

2018

Coordination of Atomic Co-Pt Coupling Species at Carbon Defects as Active Sites for Oxygen Reduction Reaction

Longzhou Zhang
Griffith University

Julia Fischer
University of Queensland

Yi Jia
Griffith University

Xuecheng Yan
Griffith University

Wei Xu
Chinese Academy Of Sciences, Jilin University

See next page for additional authors

Publication Details

Zhang, L., Fischer, J. Melisande Theresa Agatha, Jia, Y., Yan, X., Xu, W., Wang, X., Chen, J., Yang, D., Liu, H., Zhuang, L., Hankel, M., Searles, D. J., Huang, K., Feng, S., Brown, C. L. & Yao, X. (2018). Coordination of Atomic Co-Pt Coupling Species at Carbon Defects as Active Sites for Oxygen Reduction Reaction. *Journal of the American Chemical Society*, 140 (34), 10757-10763.

Coordination of Atomic Co-Pt Coupling Species at Carbon Defects as Active Sites for Oxygen Reduction Reaction

Abstract

Platinum (Pt) is the state-of-the-art catalyst for oxygen reduction reaction (ORR), but its high cost and scarcity limit its large-scale use. However, if the usage of Pt reduces to a sufficiently low level, this critical barrier may be overcome. Atomically dispersed metal catalysts with high activity and high atom efficiency have the possibility to achieve this goal. Herein, we report a locally distributed atomic Pt-Co nitrogen-carbon-based catalyst (denoted as A-CoPt-NC) with high activity and robust durability for ORR (267 times higher than commercial Pt/C in mass activity). The A-CoPt-NC shows a high selectivity for the 4e-pathway in ORR, differing from the reported 2e-pathway characteristic of atomic Pt catalysts. Density functional theory calculations suggest that this high activity originates from the synergistic effect of atomic Pt-Co located on a defected C/N graphene surface. The mechanism is thought to arise from asymmetry in the electron distribution around the Pt/Co metal centers, as well as the metal atoms' coordination with local environments on the carbon surface. This coordination results from N8V4 vacancies (where N8 represents the number of nitrogen atoms and V4 indicates the number of vacant carbon atoms) within the carbon shell, which enhances the oxygen reduction reaction via the so-called synergistic effect.

Disciplines

Engineering | Physical Sciences and Mathematics

Publication Details

Zhang, L., Fischer, J. Melisande Theresa Agatha., Jia, Y., Yan, X., Xu, W., Wang, X., Chen, J., Yang, D., Liu, H., Zhuang, L., Hankel, M., Searles, D. J., Huang, K., Feng, S., Brown, C. L. & Yao, X. (2018). Coordination of Atomic Co-Pt Coupling Species at Carbon Defects as Active Sites for Oxygen Reduction Reaction. *Journal of the American Chemical Society*, 140 (34), 10757-10763.

Authors

Longzhou Zhang, Julia Fischer, Yi Jia, Xuecheng Yan, Wei Xu, Xiyang Wang, Jun Chen, Dongjiang Yang, Hongwei Liu, Linzhou Zhuang, Marlies Hankel, Debra J. Searles, Keke Huang, Shouhua Feng, Christopher L. Brown, and Xiangdong Yao

Coordination of atomic Co-Pt coupling species at carbon defects as electrochemical Janus active sites

Longzhou Zhang^{1†}, Julia Melisande Theresa Agatha Fischer^{2†}, Yi Jia^{1*}, Xuecheng Yan¹, Wei Xu³, Xiyang Wang⁴, Jun Chen⁵, Dongjiang Yang¹, Hongwei Liu⁶, Marlies Hankel², Debra J. Searles^{2,7}, Keke Huang⁴, Shouhua Feng⁴, Christopher L Brown¹ and Xiangdong Yao^{1,4*}

¹School of Natural Sciences and Queensland Micro- and Nanotechnology Centre, Griffith University, Nathan Campus, Queensland 4111, Australia.

²Centre for Theoretical and Computational Molecular Science, Australian Institute for Bioengineering and Nanotechnology, The University of Queensland, Queensland 4072, Australia.

³Beijing Synchrotron Radiation Facility, Institute of High Energy Physics, Chinese Academy of Sciences, Beijing 100049, P. R. China.

⁴State Key Laboratory of Inorganic Synthesis and Preparative Chemistry, College of Chemistry, Jilin University, Changchun 130012, P. R. China.

⁵Intelligent Polymer Research Institute, ARC Centre of Excellence for Electromaterials Science, AIIM Facility, University of Wollongong, Innovation Campus, Wollongong, New South Wales 2522, Australia.

⁶Australian Centre for Microscopy & Microanalysis (ACMM), the University of Sydney, Sydney, NSW 2006, Australia.

⁷School of Chemistry and Molecular Biosciences, The University of Queensland, Queensland 4072, Australia.

E-mail: y.jia@griffith.edu.au and x.yao@griffith.edu.au

[†]L. Z. Zhang and J. M. T. A. Fischer contributed equally.

Abstract

Platinum (Pt) is the state-of-the-art catalyst for oxygen reduction (ORR) and hydrogen evolution (HER), but its high cost and scarcity limit its large-scale use. However, if the usage of Pt can be reduced to a sufficiently low level ($0.125\text{mg}/\text{cm}^2$ according to the Department of Energy (DOE) target), this critical barrier may be overcome. Atomically dispersed metal catalysts with high activity and high atom efficiency make them possible to achieve this goal. Herein, we report a locally distributed atomic Pt-Co nitrogen-carbon based catalyst (denoted as A-CoPt-NC) with high activity and robust durability for ORR (267 times higher than commercial Pt/C in mass activity) and HER (much superior to Pt/C). The A-CoPt-NC shows a high selectivity for the $4e^-$ pathway in ORR, differing from the reported $2e^-$ pathway characteristic of atomic Pt catalysts. Importantly, the A-CoPt-NC catalyst in this experiment only contains $\sim 0.0005\text{ mg}/\text{cm}^2$ of Pt and the activity is about 20 times the DOE target at 0.9V in alkaline solution. Density functional theory (DFT) calculations suggest that this high activity originates from the synergistic effect of atomic Pt-Co located on a defected C/N graphene surface. The mechanism is thought to arise from asymmetry in the electron distribution around the Pt/Co metal centres, as well as the metal atoms coordination with local environments on the carbon surface. This coordination results from N₈V₄ vacancies (where N₈ represents the number of nitrogen atoms, V₄ indicates the number of vacant carbon atoms) within the carbon shell that enhances the oxygen reduction reaction *via* the so-called synergistic effect. DFT calculations suggest the HER activity arises from atomic Pt-Co coupling species locating at N₆V₄ vacancies within the graphitic shell causing the A-CoPt-NC catalyst to show extremely high HER activities in both acid and alkaline solutions.

Platinum (Pt) is the benchmark electrocatalyst for the oxygen reduction reaction (ORR)¹⁻⁵ and hydrogen evolution reaction (HER)^{6, 7}, exhibiting high activities. However, the high cost and the natural scarcity of Pt still hamper its industrial implementation. Downsizing the Pt particles to expose more Pt atoms on the surface (rendering higher atom efficiency) is a viable strategy to enable Pt based catalysts more affordable. Generally, when particle sizes are further reduced to the nanoscale, quantum size effects will be induced in the catalysts, which not only alter the surface energy due to the unsaturated coordination, but change the *d* state energy of metal atoms leading to spatial electron localization^{8, 9}. This size-induced change of electronic structures at active sites will subsequently tailor the binding capability with the diverse species of reactants (e.g. O₂ in ORR or H⁺ in HER), thus increases in the activities of catalysts of the electrocatalytic reactions are attainable.

Recently, the so-called single-atom catalyst (SACs) have sparked new interests in heterogeneous catalysis, maximizing the atom efficiency and demonstrating excellent catalytic performance in CO oxidation¹⁰⁻¹², water-gas shift (WGS) reaction^{13, 14}, and electrochemical/photoelectrochemical reactions¹⁵⁻¹⁷. However, the development of SACs in electrocatalysis is still in its infancy, as several issues are urgently to be addressed. Firstly, SACs are not really “atoms” (i.e. zero valence state), as the atomic metal species interact strongly with the neighbouring atoms on support and exhibit valence states. This pivotal fact pushes us to reconsider that the active sites in SACs are originated from the unique coordination structures between the single metal atoms and surround non-metallic atoms of the support. Secondly, optimization of the electronic structure on the active sites by modulating the coordination environment of the metal atoms (MAs) has become a critical route to enhance the reactivity of the active sites. Take the MA-N-C (MA = Fe or Co) coordination structure for an example, the MA-N₂ moieties as active centres have been reported more efficient for ORR than those of the MA-N₄ moieties. This is ascribed to the more suitable interaction of MA-N₂

moieties with $*O_2$ and $*OH$ intermediates^{15, 18}. Thirdly, selectivity in SACs for specific electrochemical reactions is important. For instance, the isolated atomic Pt species (denoted as IA-Pt) were reported to possess a high selectivity for the production of H_2O_2 via a $2e^-$ pathway ($O_2 + 2H^+ + 2e^- \rightarrow H_2O_2$) rather than H_2O via a $4e^-$ pathway ($O_2 + 4H^+ + 4e^- \rightarrow 2H_2O$). This is because the breaking of the O-O bond is not energetically favourable on the isolated atomic Pt sites¹⁹⁻²¹. However, it is possible to alter the reaction to a $4e^-$ pathway for ORR by putting another Pt (or different metal) atom at a certain distance. Fourthly, the research on the atomic interaction between metallic atoms in a certain local environment may provide insightful understanding of “synergetic effect”. The so-called “synergetic effect” has been extensively used, but it is very rare to studies on its origination, especially at the atomic level. Therefore, it is imperative to get an in-depth insight into the interactions between the atomic metal species and local environment on the support (e.g. atomic metal-nonmetal coordination), which may direct us to design the new generation of atomic metal catalysts with high activity and selectivity. Very recently, our group developed a defective graphene (DG) with a high density of structural defects²². Besides the defects themselves activating the electrochemical reactions according to the proposed defect mechanism²², defects are highly likely to provide unique sites for trapping metallic species^{23, 24}. Due to the different structures and sizes of the defects, one or more metal atoms might be trapped into one specific defect, providing the possibility of studying the interaction between the single metal atom and neighboring nonmetal atoms, the interaction between the metal atom pairs (in this case atomic Pt-Me (Me=Co/Pt) coupling species) and the interaction between the metal atom pair and the neighboring nonmetal atoms. Additionally, compared to the metal oxide supports, defective carbon presents a high tolerance in electrolyte environments at a wide range of pH, which enables the catalyst to work in alkaline, neutral and acid electrolytes. This is particularly relevant to the HER in this study.

Herein, we report a new class of atomic Co-Pt carbon/N based catalyst (denoted as A-CoPt-NC) that directly utilized the induced defects in the shell of carbon capsules to form atomic Co-Pt-N-C coordination structures as active sites through electrochemical activation. According to the analysis of X-ray absorption near-edge structure (XANES), the atomic configurations between Co/Pt and N/C can be deduced. Direct observation from HAADF-STEM image clearly demonstrates that the atomic metals (Co/Pt) are trapped into a vacancy type defect to form integrity of atomic Co-Pt-N-C coordination structures. Experimentally, the obtained A-CoPt-NC catalyst exhibited very high activity and robust stability for the ORR in alkaline solution, delivering the specific (electrochemical active surface area (ECSA) normalized to Pt mass) and mass activities of 85 and 267 times greater than those of the commercial Pt/C catalyst, respectively. Meanwhile, the activity had no obvious loss after a 240 min electrochemical durability test. The atomic Pt shows high selectivity for the $4e^-$ pathway in ORR, which is different from the counterpart reported in former literatures¹⁹⁻²¹. Furthermore, A-CoPt-NC exhibits extremely high activities in HER under all pH (acidic, neutral and alkaline) conditions. Density functional theory (DFT) calculations on model structures developed based on the observed and other possible Co-Pt-N-C configurations reveal that atomic Pt-Me (Me=Co/Pt) coupling at the carbon defects (denoted as a(Pt-Me)@NXVY where X is the number of nitrogens surrounding the defect and Y is the number of carbon atoms removed) can significantly tailor the electronic structure of the metal atoms and alter the charge distribution at the coordination structures, thereby enhancing the specific electrocatalytic performances (a(Pt-Co)@N8V4 for ORR and a(Pt-Co)@N6V4 for HER).

Preparation and characterization of A-CoPt-NC electrocatalyst

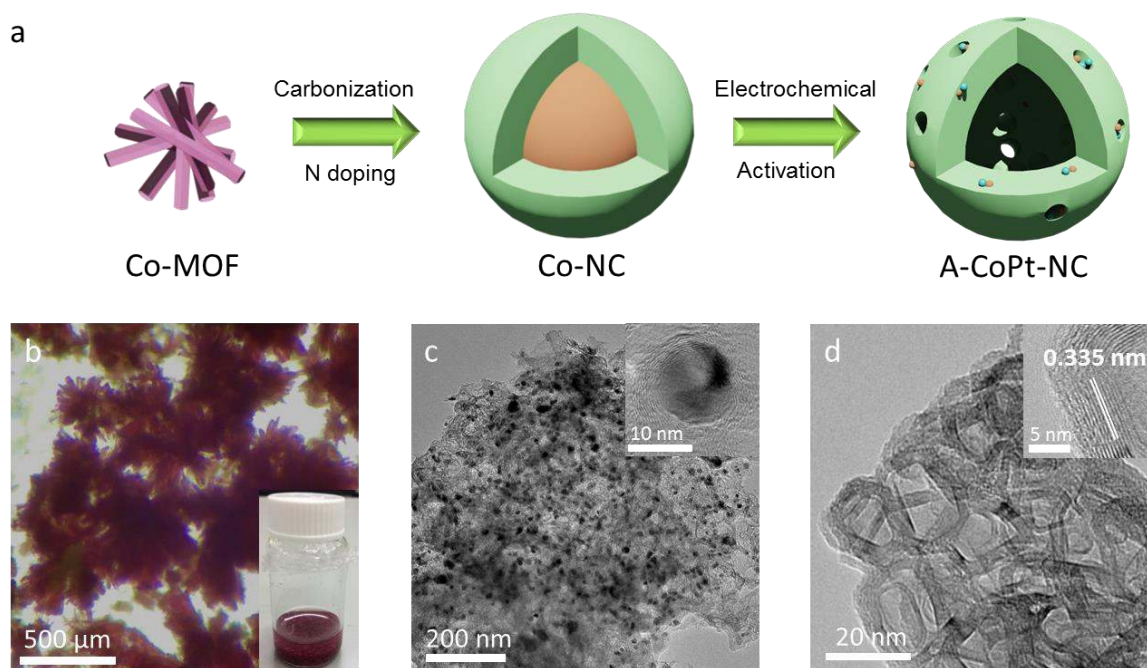


Figure 1 | Preparation and morphology characterization of A-CoPt-NC. **a**, Schematic illustration of the synthesis procedure of A-CoPt-NC. **b**, An optical photo of the Co-MOF crystal. The inset image shows Co-MOF in the DMF solution. **c**, TEM images of the Co-NC composite. The inset TEM image demonstrates the core-shell structure. **d**, TEM images of A-CoPt-NC. The inset TEM image demonstrates the hollow graphitic shells.

A-CoPt-NC was fabricated by a facile two-step synthesis strategy with the precursor of rod-like cobalt-metal organic framework (Co-MOF) as shown in Fig. 1a. In the first step, the as-prepared Co-MOF (Fig. 1b and inset) was carbonized at 850 °C with simultaneous nitrogen doping treatment, forming the core-shell Co-NC structure (Fig. 1c and inset). We then electrochemically applied a cyclic-potential to the Co-NC electrode (a similar activation process can be referred to the literature^{25,26}). Thereby carbon based hollow nanostructures with graphitic shells were generated (Fig. 1d and inset). Although XRD patterns show that no obvious metallic crystal phases exist after the electrochemical activation (Supplementary

Fig.1), the characterization of both inductively coupled plasma atomic emission spectroscopy (ICP-AES) and energy-dispersive X-ray spectroscopy (EDS) elemental mapping (Supplementary Fig.2) demonstrate trace amounts of Co and Pt in A-PtCo-NC (the content of Co is 1.72 wt %, compared to 21.10 wt % before activation, and the content of Pt is 0.16 wt %), indicating almost all of the Co cores has been removed. These results prompted us to propose that the atomically disperse Co and Pt co-doped in NC capsules can form during the electrochemical activation process, which will be verified below.

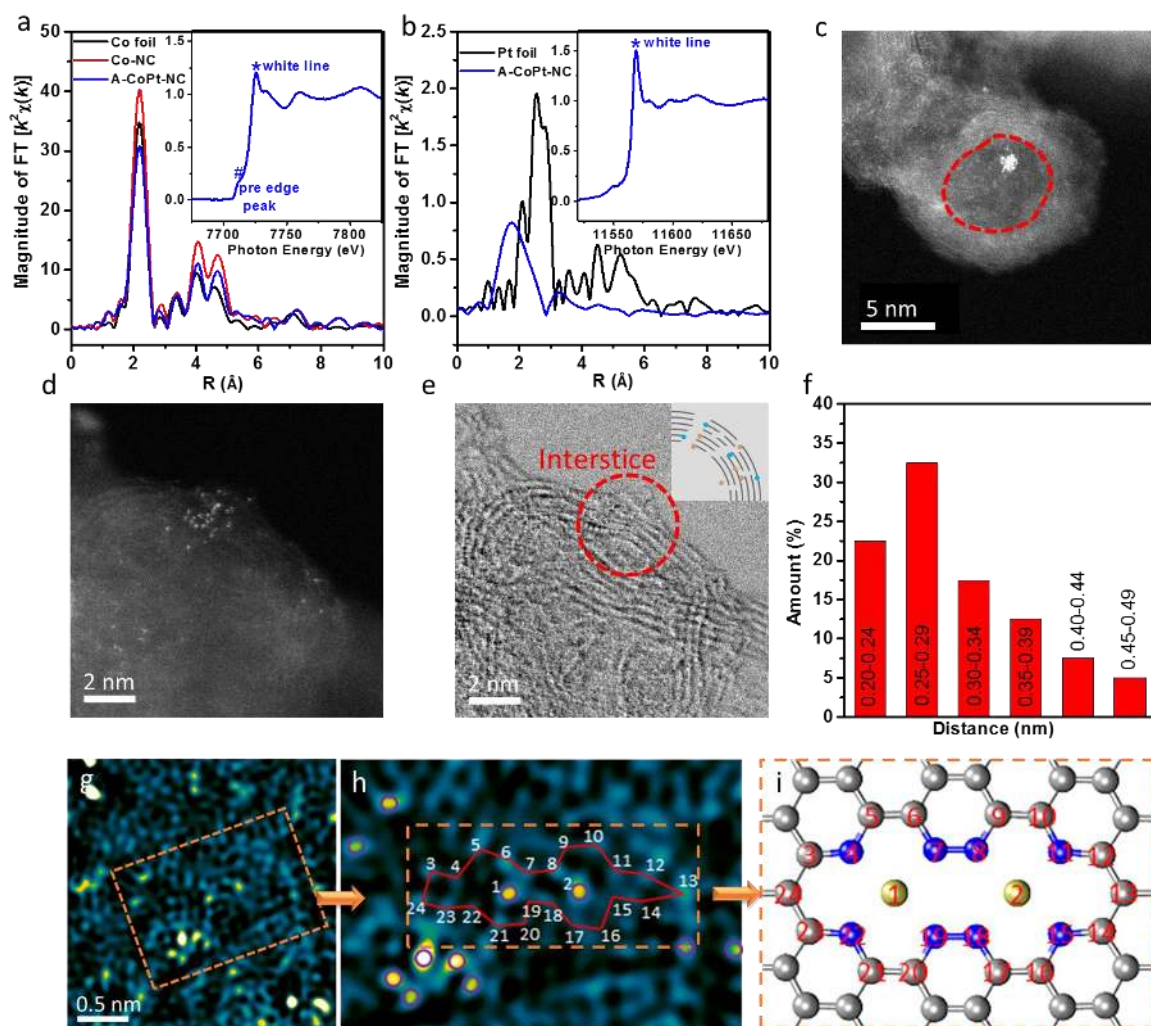


Figure 2 | The structural characterizations of A-CoPt-NC. a,b, The k^2 -weighted Fourier transform spectra of the Co and Pt EXAFS for Co-NC, A-CoPt-NC, Co foil and Pt foil, respectively. The insets are the Co and Pt XANES spectra, respectively, for A-

CoPt-NC. **c,d**, The dark-field STEM images of the A-CoPt-NC. The position of the cobalt core dissolved in the activation, is marked with the red dashed ring. **e**, The corresponding bright-field STEM image of (d) with the inset of the schematic diagram of the interstice zone. **f**, The distribution of distances of adjacent metal atoms counted from 40 neighbouring metal atom pairs. **g**, The HAADF image of A-CoPt-NC after fast Fourier Transformation (FFT) filtering. The bright yellow spots are metal atoms and the cyan spots are carbon atoms. **h**, A partially zoomed-in image of the area framed in **g**. Metal atoms are marked by purple circles. The carbon atoms adjacent to #1 and #2 metal atoms are linked with red line. **i**, Model of the configuration of the 2 metal atoms trapped in the defect, reconstructed from the observed atomic structure in **h**.

To further investigate the fine structure of Pt and Co on carbon, we performed X-ray absorption near-edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) spectrometry. The intensity of Co-Co peak around 2.2 Å decays after the activation, revealing a change in the Co local environment (Fig. 2a)^{15, 27}. The dissolution of Co cores during activation disrupts the majority of the original Co-Co coordination and the residual Co clusters contribute to the declined Co-Co peak (there may also exist some undissolved Co cores although we did not observe any in searching domain by TEM). This is further confirmed by the aberration-corrected scanning transmission electron microscopy (STEM) images. In Fig. 2c, the area of the initial Co core before activation is marked with a red dashed ring. It can be seen that the Co core was removed and only a small Co cluster sized ~ 1 nm remained, which agrees with the EXAFS analysis. The high intensity of the white line of A-CoPt-NC in the XANES (inset of Fig.2a) indicates an oxidized electronic structure of Co, which is due to the Co-N and/or Co-C and/or Co-C/N coordination newly formed during the activation process¹⁵. It is suggested that the pre-edge peak of A-CoPt-NC at 7712 eV is the fingerprint of the Co-N₄ square planar structure^{28, 29} due to the dipole forbidden 1s → 3d transition with dominantly

quadrupole coupling, which is analogous to the pre-edge peak of cobalt phthalocyanine³⁰. This finding is also consistent with the previous report of atomic Co catalyst used for HER²⁷. It is very difficult to distinguish the bond of the Co-N₄ and Co-C₄ due to the very small difference of the bonding energy. However, it is reasonable to suppose that the coordination is Co-C/N₄ because of the large amount of N in carbon shell.

In the R space spectra of Pt (Fig. 2b), the scattering peak derived from Pt-Pt coordination at 2.6 Å is not observed, in contrast to the Pt foil, indicating atomic dispersion of Pt species in the N-doped carbon capsules. The predominant peak around 1.8 Å can be ascribed to Pt-C or Pt-N coordination^{20, 31}. The XANES also indicates an oxidized electronic structure of Pt (inset of Fig.2b). Further study shows that A-CoPt-C without N doping, as a control experiment, exhibits a high amount of metallic Pt-Pt coordination (Supplementary Fig.3). The corresponding Pt particles are also observed in A-CoPt-C by the TEM images in Supplementary Fig.4. These findings suggest that N plays a critical role in trapping single Pt atoms and a Pt-N-C coordination structure is highly possible.

Fig. 2d shows the distribution of single metal atoms. Interestingly, these single metal atoms do not distribute uniformly on the carbon, but predominantly locate at the interstice of the graphitic layers. Fig. 2e is the bright field image focussing on the same area shown in Fig. 2d, clearly illustrating the opening of graphitic layers. Since amorphous carbon is less stable during the electrochemical activation and more easily oxidized³², we can deduce that those openings were created through the corrosion of partially existing amorphous carbon in the shell during the activation process, which is evidenced by increased graphitization degree of A-CoPt-NC at 26.2° in the XRD pattern (Supplementary Fig.1). With the removal of amorphous carbon forming the interstices, the cross-sections of the graphitic layers were exposed and distorted, thereby increasing the disorder of the lattice. The acidic solution subsequently permeated through these openings and dissolved the cobalt cores. Simultaneously, the Co and

Pt ions near the interstices were captured by the exposure of lattice defects in the carbon with the assistance of the freshly generated dangling carbon and nitrogen bonds. These contribute to the atomic metal species being distributed locally at the graphitic layer openings. The statistical distribution¹¹ of 40 pairs of two adjacent metal atoms (denoted as Me-Me, Me = Co or Pt) shows that the Me-Me distances are in the range from 0.2 nm to 0.5 nm with most in the interval of 0.25-0.29 nm. This phenomenon provides the experimental basis for structures used in modelling, as discussed later.

Benefiting from fast Fourier Transformation (FFT) and inverse FFT, noise was filtered and the HAADF image of A-CoPt-NC with strong contrasts was gained from a less damaged region of interest (ROI). This is shown in Fig. 2g with a clear view of the local coordination environment of the metal atoms. In Fig. 2h, the metal atoms are marked with purple cycles and the adjacent carbon/nitrogen atoms are linked. As a result, the configuration of the metal atoms (Me-#1 and Me-#2) can be identified. The structure observed in the STEM image is simulated in Fig. 2i. The same arrangement of the numbered atoms in Fig. 2h are used to obtain insight into the structure-property correlation in A-CoPt-NC catalyst through computer simulations.

Electrocatalytic ORR activity of A-CoPt-NC

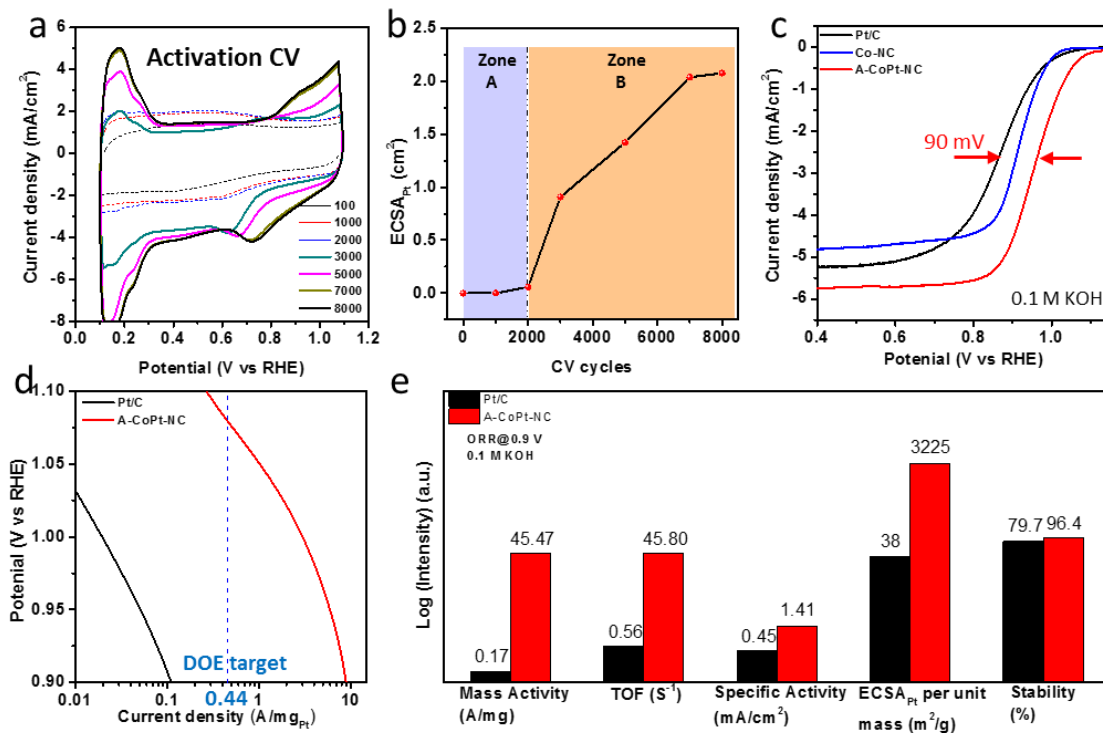


Figure 3 | Electrochemical oxygen reduction activities. **a**, Cyclic voltammetry (CV) curves after different activation cycles in 0.5 M H₂SO₄ electrolyte. **b**, The evolution of the ECSA_{Pt} during activation. **c**, ORR linear sweep voltammetry (LSV) curves of Co-NC, A-CoPt-NC and Pt/C in 0.1 M KOH electrolyte. **d**, Mass activity Tafel plot for A-CoPt-NC and Pt/C. **e**, Comparison of mass activity, TOF, specific activity, ECSA_{Pt} per unit Pt mass and stability of the A-CoPt-NC and Pt/C for ORR.

The electrochemical activation was performed with CV cycles from 0.1V to 1.1V vs RHE in 0.5 M sulphuric acid. Fig. 3a shows the evolution of the CV curves during activation. At the beginning of activation, no typical hydrogen adsorption or desorption peaks in the range of 0.1V to 0.38V (signifying the presence of Pt) were observed. The hydrogen adsorption peak appeared after 2000 CV cycles, indicating Pt started to load onto the carbon shell. The loading amount of Pt increased with the additional CV cycles until 7000 cycles. From 7000 to 8000

cycles, the CV curves nearly overlapped demonstrating the saturation of Pt loading. The corresponding ECSA analysis of the Pt also reveals that in the first 2000 CV cycles (zone A in Fig. 3b), there was little Pt loading on the shell (the $ECSA_{Pt}$ is nearly zero) due to the rarity of adsorption sites in the carbon. During this stage, the amorphous carbon was oxidised and the graphitic shell cracked. With the continued oxidation, an increasing number of adsorption sites at the openings were generated and the Pt atoms were subsequently anchored (shown as zone B).

Linear sweep voltammetry (LSV) curves in 0.1 M KOH electrolyte (Supplementary Fig.5) shows the current density increased with the rotation rate from 400 to 2500 rpm, indicating a defined mass transfer controlled process. The Koutecky-Levich plot shown in the inset implies a $4e^-$ transfer pathway for the ORR. The rotating ring-disk electrode (RRDE) voltammogram of A-CoPt-NC is also performed in O_2 -saturated 0.1 M KOH electrolyte at a rotation rate of 1600 rpm to accurately determine the electron transfer number. Based on the ring and disk currents, the electron transfer number is calculated to be larger than 3.6 over the potential range from 0.4 to 1.0 V vs RHE, and the H_2O_2 yield remained below 17% (Supplementary Fig.6), indicating that $4e^-$ transfer pathway is predominant in A-CoPt-NC for ORR in 0.1 M KOH electrolyte. The polarization curves in Fig. 3c show that the Co-NC exhibited higher half-wave potential (0.92 V vs RHE) but lower limit current density (4.8 mA/cm^2) compared to those of commercial Pt/C catalyst, due to the metal-N-C catalysis³³. Notably, the A-CoPt-NC exhibited much better performance than those of Co-NC and Pt/C as well as a robust stability (as shown in Supplementary Fig.7, after 4 hours reaction, A-CoPt-NC still retains the 96.4% of the initial activity whereas the commercial Pt/C catalyst only preserves 79.7%). The half-wave potential of A-CoPt-NC is 0.96 V vs RHE, 90 mV superior to that of Pt/C. The mass activity Tafel plot (Fig. 3d) shows that the A-CoPt-NC can deliver 20 times higher mass activity than the 2017 target set by the DOE (a current density of 0.44 A/mg_{Pt} at

0.90 V, highlighted by blue dash line in Fig. 3d). The A-CoPt-NC can deliver the DOE targeted mass activity at 1.08 V vs RHE, thus reducing the overpotential by 0.18 V. Impressively, the ECSA_{Pt} per unit Pt mass (3225 m²/g) of A-CoPt-NC is the highest among the Pt contained ORR catalysts reported so far (Fig. 3e), which is attributed to the high atom efficiency^{1, 2, 4}. With this advantage, the A-CoPt-NC also presents a much higher mass activity of 45.47 A/mg compared to those of Pt/C and other Pt contained catalysts (Supplementary Table 1). More importantly, the specific activity of A-CoPt-NC, which normalizes the performance to the ECSA, is 3-fold greater than that of Pt/C, indicating that the individual active site in A-CoPt-NC is more energetically favourable for ORR than that of Pt/C (Fig. 3e). As well as in the alkaline media, the A-CoPt-NC presents a very good ORR performance in the acidic electrolyte (0.1 M HClO₄), with a mass activity 5.6 times higher than that of Pt/C (Supplementary Fig.8).

Catalytic mechanism of ORR in A-CoPt-NC

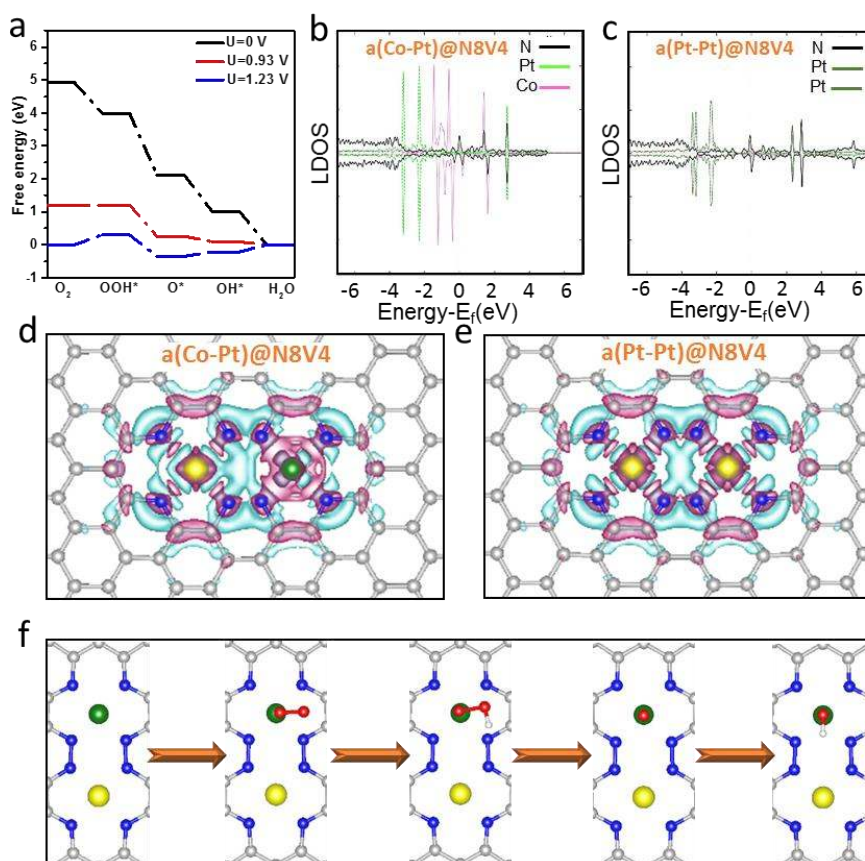


Figure 4 | Mechanistic study of ORR in A-CoPt-NC. **a**, The ORR free energy profiles of a(Co-Pt)@N8V4 at the equilibrium potential ($U=1.23$ V), onset potential and zero potential. **b,c**, The local densities of states of a(Co-Pt)@N8V4 and a(Pt-Pt)@N8V4. **d,e**, The top view of the charge densities of a(Co-Pt)@N8V4 (d) and a(Pt-Pt)@N8V4 (e). Pink and aqua iso-surfaces with an isosurface level of $0.0025 e/a_0^3$ represent electron accumulation and depletion areas respectively. **f**, An illustration of the ORR reaction pathway on a(Co-Pt)@N8V4.

According to the published literature, the atomic Pt based catalysts have a low selectivity for the $4e^-$ transfer pathway for ORR (preferring to produce H_2O_2 rather than H_2O), due to the need of a synergistic effect from Pt sites within a desirable distance to break the O-O bond¹⁹⁻²¹. Therefore, we hypothesize that the synergetic effect of atomic Pt-Me (Me=Co/Pt) coupling species at the carbon defects (denoted as a(Pt-Me)@Defects) in A-CoPt-NC can enhance the selectivity of $4e^-$ transfer pathway and the overall activity by modulating the electronic structure of metal atoms and altering the charge distribution at the coordination structures. To investigate the synergistic effect, we performed DFT calculations on five different coordination structures (each of them contains Pt-Pt and Co-Pt coordination as shown in Supplementary Fig.9), which were selected according to the analysis of the distribution of adjacent metal atomic interdistances (Fig. 2f) and STEM observations (Fig 2g-2i). The distances between the two adjacent metal atoms in these 5 models are in the range from 0.227 nm to 0.504 nm, which are in accord with the experimentally measured distances (Fig. 2f). Notably, the structure N8V4 is directly observed from the STEM image in Fig. 2h. The binding energy of the atomic metals on the 5 models were calculated as shown in Supplementary Table 2. It is shown that atomic metals on N6V6 and N8V10 are not thermodynamically stable, thereby they are excluded for further calculations. Accordingly, six ORR energy profiles are obtained as shown

in Supplementary Fig.10 and the corresponding onset potentials are summarized in Supplementary Table 3. The most energetically favourable configuration is a(Co-Pt)@N8V4 with a low overpotential of 0.30 V. As shown in Fig. 4a, the first protonation step ($O_2 \rightarrow OOH^*$) determined the onset potential, because the magnitude of change in free energy (0.3 eV) for this step is the largest at equilibrium. Meanwhile, the potential determine step for a(Pt-Pt)@N8V4 is the same with a(Co-Pt)@N8V4 (Supplementary Fig.10), whereas requires an overpotential as large as 1.07 V (Supplementary Table 3). These results reveal that heterogeneous atomic metals (Co and Pt) on N8V4 precede the homogenous atomic metals (sole Pt) on N8V4, due to the stronger binding effect between a(Co-Pt)@N8V4 and O_2 .

To investigate the underlying origin of the interactions between different active sites and the adsorbates, the density of the states of a(Co-Pt)@N8V4 and a(Pt-Pt)@N8V4 were simulated (Fig. 4b and 4c). Since the *d* orbitals of noble/transition metal atoms and the 2*p* orbitals of oxygen atoms participate in orbital coupling and form the new molecular orbitals during the adsorption, we concentrate on the states of *d* orbitals of Pt and Co. According to the *d* band centre theory proposed by Nørskov et al., the up-shifted *d* orbital relative to the Fermi level will result in a strong binding between the catalyst and the adsorbate, and vice versa³⁴. Here, the energy of the Co 3*d* orbital in a(Co-Pt)@N8V4 is much closer to the Fermi level than that of the Pt 5*d* orbital in a(Pt-Pt)@N8V4, indicating the strong binding between the a(Co-Pt)@N8V4 and oxygen. This difference in binding strength will further affect the ORR activities of these two active sites, which is supported by the energy profiles (Fig. 4a). In addition, the charge distribution patterns (Fig. 4d and 4e) show a strong electron accumulation (pink area) around the Co atom in a(Co-Pt)@N8V4, but weak electron accumulation or depletion around Pt in a(Pt-Pt)@N8V4, which can be attributed to the asymmetric deployment of Pt and Co in a(Co-Pt)@N8V4, polarizing the surface charges near the active sites. The electrons near Co will enable O_2 to be transformed to H_2O , thus enhancing the ORR

performance. Fig. 4f shows the ORR reaction associated with the $4e^-$ pathway on a(Co-Pt)@N8V4, which involves four protic hydrogen and electron transfer steps: (i) the adsorbed O_2 transfers into OOH^* ; (ii) desorption of H_2O and formation of O^* ; (iii) OH^* is formed and (iv) the OH^* further associates with a protic H and an electron to generate H_2O .

It is worth noting that the elementary steps from O_2 to O^* can also proceed through a dissociative $4e^-$ pathway, which is also energetically downhill in the energy profile (Supplementary Fig. 11) with an even lower overpotential of only 0.21 V. In that case, the last H_2O desorption step is the reaction determining step. The reaction mechanism for the dissociative $4e^-$ pathway separates the oxygen bond in the first protonation step producing O^* and OH^* instead of forming OOH^* . Thermodynamically, this pathway is favourable, but the O-O dissociation has a kinetic barrier with an activation energy of 0.56 eV. This is less than half the barrier on single Pt atoms on a similar substrate³⁵, and could be further influenced due to solvent effects. Both these two pathways are different from that of IA-Pt catalysts^{21, 35}, which follow a $2e^-$ pathway and has H_2O_2 as an intermediate product. Since O-O dissociation is the key step of the $4e^-$ pathway, it is suggested that the A-CoPt-NC exhibits a different reaction pathway compared to the reported IA-Pt catalysts^{21, 35}. Here, we attribute the high $4e^-$ pathway selectivity of A-CoPt-NC to the specific configuration structure of a(Co-Pt)@defect and the synergistic effect between the atomic Co and Pt sites, which results in the up-shifting of the d orbital and the charge polarization on the active site (a(Co-Pt)@N8V4). These factors eventually alter the dissociation energy of O-O bond and endow the $4e^-$ ORR pathway in A-CoPt-NC.

Electrocatalytic HER activity of A-CoPt-NC in wide pH range

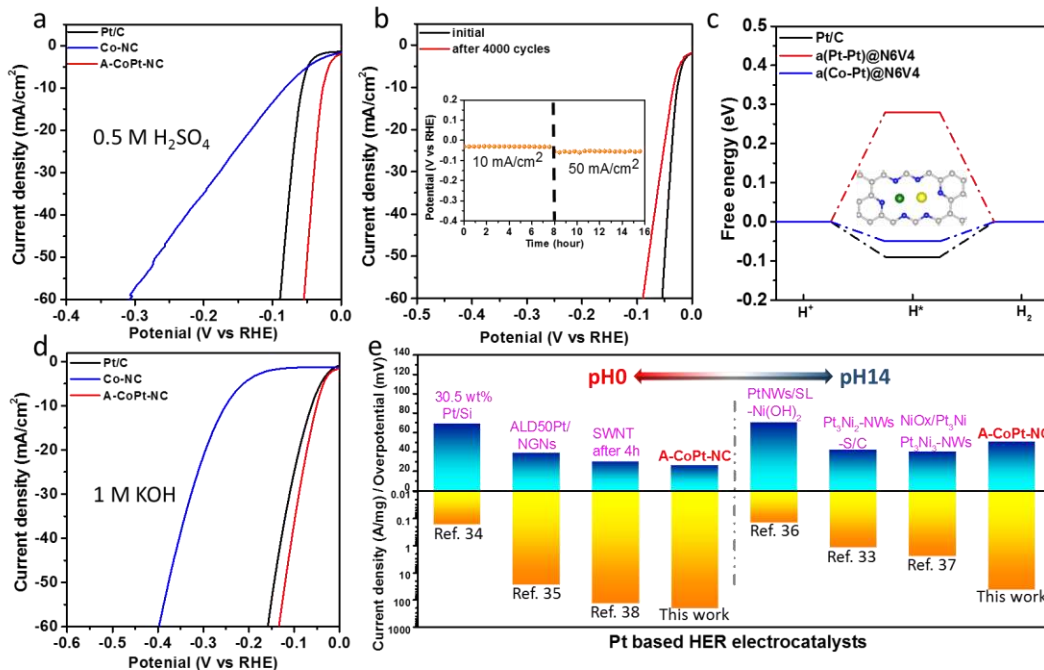


Figure 5 | Electrochemical hydrogen evolution activities. **a**, HER LSV curves of Co-NC, A-CoPt-NC and Pt/C in 0.5 M H₂SO₄ electrolyte. **b**, The durability test of A-CoPt-NC for HER. The polarization curves were recorded initially and after 4000 CV sweeps at a rate of 100 mV/s. The inset contains chronopotentiometry curves with a current density of 10 mA/cm² and 50 mA/cm², respectively. **c**, The HER free energy profiles of the Pt/C, 2H/a(Pt-Pt)@N6V4 and 2H/a(Co-Pt)@N6V4. Top view of the model a(Co-Pt)@N6V4 are inset. **d**, HER LSV curves of Co-NC, A-CoPt-NC and Pt/C in 1 M KOH electrolyte. **e**, The comparison of the overpotential needed to reach a current density of 10 mA/cm² (up) and the current density normalized to Pt mass at an overpotential of 70 mV (down) of various HER catalysts in the wide pH range. The data were collected from ref.³⁶⁻⁴¹.

As demonstrated above, electron accumulation on Co will benefit for the electrocatalytic reduction reactions, so it is natural to consider the A-CoPt-NC as an ideal catalyst for the HER.

Fig. 5a shows the LSV curves of Co-NC, A-CoPt-NC and Pt/C in the 0.5 M H₂SO₄ solution. The overpotentials at a current density of 10 mA/cm² were measured to be 27 mV for A-CoPt-NC and 59 mV for Pt/C, respectively (Supplementary Fig.12). A-CoPt-NC also exhibits a similar Tafel slope of 31 mV/dec to Pt/C, indicating the Volmer-Tafel pathway. Moreover, the durability evaluation (Fig. 5b) shows that after 4000 CV cycles, only a slight decay can be observed in the LSV curve and the chronopotentiometry curves retain stable for 8 hours reaction at the current densities of 10 mA/cm² and 50 mA/cm², respectively, indicating good stability of the A-CoPt-NC during the long-term HER reaction in an acidic environment. To further investigate the reaction mechanism of A-CoPt-NC for HER, we determined the energy profiles using DFT calculations. As shown in Fig. 5c, the two optimized configurations with the lowest $|\Delta G_H|$ are a(Co-Pt)@N6V4 and a(Pt-Pt)@N6V4, differing from that in ORR process. A $|\Delta G_H|$ value of zero represents the ideal interacting energy between the adsorbed hydrogen and the catalyst. The value of $|\Delta G_H|$ for a(Co-Pt)@N6V4 is 0.05 eV, lower than those of Pt/C and a(Pt-Pt)@N6V4. The local densities of states (Supplementary Fig.13) of a(Co-Pt)@N6V4 show an obvious higher state density than a(Pt-Pt)@N6V4 near the Fermi level, thereby increasing the coupling strength of the H* and catalysts. Furthermore, the charge distribution patterns (Supplementary Fig.14) demonstrate that the a(Co-Pt)@N6V4 possesses higher electron density around Co atom than that around Pt atoms in a(Pt-Pt)@N6V4, implying the strong bonding capability with H* at Co site. In desorption step of H*, for a(Co-Pt)@N6V4, the coupling is not strong enough, while for Pt/C, the interactions are stronger and thus the desorption of the H₂ produced requires more energy than for a(Co-Pt)@N6V4.

Considering that different HER devices in the realistic applications may be operated in a various range of pH, a wide pH range tolerance is significant for the HER catalyst. Moreover, the wide pH range tolerance also endows the catalyst to be multifunctional with OER or/and ORR in their specific pH ranges. Fig. 5d indicates that A-CoPt-NC outperforms Pt/C in alkaline

media for HER. Due to the ultralow loading of Pt (0.16 wt %) in A-CoPt-NC, it can be a promising alternative to commercial Pt/C catalyst and other Pt contained catalysts in the wide pH range of HER applications. Fig. 5e and Supplementary Table 4 show η_{10} (the overpotential required to reach a current density of 10 mA/cm²) and $J_{m,70}$ (activity normalized to Pt mass at an overpotential of 70 mV) of various reported catalysts and A-CoPt-NC in the wide pH range (A-CoPt-NC is from this work and the other data were collected from ref³⁶⁻⁴¹). The A-CoPt-NC exhibits the lowest η_{10} in acidic media and a competitive η_{10} in alkaline media for HER compared to other Pt based electrocatalysts. Considering the advantage of ultralow loading, the A-CoPt-NC achieves the highest $J_{m,70}$ in both acidic and alkaline media, which are 224 and 45 A/mg, respectively. Even in the neutral media, A-CoPt-NC still exhibits a comparable activity to Pt/C catalyst (Supplementary Fig.15), implying the A-CoPt-NC could be utilized over the full pH range from acid to alkaline. Supplementary Fig.16 and Supplementary Table 5 summarized the η_{10} of all pH range functioned HER catalysts without Pt and the A-CoPt-NC in this work (the Pt content is extremely low, with a ~ 0.0005 mg/cm² loading). Compared to these reported catalysts, A-CoPt-NC presents the highest HER activities in acidic and alkaline media and comparable activity in neutral media.

Conclusions

Utilizing the strategy of the electrochemical activation, Co cores were removed from the stable Co/C core-shell structures producing nitrogen doped defective carbons with atomic metal species. The activation process enables the removal of amorphous carbons, creating channels in graphitic carbon shells to allow ingress of acidic solvent and the gradual removal Co cores. Some of the atomic Co species were captured in the carbon shell. When using a Pt electrode during the activation, atomic Pt species from the dissolution in electrolyte can also be co-captured by the N-doped carbon shell. Accordingly, a carbon-based catalyst decorated by Co/Pt at an atomic scale is synthesized. The resulting catalyst (denoted as A-CoPt-NC) only

contains a small amount of Co (~1.72 wt%) and a very little Pt (~0.16 wt%), but shows extremely high activities for both ORR and HER. The ORR mass activity is as high as 267 times of the commercial Pt/C and 20 times of the US DOE target at 0.90 V in alkaline. The catalyst also exhibits considerably high ORR activity in acid but requires further improvement, which is being considered in our further research. Moreover, the HER of this catalyst is much superior to the commercial Pt/C both in acid and alkaline media. DFT calculations suggested that the excellent electrocatalytic performance may originate from the charge redistribution and the *d* orbital shift resulting from the synergetic effect of the atomic Pt and Co species in the specific coordination structure (a(Co-Pt)@N8V4 for ORR and a(Co-Pt)@N6V4 for HER). It is found that the atomic interaction of Pt-Co may be responsible for the high selectivity for 4e⁻ pathway in ORR, differing from the reported 2e⁻ pathway in isolated atomic Pt-based catalysts. Thus, the appropriate coordination environment of atomic metal species by defect engineering is of importance to tune the corresponding electronic redistribution for electrocatalysis, calling for a re-thinking of accepted strategies for developing efficient electrochemical catalysts.

Methods

Synthesis of Co-MOF. In a typical synthesis, 0.2 g Co(NO₃)₂·6H₂O, 0.21 g trimesic acid (H₃BTC) and 0.02 g 4,4'-bipyridine were dissolved in the 8.47 mL dimethylformamide (DMF). Subsequently, 1 mL water and 7.18 mL diethylene glycol were added to the solution with 2 hours stirring to mix the components uniformly. Then the solution was kept at the temperature of 65 °C for 48 hours. The Co-MOF was obtained after centrifugation.

Synthesis of A-CoPt-NC. A-CoPt-NC was prepared from Co-NC. Typically, the Co-MOF was mixed with dicyandiamide (mass ratio is 1:16) and annealed at 850 °C for 2 hours with a ramp rate of 4 °C under nitrogen atmosphere. Before calcining, the system was purged for two hours with nitrogen gas to ensure the removal of oxygen from the furnace. Then 4 mg of the

Co-NC was dispersed in the 1 mL ethanol/nafion (200 μ L/80 μ L) solution for at least 30 min ultrasonication. 10 μ L of the mixture was dropped onto a polished glassy carbon electrode (4 mm in diameter).

Characterizations. Raman spectrum was recorded on a Renishaw InVia spectrometer with a model 100 Ramascope optical fibre instrument. X-ray photoelectron spectrum (XPS) data was collected from a Kratos Axis ULTRA X-ray photoelectron spectrometer, and the binding energy of the C 1s peak at 284.8 eV was used as an internal reference. Co and Pt K edge X-ray absorption fine structure (XAFS) data were collected at Hard X-ray micro analysis beamline (HXMA, 06ID). Transmission electron microscopy (TEM) images were collected from TECNAI 12 with acceleration voltages of 120 kV. Scanning transmission electron microscopy-Energy-dispersive X-ray spectroscopy (STEM-EDS) elemental mapping images were obtained from TECNAI G2 F20 with acceleration voltages of 200 kV. High angle annular dark field (HAADF) images and BF images are collected from probe-corrected JEOL ARM200F with acceleration voltages of 80 kV.

Electrochemical measurements. All the electrochemical tests were performed in a conventional three-electrode system at an electrochemical station (CHI 760E), using Ag/AgCl (saturated KCl solution) electrode as the reference electrode, graphitic carbon rod as the counter electrode and glassy carbon (GC) electrode as the working electrode. All potentials were referred to the reversible hydrogen electrode (RHE) by following calculations: E (vs RHE) = E (vs Ag/AgCl) + 0.197 + 0.059pH. 4 mg of sample and 80 μ l of 5 wt. % Nafion solution were dispersed in 1 ml of 4:1 v/v water/ethanol by at least 60 min sonication to form a homogeneous solution. Then 5 μ l of the solution was loaded onto the GC electrode of 3 mm in diameter. The final loading for all catalysts and commercial Pt/C electrocatalysts on the GC electrodes is about 0.262 mg/cm². Linear sweep voltammetry with a scan rate of 5 mV/s was conducted in

1M KOH. Chronopotentiometry measurement ($j = 5 \text{ mA/cm}^2$ and 10 mA/cm^2) was performed to evaluate the long-term stability.

Calculations. All calculations were performed with the Vienna Ab initio Simulation Package (VASP)⁴² using density functional theory (DFT). The projector augmented-wave (PAW)⁴³ method, the revised Perdew-Burke-Ernzerhof (RPBE)⁴⁴ functional and the dispersion correction by Grimme (DFT-D3)⁴⁵ were used to describe the electronic interactions. Different structures were constructed to model the experimental systems to match the metal-metal distance. Pores were formed in a supercell of 8×8 graphene unit cells by removing carbon atoms. The different structures are illustrated and labelled in Supplementary Fig.8 (V – number of missing carbon atoms) and some carbon atoms at the edges were exchanged with nitrogen (N – number of nitrogen atoms). The unit cell size was $19.74 \times 17.09 \times 20 \text{ \AA}$ with a $2 \times 2 \times 1$ gamma-centred k-point grid and a cut off energy of 450 eV. These parameters were selected after carrying out preliminary tests. The free energies were calculated using a method similar to the approaches by Nørskov et al.^{46, 47} (see supplementary information) using the zero point energy (ZPE) and change in entropy (ΔS°) for adsorption of the gases species on a zirconia surface⁴⁸. These results were used since a complete set of data was available and the results are expected to be insensitive to the surface.

References

1. Li MF, *et al.* Ultrafine jagged platinum nanowires enable ultrahigh mass activity for the oxygen reduction reaction. *Science* **354**, 1414-1419 (2016).
2. He DP, *et al.* Amorphous nickel boride membrane on a platinum-nickel alloy surface for enhanced oxygen reduction reaction. *Nature Communications* **7**, 12362 (2016).
3. Escudero-Escribano M, *et al.* Tuning the activity of Pt alloy electrocatalysts by means of the lanthanide contraction. *Science* **352**, 73-76 (2016).

4. Bu LZ, *et al.* Biaxially strained PtPb/Pt core/shell nanoplate boosts oxygen reduction catalysis. *Science* **354**, 1410-1414 (2016).
5. Huang XQ, *et al.* High-performance transition metal-doped Pt₃Ni octahedra for oxygen reduction reaction. *Science* **348**, 1230-1234 (2015).
6. Chen Z, Ye S, Wilson AR, Ha Y-C, Wiley BJ. Optically transparent hydrogen evolution catalysts made from networks of copper–platinum core–shell nanowires. *Energy & Environmental Science* **7**, 1461-1467 (2014).
7. Sheng W, Zhuang Z, Gao M, Zheng J, Chen JG, Yan Y. Correlating hydrogen oxidation and evolution activity on platinum at different pH with measured hydrogen binding energy. *Nature communications* **6**, 6848 (2015).
8. Roduner E. Size matters: why nanomaterials are different. *Chemical Society Reviews* **35**, 583-592 (2006).
9. Li L, *et al.* Investigation of catalytic finite-size-effects of platinum metal clusters. *The journal of physical chemistry letters* **4**, 222-226 (2012).
10. Qiao BT, *et al.* Single-atom catalysis of CO oxidation using Pt-1/FeOx. *Nat Chem* **3**, 634-641 (2011).
11. Moses-DeBusk M, *et al.* CO Oxidation on Supported Single Pt Atoms: Experimental and ab Initio Density Functional Studies of CO Interaction with Pt Atom on theta-Al₂O₃(010) Surface. *J Am Chem Soc* **135**, 12634-12645 (2013).
12. Qiao B, *et al.* Highly efficient catalysis of preferential oxidation of CO in H₂-rich stream by gold single-atom catalysts. *ACS Catal* **5**, 6249-6254 (2015).
13. Yang M, *et al.* Catalytically active Au-O(OH)(x)-species stabilized by alkali ions on zeolites and mesoporous oxides. *Science* **346**, 1498-1501 (2014).
14. Lin J, *et al.* Remarkable Performance of Ir-1/FeOx Single-Atom Catalyst in Water Gas Shift Reaction. *J Am Chem Soc* **135**, 15314-15317 (2013).
15. Yin PQ, *et al.* Single Cobalt Atoms with Precise N-Coordination as Superior Oxygen Reduction Reaction Catalysts. *Angew Chem Int Ed* **55**, 10800-10805 (2016).
16. Fan LL, *et al.* Atomically isolated nickel species anchored on graphitized carbon for efficient hydrogen evolution electrocatalysis. *Nat Commun* **7**, 10667 (2016).

17. Qiu HJ, *et al.* Nanoporous Graphene with Single-Atom Nickel Dopants: An Efficient and Stable Catalyst for Electrochemical Hydrogen Production. *Angew Chem Int Ed* **54**, 14031-14035 (2015).
18. Shen H, *et al.* Atomically FeN₂ moieties dispersed on mesoporous carbon: A new atomic catalyst for efficient oxygen reduction catalysis. *Nano Energy* **35**, 9-16 (2017).
19. Yang S, Tak YJ, Kim J, Soon A, Lee H. Support Effects in Single-Atom Platinum Catalysts for Electrochemical Oxygen Reduction. *ACS Catal* **7**, 1301-1307 (2016).
20. Yang S, Kim J, Tak YJ, Soon A, Lee H. Single - Atom Catalyst of Platinum Supported on Titanium Nitride for Selective Electrochemical Reactions. *Angew Chem Int Ed* **55**, 2058-2062 (2016).
21. Choi CH, *et al.* Tuning selectivity of electrochemical reactions by atomically dispersed platinum catalyst. *Nat Commun* **7**, 10922 (2016).
22. Jia Y, *et al.* Defect Graphene as a Trifunctional Catalyst for Electrochemical Reactions. *Adv Mater* **28**, 9532-9538 (2016).
23. Tang C, Wang B, Wang H-F, Zhang Q. Defect Engineering toward Atomic Co-N_x-C in Hierarchical Graphene for Rechargeable Flexible Solid Zn-Air Batteries. *Advanced Materials*, 1703185.
24. Zhang L, *et al.* Defects on graphene trapping atomic Ni species for hydrogen and oxygen evolution reactions. *Chem*, (2018).
25. Xiao P, Ge X, Wang H, Liu Z, Fisher A, Wang X. Novel molybdenum carbide-tungsten carbide composite nanowires and their electrochemical activation for efficient and stable hydrogen evolution. *Advanced Functional Materials* **25**, 1520-1526 (2015).
26. Das RK, *et al.* Extraordinary Hydrogen Evolution and Oxidation Reaction Activity from Carbon Nanotubes and Graphitic Carbons. *Acs Nano* **8**, 8447-8456 (2014).
27. Fei H, *et al.* Atomic cobalt on nitrogen-doped graphene for hydrogen generation. *Nat Commun* **6**, 8668 (2015).
28. Zitolo A, *et al.* Identification of catalytic sites for oxygen reduction in iron- and nitrogen-doped graphene materials. *Nature Materials* **14**, 937-942 (2015).
29. Liu W, *et al.* Single-atom dispersed Co-N-C catalyst: structure identification and performance for hydrogenative coupling of nitroarenes. *Chemical Science* **7**, 5758-5764 (2016).
30. Alves MCM, Dodelet JP, Guay D, Ladouceur M, Tourillon G. Origin of the electrocatalytic properties for oxygen reduction of some heat-treated polyacrylonitrile and phthalocyanine

cobalt compounds adsorbed on carbon black as probed by electrochemistry and x-ray absorption spectroscopy. *The Journal of Physical Chemistry* **96**, 10898-10905 (1992).

31. Li XG, *et al.* Single-Atom Pt as Co-Catalyst for Enhanced Photocatalytic H₂ Evolution. *Advanced Materials* **28**, 2427-2431 (2016).
32. Li L, Xing Y. Electrochemical durability of carbon nanotubes in noncatalyzed and catalyzed oxidations. *Journal of the Electrochemical Society* **153**, A1823-A1828 (2006).
33. Varnell JA, *et al.* Identification of carbon-encapsulated iron nanoparticles as active species in non-precious metal oxygen reduction catalysts. *Nature Communications* **7**, 12582 (2016).
34. Hammer B, Nørskov JK. Theoretical surface science and catalysis—calculations and concepts. *Advances in catalysis* **45**, 71-129 (2000).
35. Liu S, Huang S. Theoretical insights into the activation of O₂ by Pt single atom and Pt₄ nanocluster on functionalized graphene support: Critical role of Pt positive polarized charges. *Carbon* **115**, 11-17 (2017).
36. Wang PT, *et al.* Precise tuning in platinum-nickel/ nickel sulfide interface nanowires for synergistic hydrogen evolution catalysis. *Nature Communications* **8**, 14580 (2017).
37. Zhu LL, *et al.* A rhodium/silicon co-electrocatalyst design concept to surpass platinum hydrogen evolution activity at high overpotentials. *Nature Communications* **7**, 12272 (2016).
38. Cheng NC, *et al.* Platinum single-atom and cluster catalysis of the hydrogen evolution reaction. *Nature Communications* **7**, 13638 (2016).
39. Yin HJ, *et al.* Ultrathin platinum nanowires grown on single-layered nickel hydroxide with high hydrogen evolution activity. *Nature Communications* **6**, 6430 (2015).
40. Wang PT, Jiang KZ, Wang GM, Yao JL, Huang XQ. Phase and Interface Engineering of Platinum-Nickel Nanowires for Efficient Electrochemical Hydrogen Evolution. *Angewandte Chemie-International Edition* **55**, 12859-12863 (2016).
41. Tavakkoli M, *et al.* Electrochemical Activation of Single-Walled Carbon Nanotubes with Pseudo-Atomic-Scale Platinum for the Hydrogen Evolution Reaction. *ACS Catalysis* **7**, 3121-3130 (2017).
42. Kresse G, Furthmüller J. Efficient iterative schemes for ab initio total-energy calculations using a plane-wave basis set. *Physical review B* **54**, 11169 (1996).
43. Blöchl PE. Projector augmented-wave method. *Physical review B* **50**, 17953 (1994).

44. Hammer B, Hansen LB, Nørskov JK. Improved adsorption energetics within density-functional theory using revised Perdew-Burke-Ernzerhof functionals. *Physical Review B* **59**, 7413 (1999).
45. Grimme S, Antony J, Ehrlich S, Krieg H. A consistent and accurate ab initio parametrization of density functional dispersion correction (DFT-D) for the 94 elements H-Pu. *The Journal of chemical physics* **132**, 154104 (2010).
46. Nørskov JK, *et al.* Origin of the overpotential for oxygen reduction at a fuel-cell cathode. *The Journal of Physical Chemistry B* **108**, 17886-17892 (2004).
47. Nørskov JK, *et al.* Trends in the exchange current for hydrogen evolution. *Journal of The Electrochemical Society* **152**, J23-J26 (2005).
48. Wang G, *et al.* A first-principle study of oxygen reduction reaction on monoclinic zirconia (111), (011) and (110) surfaces. *Catalysis Communications* **69**, 16-19 (2015).

Acknowledgements

The authors thank the financial support from Australia Research Council (ARC DP170103317). Y.J. also thanks ARC Discovery Early Career Researcher Award (ARC DE180101030), the Griffith University Postdoctoral and Research Fellowship and Griffith University New Research Grant. The authors would like to thank the Australian National Fabrication Facility (ANFF) – Materials node, University of Wollongong (UOW) Electron Microscopy Centre (EMC) and Shanghai Synchrotron Radiation Facility for equipment access. J.M.T.A.F. thanks the University of Queensland for the UQI scholarship supporting her Ph.D. We acknowledge access to the computational resources of the NCI National Facility at the Australian National University through the National Computational Merit Allocation Scheme supported by the Australian Government; and support from the Queensland Cyber Infrastructure Foundation (QCIF) and the University of Queensland Research Computing Centre.

Author contributions

X.Y. conceived and designed the project. X.Y. and Y.J. supervised the project. L.Z. prepared the samples and did the electrocatalytic performances test. L.Z., X.Y., W.X., X.W., S.F. and J.C. performed the characterizations including XRD, XPS, TEM, XAS and so on. J.M.T.A.F., M.H. and D.J.S. performed the DFT calculations. L.Z., Y.J., X.Y. and J.M.T.A.F. wrote the manuscript. L.Z. and J.M.T.A.F. contributed equally to this work. All authors discussed the results and commented on the manuscript.

Additional information

Supplementary information is available for this paper. Reprints and permissions information is available at www.nature.com/reprints. Correspondence and requests for materials should be addressed to Y.J. and X.Y.

Competing interests

The authors declare no competing financial interests.