Native ligand carbonization renders common platinum nanoparticles highly durable for electrocatalytic oxygen reduction: annealing temperature matters

Zhicheng Li  
Fudan University

Jinxiang Zou  
Fudan University

Xiangyun Xi  
Fudan University

Pengshuo Fan  
Fudan University

Ye Peng  
Foshan University

Dustin Banham  
Foshan University

Dong Yang  
Fudan University  https://orcid.org/0000-0001-6527-1272

Angang Dong (✉ agdong@fudan.edu.cn)  
Fudan University  https://orcid.org/0000-0002-9677-8778

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Native ligand carbonization renders common platinum nanoparticles highly durable for electrocatalytic oxygen reduction: annealing temperature matters

Abstract:

Monodisperse nanoparticles have attracted great attention in fundamental and applied research. Current protocols for the synthesis of monodisperse nanoparticles typically involve hydrocarbon molecules as surface-capping ligands. Utilizing platinum (Pt)-based nanoparticles for the oxygen reduction reaction (ORR), however, hydrocarbon ligands must be removed in order to expose the surface sites. Here, highly active and durable Pt catalysts are realized without removing the ligands; instead, the native surface ligands are directly converted into uniform, bilayered graphitic shells conformally coated on individual Pt nanoparticles by simple thermal annealing. Strikingly, the annealing temperature is found to be a critical factor dictating the ORR performance of Pt catalysts. Pt nanoparticles thermally treated at 500 °C show a very poor ORR activity, whereas those annealed at 700 °C become highly active along with exceptional stability. In-depth characterization reveals that thermal treatment from 500 to 700 °C triggers the subtle structural reconstruction of carbon shells through graphitization, gradually opening up the porosity without affecting the carbon shell thickness. Additionally, the ligand-derived graphitic shells can effectively prevent Pt nanoparticles from detachment, agglomeration, and dissolution while largely maintaining the accessibility of surface sites. As a result, such graphitic-shell-coated Pt catalysts can exhibit superior long-term stability, largely retaining the activity after
20,000 accelerated durability test cycles in a membrane electrode assembly. Moreover, this ligand carbonization strategy is amenable to Pt-based alloy nanoparticles and can further be extended to modify commercial Pt/C catalysts with substantially enhanced durability. Our work shows that boosting the ORR performance of common Pt nanoparticles is certainly possible by harnessing the native surface ligands, thus offering a robust approach of designing highly durable catalysts for proton-exchange membrane fuel cells.

1 Introduction

As one of the most promising energy conversion devices, proton-exchange membrane fuel cells (PEMFCs) have been attracting much attention for many years. It is known that the sluggish cathodic kinetics of the oxygen reduction reaction (ORR) greatly limits the conversion efficiency of PEMFCs. To date, Pt (or Pt-based alloys) nanoparticles (NPs) loaded on a carbon support are still the most promising ORR catalysts due to their intrinsically high activity. However, Pt catalysts typically suffer from the poor durability during long-term operation, which is one of the critical issues hampering the widespread applications of PEMFCs. Under the harsh operation conditions of PEMFCs, Pt catalysts can easily lose the electrochemical surface area (ECSA) due to multiple reasons, including the Ostwald ripening and agglomeration of Pt NPs, detachment of Pt NPs from the carbon support, dissolution of Pt, and corrosion of the carbon support.

In the past few decades, many efforts have been dedicated to addressing the stability issue of Pt catalysts. Coating Pt NPs with a porous and electrochemically
stable encapsulating layer has proven to be a promising approach to enhance the ORR durability\textsuperscript{17}, as this can effectively prevent the agglomeration of NPs while simultaneously suppressing the dissolution of Pt, due to the combined barrier and confinement effects\textsuperscript{18}. Among various encapsulating materials explored, carbon shows great promise because of its high electrical conductivity, electrochemical stability, and relatively benign corrosion products\textsuperscript{18,19}. For instance, Kwon \textit{et al.} reported the synthesis of carbon-coated Pt NPs on carbon nanofibers using Pt-aniline complexes as the precursor\textsuperscript{18}. Hyeon \textit{et al.} presented the synthesis of intermetallic ordered FePt NPs coated with N-doped carbon shells derived from polydopamine, where the carbon shell thickness can be tuned by controlling the polymerization duration\textsuperscript{20}. Indeed, these carbon-coating strategies can dramatically enhance the durability of ORR catalysts as indicated by both rotating disk electrode (RDE) and membrane electrode assembly (MEA) tests. Despite the improved durability, the presence of a protective layer at the particle surface can inevitably block some surface sites and restrict mass transport\textsuperscript{21}, thereby adversely influencing the catalytic performance of Pt catalysts. Therefore, the ability to fine-tune the thickness and microstructure of carbon shells to achieve an optimized tradeoff between activity and durability, although challenging, is critical to taking full advantage of this carbon-coating strategy\textsuperscript{19,22}. Moreover, most previously reported carbon-coating methods involve the incorporation of a foreign species (usually a polymer) as the carbon source\textsuperscript{20,21}, which does not necessarily have a preferential affinity to Pt, thus making it challenging to achieve complete and uniform carbon shells coated on individual NPs.
On the other hand, exploiting monodisperse Pt NPs is an effective means to inhibit Ostwald ripening during ORR\textsuperscript{8,23-25}. To date, monodisperse NPs are typically obtained by colloidal synthesis, involving the use of hydrocarbon ligands to control the nucleation and growth kinetics of NPs\textsuperscript{24}. Typical ligands include oleylamine (OAm) and oleic acid (OA)\textsuperscript{24-26}. The as-synthesized, ligand-capped NPs are highly soluble in nonpolar solvents\textsuperscript{25}, which is beneficial for subsequent solution processing. Despite these advantages, however, these long-chain organic ligands are highly insulating and can act as a dense barrier to restrict the accessibility of NPs\textsuperscript{27}, thus having a detrimental influence on their catalytic applications. As a consequence, to harness Pt NPs as ORR catalysts, the native surface ligands have to be replaced or removed in order to expose the surface sites\textsuperscript{28,29}. To this end, a number of techniques have been developed to treat Pt NPs, such as multiple cycles of washing with non-solvents\textsuperscript{30}, UV-ozone cleaning\textsuperscript{24}, and ligand exchange with small molecular (or ionic) species\textsuperscript{25,31}. Alternatively, the hydrocarbon ligands can also be removed by calcination in air\textsuperscript{32}. Although Pt NPs become active toward the ORR upon post surface treatment, the catalytic performance depends highly on the surface treatment methods. In addition, whether the native ligands are completely removed by these treatment methods remains an open question\textsuperscript{25}, which accounts for the performance inconsistency commonly encountered.

In contrast to previous methods where attempts are made explicitly to remove or replace the surface ligands\textsuperscript{24-32}, we report herein that the native ligands can be exploited to render common Pt NPs highly active and durable for the ORR. We show that simple thermal annealing can convert the hydrocarbon ligands into uniform, ultrathin graphitic
layers conformally coated on individual Pt particles, without sacrificing the size monodispersity of NPs. We highlight that the annealing temperature has a profound influence on the ORR performance of the resulting carbon-coated Pt catalysts. Specifically, thermal treatment of Pt NPs at 500 °C leads to highly dense carbon shells severely restricting the accessibility of surface sites, whereas increasing the annealing temperature triggers the reconstruction of carbon shells through graphitization, gradually opening up the transport channels for accessing the particle surface. In particular, thermal annealing of Pt NPs at 700 °C leads to bilayered graphitic shells with optimized porosity, yielding graphitic-shell-coated Pt catalysts with a superior ORR activity and stability. Importantly, this *in situ* ligand carbonization strategy is robust and broadly applicable for all Pt or Pt-based alloy NPs and can be further extended to boost the durability of commercial Pt/C catalysts. Our work demonstrates that the native surface ligands, previously considered detrimental for catalytic applications, can be harnessed to develop highly active and stable ORR catalysts for high-performance PEMFCs.

2 Results and Discussion

2.1 Synthesis and characterization of graphitic-shell-coated Pt catalysts

*Scheme 1* illustrates the preparation of the graphitic-shell-coated Pt catalysts through *in situ* ligand carbonization. Monodisperse Pt NPs (~5 nm in diameter) are synthesized via a standard thermal decomposition approach using OAm and OA as the capping ligands. Prior to ligand carbonization, the as-synthesized Pt NPs are first loaded onto a carbon support such as Ketjenblack (KB) by sonication (Pt@KB). Subsequently, the
Pt@KB is subjected to thermal annealing in a N\textsubscript{2} atmosphere at 500 °C for 1 h, which converts the OAm/OA ligands into conformal carbon shells. To obtain graphitic-shell-coated Pt catalysts, such carbon-coated Pt NPs are further annealed at various temperatures (600-800 °C) for another 1 h, which triggers the reconstruction of carbon shells to afford porous, graphitic layers\textsuperscript{34}. The resulting catalyst is labeled with Pt-T, where T is the annealing temperature. The typical Pt loading of Pt-T catalysts is ~35 wt.%, determined by thermogravimetric analysis (TGA) and inductively-coupled plasma mass spectrometry (ICP-MS).

**Scheme 1.** Schematic illustration of the synthesis of graphitic-shell-coated Pt catalysts through native ligand carbonization.

**Supplementary Fig. 1** shows a transmission electron microscopy (TEM) image of the as-synthesized 5 nm-Pt NPs with narrow size distribution. The high-resolution TEM (HRTEM) image of Pt-500 (Fig. 1a) reveals the presence of a complete carbon shell conformally coated on each NP (Fig. 1e), presumably arising from the carbonization of the OAm/OA capping ligands. A closer inspection of the HRTEM images indicates that the carbon shells are quite uniform in thickness, with the majority comprising two carbon layers. TEM also reveals that such carbon-coated Pt NPs are
highly sintering-resistant and can maintain their size upon thermal annealing up to 700 °C (Fig. 1a-c), although apparent particle agglomeration was observed when annealed at 800 °C (Fig. 1d). Notably, no substantial variation in carbon shell thickness was observed for Pt-600 and Pt-700 (Fig. 1f and 1g), whereas many Pt particles in Pt-800 are no longer coated by carbon shells (Fig. 1h). These results demonstrate that Pt NPs supported on KB can withstand thermal annealing without sacrificing the size monodispersity when the annealing temperature is below 700 °C, which is remarkable considering the high Pt loading (~35 wt.%). Presumably, the excellent thermal stability of the Pt NPs is attributed to the ligand-derived carbon shells, which effectively prevent the coarsening of NPs during thermal annealing. Control experiments show that the same Pt NPs loaded on KB, upon ligand removal by repeated washing with ethanol, underwent severe sintering when thermally treated at 700 °C (Supplementary Fig. 2). This illustrates the important role of the ligand-derived carbon shells in preventing the sintering of Pt NPs.

![Fig. 1](image)

**Fig. 1:** (a-d) TEM images and the corresponding particle size distributions (inset) of various Pt-T catalysts. (e-h) HRTEM images of various Pt-T catalysts.
The size variation of Pt NPs with respect to the annealing temperature was further investigated by X-ray diffraction (XRD). **Fig. 2a** shows the XRD patterns of various Pt-T catalysts, where the three major peaks located at 39.8°, 46.3°, and 67.5° were ascribed to the (111), (002), and (022) planes of Pt (JCPDS# 96-101-1108)\(^{35}\). The crystalline size of Pt NPs, calculated based on the Scherrer equation\(^{36}\), was essentially unchanged from Pt-500 to Pt-700. The significantly reduced peak width of Pt-800 was consistent with the particle agglomeration at 800 °C. To reveal the characteristics of the ligand-derived carbon shells, Raman spectra of Pt-500 and Pt-700 were comparatively analyzed. As shown in **Supplementary Fig. 3**, both Pt-500 and Pt-700 show the typical D and G bands at 1300 and 1590 cm\(^{-1}\), respectively, with the \(I_D/I_G\) ratio decreasing from 1.32 to 1.01, suggesting that the graphitization degree of Pt-T increases with the increased annealing temperature\(^{37}\). However, we note that the KB support may dominate the Raman signals of Pt-T, and therefore further characterization is needed to reliably evaluate the graphitization degree of carbon shells. As shown in **Fig. 2b**, the TGA curves of both Pt-500 and Pt-700 show the major weight loss from 300 to 450 °C, which corresponds to the oxidation of the KB support. For Pt-500, an additional weight loss of ~3.8 wt.% appears in the lower temperature region between 160 and 200 °C (indicated by the arrow in Fig. 2b). This weight loss, presumably ascribed to the oxidation of the less-graphitized carbon shells\(^{38,39}\), was not observed for Pt-700. This result suggests that as the annealing temperature increased from 500 to 700 °C the graphitization degree of carbon shells was increased to a level comparable to that of the KB support.
Fig. 2: (a) XRD patterns of various Pt-T catalysts. (b) TGA curves of Pt-500 and Pt-700. The blue arrow indicates the weight loss attributed to the carbon shells in Pt-500. (c) TEM and HRTEM (inset) images of Pt@SiO$_2$-700. (d) C K-edge XANES spectra of Pt@SiO$_2$-500 and Pt@SiO$_2$-700. (e) XPS survey spectra of Pt@SiO$_2$-500 and Pt@SiO$_2$-700. (f) N 1s, (g) C 1s, and (h) Pt 4f XPS spectra of Pt@SiO$_2$-500; (i) Pt 4f XPS spectrum of Pt@SiO$_2$-700.

To verify that the carbon shell is transformed from the OAm/OA ligands rather than the unstable species in the carbon support$^{40}$, Pt NPs were also loaded onto non-carbonaceous supports such as hollow SiO$_2$ spheres followed by thermal annealing under otherwise identical conditions. The product was named as Pt@SiO$_2$-T, depending on the annealing temperature (i.e., 500-700 °C). Similar to the Pt NPs loaded on KB, heat treatment of Pt@SiO$_2$ also led to carbon-coated Pt NPs (Fig. 2c), thus confirming that the carbon shell indeed originates from the surface ligands. Moreover, the absence
of carbon supports further allows us to disentangle the fine structural features of carbon shells with the varied annealing temperatures. **Fig. 2d** shows the C K-edge X-ray absorption near-edge structure (XANES) spectra of Pt@SiO$_2$-500 and Pt@SiO$_2$-700, where the characteristic $\pi-\pi^*$ (~285 eV) and $\sigma-\sigma^*$ (~291 eV) resonance peaks elucidate the typical graphene-like features exhibited by the ligand-derived carbon shells. For all Pt@SiO$_2$-T samples, the resonance absorption peak at 288.3 eV suggests the presence of heteroatoms (e.g., O) in carbon shells$^{41}$. In addition, a weak shoulder at around 283.2 eV is also observed in both cases. This feature is attribute to the surface defects, such as dangling bonds and amorphous carbon$^{42}$. With the increased annealing temperature, the pre-edge feature intensity of Pt@SiO$_2$-T slightly increases, suggesting that more defect sites have been created in the carbon shells$^{43}$.

The X-ray photoelectron spectroscopy (XPS) spectrum of Pt@SiO$_2$-500 shows the presence of Pt, C, O, and N (**Fig. 2e**), where N atoms (~2.3 at.%) should originate from the OAm ligands. As indicated by the high-resolution N 1s spectrum (**Fig. 2f**), N mainly exists in the forms of pyridine N (~398 eV) and pyrrole N (~400 eV). The high-resolution C 1s spectrum indicates the presence of C-N species in carbon shells (**Fig. 2g**)$^{44}$, indicating that the native OAm ligands were converted into N-doped carbon shells upon thermal annealing at 500 °C. Notably, the N signals of Pt@SiO$_2$-700 were essentially undetectable by XPS (**Fig. 2e**), suggesting that thermal annealing at 700 °C nearly completely removed N atoms from carbon shells, in addition to increasing the graphitization degree. For both Pt@SiO$_2$-500 and Pt@SiO$_2$-700, the Pt 4f signals can be deconvoluted into two doublets assigned to Pt$^0$ and Pt$^{2+}$ states (**Fig. 2h** and **2i**),
respectively, indicating that Pt NPs remained unchanged during thermal annealing\textsuperscript{20,45}.

### 2.2 Electrochemical tests of graphitic-shell-coated Pt catalysts

The ORR performance of various Pt-T catalysts was first investigated using RDE in O\textsubscript{2}-saturated 0.1 M HClO\textsubscript{4}. For comparison, commercial Pt/C (Johnson Matthey, particle size: \textasciitilde3 nm, Pt loading: \textasciitilde35 wt.\%) was tested as a standard (Supplementary Table 1)\textsuperscript{46}. As expected, the as-synthesized, ligand-capped Pt NPs (Pt@OAm) loaded on KB were essentially inactive as indicated by linear sweep voltammetry (LSV, Fig. 3a). The ORR inactivity of Pt@OAm was presumably caused by the surface hydrocarbon ligands, which effectively block the accessibility of surface sites, consistent with previous results\textsuperscript{25}.

Following the thermal annealing, all of the Pt-T catalysts demonstrated ORR activity, with the degree of activity varying depending on the annealing temperature that was used. Specifically, both Pt-500 and Pt-600 exhibited a relatively poor ORR activity, though the latter is slightly better in terms of the onset and half-wave potentials (Fig. 3a). Strikingly, when the annealing temperature reaches 700 °C, the resulting Pt-700 catalysts became highly active (Fig. 3a), with the half-wave potential higher than that of commercial Pt/C (0.902 V vs. 0.892 V). Further increasing the annealing temperature to 800 °C decreased the ORR activity. These results suggest that increasing the annealing temperature from 500 to 700 °C effectively activates Pt NPs. This annealing temperature-dependent ORR activity was also evidenced by cyclic voltammograms (CV). As shown in CV curves (Fig. 3b), Pt-700 showed the well-defined peaks of the monolayer hydrogen adsorption and desorption, whereas these
features were barely observable for other three catalysts. Likewise, the ECSA of Pt-700 (35.9 m$^2$ g$^{-1}$Pt) is much higher than that of Pt-500 (3.1 m$^2$ g$^{-1}$Pt) and Pt-600 (8.8 m$^2$ g$^{-1}$Pt) (despite all three having similar Pt particle sizes), indicating the greater exposure of surface sites as the annealing temperature increased from 500 to 700 °C$^{21}$.

Fig. 3: (a) ORR polarization curves and (b) CV curves of Pt@OAm and various Pt-T catalysts. (c) ORR polarization curves of Pt-700 before and after ADT cycles. (d) ORR polarization curves of Pt/C before and after ADT cycles. (e) Changes in ECSA for Pt/C and Pt-700 during the ADT. (f) Changes in MA (at 0.9 V) and SA for Pt/C and Pt-700 during the ADT.
More importantly, in addition to the excellent ORR activity, Pt-700 also exhibited superior long-term stability. Fig. 3c shows the polarization curves of Pt-700 according to the number of accelerated durability test (ADT) cycles. The half-wave potential of Pt-700 was nearly unchanged after 20k ADT cycles, indicative of the negligible activity degradation, which was also verified by CV (Supplementary Fig. 4). The ECSA, mass activity (MA, at 0.9 V), and specific activity (SA) of Pt-700 were largely retained as well after 20k ADT cycles (Fig. 3e and 3f), further confirming the negligible decrease of the accessible active sites. In comparison, the half-wave potential of commercial Pt/C was decreased by ~50 mV after 20k ADT cycles (Fig. 3d), along with an intensive degradation in ECSA, MA, and SA (Fig. 3e and 3f).

The poor stability of commercial Pt/C is typical for Pt-based catalysts, which is presumably caused by the agglomeration and/or Ostwald ripening of Pt NPs during ORR. To further explore the degradation mechanism, we investigated the structural change of the same NPs during cycling by ex situ TEM by employing the positioning gold grids (see Supplementary information for details). TEM shows that after 20k ADT cycles some irregular large particles emerged for commercial Pt/C (Fig. 4a and 4b), whereas Pt-700 remained essentially unchanged with no particle agglomeration and detachment (Fig. 4c and 4d). In addition, we also evaluated the dissolution of Pt NPs by analyzing the Pt species dissolved in electrolytes after 20k ADT cycles by ICP-MS, which showed that the dissolved Pt content for Pt-700 was substantially lower than that for Pt/C (0.232 ng vs. 40 ng). The Pt particles from the Pt/C catalyst are ~1.7x smaller than the Pt particles from the Pt-700 catalyst, and it is well known that smaller
Pt particles are more prone to dissolution than larger ones\textsuperscript{50-52}. However, the anti-dissolution improvement of ~170x far exceeds what would be expected based purely on differences in Pt particle size, and suggests that a further electronic effect may be largely responsible for the difference in Pt dissolution. Nonetheless, these results emphasize the great role of the ligand-derived graphitic shells in preventing migration, agglomeration, and dissolution of Pt NPs during the ADT.

Fig. 4: Morphological variations of Pt/C (a) before and (b) after 20k ADT cycles. Red cycles in (b) indicate the large particles formed during the ADT cycles. Morphological variations of Pt-700 (c) before and (d) after 20k ADT cycles.

2.3 Mechanistic understanding of the annealing temperature-dependent ORR activity

Having established the importance of ligand carbonization for improving the ORR activity and stability of Pt NPs, we turn our attention to investigating the annealing temperature-dependent catalytic activity exhibited by Pt-T catalysts. As
mentioned above, Pt-800 showed an inferior ORR activity, which is expected considering the severe agglomeration of Pt NPs at 800 °C. However, the drastically different ORR activity exhibited by Pt-500, Pt-600, and Pt-700 is somewhat surprising, because these three catalysts comprise the same Pt NPs coated with carbon shells having a similar thickness (i.e., bilayer carbon)\textsuperscript{53}. The fact that Pt-500 has a very small ECSA (~3.1 m\textsuperscript{2} g\textsuperscript{-1}Pt) despite no apparent change in Pt particle size (Fig. 1a) indicates that the carbon shell resulting from annealing at 500 °C is very dense in nature, which severely blocks the accessibility of surface sites despite its ultrathin thickness. Conversely, the enhanced ECSA and ORR activity exhibited by Pt-600 and Pt-700 suggests the accessibility of Pt NPs was gradually improved with the annealing temperature increasing from 500 to 700 °C. Given the essentially unchanged carbon shell thickness, we speculate that the initially dense carbon shell undergoes structural reconstruction and becomes progressively porous with the increased annealing temperature, as illustrated in Scheme 1.

Since it is difficult to directly visualize the proposed porosity in carbon shells by HRTEM (Fig. 1e-g), we designed poisoning experiments to better reveal the subtle carbon-shell difference between various Pt-T catalysts\textsuperscript{20}. Polystyrene with a terminal thiol group (denoted as PS-SH), with a molecular weight of 12600 (the molecular length of PS-SH is estimated to be ~28 nm)\textsuperscript{54}, was synthesized as a molecular probe. The thiol group is a well-known deactivator of ORR due to its strong binding affinity to Pt\textsuperscript{55}. The three catalysts (Pt/C, Pt-500 and Pt-700) were treated with PS-SH in a tetrahydrofuran (THF) solution for 12 h. As reference, the small thiol-containing molecule,
dodecanethiol (C_{12}H_{25}SH), was also used to treat catalysts under the same conditions. As expected, for Pt/C without carbon coating, both C_{12}H_{25}SH and PS-SH can easily poison Pt NPs as indicated by the fully deactivated ORR activity (Fig. 5a). Likewise, both Pt-500 and Pt-700 were completely deactivated upon treatment with C_{12}H_{25}SH (Fig. 5b and 5c), indicating that their carbon shells are permeable to C_{12}H_{25}SH molecules under the aforementioned treatment conditions. In the case of poisoning with PS-SH, however, Pt-500 and Pt-700 showed quite different anti-deactivation behaviors. Specifically, Pt-500 largely retained the ORR activity after PS-SH treatment (Fig. 5b), indicating that the carbon shells are impermeable to the bulky PS-SH molecules. In contrast, treatment of Pt-700 with PS-SH led to an apparent degradation in the catalytic activity (Fig. 5c). This implies that the carbon shells of Pt-700 have wider porous channels compared to Pt-500, which allow the partial penetration of PS-SH molecules.

These results not only qualitatively reveal the porosity difference of carbon shells between various catalysts, but also demonstrate that the porosity of the ligand-derived carbon shells, which is a key factor dictating the ORR activity of Pt NPs, can be tailored by simply controlling the thermal annealing temperature. It is likely that with the increasing of the annealing temperature the graphitic and rigid carbon shells can no longer maintain a complete and continuous spherical geometry due to the high curvature, with multiple discontinuous segments becoming favorable as depicted in Fig. 5d. Such fractures and defects could provide porous channels for accessing the encapsulated Pt NPs, which accounts for the high ORR activity exhibited by Pt-700.
Fig. 5: ORR polarization curves of (a) Pt/C, (b) Pt-500, and (c) Pt-700 before and after treatment with C_{12}H_{25}SH and PS-SH. (d) Schematics of the surface accessibility of Pt/C, Pt-500, and Pt-700, respectively, by C_{12}H_{25}SH and PS-SH molecules.

2.4 Membrane electrode assembly (MEA) tests

To test the durability of the graphitic-shell-coated Pt catalysts under the realistic operating conditions for PEMFCs, MEAs were fabricated and put through a Pt-dissolution ADT 20k cycles\textsuperscript{56, 57}. As shown in Fig. 6a, the performance of the MEA with Pt/C degraded rapidly with the increased ADT cycles, whereas the Pt-700 cell demonstrated an exceptional durability (Fig. 6b), consistent with RDE tests\textsuperscript{58}. Interestingly, Pt-700 showed slightly improved performance after the first 5k ADT cycles (both under air and 100% O\textsubscript{2}). While it may be anticipated that this was due to oxidation of the carbon shell leading to more exposed Pt sites\textsuperscript{18}, there was no noticeable increase in ECSA following 5k cycles. Thus, the increase in performance is more likely
a result of surface oxidation leading to enhanced proton transport within the catalyst layer. The variation of the maximum power density during the ADT was also compared. While the Pt-700 cell displayed a smaller initial maximum power density relative to Pt/C due to the lower ECSA (Supplementary Table 1), Pt-700 could maintain a more stable power density, which started to exceed that of Pt/C after 10k ADT cycles (Fig. 6c). Furthermore, the Pt-700 cell exhibited a small voltage loss of only 11 mV at an operating current density of 0.8 A/cm² after 20k ADT cycles (Fig. 6d), whereas the cell voltage of Pt/C decreased by 75 mV under the same cycling conditions. The polarization curves under 100% O₂ are also shown in Supplementary Fig. 5, which help to highlight that the performance loss is kinetic (as would be expected from this particular ADT) for both Pt-700 and Pt/C, and the superior durability of Pt-700 vs. Pt/C.

Fig. 6: Polarization curves of the MEAs with (a) Pt/C and (b) Pt-700 before and after ADT cycles. (c) Changes of the maximum power density with the ADT cycles for Pt/C and Pt-700 cells, respectively. (d) Voltage loss (at 0.8 A cm⁻²) after 20k cycles for Pt/C and Pt-700 cells, respectively.
2.5 Generality of the ligand carbonization approach

It is well known that decreasing the size of Pt NPs will dramatically increase the ECSA, which is advantageous for enhancing the ORR activity\textsuperscript{52}. However, as the particle size decreases below \( \sim 3 \) nm, the catalyst particles begin to demonstrate unfavorable ORR kinetics as a result of unfavorable crystal structures dominating the surface (‘particle size effect’). Unfortunately, Pt particles of this size are also still prone to rapid dissolution\textsuperscript{50}. This reality has historically meant that there is an unfortunate inverse relationship between activity and stability that must be balanced by MEA designers when selecting a catalyst. To determine whether the ligand carbonization strategy presented here can break this relationship, the OAm-capped Pt NPs with a diameter of \( \sim 3 \) nm were synthesized (\textbf{Supplementary Fig. 6a and 6b}). Using the same protocol developed for 5 nm-Pt NPs, we obtained 3 nm-Pt-700 catalysts by thermally treating 3 nm-Pt NPs supported on KB (Pt loading: \( \sim 20 \) wt.\%) at 700 °C under N\textsubscript{2} protection. Despite the decreased particle size, no apparent agglomeration of 3 nm-Pt NPs was observed upon thermal annealing, as evidenced by TEM (\textbf{Fig. 7a}) and XRD (\textbf{Supplementary Fig. 6c}). HRTEM reveals that the 3 nm-Pt NPs are homogeneously coated with approximately two graphic layers derived from the OAm ligands (Fig. 7a, inset). Importantly, these 3 nm-Pt-700 catalysts exhibited a high ORR activity and stability as indicated by the polarization curves (\textbf{Fig. 7b}) and CV (\textbf{Supplementary Fig. 6d}). As expected, the ECSA (\textbf{Fig. 7c}) of 3 nm-Pt-700 is much higher than that of its 5 nm-Pt counterpart (66.8 m\textsuperscript{2} g\textsuperscript{-1}Pt vs. 35.9 m\textsuperscript{2} g\textsuperscript{-1}Pt, Supplementary Table 1), due to the smaller size of Pt NPs. Moreover, 3 nm-Pt-700 only showed small degradation in ECSA
and MA (at 0.9 V) after 20k ADT cycles (Fig. 7c). It is thus confirmed that our in situ ligand carbonization strategy is general and can be exploited to boost the ORR stability of highly active Pt catalysts, which could greatly expand the design abilities for commercial MEA developers.

Fig. 7: (a) TEM and HRTEM (inset) images and (b) ORR polarization curves of 3 nm-Pt-700 catalysts. (c) Changes of ECSA and MA (at 0.9 V) for 3 nm-Pt-700 according to the number of ADT cycles. (d) TEM and HRTEM (inset) images and (e) ORR polarization curves of 3 nm-Pt-600N catalysts. (f) Changes of ECSA and MA (at 0.9 V) for 3 nm-Pt-600N according to the number of ADT cycles.

It should be noted that increasing the Pt loading of 3 nm-Pt NPs to ~35 wt.\% could lead to severe particle agglomeration when thermally treated at 700 °C (Supplementary Fig. 7), which is expected considering the ultrasmall particle size. Lowering the annealing temperature to 600 °C could mitigate the sintering problem (Supplementary Fig. 8a), yet the resulting 3 nm-Pt-600 catalyst showed a similarly poor ORR activity as its 5-nm counterpart (Supplementary Fig. 8b), presumably because of the dense nature of the carbon shells resulting from thermal annealing at 600 °C.
°C. Thus, we introduced N atoms during thermal annealing to afford N-doped carbon shells, as N doping has been demonstrated to be capable of enhancing the ORR activity of carbon-coated Pt NPs. N doping was realized by conducting thermal annealing at 600 °C in the presence of urea, yielding 3 nm-Pt-600N catalysts with an N content of ~3.7 at.%. No particle aggregation was observed during N doping (Fig. 7d), and more importantly, the resulting 3 nm-Pt-600N showed a significantly improved ORR activity (Fig. 7e), while simultaneously maintaining the superior long-term stability (Fig. 7f). The enhanced ORR activity of 3 nm-Pt-600N relative to 3 nm-Pt-600 (Supplementary Fig. 8b) indicates the greater exposure of surface-active sites. We speculate that in addition to altering the electronic structure of Pt which favors the ORR, N doping also facilitates the widening of porous channels in carbon shells. These results thus establish an alternative route of obtaining highly efficient ORR catalysts based on ultrasmall Pt NPs. In addition to Pt, this in situ ligand carbonization strategy is also applicable to alloy NPs. As shown in Supplementary Fig. 9, thermal annealing of PtPd NPs (~ 4 nm in diameter) supported on KB at 700 °C produced graphitic-shell-coated PtPd catalysts with an excellent ORR activity and stability.

Moreover, this ligand carbonization strategy can further be applied to improve the durability of commercial Pt/C catalysts. Surface modification was realized by mixing Pt/C with OAm ligands in isopropanol followed by stirring overnight. Considering the small size of Pt NPs (~3 nm), subsequent ligand carbonization was conducted at 600 °C in the presence of urea to produce Pt/C-600N catalysts. TEM (Fig. 8a and 8b) and XRD (Supplementary Fig. 10) verify that the size of Pt NPs was unchanged upon
annealing, indicative of the high sintering resistance of Pt NPs. Meanwhile, HRTEM reveals that the original bare Pt NPs was uniformly coated with bilayered graphitic shells derived from the OAm ligands (Fig. 8b, inset). The ECSA of Pt/C was reduced by ~15% upon ligand carbonization (77.4 m²/gPt vs. 65.8 m²/gPt, Supplementary Table 1), indicating that over-coating Pt NPs with bilayered graphitic shells only blocked a fraction of surface sites. Moreover, compared with pristine Pt/C (Fig. 3d), Pt/C-600N exhibited significantly improved stability as indicated by the negligible activity degradation after 20k ADT cycles (Fig. 8c and 8d). The substantially enhanced ORR durability was also manifested in the MEA tests (Fig. 8e), where the Pt/C-600N cell showed a voltage loss of only 13 mV at 0.8 A/cm² after 20k ADT cycles (Fig. 8f). The percent ECSA loss of Pt/C-600N was also provided in Supplementary Fig. 11, where it is apparent that the graphitic shell greatly reduces the rate of loss in ECSA for both Pt/C-600N and Pt/C-700 vs. the untreated Pt/C.

Fig. 8: TEM images of (a) Pt/C and (b) Pt/C-600N. Insets are the corresponding HRTEM images. (c) ORR polarization curves and (d) CV curves of Pt/C-600N. (e) Polarization curves of the MEAs with Pt/C and Pt/C-600N before and after 20k cycles. (f) Voltage loss (at 0.8 A cm⁻²) after 20k cycles for Pt/C and Pt/C-600N cells,
respectively.

3 Conclusion

In this work, we have developed a facile and effective method for preparing stable graphitic-shell-coated Pt catalysts by harnessing the native ligands tethered to the surface of colloidal Pt NPs. Simple thermal annealing of Pt NPs converts the hydrocarbon ligands into uniform, bilayered graphitic shells, which are sufficiently robust to protect Pt NPs against migration, agglomeration, and dissolution while largely maintaining the accessibility of surface sites. The ORR activity of the resulting carbon-coated Pt catalysts depends strongly on the thermal annealing temperature, which dictates the microstructure of graphitic shells. In particular, the graphitic-shell-coated Pt catalysts resulting from thermal annealing at 700 °C show superior long-term durability to those of commercial Pt/C catalysts. Importantly, this in situ ligand carbonization strategy is general and can also be applied to improve the stability of commercial Pt/C, thus opening a new route of developing highly durable ORR catalysts for PEMFCs.

Methods

Synthesis of colloidal Pt and PtPd NPs.

Monodisperse Pt NPs with a diameter of ~5 nm were synthesized according to a literature method with slight modifications. A precursor mixture was prepared by dissolving 0.16 g of Pt(acac)₂ in 20 mL of benzyl ether, 14.72 mL of OAm, and 2.5 mL of OA. The mixture was degassed under vacuum at 120 °C for 30 min, and was then
heated to 160 °C under a N₂ atmosphere, into which an additive solution of manganese carbonyl (0.16 g in 0.5 mL of chloroform) was injected. The reaction mixture was then heated to 200 °C and kept at this temperature for 30 min. The as-synthesized Pt NPs were precipitated from the crude reaction mixture by addition of ethanol. After centrifugation, the precipitated NPs were re-dispersed in chloroform with a concentration of 10 mg mL⁻¹.

For the synthesis of monodisperse Pt NPs with a diameter of ~3 nm, 50 mg of Pt(acac)₂ and 10 mL of OAm were added into a three-neck flask with vigorous stirring. After degassing, the reaction mixture was heated to 100 °C. After that, a mixture containing 0.2 g of borane tert-butylamine complex and 5 mL of OAm was quickly injected into the flask. Subsequently, the mixture was heated up to 120 °C and kept for 1 h. The purification of 3 nm-Pt NPs was conducted similarly as 5 nm-Pt NPs mentioned above.

Monodisperse PtPd NPs (~4 nm in diameter) were obtained by a literature method with minor modifications. Briefly, 16.2 mg of Pt(acac)₂, 12.8 mg of Pd(acac)₂, 825 μL of OAm, and 400 μL of OA were added to a three-neck flask together with 20 mL of ODE as solvent. The mixture was evacuated for 15 min and then 140 μL of TOP was added. After degassing for another 60 min at 50 °C, the reaction mixture was heated to 320 °C with N₂ protection under vigorous stirring. After 30 min of reaction, the solution was allowed to cool to room temperature. Subsequent purification of PtPd NPs was conducted similarly as Pt NPs mentioned above.

**Synthesis of graphitic-shell-coated Pt catalysts.**
A chloroform solution (2 mL) containing the as-synthesized Pt NPs was mixed with 15 mg of KB followed by sonication for 30 min. After drying at 60 °C overnight, the Pt@KB powder was annealed at 500 °C in a N₂ atmosphere for 1 h to induce the ligand carbonization. Graphitic-shell-coated Pt catalysts were obtained upon further annealing at different temperatures (500-800 °C) for another 1 h. Graphitic-shell-coated PtPd catalysts were obtained by a similar procedure, with the final thermal annealing temperature of 700 °C.

**Modification of commercial Pt/C catalysts.**

Commercial Pt/C catalysts, 200 μL of OAm, and 10 mL of isopropanol were mixed in a vial followed by stirring overnight, during which OAm ligands were attached to Pt NPs. The OAm-modified Pt/C was collected by centrifugation and was then heated at 500 °C for 1 h to carbonize the OAm ligands. Pt/C-600N catalysts were obtained by uniformly mixing urea with Pt/C with a mass ratio of 10:1 followed by annealing in a N₂ atmosphere at 600 °C for another 1 h.

**Data availability**

All relevant data that support the findings of this study are available within the article and its Supplementary Materials.

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**Competing interests**

The authors declare no competing financial interests.

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