Single atom surface engineering: A new strategy to boost electrochemical activities of Pt catalysts

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A R T I C L E  I N F O

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- Surface engineering
- Electrochemical reactions
- Atomic layer deposition
- DFT calculations

A B S T R A C T

Pt-based catalysts are widely applied in several catalytic electrochemical reactions for energy storage and conversion. The improvement of specific activity of Pt is typically achieved by introducing the transition metal to obtain the alloy structure. Different from the traditional alloy structure, herein, we report Pt catalyst modified with Co single atoms obtained by atomic layer deposition (ALD). The as-prepared catalysts show much higher mass activity and excellent stability compared to commercial Pt/C catalysts towards the hydrogen evolution reaction (HER) and oxygen reduction reaction (ORR). The atomic resolution TEM images and X-ray absorption spectroscopy (XAS) indicate the formation of atomically dispersed Co on Pt. First principle calculations reveal that the Co atom affects the electronic structure of the Pt catalysts, which resulted in the high HER and ORR performance. This work provides a new approach for the rational design of highly active and stable Pt-based catalysts, which hold great potential for application in various catalytic reactions.

1. Introduction

Pt-based catalysts have wide applications in several industrial areas due to their great electrochemical performance [1–5]. However, the widespread application of Pt is significantly hindered by its low abundance, limited supplies, and ever increasing price. Accordingly, optimizing the mass activity of Pt nanocatalysts is of great concern for minimizing the cost and achieving broader commercialization of Pt. Various methods have been developed for improving two key factors of Pt catalysts; the utilization efficiency and specific activity [6–9]. The improvement of Pt specific activity can be achieved by introducing transition metals to tune the surface electronic structure and atomic coordination. Compared with pure Pt catalysts, Pt-based multi-metallic nanocatalysts have shown great promise in enhancing the ORR [10–15] and HER [16–18] activities due to their intrinsic ligand and geometric effects. In addition to the alloy structure, it is found that the single atom-modified Pt exhibits extremely high activity compared with pure Pt and Pt-based alloy structures [19,20]. For example, Li and co-workers created Ni single atom-modified Pt nanowires through an electrochemical dealloying approach [20]. However, the aforementioned method is limited by the accuracy control of dealloying process, and it is not suitable for potential industrial application. Therefore, it remains a great challenge to develop a new strategy to obtain Pt-based catalysts with single atom modifications that can achieve good performance.

As the electrocatalytic process occurs on the surface of the catalysts, the modification of surface structure can effectively change the catalytic activity compared to the bulk part. ALD is a powerful tool to engineer the surface structure of Pt catalysts, as it can enable precisely control over the deposition of single atoms and nanoclusters [21,22]. In 2013, our group firstly reported a practical synthesis method to fabricate Pt

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single atoms on graphene nanosheets by the ALD [23]. The as-prepared Pt single atoms (SA) exhibit greatly enhanced electrochemical activities compared with commercial Pt/C catalysts. The preparation procedure of Pt SA can also be applied for the deposition of other metals. It can be expected that the deposition of single atomic transition metals will further improve the activity of Pt catalysts.

Herein, for the first time, we successfully prepared Co SA-modified Pt nanoparticles (NPs) on nitrogen-doped carbon nanotubes (NCNT) through an ALD process. The detailed structure of the Co SA-modified Pt NPs has been investigated by scanning transmission electron microscopy (STEM), X-ray absorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS). The as-prepared Co SA-modified Pt NPs showed much higher catalytic activity and stability compared to commercial Pt/C catalysts for both HER and ORR. Density functional theory (DFT) calculation results indicated that the Co atom affects the electronic structure of the Pt catalysts, which leads to the high HER and ORR performance.

2. Results and discussion

2.1. ALD preparation and characterization of Co SA-modified Pt NPs

Fig. 1a shows the synthesis route for Co SA-modified Pt NPs on NCNT with ALD. NCNTs with an average diameter of 100 nm were prepared by
ultrasonic spray pyrolysis as outlined previously [24]. Pt NPs were deposited onto the surface of NCNTs through an ALD process by using trimethyl(methylcyclopentadienyl)-platinum (IV) (MeCpPtMe$_3$) and O$_2$ as the precursors with a nitrogen (99.9995%) purge gas. As shown in Fig. S1, the SEM images indicated that Pt NPs are successfully deposited onto the substrates after 20 Pt ALD cycles. Due to the N-doped sites on NCNT, the Pt atoms can be deposited onto NCNT more easily compared with that on graphene structure. The as-prepared NCNT-Pt catalyst has also been characterized by TEM. As shown in the typical low-resolution TEM image, the Pt NPs are well dispersed on the NCNT substrates (Fig. S2a). The high-resolution TEM image clearly shows uniform Pt particles formed and the surface of the NPs is smooth (Fig. S2b). The periodic fringe spaces are confirmed to be 0.22 nm, which agree well with the d values for the (111) of Pt.

Further modification of the catalyst was conducted by deposition of Co SAs on Pt NPs by ALD using bis(ethylcyclopentadienyl)cobalt(II) as precursor. The typical SEM and HAADF-STEM images show that the size of Pt NPs is not changed after the deposition of Co (Fig. 1b and c). The (111) spacing of Co SA-modified Pt NPs is also found to be 0.23 nm (Fig. 1d). The high-resolution HAADF-STEM images clearly show that the surface of Pt NPs become rough, indicating the formation of isolated Co atoms formed on Pt NPs (Fig. 1e and f; Fig. S3). Furthermore, HAADF-STEM-EDS spectroscopy mapping profiles shown in Fig. S4 clearly reveal the homogeneous distribution of C, N, Pt and Co elements on the NCNTs. In addition, the Energy-dispersive X-ray spectroscopy (EDX) elemental mapping (Fig. 1g) reveals that most of the Co atoms are sparsely deposited on the Pt NPs. The Inductively coupled plasma optical emission spectrometer (ICP-OES) results showed that the loading of the Pt and Co loadings on NCNT are 14 and 0.9 wt%, respectively.

2.2. X-ray absorption fine structure of Co SA-modified Pt NPs

To further investigate the effect of Co SA on Pt structures, we carried out XANES and EXAFS measurements to study the electronic environment of Pt and Co in the Co-modified Pt catalysts (Fig. 2).
shows the normalized XANES spectra at the Pt L₂ and L₃ edges, respectively. Detailed examination of the spectra was conducted by qualitative and quantitative analysis of the Pt L₂ and L₃ edges white lines (WLs, the sharp intense peak lead by the rising absorption edge) [25]. It is apparent that, with the exception of Pt foil, which has an intense L₂-edge WL and a very weak L₃-edge WL due to the large spin orbit coupling of the 5d and an even distribution of the 5d 3/2 and 5d 5/2 densities of states just above the fermi level in metallic Pt, both the Pt NPs and Co modified Pt catalysts exhibit substantial WL intensity at both edges. In addition, the Co modified Pt catalysts appear to have the most intense WL compared to Pt NPs and Pt foil. It has been shown that the area under the WL peak of L₂,3-edge x-ray absorption spectra of the Pt metal is directly related to the unoccupied density of states of the Pt 5d orbitals. An increase in the L₂,3-edge WL intensity indicates a decrease in the number of electrons in the occupied d band. In addition, a small positive shift in the threshold energy E₀ can be observed for Co-modified Pt catalysts compared to Pt NPs and Pt foil. The detailed E₀ position can be determined by the first derivative of the XANES spectrum at the Pt L₃ edge (Fig. 2c). Among the three samples, the E₀ for Co modified Pt catalysts is 11.5645 eV, which is slightly higher than that of 11.564 eV of Pt NPs. This result suggests that Pt experiences a more oxidized environment when modified by Co SAs. To further explore the implication of the unoccupied densities of 5d states in Pt, quantitative WL intensity analysis was conducted based on a reported method to determine the occupancy of the 5d states in each sample. The Pt L₂- and Pt L₃-edge threshold and WL parameters are summarized in Table 1. From the analysis, the Co-modified Pt catalysts have the highest total unoccupied density of states of Pt 5d character (0.85), while the Pt foil sample had the lowest of 0.67. It has been demonstrated in literature that the vacant d-orbitals of Pt atoms play a vital role in the activity of catalysts [20,22,26].

Furthermore, to study the local structure environment of Pt, the EXAFS spectrum was studied. The magnitude of Fourier transforms (FT) of the Pt EXAFS for different samples were plotted in Fig. 2d. The EXAFS R space curve fitting attributes the FT magnitude peak at around 2.6 Å to the Pt–Pt or Pt-Co bonding. When we carefully compare the EXAFS peaks of Pt NPs and Co modified Pt NPs, we found that the peak position slightly shifted from 2.58 Å to 2.51 Å. The peaks are fitted to quantitatively obtain the coordination number (CN) and bonding length of Pt-Pt or Pt-Co bonding. When we carefully compare the EXAFS experimental data. From the Co K-edge R space fitting results, the Co atoms have the CN of 0.8 for Co-Pt, also suggesting the formation of Co single atoms on Pt surface. In addition, the bonding length of Pt-Co is 2.83 Å, which is close to that of Pt-Co (3.04 Å). The relatively longer distance of Co-Pt bond is due to the formation of Co-O bond during the ALD process.

2.3. Electrocatalytic performance of Co SA-modified Pt NPs

The HER activity of the Co SA-modified Pt NPs was firstly measured in comparison to the Pt NPs and commercial Pt/C catalysts by conducting linear sweep voltammetry measurements in 0.5 M H₂SO₄ at room temperature (Fig. 3a–c). The polarization curves show that the Co SA-modified Pt NPs exhibit better HER performance compared with Pt NPs and commercial Pt/C, as shown in Fig. 3a. The specific activity for each catalyst is calculated from the polarization curves by normalizing the current with the geometric area of the electrode (Fig. 5f). The Co SA modified Pt NPs exhibited a current density of 158 mA/cm² at an overpotential of 0.07 V. When normalized to the metal loading, the mass HER activities for the Co SA-modified Pt NPs at an overpotential of 0.07 V is 11.5 A mg⁻¹ (Fig. 3b), which is 3.5 and 16.4 times greater than that of the Pt NPs (3.3 A mg⁻¹) and Pt/C catalysts (0.7 A mg⁻¹), respectively. In addition, we prepared the Pt NPs with 5 cycles Co ALD deposition (SALD Co-Pt NPs/NCNT) and tested the electrochemical performance. As shown in Fig. 5g, the mass HER activity reduced to 4.3 A mg⁻¹ at the overpotential of 0.07 V, which indicated that increasing the Co deposition on Pt catalysts might cover the active sites on Pt surface, thus affect the electrochemical performance of the catalysts. To evaluate the durability of the as-prepared Co SA modified Pt catalysts, accelerated degradation tests (ADTs) were adopted between +0.4 and −0.15 V (versus RHE) at 100 mV s⁻¹ for 5000 cyclic voltammetry sweeps. As exhibited in Fig. 3c, the polarization curve of Co SA modified Pt NPs after 5000 cycles retained a similar performance to the initial test, resulting in a loss of only 17% of its initial current density at an overpotential of 0.07 V (Fig. 3e). Furthermore, we examined the HER performance of Co SA-modified Pt NPs, Pt NPs and commercial Pt/C catalysts in 1.0 M KOH at room temperature (Fig. 3d–f). Fig. S9 reveals the current densities in alkaline solution were 32.7, 13.7 and 17.8 mA/

### Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>Pt L₃, L₂ edge threshold and whiteline (WL) parameters</th>
<th>Pt L₂ edge WL</th>
<th>Pt L₃ edge WL</th>
<th>h₂/2</th>
<th>h₃/2</th>
<th>Total</th>
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<td>Eₚ (eV)</td>
<td>ΔEₚ</td>
<td>E₀ (eV)</td>
<td>Eₚ (eV)</td>
<td>ΔEₚ</td>
</tr>
<tr>
<td>Pt NPs on NCNT</td>
<td>11564</td>
<td>11,566.7</td>
<td>6.46</td>
<td>13,273</td>
<td>13,276.1</td>
<td>1.47</td>
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<td>Co SA modified Pt NPs</td>
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<td>11,567.0</td>
<td>7.15</td>
<td>13,273.3</td>
<td>13,276.3</td>
<td>4.39</td>
</tr>
</tbody>
</table>

*Position of the first inflection point of the edge jump for the corresponding Pt L₃ edge.*

*Peak position.*

*Area under the difference curve for normalized edge jump, the normalized edge jump for the Pt L₃ and L₂ edge corresponds to a value of 2.5 × 10³ cm⁻¹ and 1.16 × 10³ cm⁻¹, respectively.*
cm$^2$ on the Co SA modified Pt NPs, Pt NPs and commercial Pt/C catalysts at $-0.07$ V, respectively. In addition, the Co SA-modified Pt NPs still exhibited the best mass activity (2.7 A mg$^{-1}$ at the overpotential of 0.07 V) among these three catalysts (Fig. 3c). After 5000 cycles ADT test, the Co SA modified Pt NPs showed only a loss of only 7% of its initial current density at an overpotential of 0.07 V in 1.0 M KOH solution. It can be found that the electrocatalytic abilities of the Co SA modified Pt NPs exhibited superior HER performance in both acid and alkaline solution compared to the commercial Pt/C catalyst.

We also evaluated the ORR performance of the Co SA-modified Pt NPs, Pt NPs and commercial Pt/C catalysts. Cyclic voltammograms (CV) were recorded in 0.10 M aqueous HClO$_4$ at a scanning rate of 50 mV s$^{-1}$(Fig. 4a). The ECSA for the Co SA modified Pt NPs, Pt NPs and Pt/C catalysts were 63.6, 70.3 and 34.3 m$^2$/g, respectively. The polarization ORR curve showed that the Co SA modified Pt NPs exhibited better activity than that of the Pt NPs and Pt/C (40%) (Fig. 4b). In order to compare the activity for different catalysts, we normalized the kinetic current to the Pt mass (Fig. 4c and d). The kinetic current density ($j_k$) was derived from the Randles-Sevcik equation [29]. As shown in Fig. 4e, the Co SA modified Pt NPs exhibited greatly improved specific activity, with $j_k$ specific values of 0.77 mA cm$^{-2}$ based on the ECSA at 0.9 V vs. RHE, which was 1.5 and 3.8 times greater than that of the Pt NPs (0.51 mA cm$^{-2}$) and Pt/C catalyst (0.20 mA cm$^{-2}$), respectively. The mass activities of the Co SA modified Pt NPs, Pt NPs and Pt/C catalysts were 0.49, 0.26 and 0.11 A/mg, respectively, showing similar trends to that of the specific activities. As shown in Fig. 4f, the Co SA-modified NPs had obviously improved $j_k$, mass relative to the Pt/C catalyst. These results indicate that the shortened Pt-Pt bond distance by the Co modification can greatly improve the performance of the catalysts for ORR. It should be pointed that the increase of Co ALD cycles also affected the ORR performance of the catalysts. As shown in Fig. S10a, the ECSA of the catalysts significantly reduced due to the high coverage of Co on Pt surface. In addition, the mass activity reduced to 0.17 A/mg (Fig. S10b-d), indicating the significant role of Co single atom modification. We also tested the long-term stability of the catalysts through ADT between +0.6 and +1.1 V (vs. RHE) at 100 mV s$^{-1}$ for 10,000 cyclic voltammetry sweeps. For the mass ORR activities at 0.9 V, the Co SA-modified Pt NPs, which achieved the best activity towards ORR, only showed a 20.8% loss in mass activity after 10,000 cycles (Fig. S11). In addition, the TEM images of post-testing samples indicated that the particles are still well-dispersed on the NCNT (Fig. S12a and b). EDX mapping indicated that Co and Pt are not dissolved in the solution during the ADT test (Fig. S12c). These results indicate that the Co SA-modified Pt NPs exhibit good durability.

### 2.4. The enhanced mechanism elucidated by DFT calculations

Density functional theory (DFT) calculations were conducted to further elucidate the mechanism behind the improved HER and ORR performance enabled by Co SA-modified Pt NPs. Pt (111) and Co SA modified Pt (111) were used for the calculation models (Fig. 5a) based on the HAADF-STEM and EXAFS results discussed above (Fig. 1, Tables S1 and S2). The hydrogen adsorption free energies ($\Delta G_{\text{H}}$) and all potential adsorption sites for hydrogen are first calculated to probe the activity of HER. Furthermore, in order to get simulation results closer to the real reaction conditions, a volcano-type kinetic model, which expresses the experimental exchange current $i_0$ as a function of $\Delta G_{\text{H}}$ (Fig. 5b) was employed [30]. An obvious volcano plot is shown in Fig. 5b. On the left part of the volcano plot, hydrogen bond energy is strong, leading to the difficulty in H$_2$ generation. On the contrary, the right part of the volcano means proton transfer becomes rough, which result in the weak hydrogen bond. Generally, hydrogen-binding on Pt (111) is strong and concentrated on the left side of the volcano. In addition, the calculations for the Co SA-modified Pt(111) show that $\Delta G_{\text{H}}$ decreases to region of optimal HER activity. Moreover, the d band center of Pt(111) (−1.94 eV) and Co SA-modified Pt(111) (−2.04 eV, −2.30 eV) (Fig. S13) also demonstrate that Co SA species can modify the electronic structure of the surface Pt atoms by lowering of the d band.

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**Fig. 3.** (a) The HER polarization curves recorded on Co SA modified Pt NPs, regular Pt NPs and commercial Pt/C catalysts in 0.5 M H$_2$SO$_4$ at room temperature. (b) The current mass activity at $-0.07$ V in acid solution. (c) Durability measurement of the Co SA modified Pt NPs catalysts. (d) The HER polarization curves recorded on Co SA modified Pt NPs, regular Pt NPs and commercial Pt/C catalysts in 1.0 M KOH at room temperature. (e) The current mass activity at $-0.07$ V in alkaline solution. (f) Durability measurement of the Co SA modified Pt NPs catalysts in alkaline solution.
Therefore, the adsorption strength of hydrogen is weakened, leading to the optimal HER activity.

To further explore the ORR performance between Pt(111) and Co SA modified Pt(111), theoretical calculations for the reaction mechanism were explored. The free energy diagrams of the ORR on Pt(111) and Co SA modified Pt(111) are shown in Fig. 5c. In the case where the electrode potential is zero ($U = 0$ V), the elementary reactions of ORR on both surfaces are exothermic, and the adsorption of O* and OH* is weakened on Co SA modified Pt(111). Moreover, previous studies [31, 32] on the ORR mechanism indicate that O protonation to OH is the rate-determining step, and thus catalysts with weakly binding oxygen species have better ORR activity. Therefore, the Pt atoms electronically modified by Co SA species are beneficial for the ORR performance. Additionally, when the electrode potential is 1.23 V, OH protonation on Co SA modified Pt(111) is found to be more facile than Pt(111). This further suggests that the Co SA modified Pt(111) could provide more available sites for ORR, which is consistent with the stability of Pt(111) and Co-SA modified Pt(111) found experimentally [33–35].

3. Conclusion

In conclusion, we have successfully synthesized Co SA-modified Pt catalysts on NCNTs by ALD. The obtained Co SA-modified Pt NPs showed significantly improved activity and excellent stability compared to commercial Pt/C catalysts for both HER and ORR. X-ray absorption spectroscopy indicates that the structure model of Co SA-modified Pt NPs contain one Co-Pt bonding configuration. Furthermore, the DFT calculation results reveal that the Co SA-modified Pt(111) show decreased $\Delta G^*$ to the value of the optimal HER activity region. During the ORR process, the adsorption of O* and OH* weakened on Co SA modified Pt(111), which is beneficial for the ORR performance. This work paves a new way for the rational design of bimetallic catalysts, which have great potential for application in various catalytic reactions.

CRediT authorship contribution statement

L. Zhang conceived and designed the experimental work and prepared the manuscript; M. Norouzi Banis, Y. Sun and J. Li helped with ALD characterization; Q. Wang and M. Gu performed STEM characterization; Z.-J. Zhao and L. Li performed the DFT calculations; R. Li, and K. Adair participated in the discussion of the data; X. Sun supervised the overall project. All authors have given approval to the final version of the manuscript.

Declaration of Competing Interest

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

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.nanoen.2021.106813.

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