Tailoring three-phase microenvironment for high-performance oxygen reduction reaction in PEMFCs

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Proton exchange membrane fuel cells (PEMFCs) offer an attractive zero-emission mobile power source. However, the requirement of excessive platinum group metal (PGM) catalysts to facilitate the sluggish oxygen reduction reaction (ORR) in PEMFCs has prevented their widespread adoption. Despite tremendous progress in catalyst development with greatly increased catalytic activities, the reduction of PGM loading in practical PEMFCs remains a significant challenge. The ORR in PEMFCs occurs at a catalyst-electrolyte-gas three-phase interface, with multi-faceted challenges involving the activity of the catalysts, available active sites, and concerted transport of the reactants (oxygen, protons) to and removal of the product (water) from the active sites. The reduction of PGM loading reduces the number of catalytic sites and requires a higher reaction rate on each site to sustain the overall power output, which necessitates a faster delivery of the reactants to and removal of the product from each active site. A desirable interface must allow efficiently feeding oxygen and protons to the catalytic sites without starving the reaction and timely removing water to avoid interface flooding. Herein we report the design of the three-phase microenvironment in PEFMCs by tailoring the interactions between the carbon supports and the electrolyte ionomers. We show that the carbon surface with 2.6% oxygen interacts with the ionomers through both its hydrophilic and hydrophobic regions, creating favorable transport paths for rapid delivery of both oxygen and protons, and timely removal of water. Such an elaborated interfacial design allows reducing costly platinum catalysts together with state-
of-the-art performance, for the first time enabling PEMFCs with all key ORR catalyst performance metrics, including mass activity, rated power and durability, surpassing the U.S. DOE targets.

Proton exchange membrane fuel cells (PEMFCs) are of immense interest as a zero-emission, quiet mobile power source for automobiles, drones, submarines, and autonomous underwater vehicles\textsuperscript{1-3}. Platinum (Pt) group metals (PGMs), particularly Pt, are typically used as the catalysts in commercial PEMFCs, especially as the cathode catalyst for boosting the sluggish oxygen reduction reaction (ORR)\textsuperscript{3}. To reduce PGM usage in PEMFCs is essential for lowering the cost to enable more widespread adoption\textsuperscript{4,5}. To this end, intensive efforts have been devoted to developing new generations of electrocatalysts with greatly increased mass activities\textsuperscript{6-9}. However, such high mass activities are usually only achieved in half-cell rotating disc electrode (RDE) test where the mass transport limitation is compensated using Koutecký–Levich equation, and only represent the fundamental material limit of these new generations of electrocatalysts. To date, the performance advancements achieved in the half-cell RDE test can hardly be captured in practical PEMFCs, and the power density target set by the DOE for PEMFCs is rarely met or addressed. Overall, despite tremendous progress in catalyst development, the reduction of PGM loading in practical PEMFCs (e.g., to meet the United States DOE target of cathode loading level ≤ 0.1 mg\textsubscript{PGM}/cm\textsuperscript{2}) has met with much less success\textsuperscript{5}.

The ORR in a practical PEMFC occurs at a \textit{catalyst-electrolyte-gas} three-phase interface. The challenge is multifaceted, involving the activity of catalysts, available active sites, and concerted transport of oxygen and proton through the polymer electrolyte (ionomers) to reach the active sites and the removal of water from the active sites\textsuperscript{10}. The reduction of PGM loading reduces the number of catalytic sites and requires a higher reaction rate for each catalytic site to sustain the overall power output, which in turn requires faster delivery of the reactants to and removal of the product from each active site. Therefore, the mass transport resistance starts to play an increasingly limiting role in the electrodes with lower PGM loading\textsuperscript{5,11}. To this end, an elaborated \textit{catalyst-electrolyte-gas} three-phase interface design is necessary to facilitate the relevant mass transport processes for efficient ORR. Beyond a highly active and durable Pt-based catalyst\textsuperscript{12-16}, the gas permeation\textsuperscript{17-19}, proton conduction\textsuperscript{20,21}, and water removal\textsuperscript{22} are all equally important. Therefore, a desirable three-phase interface must include an efficient ORR catalyst with high intrinsic activity, while simultaneously allowing for rapidly feeding reactants (O\textsubscript{2} and proton) without starving the reaction and timely removing the product (water) to avoid interface flooding. This rarely addressed issue is the subject of this study.

Recognizing that the ionomers are mostly in contact with the carbon support, here we focus on chemical modification the carbon support to tailor carbon-ionomer interactions to create a favorable three-phase interface for facilitating the ORR in PEMFCs. Specifically, we used a series of carbon materials with different surface oxygen contents (2.6, 3.9, 12.1%, respectively) as the catalyst support to prepare cathode catalysts in membrane electrode assemblies (MEA) and evaluate their performance. Our systematic studies demonstrate that the catalysts supported on carbon with 2.6% surface oxygen display a state-of-the-art mass activity among all PGM-based (with PGM as the only active sites) MEAs reported to date, for the first time enabling PEMFCs with all key ORR catalyst performance metrics, including mass activity, rated power and durability, exceeding the targets set by the DOE. Our molecular dynamics (MD) simulations suggest that an optimal surface oxygen ratio on the carbon surface leads to favorable carbon-ionomer interactions\textsuperscript{23,24} and a microenvironment comprised of both hydrophobic and hydrophilic
paths, which simultaneously allow the efficient shuttle of oxygen molecules and protons to, and timely removal of water from the catalytic sites, providing a desirable microenvironment for highly efficient ORR$^{25}$.

To explore the impact of surface oxygen in carbon supports on ORR, we prepared catalysts on three types of carbon materials, each with different surface oxygen ratio (labeled as C1, C2, and C3). We employed synchrotron-based near-edge X-ray absorption fine structure (NEXAFS) spectroscopy to determine the ratio of surface oxygen on the surface of each carbon support. The carbon K-edge intensities are comparable for all samples (Fig. 1a), suggesting the identical carbon loading, whereas the oxygen K-edge intensity indicates that the oxygen/carbon (O/C) ratio qualitatively follows a trend of C1 > C2 > C3 (Fig. 1b). In addition, components corresponding with C=O, COOH, C-OH vibration modes are identified in both carbon K-edge and oxygen K-edge spectra, suggesting the presence of various oxygen forms on the surface$^{26-28}$. The oxygen-containing surface functional groups are also evidenced by attenuated total reflection (ATR) Fourier-transform infrared spectroscopy (FTIR). The band at ~1750 cm$^{-1}$ can be assigned to C=O, while the band from 1000 to 1300 cm$^{-1}$ can be assigned to C-O. We find that the sum of both intensities follows the trend of C1 > C2 > C3 (Fig. 1c, full spectra presented in Supplementary Fig. 1)$^{29-31}$.

![Fig. 1](image-url) **Fig. 1** | Characterization of carbon materials. a, b, synchrotron-based NEXAFS spectra for carbon K-edge (a) and for oxygen K-edge (b). The intensity is normalized by sample loading. c, ATR-FTIR spectra. d, e, synchrotron-based XPS spectra for carbon 1s (d) and oxygen O 1s (e). f, the surface oxygen ratio for three carbon samples evaluated; C1, C2, and C3.

The quantitative ratio of surface oxygen (O/C ratio) was obtained via synchrotron-based X-ray photoelectron spectroscopy (XPS). The incident beam energy was fixed at 620 eV, making the
inelastic mean free path (IMFP) about 0.6 nm and 1.2 nm for the oxygen 1s and carbon 1s photoelectrons generated, respectively. After normalization by photoionization cross-section, the surface oxygen ratios were determined to be 12.1%, 3.9%, and 2.6% for C1, C2, and C3, respectively (Fig. 1d-f). Except for the surface oxygen ratio, other key properties of all three carbon materials are comparable. Transmission electron microscopy (TEM) images reveal that C1, C2, and C3 show similar morphology of carbon spheres with an average diameter of 30 nm (Supplementary Fig. 2a). The N2 adsorption/desorption tests also reveal similar Brunauer–Emmett–Teller (BET) surface areas in the range of 755-766 m²/g and comparable pore size distribution for C1, C2, and C3 (Supplementary Fig. 2b, c, Table 1). X-ray diffraction (XRD) patterns also show identical features, suggesting that these carbon materials share the same layered carbon structure and interplane distance (Supplementary Fig. 2d, Table 1).

To investigate the impact of these carbon supports on fuel cell performance, we prepared PtNi ORR catalysts on these carbons (PtNi/C1, PtNi/C2, PtNi/C3). Specifically, PtNi nanoparticles were directly grown on the respective carbon supports (C1, C2, C3) through a solution-phase synthesis process. They were then annealed in a mixture of Ar and H2, followed by acid wash and further annealing in Ar and H2 (details noted in method part). The TEM studies reveal that the PtNi nanoparticles on three different carbon supports exhibit a similar spherical morphology with a comparable average diameter of 4.9-5.0 nm (Supplementary Fig. 3, 4). The X-ray powder diffraction (XRD) patterns of PtNi/C1, PtNi/C2, and PtNi/C3 are also comparable, which matches well with the comparable Pt ratios (72-74%) revealed by energy dispersive spectroscopy (EDS) (Supplementary Fig. 4, Table 2). In addition, PtCo/C3 was also synthesized with a similar catalyst size for comparison (Supplementary Fig. 3-4, Tables 2). These catalysts were tested by a rotating disk electrode (RDE) system for preliminary ORR performance screening (Supplementary Fig. 5, Table 3), in which they showed comparable half-cell performance.

The membrane electrode assembly (MEA) was prepared by coating the prepared catalysts directly on the proton exchange membrane as the cathode layer using an ultrasonic spray system. The catalyst coated membrane (CCM), gas diffusion layer (GDL), and gasket were assembled in a fuel cell fixture (details noted in method part). The Pt loading in all prepared MEAs is fixed at 0.10 mgPGM/cm² and 0.05 mgPGM/cm² for cathode and anode, respectively. For a fair comparison, all materials are the same in all prepared MEAs except for the cathode layer catalyst. Thus, the MEAs are labeled by the type of cathode catalysts. All performance metrics in MEA, including ORR mass activity (MA), rated power, and durability, etc. are tested and evaluated following the published DOE guidelines.

First, the MA of ORR catalysts in MEA was evaluated by normalizing the iR-free currents at 0.9 V in the H2/O2 test by the cathode PGM mass. The beginning of test (BOT) MAs are 0.44, 0.54, 0.68, A/mgPGM for PtNi/C1, PtNi/C2, PtNi/C3, respectively (Fig. 2a, Supplementary Table 4). It is important to note, although the PtNi nanocatalysts used in these different MEAs showed comparable ORR MA in RDE half-cell test (Supplementary Fig. 5, Table 3), the ORR MAs observed in MEAs (Fig. 2a) showed notable differences, indicating the mass transport in these MEAs rather than intrinsic catalytic activity plays a dictating role in the overall performance. Therefore, the observed trend of MA in PtNi/C1 < PtNi/C2 < PtNi/C3 indicates more efficient mass transport in PtNi/C3 than PtNi/C2 and PtNi/C1. Notably, the beginning of test (BOT) MAs of the PtNi/C2, PtNi/C3, PtCo/C3 all surpass the DOE target for MA (0.44 A/mgPGM). Impressively, the MA of PtCo/C3 (0.74 A/mgPGM) is 2.2 times of the MA (0.33 A/mgPGM) of the
benchmark Tanaka Kikinzoku Kogyo (TKK) Pt/C and represents the state-of-the-art MA for PGM-based ORR catalysts achieved in MEA tests to date (Supplementary Table 4).

The durability of the MEA performance is critical for long term stable operation of practical PEMFCs. To test the practical durability of MEA performance with reasonable experimental time in a laboratory setting, the US DOE has recently suggested a standard accelerated durability test (ADT) protocol, in which a square wave potential between 0.60 and 0.95 V (3-s hold at each voltage) is applied to the MEA for 30,000 cycles with a targeted end of test (EOT) MA above 60% of BOT MA, and a minimum value above 0.26 A/mgPGM. Although an ADT protocol with a triangle wave between 0.60 and 1.0 V (50 mV/s sweep rate) was commonly used in previous studies, the newly suggested square wave protocol shows five times acceleration factor over the triangle wave protocol and is proven to be much harsher than triangle wave protocol to better reflect the long-term durability of the MEA in practical devices. Notably, PtNi/C2, PtNi/C3, PtCo/C3 MEAs all retain more than 60% of initial MA (DOE target) after 30000 cycles square wave ADT, and with EOT MA exceeding the DOE target. To the best of our knowledge, this represents the first time that the MEA durability and EOT performance exceeds the DOE target after the suggested square wave ADT, highlighting the long-term durability of our MEAs for practical PEMFCs.

![Fig. 2](image_url)

**Fig. 2** | **Membrane electrode assembly (MEA) test.** a, Comparison of mass activities obtained in the H₂/O₂ test at the beginning of test (BOT; before ADT) and end of test (EOT; after ADT). b-e, H₂/Air test. b, polarization plots at BOT. c, power densities plots at BOT. d, polarization plots at EOT. e, power densities plots at EOT. f, Comparison of peak power densities. The MEA test was performed at 80 ºC and 150 kPa_{abs} pressure unless specifically noted.
To further evaluate the performance of these MEAs in fuel cell working environment, we next obtained the current/power polarization plot using air as an oxygen source (Fig. 2b). It was observed at low current densities, all our catalysts perform rather similarly and show higher performance than Pt/C. At higher current density where mass transport starts to play an increasingly important role, the differences start to show up for C1, C2, C3 supported catalysts: a much obvious voltage drop was observed in PtNi/C1 while PtNi/C2, PtNi/C3, and PtCo/C3 better maintained their performance in this mass-transport region (Fig. 2b). Such differences can be more clearly seen in the power density plots (Fig. 2c). Power density values, especially at high current densities, are good indicators for PEMFC performance related to the mass transports, including proton, oxygen transport, and water removal. The peak power densities of the prepared MEAs follow the trend of PtNi/C1 << PtNi/C2 < PtNi/C3 ≈ PtCo/C3 (Fig. 2c, f) which is similar to the trend of MAs (Fig. 2a), again supporting that mass transport is more efficient in C3-based catalyst layer than that of C2-based and much more than C1-based. Thus, the performance differences observed in both H2/O2 and H2/Air tests can all be attributed to the different mass transport limitations related to the different carbon supports used. We have further characterized the EOT performance following the aggressive square wave ADT protocol suggested by DOE to evaluate the durability of the MEAs. At the EOT, all MEAs based on our catalysts are significantly better than that using Pt/C (Fig. 2d, e). Compared to BOT, the peak power density follows the same trend of PtNi/C1 < PtNi/C2 < PtNi/C3, though the difference between PtNi/C3 and PtNi/C2 is smaller (Fig. 2e, f).

Furthermore, we have also evaluated the rated power of the MEAs. The rated power of an MEA takes into account of heat rejection, which is an important aspect during practical fuel cell operation as the mass transport induced power loss at high current density is converted to waste heat. Thus, the DOE sets a target of Q/ΔT < 1.45 kW/°C for heat rejection, which translates to the targeted rated power > 1 W/cm² measured at 0.67 V when working temperature is 94 °C (17,33,36) (see Methods for details). Thus, the rated power is an important figure of merit directly reflecting the performance of practical fuel cells at working conditions, which is, however, rarely reported. Significantly, our PtNi/C3 and PtCo/C3 MEAs deliver a rated power of 1.21 and 1.16 W/cm², respectively, both above the DOE target (1W/cm²) (Fig. 3). Furthermore, the EOT rated power of both PtNi/C3 (1.05 W/cm²) and PtCo/C3 (1.03 W/cm²) remained above the DOE target after the square wave ADT (Fig. 3b) (33), showing significant promise for long term stable operation. It is important to note that the rated power corresponds directly to the working performance of practical PEMFCs. The achievement of rated power target both before and after ADT clearly highlights the practical potential of our MEAs.

The voltage loss at a fixed current density from the BOT to EOT is another important measure characterizing the durability of the PEMFC operation. Significantly, the voltage losses for the PtNi/C3 and PtCo/C3 MEAs at 0.8 A/cm² after the square wave ADT is only 16.2 and 16.9 mV, respectively, far below the U.S. DOE target (< 30 mV) (Fig. 3c) (33). These evaluations attest the superior performance, especially the sustained high MEA performance after ADT, of these designed catalysts compared to previously reported catalysts (Supplementary Table 5) (17,37). Although our studies have focused only on the catalyst layer, there are excellent examples in the literature to improve MEA rated power by optimizing membrane, ionomer, gas diffusion layer, or gas flow pattern design (38–40). For instance, we expect further enhancement may be achieved if the state-of-the-art Gore ultrathin membrane is used (41).
**Fig. 3** Rated power of representative MEAs and EDS map analysis of cathode catalysts. a, H₂/Air polarization plots obtained at the temperature of 94 °C and pressure of 250 kPaₐbs for rated power evaluation. b, Comparison of rated power densities at BOT and EOT in for PtNi/C3, PtCo/C3. c, Voltage loss from BOT to EOT at a fixed current density of 0.80A/cm². d-g, STEM image and EDS elemental analysis of representative Pt-based nanoparticles. PtNi/C3 samples before MEA test d, at EOT e. PtCo/C3 samples before MEA test f, and at EOT g. Each panel consists of the STEM image, mapping image of individual elements, and combined mapping of both elements.

Furthermore, we performed TEM, scanning transmission electron microscopy (STEM), and EDS analysis on catalysts at EOT. The Pt/C exhibited significant size coarsening, with average size increasing from 3.8 ± 1.1 nm (before the test) to 6.3 ± 1.5 nm (EOT, after ADT), which can explain the 67% MA loss in H₂/O₂ test and the large voltage loss in the H₂/Air test (Fig. 2, Supplementary Fig. 6, Table 4). We also observed size coarsening in TEM images for the Pt-alloy nanocatalysts, but to a much less degree. The sizes of the original PtNi nanocatalysts prepared on C1, C2, C3 are 4.9 ± 1.0 nm, 4.9 ± 1.0 nm, 4.9 ± 1.1 nm, respectively, while the sizes of the corresponding PtNi nanoparticles after ADT are 6.1 ± 1.7 nm, 6.1 ± 1.6 nm, 6.1 ± 1.6 nm, respectively (Fig. 2, Supplementary Fig. 3). The size of PtCo nanoparticles grown on C3 is 5.0 ± 0.7 nm (before test and), and 6.0 ± 1.5 nm at EOT (Fig. 2, Supplementary Fig. 3). The coarsening of nanocatalysts leads to a decrease in surface active sites, which contributes to the observed performance degradation at EOT.

In addition, we also observed a loss of non-noble transition metal in the evaluated catalysts at EOT. The atomic ratio of Ni drops from 28.1% to 11.2%, 26.7% to 10.8%, 25.8% to 10.4% for PtNi/C1, PtNi/C2, PtNi/C3, respectively. The atomic ratio of Co decreases from 17.9% to 10.7% for PtCo/C3 (Supplementary Table 2). More interestingly, these nanocatalysts demonstrated excellent structural durability, as revealed in EDS elemental maps. The representative PtNi catalyst demonstrates a Pt-rich shell structure (Fig. 3d), which has been suggested to be important for high durability in Pt alloy nanocatalysts. The Pt-rich shell was retained at EOT but with increased...
thickness, as evidenced by the EDS map (Fig. 3e). Compared to the PtNi, PtCo shares a similar core-shell structure with a Pt-rich shell, but the distribution of Co is more uniform in the PtCo nanocatalysts both before the MEA test and at EOT, indicating that more Co remained near the surface after ADT. The loss of Co is indicated by a comparison of the relative intensity change of Co in the EDS maps (Fig. 3d, g). It is noted that the leached metal cations may also poison the ionomers and also contribute to the performance degradation after ADT (Fig. 2d-f)\textsuperscript{43}.

It is recognized that the three-phase interface is critical for MEA performance\textsuperscript{17}. However, due to the complexity of the interface and lack of reliable characterization techniques, there is little atomic-scale insight on the three-phase interface, especially the ionomer/substrate interface. For instance, the electron-beam based techniques suffer from beam-induced damage on ionomer\textsuperscript{44,45}. Soft X-ray based low radiation damage techniques provide the best resolution at sub-15 nm scale, which is far from the resolution needed for capturing the nanometer scale feature of the ionomer structure and distribution at the three-phase interface\textsuperscript{44,46}. Therefore, we performed molecular dynamics (MD) simulation to investigate the effect of the surface oxygen ratio in carbon support on the interaction between the support and the perfluorosulfonic acid (PFSA) ionomers at the atomic scale. We also studied the impact of the above interactions on fuel cell performance.

To build the carbon support model, we equilibrated the amorphous carbon portion in the model using Reactive Force Field (ReaxFF) based molecular dynamics calculations (using LAMMPS) (Supplementary Fig. 7a). Then, we added three top layers of graphitic carbon on the top of pre-equilibrated amorphous carbon to simulate the surface structure of carbon, as noted in the literature\textsuperscript{47,48}. This core-shell carbon model leads to a predicted density of 2.15 g/cm\textsuperscript{3}, with a predicted X-ray diffraction pattern with the (002) peak at 26° and (100) peaks at 42° diffraction angle, which are in good agreement with experimental observation (Fig. 1, Supplementary Fig. 7b). Then, the top graphitic surface was oxidized at four different surface oxygen ratios (0%, 2.5%, 4.0%, and 12.0%) with the distribution of C=O, C-OH, and COOH oxygenated functional groups based on experimental results. These sites are evidenced by corresponding bonds detected in NEXAFS (Fig. 1a, b)\textsuperscript{26,28}. Then, we equilibrated the predicted structure for 100 ps at 300 K (Supplementary Fig. 8) using ReaxFF reaction dynamics.

To describe the electrolyte, we used hydrated PFSA ionomers and equilibrated at 600 K for 200 ps. The PFSA ionomer includes two portions, the polar or hydrophilic portion composed of sulfonic groups (-SO\textsubscript{3}\textsuperscript{-}), and the non-polar or hydrophobic PTFE backbone. To determine the interaction between the PFSA ionomers and the carbon surface, we used ReaxFF MD to equilibrate the hydrated PFSA ionomer structures on top of four different carbon surfaces at 300K for 200 ps and then predicted their interaction energy (Supplementary Fig. 9)\textsuperscript{40}. The carbon surface with 2.5% oxygen shows stronger interaction compared to other carbon surfaces (0%, 4.0% or 12.0% oxygen) (Fig. 4a, Supplementary Table 6). It is also noted that the carbon with 2.5% surface oxygen demonstrates the largest number of ionomer atoms within 0.3 nm distance from the carbon surface, which reflects the interface energy since those atoms contribute to the bonding between the ionomers and the carbon surface (Fig. 4b). The stronger interaction between the hydrated PFSA ionomers and carbon surface (2.5% oxygen) leads to a more uniform distribution of the ionomers, which in turn alleviates the known mass transfer issue in the catalyst layer due to uneven distribution of the ionomers\textsuperscript{23,24}. Indeed, the PtNi/C3 (C3 contains 2.6% surface oxygen) MEA displays the smallest pressure independent oxygen transport resistance (R\textsuperscript{P-ind}) (Fig. 4d, Supplementary Fig. 10).
The interfacial interactions between the carbon surface and hydrated PFSA ionomers. a. The image of ionomer, water distribution on carbon surfaces after MD simulation. The bottom caption shows interface energy between the ionomers and the carbon surface with different oxygen contents (0%, 2.5%, 4.0%, and 12.0%). b. The number of atoms belongs to the ionomer distributed within 0.3 nm distance above carbon surfaces in simulation models (model size: 3.4 nm in width, 3.4 nm in depth). c. The average distance of water molecules within 0.3 nm above carbon surfaces. d. Pressure independent parts of oxygen transport resistance measured at 80 °C, RH 65% (details noted in Method part).

Additionally, our simulation shows that the water molecules are repelled further away from the carbon surface as the surface oxygen ratio decreases, as indicated by the average distance of water molecules above the carbon surface (within 0.3 nm) (Fig. 4c). Thus, the simulation suggests that the carbon with lower surface oxygen ratio may have a lesser problem of interface water flooding MEA (Fig. 4c)\(^{22}\). The calculated results show that the carbon surface with 0% oxygen interacts only with the hydrophobic PTFE part of the PFSA ionomer (via C-F and C-C interactions) (Fig. 4A, Supplementary Fig. 9). In contrast, the 12.0% oxygenated surface is highly hydrophilic and allows only the sulfonic group to bind on its highly oxygenated surface, with very few non-polar groups on the carbon sites (Fig. 4a, Supplementary Fig. 9). In contrast, 2.5% and 4.0% oxygenated surfaces have both hydrophilic (sulfonic-oxygenated carbon) and hydrophobic (C-F, C-C) interactions within the interface (Fig. 4a, Supplementary Fig. 9). Together, the C3 shows strongest interactions with both hydrophilic and hydrophobic parts of the ionomers, which simultaneously promotes both O\(_2\) and proton transfer: i.e., O\(_2\) prefers a hydrophobic region (PTFE backbone) and diffuses through it quickly while the proton (H\(_3\)O\(^+\)) and water prefer hydrophilic part (sulfonic acid side chain) for fast diffusion\(^{49}\). Thus the most desirable three-phase interface should involve a combination of both the hydrophilic and hydrophobic interface near or next the catalysts facilitating O\(_{ad}\), OH\(_{ad}\), and H\(_2\)O\(_{ad}\) formation\(^{25}\).

Together, this work demonstrates that carbon surface chemistry plays a critical role in the MEA performance of Pt-based catalysts in practical PEMFCs. We showed that the Pt alloy nanocatalysts on carbon support with an optimal surface oxygen ratio lead to state-of-the-art MEA performance, with all the key ORR performance metrics including mass activities, rated power, and durability, exceeding DOE targets (Table 1). We attribute this outstanding performance to the favorable
interaction between the carbon substrate and the ionomers, which results in a more uniform distribution of ionomers in the catalyst layer and a desirable microenvironment involving both hydrophilic and hydrophobic paths that facilitate the balanced supply of protons and oxygen molecules to the catalytic sites, as well as the timely removal of water molecules from the interface. These findings highlight the critical role of the three-phase microenvironments in facilitating catalytic reactions in a complex system, and opens a new pathway to greatly boost the performance of practical PEMFCs.

**Table 1.** Summary of DOE technical targets and the performance achieved in our MEAs with a tailored three-phase microenvironment. The cathode loading is maintained at 0.10 mg\textsubscript{PGM}/cm\textsuperscript{2} for all MEAs.

<table>
<thead>
<tr>
<th>Performance Metrics</th>
<th>Unit</th>
<th>DOE Target</th>
<th>PtNi/C3</th>
<th>PtCo/C3</th>
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<td>Mass Activity</td>
<td>A/mg\textsubscript{PGM} @0.9V iR free</td>
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<td>38%</td>
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<td>Rated Power</td>
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<td>1.21</td>
<td>1.16</td>
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<tr>
<td>Loss in voltage at 0.8 A/cm\textsuperscript{2} (EOT vs. BOT)</td>
<td>mV</td>
<td>&lt;30.0</td>
<td>16.2</td>
<td>16.9</td>
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</tbody>
</table>

**Methods**

**Chemicals and Materials**

Platinum (II) acetylacetonate [Pt(acac)\textsubscript{2}], nickel(II) acetylacetonate [Ni(acac)\textsubscript{2}], nickel(II) acetate tetrahydrate [Ni(Ac)\textsubscript{2}\cdot4H\textsubscript{2}O], cobalt (II) acetate tetrahydrate [Co(Ac)\textsubscript{2}\cdotH\textsubscript{2}O], benzyl acid (BA), Nafion dispersion, Aquion D83-06A ionomer dispersion were purchased from Sigma Aldrich. Bis(triphenylphosphine)dicarbonylnickel [(C\textsubscript{6}H\textsubscript{5})\textsubscript{3}P]\textsubscript{2}Ni(CO)\textsubscript{2}) was purchased from Alfa Aesar. N, N-Dimethylformamide (DMF), acetone, isopropanol (IPA) were purchased from Fisher Scientific. Ethanol was purchased from Decon Labs, Inc. SGL-29BC gas diffusion layer (GDL), polytetrafluoroethylene (PTFE) gasket were purchased from The Fuel Cell Store. Carbon black (Ketjenblack EC-300J) was obtained from Fitz Chem LLC. Water used was Ultrapure Millipore (18.2 MΩ-cm).

**Preparation of Carbon Materials (C1, C2, C3)**

C1 is prepared by annealing Ketjenblack EC-300J in the air at 400℃ for 3 hours. C2 is the original Ketjenblack EC-300J. C3 is prepared by annealing Ketjenblack EC-300J in a gas mixture of argon (Ar) and hydrogen (H\textsubscript{2}) at 800℃ for 10 hours.

**Synthesis of PtNi/C (C can be either C1, C2, C3)**

1. **Solution phase synthesis**

140 mg carbon black (C1, C2, or C3) and 28 mg [(C\textsubscript{6}H\textsubscript{5})\textsubscript{3}P]\textsubscript{2}Ni(CO)\textsubscript{2} were dispersed in 135 mL DMF under ultrasonication for 30 mins in a 325 mL pressure bottle (sealed). 190 mg Pt(acac)\textsubscript{2}, 110 mg Ni(ac)\textsubscript{2}\cdot4H\textsubscript{2}O, and 1030 mg benzoic acid were dissolved in 25 ml DMF in a 20 mL vial. Then the prepared solution noted above was added into the 325 ml pressure bottle and mixed with carbon black and [(C\textsubscript{6}H\textsubscript{5})\textsubscript{3}P]\textsubscript{2}Ni(CO)\textsubscript{2} solution. After ultrasonication for 5 mins, the sealed
pressure bottle was directly put into 140 °C oil bath and then slowly heated to 160 °C. The pressure bottle was then kept at 160 °C for 48 hours (hrs). After reaction finished, the catalyst was collected by centrifugation, then dispersed and washed with isopropanol and acetone mixture. The centrifugation and re-dispersion steps were repeated for three times. Then the catalysts were dried in vacuum at room temperature.

2. Pre-Acid Wash Annealing

The dried catalyst was then annealed at 250 °C for 2 hrs in Ar and H\textsubscript{2} mixture (the volume ratio of Ar/H\textsubscript{2} =500/1).

3. Acid Wash

About 200 mg of annealed catalyst is dispersed in 20 ml 0.05 M H\textsubscript{2}SO\textsubscript{4} in a 25 mL vial. The dispersion is then purged with N\textsubscript{2} till N\textsubscript{2} saturation. Then the vial is kept in 85 °C for 6 hours. The catalyst is collected through centrifugation when the acid wash is finished, then dispersed and washed with water. The centrifugation and re-dispersion steps were repeated for three times. Then the catalysts were dried in vacuum at room temperature.

4. Post-Acid Wash Annealing

The dried catalyst was then annealed at 200 °C for 2 hrs in Ar and H\textsubscript{2} mixture (the volume ratio of Ar/H\textsubscript{2} =500/1).

Synthesis of PtCo/C3

1. Solution phase synthesis

140 mg carbon black (C3) was dispersed in 135 mL DMF under ultrasonication for 30 mins in a 325 mL pressure bottle (sealed). 190 mg Pt(acac)\textsubscript{2}, 140 mg Co(ac)\textsubscript{2}·4H\textsubscript{2}O, and 1030 mg benzoic acid were dissolved in 25 ml DMF in a 20 mL vial. Then the prepared solution noted above was added into the 325 ml pressure bottle and mixed with carbon black solution. After ultrasonication for 5 mins, the sealed pressure bottle was directly put into 140 °C oil bath and then slowly heated to 160 °C. The pressure bottle was then kept at 160 °C for 48 hours (hrs). After reaction finished, the catalyst was collected by centrifugation, then dispersed and washed with isopropanol and acetone mixture. The centrifugation and re-dispersion steps were repeated for three times. Then the catalysts were dried in vacuum at room temperature.

2. Pre-Acid Wash Annealing

The dried catalyst was then annealed at 400 °C for 8 hrs in Ar and H\textsubscript{2} mixture (the volume ratio of Ar/H\textsubscript{2} =500/1).

3. Acid Wash

About 200 mg of annealed catalyst is dispersed in 20 ml 0.05 M H\textsubscript{2}SO\textsubscript{4} in a 25 mL vial. The dispersion is then purged with N\textsubscript{2} till N\textsubscript{2} saturation. Then the vial is kept in 85 °C for 6 hours. The catalyst is collected through centrifugation when the acid wash is finished, then dispersed and washed with water. The centrifugation and re-dispersion steps were repeated for three times. Then the catalysts were dried in vacuum at room temperature.

4. Post-Acid Wash Annealing

The dried catalyst was then annealed at 200 °C for 2 hrs in Ar and H\textsubscript{2} mixture (the volume ratio of Ar/H\textsubscript{2} =500/1).
**Characterization**

Transmission electron microscopy (TEM) images were taken with an FEI T12 transmission electron microscope operated at 120 kV. Atomic resolution high angle annular dark-field (HAADF) images, as well as energy-dispersive spectroscopy (EDS) maps, were taken using a JEOL Grand ARM300CF scanning/transmission electron microscopy (S/TEM) operated at 300 kV. HAADF images were also taken using an FEI TITAN operated at 200 kV at STEM mode. The TEM/STEM sample grids were prepared by dispersing the sample in a mixture of water and ethanol, then dripping the dispersion onto carbon-coated aluminum grids (Ted Pella, Redding, CA) using a pipette and drying under ambient condition. X-ray powder diffraction (XRD) patterns were collected with a Panalytical X’Pert Pro X-ray Powder Diffractometer with Cu-Kα radiation. The concentration of catalysts was determined by inductively coupled plasma atomic emission spectroscopy (ICP-AES, Shimadzu ICPE-9000) as well as by EDS coupled in FEI TITAN TEM. The data for the Brunauer–Emmett–Teller (BET) surface areas and pore distributions were acquired with a Micromeritics Tristar II 3020 surface area and porosity analyzer. ATR-FTIR spectra were recorded by using a horizontal reflection ATR accessory (PIKE Technology, MIRacle, ZnSe crystal) and a DTGS detector coupled to an FTIR spectrometer (Bruker, Tensor 27). The spectra are the result of averaging data from 1024 scans, taken with a 4 cm⁻¹ resolution. The synchrotron radiation based high-resolution XPS measurements were carried out at Taiwan Light Source beamline 09A1 of National Synchrotron Radiation Research Center (NSRRC), Taiwan. The measurements were conducted at room temperature. The photon energy for XPS measurements was set at 620 eV. The photon energies were calibrated by the Au 4f core level signal emitted from a clean gold foil electrically connected with the samples. The near-edge X-ray absorption fine structure (NEXAFS) spectroscopy measurements on Carbon and Oxygen K-edge were carried out on beamline 7.3.1 at the Advanced Light Source, Lawrence Berkeley National Laboratory, CA. The energy resolution on Carbon and Oxygen K-edge was set to 0.2 eV and recorded in total electron yield mode with careful energy calibration before and after the experiment.

**Electrode Preparation and Electrochemistry Test**

A typical catalyst ink was prepared by mixing 2 mg of catalyst powder (PtNi/C1, PtNi/C2, PtNi/C3, PtCo/C3) with 2 mL of ethanol solution containing 16 μL of Nafion (5 wt%) with 5 min ultrasonication time. The benchmark Pt/C ink was prepared by mixing 1.5 mg of Pt/C powder with 2 mL of ethanol solution containing 20 μL of Nafton (5 wt%) with 5 min ultrasonication time. Then, 10 μL of prepared ink was dropped onto a 5 mm diameter glassy carbon electrode (Pine Research Instrumentation). Estimation of platinum group metal (PGM) loading is based on the overall PGM ratio within catalyst determined by ICP-AES. The ink was dried under an infrared lamp. Then the electrode was ready for the electrochemical test.

A three-electrode cell was used to carry out the electrochemical measurements. The working electrode was a catalyst coated glassy carbon electrode. An Ag/AgCl electrode was used as the reference electrode. Pt wire was used as the counter electrode. Cyclic Voltammetry (CV) measurements were conducted in an N₂ saturated 0.1 M HClO₄ solution between 0.05 to 1.1 V vs. reversible hydrogen electrode (RHE) at a sweep rate of 100 mV/s. Electrochemical active surface area (ECSA) is measured by integrating the hydrogen underpotential deposition (Hupd) peak in CV, assuming 210 μC/cm²PGM. Oxygen reduction reaction (ORR) measurements were conducted in an O₂ saturated 0.1 M HClO₄ solution between 0.05 to 1.05 V vs. RHE at a sweep rate of 20 mV/s. Accelerated durability test (ADT) was performed in an oxygen saturated 0.1 M HClO₄ solution by
applying square wave potential sweeps between 0.6 to 0.95 V vs. RHE (0.6 V 3s and 0.95 V 3s) for 30000 cycles.

**Membrane Electrode Assemblies (MEA) Fabrication and Test**

A series of catalysts were investigated as the cathode catalyst layers in MEA testing. The catalyst inks were made by mixing the catalysts with the ionomer solution (Aquivion D83-06A, ionomer to carbon ratio = 1.0) and water-IPA solvent mixture, followed by sonicitating the dispersion in ice water for 1.5 hours. The fresh inks were then spray-coated onto the Johnson Matthey (JM) half catalyst-coated membrane (CCM) by using a Sono-Tek ultrasonic spray system. The anode catalyst layer on the JM half-CCM had a platinum loading of 0.05 mg\text{PGM}\ cm^{-2}, and the membrane thickness was 18 µm. The cathode catalyst loading was controlled to be 0.10 mg\text{PGM}\ cm^{-2}, which was confirmed by the ICP measurements. The fabricated CCM was dried in a vacuum oven at 80 °C for 24 hours to evaporate the solvents completely. Two 3-mil PTFE sheets were used as the gaskets for both the anode and cathode electrodes. The SGL-29BC gas diffusion layer (GDL), PTFE gaskets, and prepared CCM were used to make the MEA. Then, MEAs were loaded in Fuel Cell Technology 5 cm² single-cell hardware and tested in the Scribner 850E fuel cell test stand under 80 °C, 150 kPa\text{abs} (abs: absolute; all pressure noted in this work refer to the absolute pressure) and 100% relative humidity (RH), following the U.S. Department of Energy (DOE) fuel cell test protocol. The gas flow rate is 1500/2000 sccm for anode/cathode, respectively. The ADT for Pt-based catalysts was performed using a square wave voltage from 0.6 V to 0.95 V with a duration of 3 seconds for each voltage level, according to the U.S. DOE MEA ADT protocol for PGM based catalysts. Each test was run up to 30,000 cycles at 80 °C, 150 kPa\text{abs}, 100% RH with H₂/N₂ flow 100/100 sccm for anode and cathode, respectively. The MEA’s beginning of test (BOT) and end of test (EOT) performance metrics, such as polarization curves, mass activity, specific electrochemical surface areas, high-frequency resistance (HFR) by alternating current (AC) impedance and H₂ crossover (H₂x-over), were recorded for analysis. ECSA of the cathode catalyst in MEA is measured by the potentiosat associated with the 850e fuel cell station using a CV scan. The ECSA is determined by integrating the hydrogen underpotential deposition (H upd) peak in CV, assuming 210 µC/cm²\text{PGM}.

**Rated Power Test for MEA**

MEAs employing PtNi/C3 or PtCo/C3 as a cathode catalyst were prepared in a similar way as noted above. The BOT rated power of prepared MEA was tested in Scribner 850E fuel cell test stand under 94 °C, 250 kPa\text{abs}, and 100% RH. The H₂/Air gas flow rate is 835/2000 sccm for anode/cathode, respectively. Then, the durability test, which involves 30,000 cycles of square wave (0.6 V maintained 3s, 0.95 V maintained 3s) at 150 kPa\text{abs}, 80 °C, is performed for the tested MEA. The H₂/N₂ flow (100% RH) is 100/100 sccm for anode and cathode, respectively. Then EOT rated power of the MEA was tested under 94 °C, 250 kPa\text{abs}, and 100% RH. The H₂/Air gas (100% RH) flow rate is 835/2000 sccm for anode/cathode, respectively.

The U.S. DOE assumes 90 kW stack gross power (P stack) required for 80 kW net power and sets a heat rejection target, \(\frac{Q}{\Delta T_i}\) of <1.45. The rated power is measured at rated voltage (V\text{rated}), which can be defined by the following equation 1 (Eq. 1)

\[
\frac{Q}{\Delta T_i} = \frac{P_{\text{stack}}(1.25 - V_{\text{rated}})}{V_{\text{rated}}(T_{\text{stack}} - T_{\text{ambient}})}
\]  

Eq. 1
The U.S. DOE protocol assumes $P_{\text{stack}} = 90$ kW and $T_{\text{ambient}} = 40\, ^\circ\text{C}$, if $T_{\text{stack}} = 94\, ^\circ\text{C}$, $Q/\Delta T_i = 1.445$, which meets the target ($Q/\Delta T_i$ of $\leq 1.45$), the $V_{\text{rated}}$ will be around 0.67 V. Thus, the rated power will be measured at 0.67 V.

**Oxygen Transport Resistance Test for MEA**

The MEA prepared by Ford is tested at UCLA for oxygen transport resistance. The MEA is tested at 80 °C with 65% RH for both anode and cathode. Ultrahigh pure hydrogen was applied in the anode. The mix-gas of O$_2$ and N$_2$ was pre-mixed by Airgas, Inc., and the mix-gas cylinders were used as a gas source. The dry O$_2$ mole ratios for pre-mixed gases are 0.981%, 1.491%, and 2.001%, which is determined by Airgas, Inc. In order to identify the pressure independent part of the total oxygen transport resistance, the test was performed at a total pressure of 110, 150, 190, 230, and 270 kPa$_{\text{abs}}$. The total oxygen transport resistance can be approximately obtained by the following Eq. 2.

$$R_{\text{total}} = \frac{4F C_{\text{O}_2}}{i_{\text{lim}}} = \frac{4F}{i_{\text{lim}}} \times \frac{P_{\text{abs}} - P_{H_2O}}{RT} \times x_{\text{O}_2-\text{dry}}$$

**Eq. 2**

$R_{\text{total}}^O_2$ presents the total oxygen transport resistance. $C_{\text{O}_2}$ represents oxygen concentration. $P_{\text{abs}}$ is the total pressure. $P_{H_2O}$ is the pressure of water vapor at testing temperature. $x_{\text{O}_2-\text{dry}}$ is the dry O$_2$ ratio. $i_{\text{lim}}$ represents the limit current density.

The limit current density is identified as the largest current density (considering the absolute value) obtained through linear voltage sweep (0.12 to 0.41 V, 2 mV/s) using the 885 potentiostat attached to the 850e fuel cell station. The total oxygen transport resistance can be separated into pressure dependent part ($R_{\text{P-d}}$) and pressure independent part ($R_{\text{P-ind}}$), which represents the Knudsen diffusion and diffusion of oxygen through the ionomer layer $^{50,51}$. The relationship of total oxygen resistance ($R_{\text{total}}$), pressure dependent part of oxygen transport resistance ($R_{\text{P-d}}$), and pressure independent part of oxygen transport resistance ($R_{\text{P-ind}}$) can be expressed in the following Eq. 3.

$$R_{\text{total}} = R_{\text{P-d}} + R_{\text{P-ind}}$$

**Eq. 3**

**Computational details**

The calculations were performed by using Reactive Force Field (ReaxFF) implemented in the molecular dynamics package LAMMPS $^{52,53}$, which differs from traditional unreactive force fields by

- Describing the electrostatic interactions using a distributed atom size charge on each atom that is allowed to change on every step of MD, allowing the electrostatic to change as reactions occur.
- Allowing all valence interactions to break, going to zero at large separations
- Describing the bond breaking using a bond distance dependent bond order and a bond order dependent bond energy.
- Including electrostatic and van der Waals interactions between all atoms (even bonded ones) since the valence bonds can break. Since the charges are distributed over the size of the atoms Coulomb interactions are shielded, going to a constant at short distances (point charges lead to infinities)
Thus, during MD simulations of reactive systems, many bonds may form or break.

All ReaxFF MD simulations were performed with 0.2 fs time step to allow charges and bonds to be described properly as bonds are broken and formed.

We used 2.15 g/cm$^3$ carbon density in a three-dimensional (3D) periodic MD simulation box consisting of around 3000 carbon atoms. This system was melted by heating to 6000 K for 5 ps and ten quenched to 300 K over 5.7 ps (1000 K/ps), followed by equilibrated at 300K for 100 ps to obtain amorphous carbon. Then we introduced 3 graphitic layers top of the amorphous carbon, heated to 3500 K for 100 ps for equilibration, and quenched again to 300K at 100 K/ps rate.

The surface layer was oxidized with 0%, 2.5%, 4.0%, and 12.0% oxygen to carbon ratio to simulate experimental observation. Here we used the ratios of functionalized C from experimental results. The hydrated ionomer was heated and equilibrated at 600 K for 100 ps and then cooled down and further equilibrated at 300 K for 100 ps. Then the hydrated PFSA ionomer was placed on the top of the carbon surfaces having different oxidized graphitic surfaces and equilibrated for 200 ps at 300K (12 H$_2$O/sulfonic acid [-SO$_3$] at ionomer and carbon interface).

The interaction between hydrated perfluorosulfonic acid (PFSA) ionomer (based on the molecular structure of Nafion from previous simulations \textsuperscript{49}) and the carbon surface was calculated via energy minimization and equilibration. The interface energies were calculated by using the Eq. 4.

$$\text{Interface energy (kcal/area)} = \frac{(E_{\text{Total}}(\text{carbon+Nafion}) - E_{\text{carbon}}(\text{carbon subs}) - E_{\text{Nafion}}(\text{Nafion}))}{\text{Interface area}} \quad \text{Eq. 4}$$

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at [www.nature.com](http://www.nature.com).

**Reference:**

11 Inoue, G. & Kawase, M. Effect of porous structure of catalyst layer on effective oxygen diffusion coefficient in polymer electrolyte fuel cell. J. Power Sources 327, 1-10, (2016).

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**Author Contributions** Z.Z. and Y.H. conceived the idea and designed the experiments. Z.Z. conducted the materials preparation, characterization, fuel cell test at UCLA. W.X. Zeyan L. J.Z. contributed to the experiments at UCLA. M.D.H., B.M. conducted the MD simulations while Zhengtang L., W.A.G. supervised the simulations. C.X. Zijie L. J.Y. conducted the fuel cell test in Ford. Y.L. conducted synchrotron-based NEXAFS experiment, and J.G. supervised the experiment. S.H. performed the synchrotron-based XPS test. I.L. conducted the ATR-FTIR test, and F.Z. supervised the work. W.G. conducted the EDS mapping analysis, and X.P. supervised the work. Z.Z., X.D., Y.H. wrote the paper with substantial input from other authors. Y.H. supervised the whole project.

**Competing interests** A U.S. provisional patent, “High performance platinum-based catalyst combined with carbon support engineering” (application No. 62/905,564), has been filed by UCLA. The processing conditions and related characterization results of the carbon materials have been submitted for a patent application. The rest of the data and figures in this work have not been presented or published elsewhere.
Data availability All data needed to evaluate our conclusions are found in the main text and the Supplementary Information. Further data related to the paper are available from the corresponding authors on reasonable request.

Additional information

Supplementary Information is available for this paper online

Correspondence and requests for materials should be addressed to Y. H.

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Supplementary Information

Tailoring three-phase microenvironment for high-performance oxygen reduction reaction in PEMFCs
Supplementary Fig. 1 | ATR-FTIR spectra of our three carbon materials with surface oxygens.
Supplementary Fig. 2 | Characterization of carbon materials. a, TEM images. b, XRD spectra. c, N₂ adsorption-desorption isothermal plots. d, pore size distribution calculated from density function theory (DFT) model using nitrogen isothermal plots in c.
Supplementary Fig. 3 | TEM images of catalysts. Catalysts before MEA test a, PtNi/C1, b, PtNi/C2, c, PtNi/C3, d, PtCo/C3. Catalysts after 30k square wave cycles in MEA test e, PtNi/C1, f, PtNi/C2, g, PtNi/C3, h, PtCo/C3.
Supplementary Fig. 4 | Characterization of prepared catalysts. 

a, The diameter of nanoparticles based on TEM images in Supplementary Fig. 3. 

b, EDS composition. 

c, XRD spectra of prepared catalysts.
Supplementary Fig. 5| Rotating disk electrode (RDE) test of catalysts. a, cyclic voltammetry curves of tested in N₂ saturated 0.1 M HClO₄. b, ORR polarization curves in O₂ saturated 0.1 M HClO₄ (1600 rpm). c, comparison mass activities (MA) and specific activities (SA) d, comparison of retention ratios of mass activities.
Supplementary Fig. 6 | TEM images of commercial Pt/C catalyst. a, before MEA test. b, after 30k square wave cycles in the MEA test.
Supplementary Fig. 7 | Predicted Core-shell like carbon substrate structure with its predicted XRD pattern. **a**, The predicted carbon substrate has a structure similar to that obtained by experimentally. It consists of three layers of graphitic carbon on top of amorphous carbon on the bottom. We heated the amorphous carbon to 6000 K and quenched to 300 K, followed by heating to 3500 K temperature followed by quenching at 300 K at 100 K/ps rate. The final carbon density in our structure is 2.15 g/cm$^3$. **b**, The predicted X-ray diffraction pattern (XRD), which is similar to experimental data as shown in Supplementary Fig. 2b. The XRD used Cu source radiation with powder diffraction technique at 0.05 step size.
Supplementary Fig. 8| The carbon surface oxidation patterns used in the MD simulations. a, pure graphitic surface (0% oxygen). b, c, and d are 2.5%, 4.0% and 12.0% oxygen to carbon ratios. All structures were minimized and equilibrated for 100 ps at 300K temperature and 0.2 fs time step. (coffee: carbon, red: oxygen, off-white: hydrogen).
**Supplementary Fig. 9** Predicted structures from ReaxFF MD. The Hydrated PFSA ionomer structure contains non-polar hydrophobic parts (polytetrafluoroethylene [PTFE] backbone) and hydrophilic sulfonic acid (-SO$_3^-$) parts. **a,** The side view of PFSA ionomer molecules with a cluster of waters (12 H$_2$O/-SO$_3^-$) at the carbon ionomer interface. **b,** The top view of the panel **a.** The structures were equilibrated at 600 K for 200 ps and cooled down to 300 K at 0.2 fs time step (gray: carbon; light cyan: Florine; red: oxygen; white: hydrogen; yellow: sulfur).
Supplementary Fig. 10 | The measured total oxygen transport resistances in MEAs plotted with the total pressure during the measurement. a, PtNi/C1, b, PtNi/C2, c, PtNi/C3. The total oxygen transport resistance can be separated into pressure dependent part ($R_{P-d}$) and pressure independent part ($R_{P-ind}$), which represents the Knudsen diffusion and diffusion of oxygen through the ionomer layer in the catalyst layer\textsuperscript{50,51}. Thus, the pressure independent part is critical for oxygen transport in an operating fuel cell. The relationship of total oxygen resistance ($R_{total}$), pressure dependent part of oxygen transport resistance ($R_{P-d}$), and pressure independent part of oxygen transport resistance ($R_{P-ind}$) can be expressed in the following Eq. S1 (noted in the Method part “Oxygen Transport Resistance Test for MEA”).

$$R_{total} = R_{P-d} + R_{P-ind}$$  \hspace{1cm} \text{Eq. S1}
Supplementary Table 1 | BET surface area and XRD interlayer distance of carbon materials.

<table>
<thead>
<tr>
<th>Sample</th>
<th>BET surface area (m$^2$/g)</th>
<th>XRD interlayer distance (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>766</td>
<td>0.355</td>
</tr>
<tr>
<td>C2</td>
<td>761</td>
<td>0.355</td>
</tr>
<tr>
<td>C3</td>
<td>755</td>
<td>0.355</td>
</tr>
</tbody>
</table>
**Supplementary Table 2** | EDS composition analysis of catalysts before the MEA test and at the end of test (EOT) (after 30000 square wave ADT cycles in MEA test).

<table>
<thead>
<tr>
<th></th>
<th>Atomic Ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pt</td>
</tr>
<tr>
<td>PtNi/C1</td>
<td>71.9</td>
</tr>
<tr>
<td>PtNi/C2</td>
<td>73.3</td>
</tr>
<tr>
<td>PtNi/C3</td>
<td>74.2</td>
</tr>
<tr>
<td>PtCo/C3</td>
<td>82.1</td>
</tr>
<tr>
<td>PtNi/C1 MEA EOT</td>
<td>88.8</td>
</tr>
<tr>
<td>PtNi/C2 MEA EOT</td>
<td>89.2</td>
</tr>
<tr>
<td>PtNi/C3 MEA EOT</td>
<td>89.6</td>
</tr>
<tr>
<td>PtCo/C3 MEA EOT</td>
<td>89.3</td>
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</tbody>
</table>
**Supplementary Table 3** Specific activity (SA), mass activity (MA), and electrochemical active surface area (ECSA) comparison of catalysts based on RDE test results.

<table>
<thead>
<tr>
<th></th>
<th>SA (mA/cm$^2$)</th>
<th>MA (A/mg$_{PGM}$)</th>
<th>ECSA (m$^2$/g$_{PGM}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt/C</td>
<td>0.50</td>
<td>0.26</td>
<td>52.0</td>
</tr>
<tr>
<td>PtNi/C1</td>
<td>3.15</td>
<td>1.22</td>
<td>38.7</td>
</tr>
<tr>
<td>PtNi/C2</td>
<td>3.87</td>
<td>1.47</td>
<td>38.0</td>
</tr>
<tr>
<td>PtNi/C3</td>
<td>3.81</td>
<td>1.45</td>
<td>38.1</td>
</tr>
<tr>
<td>PtCo/C3</td>
<td>3.94</td>
<td>1.42</td>
<td>36.0</td>
</tr>
</tbody>
</table>
**Supplementary Table 4** | Comparison of the mass activity of tested catalysts and several representative catalysts reported in recent literature. The comparison includes the mass activity at the beginning of test (BOT, before ADT), the mass activity at the EOT, which is after ADT cycles, the MA retention ratio, ADT cycle protocol, and the cathode PGM loading.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Cathode Loading (mg_{PGM}/cm^2)</th>
<th>Mass Activity (A/mg_{PGM})</th>
<th>ADT cycle number</th>
<th>ADT test condition</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>BOT</td>
<td>EOT</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DOE Target</td>
<td>≤0.1</td>
<td>&gt;0.44</td>
<td>&gt;0.264</td>
<td>&gt;60%</td>
<td>30000</td>
</tr>
<tr>
<td>Pt/C</td>
<td>0.10</td>
<td>0.33</td>
<td>0.11</td>
<td>33%</td>
<td>30000</td>
</tr>
<tr>
<td>PtNi/C3</td>
<td>0.10</td>
<td>0.68</td>
<td>0.50</td>
<td>74%</td>
<td>30000</td>
</tr>
<tr>
<td>PtCo/C3</td>
<td>0.10</td>
<td>0.74</td>
<td>0.46</td>
<td>62%</td>
<td>30000</td>
</tr>
<tr>
<td>PtNi (P2-SA)</td>
<td>0.1</td>
<td>0.64*</td>
<td>0.52*</td>
<td>81%</td>
<td>30000</td>
</tr>
<tr>
<td>Pt ML/Pd</td>
<td>0.134</td>
<td>0.17</td>
<td>0.11*</td>
<td>65%</td>
<td>20000</td>
</tr>
<tr>
<td>D-PtCo</td>
<td>0.2</td>
<td>0.490</td>
<td>0.330</td>
<td>67.3%</td>
<td>30000</td>
</tr>
<tr>
<td>L1_{0}-FePt</td>
<td>0.103</td>
<td>0.210</td>
<td>0.230</td>
<td>109.5%</td>
<td>30000</td>
</tr>
<tr>
<td>L1_{0}-CoPt</td>
<td>0.105</td>
<td>0.560</td>
<td>0.450</td>
<td>80.4%</td>
<td>30000</td>
</tr>
</tbody>
</table>

\*Data extracted from plots.

# The square wave ADT represents the newly suggested protocols for long term durability evaluation \(^33,34\), and has five times acceleration than the typical triangle wave ADT protocol used previously \(^34\).
Supplementary Table 5 | Rated power reported in representative literature. Measurements are performed at 250 kPa\textsubscript{abs}, 94 °C.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Rated Power (W/cm\textsuperscript{2})</th>
<th>Loss in Voltage at 0.8 A/cm\textsuperscript{2} (mV)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>BOT</td>
<td>EOT</td>
<td></td>
</tr>
<tr>
<td>DOE Target</td>
<td>&gt; 1.00</td>
<td>&lt; 30.0</td>
<td>Ref.\textsuperscript{33}</td>
</tr>
<tr>
<td>PtCo/HSC-e</td>
<td>1.21</td>
<td>NA</td>
<td>Ref.\textsuperscript{17}</td>
</tr>
<tr>
<td>PtCo/HSC-f</td>
<td>1.26</td>
<td>NA</td>
<td>Ref.\textsuperscript{17}</td>
</tr>
<tr>
<td>PtNi/C3</td>
<td>1.21</td>
<td>1.05</td>
<td>this work</td>
</tr>
<tr>
<td>PtCo/C3</td>
<td>1.16</td>
<td>1.03</td>
<td>this work</td>
</tr>
</tbody>
</table>

NA: not available
Supplementary Table. 6| Interface energy predicted from ReaxFF MD for the hydrated PFSA ionomer on different carbon surfaces. (12 H₂O/−SO₃⁻ at the interface, Fig. 4).

<table>
<thead>
<tr>
<th>Percentage (%) of Surface Oxygen</th>
<th>Interface Energy (kcal/Å²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>-48.17</td>
</tr>
<tr>
<td>2.5</td>
<td>-50.69</td>
</tr>
<tr>
<td>4.0</td>
<td>-50.54</td>
</tr>
<tr>
<td>12.0</td>
<td>-48.53</td>
</tr>
</tbody>
</table>
Reference: