with unprecedentedly high ECSA. Our investigations reveal that the unprecedentedly high ECSA is achieved by a Pt-CoO bone structure with a diameter of only a few nm. Using this concept, we achieved unprecedentedly high mass activity values for self-supported ORR catalyst layer. This material concept promises stable and low platinum fuel cell operation at high temperature, high current density and low humidification.

**Graphical Abstract** 

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The continuing emission of anthropogenic CO<sub>2</sub> is perpetuating man-made climate change. Green technologies such as fuel cells and electrolyzers for a hydrogen economy using renewable energy are necessary to maintain the global standard of living. Research and development of proton exchange membrane fuel cells (PEMFCs) have made substantial progress in recent years with PEMFCs reaching industrial maturity in serial production of fuel cell cars by several automotive companies. This advancement is partly due to rapid progress in engineering, but also due to more efficient fuel cell catalysts<sup>1–3</sup>. Roughly 25 g<sub>Pt</sub> are required using state of the art catalysts for a 90 kW fuel cell car<sup>4</sup>.

At a research level, several catalysts have already shown catalytic activities capable of further reducing the amount of Pt by a factor five to ten<sup>5</sup>, which is required for large-scale applications. Most catalyst concepts reach their high mass activity (MA) by increasing the specific activity (SA) of Pt-based catalysts at the extent of a reduced electrochemical surface area (ECSA) as compared to standard Pt/C catalysts<sup>5–</sup><sup>10</sup>. However, a reduced ECSA constitutes a barrier for their use in automotive applications.

The essential reaction in a PEMFC is the oxygen reduction reaction (ORR) to water:

$$O_2 + 4 H^+ + 4 e^- \to 2 H_2 0 \tag{1}$$

Since the oxygen reduction is kinetically inhibited, an overpotential is needed to start the reaction. At a relatively low overpotential of 0.3 V, where performance testing typically takes place, the reaction rate is limited by one of the charge transfer reactions, i.e. for Pt the reduction of the different intermediates O\*, HO\* and HHO\*<sup>11</sup>:

$$H0^* + H^+ + e^- \to H_20 \tag{2}$$

$$0^* + 2 H^+ + 2 e^- \to H_2 0 \tag{3}$$

$$00H^* + 3H^+ + 3e^- \to 2H_20 \tag{4}$$



Figure 1: Volcano Plot showing experimental activity data relative to Pt as a function of the hydroxyl intermediate binding energy  $\Delta G_{HO}$ ; Data for Pt/Au(111), Pt<sub>3</sub>Y, Pt<sub>3</sub>Ni, dealloyed PtCu, Pt<sub>3</sub>Fe, Pt<sub>3</sub>Co and Pt/Ir(111) is taken from Stephens *et al.*<sup>11</sup>(black); Data for Pt/C and Pt<sub>3</sub>Co/C is as used in this publication assuming the same hydroxyl binding energy as bulk Pt or Pt<sub>3</sub>Co respectively (grey); Data for Pt-CoO is taken from DFT calculations and electrochemical activity evaluation.

As the binding energies of the individual reaction intermediates are not independent of each other but are connected via the scaling relation<sup>11</sup>, the discussion is usually based on a single reaction intermediate, e.g. the hydroxyl binding energy. If the hydroxyl binding energy is too small, the reaction is limited by the OH\* reduction, if it is too large, the reaction is limited by the HHO\* formation. As a consequence, the experimental (and theoretical) data for the ORR activity display a volcano-type dependence on the hydroxyl intermediate binding energy, see Figure 1. In order to optimize the intermediate binding energies, the electronic structure of Pt has to be slightly changed. Main principles for achieving this and to design Pt-based ORR electrocatalysts are the strain and the ligand effects<sup>12</sup>, which are discussed later on in more detail.

Automotive fuel-cell systems, however, operate at high current densities and thus relatively large overpotentials. As the overpotential increases, the nature of the rate determining step changes and mass transfer becomes crucial for the performance<sup>4,13,14</sup>. As a consequence, at low overpotentials up to 0.3 V and thus low current densities, the reduced ECSA of most catalyst concepts does not impact their performance<sup>4</sup>. However, at higher overpotential, e.g. 0.5 V, which is often reached under fuel-cell operation, the charge transfer steps are fast on an active catalyst<sup>11</sup> and mass transfer, especially the oxygen resistance, becomes more and more rate limiting<sup>4</sup>. Thus, the power density in this potential regime depends on the ECSA per mass. As a consequence, it is key to achieve improved SA and high ECSA for realizing high performance in an automotive PEMFC application. As also outlined in recent work of Kongkanand et al.<sup>4</sup>, a sole focus on optimizing the SA at low overpotentials (0.9 V<sub>RHE</sub>) disregards the oxygen mass transport resistance occurring at higher overpotentials. Additionally, it was found that ionomer-free catalysts such as nanostructured thin film (NSTF) catalysts by 3M have the lowest oxygen resistance<sup>4</sup>. For the goal of achieving high current densities of > 1.5 A cm<sup>-2</sup> in fuel-cell operation, high dispersion of the catalytically active material (high ECSA) combined with a high specific activity is therefore crucial<sup>4</sup>.

Here, we present a concept for a self-supported catalyst capable of combining high SA with an unprecedentedly high ECSA. The catalyst is inspired by developments for carbon-free ORR catalysts, such as nanostructured thin film (NSTF) catalysts introduced by 3M<sup>9</sup>. Such catalysts utilize the high SA of bulk Pt<sup>9</sup>, but perhaps equally important and technologically relevant, they circumvent inherent carbon corrosion problems of standard carbon supported catalysts<sup>9</sup>. In addition, it was recently demonstrated that, in carbon-supported catalysts, not only oxygen diffusion can be rate limiting, but also proton

transport<sup>15</sup>. Using new catalyst layer engineering methods, oxygen-related mass transport resistances can be to some extent overcome<sup>16,17</sup>. A disruptive change of the catalyst layer concept towards self-supported catalyst layers would allow new operation regimes for PEMFCs with regard to temperature and humidification. The presented catalyst layer consists of a nanoporous (np) Pt-CoO network, which is achieved by alternating magnetron sputtering of Co and Pt layers and subsequent acid leaching of the non-noble cobalt-oxide component, see methods section and supporting information (SI) for more details. This method was recently shown for synthesis of highly active self-supported iridium-based catalysts for oxygen evolution<sup>18</sup> and nanostructured Pt-Cu for ORR<sup>19</sup>, the latter presenting a low ECSA as compared to this work. Compared to complex wet-chemical approaches, the magnetron sputtering technique is easily up-scalable and allows the use of different substrates, such as glassy carbon, but also gas diffusion layers (GDLs), transfer foils or even membranes. The metal network is self-supported, i.e. it does not require any carbon support material, and is directly prepared onto such substrates. In the presented work, we used glassy carbon (GC) disk electrodes and GDLs as substrates. In Figure 2, scanning transmission electron microscopy (STEM) and scanning electron microscopy (SEM) micrographs, respectively, are shown to visualize the porous structure. In addition, a tomographic 3D movie is shown in the SI and Figure S1 – S4.



Figure 2. Morphology of self-supported nanoporous Pt-CoO networks a) Scanning transmission electron microscopy high-angle anular dark-field imaging (STEM HAADF) with elemental distribution of np Pt-CoO measured by EDX of b) Cobalt (red), c) Pt (green) and d) a combined elemental distribution graph of Pt and Co; e) Scanning electron microscopy (SEM) top-view and f) side-view

In order to investigate and optimize the np Pt-CoO network, various catalysts with different Pt:Co starting compositions as well as heat treatment steps were prepared as summarized in Table 1. It is demonstrated that, after acid leaching (see Methods section), the catalyst can still contain up to 28.4 at. % Co as indicated by energy dispersive X-ray spectroscopy (EDX). However, X-ray photoelectron spectroscopy (XPS) reveals that, in all cases, the near surface concentration of Co is very low, see Table 1; an observation that is in agreement with the formation of Pt-CoO encapsulated structures, where Co is preferably located in the core and Pt dominates at the surface. A similar conclusion is suggested by STEM HAADF micrographs, see Figure 2 d, showing the elemental distribution at nanometer scale. The presence of cobalt oxide under electrochemical conditions is confirmed by *in situ* X-ray absorption spectroscopy (XAS), see Figure 7 and SI, whereas small angle X-ray scattering (SAXS) indicates that a nanoporous structure is formed with a

bone size of about 3-5 nanometer; see Figure S5. This result is also in agreement with the TEM analysis, see Figure S2. Finally, it is found that a heating step in the synthesis leads to a shift in the Pt 4f binding energy in the XPS spectra towards higher energies.

Table 1: Basic properties of differently prepared nanoporous Pt-CoO networks; determined by EDX<sup>1</sup>; XPS<sup>2</sup> and SAXS<sup>3</sup>.

	as prepared	Co content <sup>1</sup>	Co content <sup>2</sup>	Peak position <sup>2</sup>	bone
	Co:Pt weight	after leaching	after leaching	after leaching	size <sup>3</sup>
	ratio <sup>1</sup>	[at. % of Pt and Co]	[at. % of Pt and Co]	Pt 4f Pt (0)	[nm]
				[eV]	
Pt-CoO 1	6.0	17.5	10.9	71.17	3.77
Pt-CoO 2	9.1	28.4	7.4	71.10	5.16
Pt-CoO 3	14.3	10.4	17.2	71.19	4.43
Pt-CoO heat 1	14.3	21.2	6.4	71.29	4.55
Pt-CoO heat 2	14.3	26.5	3.5	71.51	4.24

Testing the electrochemical performance of the catalysts in a rotating disk electrode RDE half-cell configuration reveals remarkable oxygen reduction reaction (ORR) activity, see Figures 3 and 4. At 0.9  $V_{RHE}$ , specific ORR current densities between 3.8 and 5.5 mA cm<sup>-2</sup><sub>Pt</sub> are observed. This correlates to an increase by a factor of *ca*. 8 as compared to a state-of-the-art supported Pt/C catalyst (TEC10E50E from Tanaka TKK) and a factor of *ca*. 2-3 as compared to polycrystalline bulk Pt, which regularly serves as

benchmark for the ORR<sup>20</sup>. Such ORR specific activities are typical for highly active bulk Pt-alloy catalysts, such as Pt<sub>3</sub>Co-alloys. For instance, Stamenkovic et al. reported a specific ORR activity of around 4.5 mA cm<sup>-2</sup>Pt at 0.9 V<sub>RHE</sub> for a polycrystalline Pt<sub>3</sub>Co bulk alloy<sup>3</sup>. The high specific ORR performance can be related to an optimization of the binding energy of the catalyst surface with the reaction intermediate O, OH and OOH, which functions as descriptor in the optimization of Pt based ORR catalysts<sup>21</sup>. Alloying Pt with a second, less noble element, tunes the electronic structure of Pt and improves its ORR performance<sup>22</sup>. Furthermore, recently, it was shown that surface distortion has a positive effect on the ORR activity as well<sup>2</sup>. As mentioned above, the XPS results indicate that in particular a heat treatment of the Pt-CoO networks leads to such a change of the electronic properties of platinum. The main enhancement of the specific activity, however, is not assigned to a ligand effect. Instead, it is suggested that the observed increase in ORR activity is related to a structural effect of the Co-Oxide core on the Pt skin layer: i.e. the underlying Co-Oxide core causes a compressive strain on the outer Pt layer, which is generally known to increase the ORR activity<sup>23</sup>. Before discussing further experimental and theoretical evidence of this hypothesis, our efforts to test the stability of the np Pt-CoO networks are presented.



Figure 3. Pt surface area-based specific ORR activities of different np Pt-CoO networks. The value for Pt/C (TEC10E50E 2-3nm) is prepared in the same lab and taken from literature<sup>24</sup>. Due to the high reduction current at 0.9  $V_{RHE}$  the calculated SA is less reliable<sup>5</sup>, see also experimental data shown in Figure S8 – S11. Therefore, we include the SAs determined at 0.95  $V_{RHE}$  as well. The error bars are determined from three independent measurements. The label heat indicates that sample Pt-CoO 3 was heated to 573 K for 8 or 16 min, respectively Pt-CoO heat 1 and Pt-CoO heat 2; see also methods section.

Stability measurements were performed at 80°C applying the accelerated stress (AST) protocols proposed by the FCCJ<sup>25</sup> which are described in detail in the SI. A commercial Pt/C catalyst (TEC10E50E, 2-3nm) was used as benchmark. Applying an AST simulating load-cycles ( $0.6 V_{RHE} - 1 V_{RHE}$ ), a similar *ca*. 45 % loss in the ECSA is found for the np Pt-CoO networks as for the commercial Pt/C catalyst after 3,600 cycles. Nevertheless, the observed trend in ECSA loss during the AST treatment is different. While under the applied conditions for the Pt/C catalyst a more or less continuous "exponential decrease" in ECSA is observed, the main ECSA loss of the np Pt-CoO networks occurs in the first 600 AST cycles, see Figure S12. Thereafter, the decrease in ECSA becomes significantly slower. Investigating the degraded np Pt-CoO networks by TEM it is found that the degradation is related to an increase in the average bone size of the networks, most likely originating from the dissolution of platinum followed by re-deposition on the bones as well as dealloying, see Figure S13. Such an effect is in good agreement with previous reports stating that an AST treatment simulating load-cycles (AST 1) mainly induces Pt dissolution and electrochemical Ostwald ripening<sup>26</sup>. It should be further noted that the TEM micrographs indicate no sign of changes in the structural morphology of the np networks. Applying an AST treatment simulating start-stop cycles (1  $V_{RHE} - 1.5 V_{RHE}$ ; AST 2) the advantage of the np Pt-CoO networks over conventional catalyst concepts becomes evident, in particular the absence of a carbon support. While under the applied conditions the Pt/C benchmark degrades very rapidly, 53% of the ECSA is lost after 800 cycles, the self-supported np Pt-CoO network catalyst is significantly more stable and loses only 15% of its surface area, see Figure S14 and S15. Furthermore, the decrease in specific ORR activity is relatively moderate; after the AST 2 the SA is still 3.6 mA cm<sup>-2</sup><sub>Pt</sub> as compared to a SA of 2.4 mA cm<sup>-2</sup><sub>Pt</sub> after AST 1, see Figure S16.

Comparing the electrocatalytic mass activity to previous efforts to optimize ORR catalysts, it is essential to note that with the presented catalysts the high SA of the np Pt-CoO network is not achieved at the expense of a reduced ECSA, see Figure 4. Typically, going from bulk surfaces to nanoparticle-based catalysts, the specific ORR activity decreases with a decrease in active surface area (decrease in particle size<sup>27</sup>). In contrast, for the np Pt-CoO networks, the achieved surface area is comparable with highly dispersed carbon-supported Pt nanoparticles, i.e. > 100 m<sup>2</sup> g<sup>-1</sup><sub>Pt</sub>. Therefore, an initial loss of active surface area during "load cycle treatments" is difficult to avoid. However, also after degradation the catalyst remains superior to standard catalysts. The high surface area of the Pt-CoO networks can be correlated to their nanoporous nature and fine Pt-CoO "bone" structure with chain diameters of approximately 4 nm as measured by SAXS and TEM, see Table 1 and Figures S1 – S4. The pore diameter was averaged to be

approximately 16 nm, see Figures S1 and S3. An estimation based on TEM Tomography indicates that the mean volume of the bone structure is 20.4%, see Figure S4. This combination of a fine bone structure and relatively large pore volume allows high oxygen transport and high electronic conductivity. Furthermore, the presence of Nafion-ionomer in the catalyst layer of a fuel cell is expected to be less crucial as compared to standard Pt/C or Pt-alloy/C catalysts as the proton transport can be established via proton diffusion on the surface of the catalyst or via adsorbed water as emphasized by Debe et al.<sup>28</sup> for ionomer-free catalyst layers.



Figure 4. Comparison of ORR performance at 0.9 V<sub>RHE</sub> for different standard and model catalysts with the best performing np Pt-CoO network. a) Specific ORR activity (bar) and electrochemically available surface area (red cross), b) Pt mass related ORR activity calculated from a). The values for Pt/C (TEC10E50E 2-3nm), Pt<sub>3</sub>Co/C, Pt (polycrystalline, MA n/a) and Pt<sub>3</sub>Co (polycrystalline, MA n/a) are taken from literature: Pt and Pt/C<sup>24</sup>, Pt<sub>3</sub>Co<sup>3</sup>, Pt<sub>3</sub>Co/C<sup>29</sup>.

As self-supported nanoporous Pt-CoO networks are a new concept for ORR catalysts, we have extensively characterized them further by both ex situ and in situ XAS, ex situ pair distribution function analysis as well as computational modelling with density functional theory (DFT). The aim is to gain a further

knowledge on the structure-activity relationship as well as understanding better the nature of the active phase.



Figure 5. Bottom: Fourier Transform (FT) magnitudes of the  $k^2$ -weighted extended X-ray absorption fine structure (EXAFS) data of the leached np Pt-CoO networks, see color coding for the individual samples. A pure Pt foil (grey line) and Pt nanoparticles supported on carbon (Pt/C) with a mean particle size of 3-4 nm (dark grey line) serve as reference. Dashed lines indicate the typical atomic Pt-Pt distance for 3-4 nm Pt/C and Pt foil. It is noted, that the mean diameters of the Pt-CoO bones are similar like the mean diameter of the nanoparticles. The k-range for the Fourier-transformed EXAFS spectra at the Pt L<sub>3</sub> edge was from 3 to 13 Å<sup>-1</sup>.

Ex situ X-ray absorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) spectroscopy of the leached Pt-CoO networks at the Pt  $L_3$  edge and Co K edge, see Figure 5 and Figure 517—S19 and Table S1, indicate that the Pt is partially oxidized, while the Co largely appears as Co oxide,

see Figure S18. The atomic Pt-Pt distances of the network catalyst are significantly reduced as compared to those of a Pt foil or even Pt nanoparticles supported on carbon, indicating a mean Pt-Pt lattice strain, see Table S1. It should also be noted that no Pt-Co alloy is formed; the EXAFS spectra for the np Pt-CoO networks reveal the absence of hetero-metallic Pt-Co pairs. That is, the alternated magnetron sputtering and dealloying largely leads to mono-metallic Pt-CoO bone structures without alloy formation. PDF analysis of x-ray total scattering data (see Figure 6 and table S4 and S5) show that the leached, np Pt-CoO networks exhibit mainly an *fcc* structure corresponding to a strained Pt nanostructure with a smaller unit cell parameter compared to that of bulk Pt, confirming the EXAFS results. The structure is highly strained: While all peaks in the PDF can be assigned to an *fcc* structure, the apparent unit cell parameter changes as structure modelling are done over different ranges of the PDF (table S54 and S5). Such behavior may be explained by the presence of a cobalt or cobalt oxide core in the structures or a curvature of the thin bones in the structure. Note that the lower X-ray scattering power of Co compared to Pt and low concentration of Co makes it difficult to establish the structure of Co-containing phases from the current X-ray total scattering data.



Figure 6. A) Fit of the fcc structure to the experimental PDF from a leached Pt-CoO sample. The fcc structure fits to the most intense peaks in the low r-region. B) Zoom of the fit from (A) at the r-range from 18-40 Å, where the model fails to fit the experimental data. C) The high r data can be fitted by a fcc model, although with a smaller unit cell parameter.

The catalyst structure is also investigated by DFT calculations to model the strain effect of CoO core on the thin Pt skin, see Figure S20 and S21 and discussion in the SI. The calculations indicate that the strain (- 1.28%) weakens the adsorption of OH by 0.038 eV leading to an estimated reduction of around 38 mV in overpotential as compared to an unstrained Pt layer. The relative strain is in agreement with the one of electronically modified electrocatalysts to Pt as found for  $Pt_x\gamma^{30}$  and dealloyed Pt-Cu<sup>31</sup>. The calculations are in line with in situ EXAFS measurements, see Figure 7 and SI, which show that indeed on the np Pt-CoO networks the Pt-O coordination as function of the applied potential is significantly lower than on a Pt/C benchmark sample. As no contribution of Co-Pt pairs is observed from the ex-situ XAS data, we assumed that the coordination number of the Pt is altered by the anodic potential applied during the ORR, where oxygen species adsorb at the electrode surface. Therefore, the high specific ORR activity of the self-supported np Pt-CoO networks can be understood as a result of local strain due to decreased Pt-Pt distances in the Pt shell.



Figure 7: Potential dependent N(Pt-O)/N(Pt-Pt) ratio measured with in-situ X-Ray Absorption Spectroscopy. The coordination number of Pt-O and Pt-N are acquired by fitting the acquired EXAFS spectra. Pt-CoO (blue) and Pt/C 30wt% (black) shown at the measured potentials with the sweeping direction indicated by the arrows.

The combination of high SA with high ECSA leads to extremely high mass activities (MA) for the ORR. As summarized in Figure 8 and Table 2, the highest achieved MA reaches 8 A mg<sup>-1</sup><sub>Pt</sub> at 0.9 V<sub>RHE</sub>. This exceeds the performance of conventional, carbon supported Pt-alloy catalysts by a factor of *ca*. 15, see Figure 4. To the best of our knowledge, the mass activity of the here presented self-supported Pt-CoO are far superior to previously reported carbon-free ORR catalysts. Similar or higher mass activity was previously observed only for systems that require a carbon support, such as reported Pt-Ni nanoframes (Nanoframes)<sup>5</sup> or ultrafine jagged Pt nanowires (J-PtNWs/C)<sup>32</sup>.



Figure 8. Comparison of ORR mass activity (MA) at 0.9  $V_{RHE}$  and electrochemical active surface area (ECSA) for different supported and self-supported (underlined) catalysts and np Pt-CoO network. The inserted values are for Nanocages<sup>6</sup>, Nanoframes<sup>5</sup>, Pt<sub>2.5</sub>Ni<sup>7</sup>, Pt<sub>x</sub>Gd<sup>8</sup>, NSTF<sup>9</sup>, Pt/C<sup>24</sup>, J-PtNWs/C<sup>32</sup>, Pt<sub>x</sub>Y<sup>30</sup>, <u>PtCu<sup>19</sup></u>, PtCo NW<sup>33</sup>, <u>Aerogel<sup>34</sup></u>, Mo-Pt<sub>3</sub>Ni/C<sup>35</sup>, LF@PF<sup>36</sup>, Nanoplate<sup>37</sup> and PALD<sup>38</sup>.

	SA	SA	ECSA	ECSA	МА
	@0.9V <sub>RHE</sub>	@0.95V <sub>RHE</sub>	CO-Stripping	H-UPD	@0.9V <sub>RHE</sub>
	[mAcm <sup>-2</sup> Pt]	[mAcm <sup>-2</sup> Pt]	[m <sup>2</sup> g <sup>-1</sup> Pt]	[m <sup>2</sup> g <sup>-1</sup> Pt]	[Amg <sup>-1</sup> Pt]
Pt-CoO 1	4.57 ± 0.67	0.52 ± 0.09	101 ± 6	85 ± 10	4.60 ± 0.96
Pt-CoO 2	4.52 ± 0.30	0.51 ± 0.02	115 ± 4	111 ± 28	5.19 ± 0.52
Pt-CoO 3	3.77 ± 0.25	0.49 ± 0.06	152 ± 10	147 ± 28	5.74 ± 0.76
Pt-CoO heat 1	4.59 ± 0.17	0.59 ± 0.02	147 ± 6	134 ± 25	6.75 ± 0.51
Pt-CoO heat 2	5.38 ± 0.21	0.62 ± 0.01	156 ± 4	156 ± 27	8.37 ± 0.55

Table 2: Electrochemical properties of differently prepared Pt-CoO networks as stated in Table 1.

To conclude, the combination of high ORR SA and high ECSA leads to high MA values of the self-supported np Pt-CoO networks. The higher SA could be assigned to a slightly strained platinum surface which is caused by the cobalt-oxide core. Although being a model catalyst, this new type of self-supported catalyst is a significant step towards applications of PEMFCs because the key property of improved SA and high ECSA is not only important for obtaining high MA. It is expected to be also crucial for realizing such performance in a PEMFC application. Optimizing these kind of catalyst layers for PEMFCs will require larger catalyst layers that enable extensive MEA testing, as it has been done with carbon supported catalysts for decades. For more realistic benchmarking test, we also measured the np Pt-CoO network catalyst in a gas diffusion electrode half-cell setup<sup>14,39</sup>. As a promising result, it is confirmed that the activity of this catalyst is indeed superior to Pt/C at low temperature and dry conditions. However, under humidified conditions and high temperature such as 80 °C the catalyst layer has to be further optimized to avoid flooding. Nevertheless, the target in automotive industry is to develop PEMFCs operable above 100 °C to reduce problems in heat management, conditions at which flooding should not be a problem but carbon corrosion is expected to increase<sup>40</sup>. At such conditions, the carbon-free self-supported catalyst

will be a particularly interesting concept. The presented technology is industrially scalable and can be applied on all planar surfaces as gas diffusion layers, transfer foils or membranes. Thus, it combines all necessary ingredients to enable a further decrease in the Pt content of future PEMFC technology for mobile applications.

#### Methods

Sample preparation. Glassy carbon electrodes (3 mm and 5 mm diameter, HTW, Germany) were polished and ultrasonicated in acetone, isopropanol, ethanol and ultrapure  $H_2O$  (Millipore 18.2 M $\Omega$ cm resistance and 5 ppb TOC). The electrodes were thereafter transferred to a vacuum chamber and for 3 minutes, plasma-activated at a pressure of 20 Pa in  $O_2$  atmosphere and a radio frequency (RF) power of 300 W. Samples were then transferred to the deposition chamber. To prepare the Pt-Co catalysts, the reactor chamber was evacuated to a base pressure of  $5 \cdot 10^{-3}$  Pa. An argon plasma was ignited in the chamber at a working pressure of 5 Pa. First, a 15 nm Ti (99.9 % Mateck GmbH, Germany) interlayer was deposited onto the glassy carbon by magnetron sputtering. For the Pt-Co film deposition, two magnetrons were equipped with planar targets of Co (99.9 %, Mateck GmbH, Germany) and Pt (99.95 %, Junker Edelmetalle, Germany). They were located at the superior part of the recipient. The RF generators (Advanced Energy) had a driving frequency of 13.56 MHz. The recipient (Neoplas GmbH, Germany) was configurated in a way that the substrate holder is turned automatically towards the respective magnetron with the sputtering being initiated when the sample is in position below the magnetron. To obtain the nanoporous metal film, Co and Pt sputtering were alternated with the respective loading deposition of Pt and Co. The alternating process was repeated 11 times so that the total Pt loading of the as-deposited electrode was 22 µg<sub>Pt</sub> cm<sup>-</sup>  $^{2}_{eeo}$ . We varied the deposition time of Co to obtain different Co:Pt compositions, respectively 6:1, 9:1, 14:1 by weight. Directly after preparation Pt-CoO heat 1 and Pt-CoO heat 2 metal films were treated in the vacuum chamber at 573 K for 8 (heat 1) and 16 minutes (heat 2). After deposition of the Pt-CoO networks, the glassy carbon electrode was inserted into a Teflon holder. Then samples were electrochemically leached by inserting them under potential control into the electrochemical cell and cycling them between 0.05 V<sub>RHE</sub> and 1.00 V<sub>RHE</sub> with a scan rate of 50 mV s<sup>-1</sup>, directly after immersion into a de-aerated 0.1 M HClO<sub>4</sub> electrolyte until a stable cyclic voltammogram (CV) was achieved. As control the Pt and Co contents of the metal films were determined by EDX before and after leaching. Each sample was prepared and tested at least three times.

*Electrochemical investigations*. The tips were mounted as working electrodes into the rotating disk electrode and placed in a separate three-compartment electrochemical Teflon cell. The reference electrode was separated by a Nafion membrane from the working electrode<sup>41</sup>. All measurements were performed at room temperature using a Nordic-EC potentiostat, a saturated Calomel reference electrode and a carbon rod as the counter electrode. Prior to each experiment the potential was calibrated to the reversible hydrogen potential (RHE). The solution resistance was electronically compensated by a positive feedback scheme. The potentials in this study are given with respect to the RHE. 0.1 M HClO<sub>4</sub> was used as the electrolyte in all measurements. The electrolytes were made from MilliQ water (> 18.2 M $\Omega$ cm, < 5 ppb TOC) and ultrapure 70% HClO<sub>4</sub> (Suprapur, Merck). Specific activity (SA) was defined as the quotient of the kinetic activity as calculated by the Koutecky-Levich equation divided by the electrochemically active surface area (ECSA) determined by the CO stripping method. The ECSA was defined as the surface area determined by CO stripping divided by the measured Pt mass with ICP-MS after leaching and ORR testing. The mass activity (MA) was defined as specific activity multiplied by the ECSA. For the calculation of the ECSA we used the loading as measured by ICP-MS after leaching because more than half of the Pt was lost during leaching. For further information see Supporting Information.

SEM and TEM characterization. The SEM (JSM7500F, Jeol) was employed with a field-emission gun, a semi in-lens conical objective lens and a secondary electron in-lens detector for high-resolution and highquality imaging of structural features. The maximum specified resolution was 1.0 nm at 15 keV. Glassy carbon electrodes were directly positioned on the substrate holder without any preparative coating. The TEM (Jeol 2100) was operated at 200 kV. Samples were characterized by recording images with a minimum of 3 different magnifications in at least 5 different areas of the TEM grid. The size analysis was performed by measuring the size of the skeleton. The TEM grids were prepared as follows. The glassy carbon disc with the nanoporous catalyst layer was sonicated in ca. 200 – 500 µl isopropanol for 5 minutes until the isopropanol turned slightly grey indicating the dispersion of the sample. This dispersion was then drop casted onto a holey carbon support film of Cu 300 mesh grids (Quantifoil). In addition, we deposited Pt-Co templates directly onto TEM grids and leached them in 0.1 M perchloric acid prior to TEM investigations. The size of the different structures in the TEM images was analyzed using imageJ software by manually measuring the size of the different structures and classifying them into pore, bone and agglomerate.

Scanning Transmission Electron Tomography. The electron tomography data was acquired in HAADF STEM mode on a Thermo Fisher Titan (S)TEM instrument fitted with a field emission electron gun and operated at 300 kV. The electron probe size and convergence angle were ~ 1.5 Å and 17 mrad, respectively, and the HAADF collection angle was 50 mrad. Tilt series in the angular range -68° to +68° were acquired with 2° increment steps. The shift between the images in the tilt series and their rotation axis were corrected using the Thermo Fisher Inspect3D program. The SIRT algorithm was used to reconstruct the 3-dimensional (3D) structure from the aligned tilt series. The 3D visualization and volume rendering was performed in the program Aviso Lite 9.0.1.

Scanning Transmission Electron Microscopy High Angle Angular Dark-Field Imaging. STEM HAADF. The samples were measured by High Resolution Transmition Electron Microscope (HRTEM) Titan G2 (FEI) with Image corrector on accelerating voltage 80 kV. Scanning TEM (STEM) images were taken with HAADF detector Model 3000 (Fishione). The qualitative EDX mapping was obtained by Super-X system with four silicon drift detectors (Bruker) in STEM mode at 80kV (Titan G2, FEI). Thereby the collected data were evaluated by Cliff-Lorimer model and averaged at lateral maps resolution of 1 nm.

*EDX and XPS characterization.* Pt to Co ratios were analyzed by EDX and XPS. EDX was performed with the SEM (JSM7500F, JEOL) using 10 keV and higher beam current. X-Ray photoelectron spectroscopy (XPS)

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was carried out using an Axis Ultra (Kratos, GB) for element analysis and determination of electronic density of states. The spectra were recorded by means of monochromatic Al K $\alpha$  excitation (1.486.6 eV) with a medium magnification (field of view 2) lens mode and by selecting the slot mode, providing an analysis area of approximately 250  $\mu$ m in diameter. A pass energy of 80 eV was used for estimating the chemical elemental composition; 10 eV was used for the high-energy resolution of the Pt 4f region to investigate Pt chemical state.

*ICP-MS measurements.* The Pt loading of the catalysts was determined by using inductively coupled plasma mass spectrometry (ICP-MS) measurements. For this, the catalysts were digested in aqua regia freshly mixed with 30 % HCl (Suprapur, Merck) and 65 % HNO<sub>3</sub> (Suprapur, Merck) in a volumetric ratio of 3:1, respectively. This process was repeated once. Then the samples were dissolved in 5-10 ml dest. Aqua. The samples were thereafter diluted 1:10 for the ICP-MS analysis. The concentration of platinum in the diluted aqua regia solution was analyzed by ICP-MS (NexION 300X, Perkin Elmer) through a Meinhard quartz nebulizer and a cyclonic spray chamber, operating at nebulizer gas flow rates of between 1.00 and 1.02 L min<sup>-1</sup> (Ar, purity grade 5.0).

SAXS analysis. Small-angle X-ray scattering (SAXS) profiles were measured using the SAXSLab instrument at the Niels Bohr Institute. The instrument uses a Rigaku 40 W micro-focused Cu-source producing Xrays with a wavelength of  $\lambda = 1.54$  Å detected by a moveable Pilatus 300 k pixel-detector allowing different *q*-ranges to be measured. Here the magnitude of the scattering vector is defined as  $q = 4\pi \sin \Theta$ / $\lambda$  with  $\Theta$  being half of the scattering angle. The *q*-calibration of the instrument was done using silver behenate. The scattering data are analyzed using the following expression:

$$I(q) = Aq^{-n} + B + \frac{C}{(1 + R^2 q^2)^2}$$

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where *A*, *B*, *C* are constants. *n* is the Porod exponent of the power law describing surface scattering, *B* is a small constant term accounting for the background scattering from the thin quartz plates the samples are mounted on, and finally, the last term describes the scattering from the metal network with the main fit parameter, the network radius *R*. The term originates from pore structure analysis<sup>42</sup>, but from Babinet's principle the scattering signal is the same whether the pore is empty, or as in this case, filled.

*XAS analysis*. X-ray adsorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) measurements were carried out at the B18 beamline, Diamond Light Source (DLS), Didcot (England) and at the beamline SuperXAS X10DA, Swiss Light Source (Switzerland). The in situ XAS experiments were performed at the ROCK beamline, SOLEIL Light Source (France). The Co edge XAS spectra were recorded in fluorescence mode, while depending on the loading the Pt edge XAS spectra were measured in transmission and fluorescence mode. A Si(111) double crystal monochromator was used in combination with a Ge 36 element or 5-element detector for fluorescence measurements, while for the transmission the ionization chambers with lengths of 15 and 30 cm were filled with N<sub>2</sub>. The averaged XAS spectra were analyzed by using the IFEFFIT software package<sup>43</sup>. In principle, the raw spectra were aligned, averaged, background corrected and normalized by the edge step. After conversion of the energy units (eV) into photoelectron wave number *k* units (Å<sup>-1</sup>), the resulting  $\chi(k)$  functions for XAS spectra were weighted with  $k^2$  and then Fourier-transformed to obtain pseudo-radial structure functions (RSFs). EXAFS spectra of the pure Pt and Co foils as references were used to estimate the amplitude reduction factors (S<sub>0</sub><sup>2</sup>).

*X-ray total scattering.* X-ray total scattering data were obtained at beamline 11-ID-B at the Advanced Photon Source (APS), Argonne National Laboratory. The data were collected from a sample dispersed on Kapton foil, and the signal from the foil was subtracted before further treatment. The data were obtained using an X-ray wavelength of 0.2112 Å, using the RA-PDF setup<sup>44</sup>. PDF analysis was done in PDFgetX3<sup>45</sup> and PDFgui<sup>46</sup>. The show that the Pt networks contain nanocrystalline Pt in the expected *fcc* structure

(Figure 6b). Due to the much higher X-ray scattering power of Pt compared to Co, as well as the lower amount of Co present in the leached sample, Pt completely dominates the PDF. The refined unit cell parameter is 3.85 Å, which is much lower than the 3.92 Å reported for bulk Pt. This agrees with the EXAFS results, where short Pt-Pt distances were also observed. Considering the fit using the simple *fcc* Pt structure in Figure 6b, the model describes the intense peaks in the low *r*-region well, while the smaller features seen in the PDF from ca. 18 Å are not described by the model. However, if fitting the *fcc* structure to the range from 18-40 Å, the structure fits these features very well, just with a lower refined unit cell parameter of 3.83 Å. Further details on the PDF modelling is given in the supplementary information.

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