Regulating the scaling relationship for high catalytic kinetics and selectivity of the oxygen reduction reaction

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The electrochemical oxygen reduction reaction (ORR) is at the heart of modern sustainable energy technologies. However, the linear scaling relationship of this multistep reaction now becomes the bottleneck for accelerating kinetics. Herein, we propose a strategy of using intermetallic-distance-regulated atomic-scale bimetal assembly (ABA) that can catalyse direct O–O radical breakage without the formation of redundant *OOH intermediates, which could regulate the inherent linear scaling relationship and cause the ORR on ABA to follow a fast-kinetic dual-sites mechanism. Using in situ synchrotron spectroscopy, we directly observe that a self-adjustable N-bridged Pt = N2 = Fe assembly promotes the generation of a key intermediate state (Pt–O–O–Fe) during the ORR process, resulting in high reaction kinetics and selectivity. The well-designed Pt = N2 = Fe ABA catalyst achieves a nearly two orders of magnitude enhanced kinetic current density at the half-wave potential of 0.95 V relative to commercial Pt/C and an almost 99% efficiency of 4-electron pathway selectivity, making it one of the potential ORR catalysts for application to the energy device of zinc–air cells. This study provides a helpful design principle for developing and optimizing other efficient ORR electrocatalysts.

The oxygen reduction reaction (ORR) is the core of renewable energy conversion technology and plays an irreplaceable role in emerging electrochemical energy devices such as metal-air batteries and fuel cells. However, the sluggish kinetics of the ORR that occurs on the cathode involves multiple-step proton-coupled electron transfer, which has a significant impact on improving the overall efficiency of energy conversion devices. Catalysts based on the noble metal platinum (Pt, US $36,084 kg⁻¹) with metal loadings up to 400 μgPt cm⁻² are the most practical for boosting the energy output efficiency of ORR-driven energy devices, but their application is still hindered by the scarcity of noble metal resources and limited stability under desirable high voltages (>0.6 V). Alloying Pt with a secondary metal can reduce the usage of Pt while improving the performance. However, the amount of Pt is still not less than 40 at.%, and the secondary metal leaches away gradually under ORR conditions, resulting in rapid performance losses, which sufficiently limits large-scale commercialization. Therefore, the design of low-loading active noble-metal ORR electrocatalysts with fast reaction kinetics and long-term durability is both desirable and essential for efficient energy conversion and storage.

According to the widely accepted conventional single-site mechanism of multiple-step proton-coupled electron transfer, the
ORR process involves multiple reaction intermediates, including 'O2, 'OOH, and 'O13,14. Of note, the binding energies of these intermediates over the single sites are closely related and follow the scaling relationship35,36. This means that the binding energy of each intermediate cannot be independently adjusted because of the insurmountable scaling relation. Consequently, the oxygen intermediates ('O2 and 'OOH) with larger binding energy directly lead to sluggish kinetics and require a high overpotential to drive the reaction37,38. As a result, the scaling relationship in the ORR now becomes the bottleneck for further improving its performance. To overcome this dilemma, an alternative route is to bypass the formation of oxygen intermediates ('OOH) and promote direct O-O radical breakage without extra reaction intermediates ('OOH) by a dual-site mechanism (M-O-M)29,30. Under this circumstance, the direct breakage of O-O on the dual-sites alleviates the complex dynamics of multiple reaction intermediates. More importantly, the dual-site mechanism is a fast 4-electron reaction path, since the sluggish kinetics of 'OOH production is overcome. Therefore, it can be anticipated that to further improve the performance and selectivity of current ORR electrocatalysis, the dual-site mechanism with a fast four-electron reaction path is an extremely ideal choice. However, the realization of this dual-sites mechanism requires stringent requirements on the geometric and electronic configuration of the active site. Intermetallic distances of too great a length cause the single-site mechanism to be accompanied by M–OOH, which is common in single-site catalysis, and distances that are too short, such as in metal nanoparticles, easily trigger the emergence of two-electron reaction paths21,22. Thus, an appropriate interatomic spacing in two adjacent metallic active sites is mandatory for the dissociation of 'O2 and the triggering of O-O radical breakage (M–O–M) to demonstrably suppress M-OOH production and limiting the two-electron reaction path selectivity during the ORR process.

Herein, to satisfy the dual-site mechanism design rule for high kinetics and selectivity of four-electron ORR, we develop a N-bridged Pt–N2 = Fe ABA (Supplementary Fig. 10a and Supplementary Table 2) catalyst is illustrated by the electron micrograph in Fig. 1a and Supplementary Fig. 5. The atomically dispersed metal sites in Pt–N2 = Fe ABA catalyst is revealed by Cs-correlated scanning transmission electron microscopy high-angle annular dark field (STEM-HAADF) images (Fig. 1b and Supplementary Fig. 6). Based on the fact that the intensity of atoms in the STEM-HAADF image highly depends on their atomic number (Z), intensity analysis was performed at four dual-site regions in STEM-HAADF images to clarify the composition of the dual-sites structure (Fig. 1c and Supplementary Fig. 6d, e)27. All the two bright spots along the arrow show different intensity, confirming that the dual-sites is composed of Pt and Fe atoms. The statistical analysis results show that the atomic distance between Pt and Fe is about 2.83–2.91 Å. And the statistical analysis based on 500 randomly selected metal sites reveals that the metal dual-sites structure dominates about 72.6% (Supplementary Fig. 7). In the X-ray diffraction (XRD) pattern (Fig. 1d), all the samples show only a major diffraction peak (2θ = 25°) of carbon, and no crystalline metal nanoparticles are observed. And the uniform distribution of Pt, Fe, N and C elements in the Pt–N2 = Fe ABA catalyst can be reflected by the energy-dispersive X-ray (EDX) mapping images (Fig. 1e, Supplementary Fig. 8).

The surface chemical states were detected by X-ray photoelectron spectroscopy (XPS). The XPS survey spectrum reveals that the Pt = N2 = Fe ABA catalyst contains Pt, Fe, N, C and O elements (Supplementary Fig. 9), and the metal contents of Pt and Fe in Pt = N2 = Fe ABA determined by XPS are 2.93 and 1.34 wt%, respectively, which are close to the inductively coupled plasma–optical emission spectrometry (ICP–OES) results (inset of Supplementary Fig. 9). The O 1 s XPS spectrum in Supplementary Fig. 10b shows the absence of M-O bond, suggesting the inexistence of metal oxide species or O-coordinated monoatomic metals in Pt = N2 = Fe ABA sample32,33. The N 1 s XPS spectrum was fitted into a typical peak at 399.35 eV, which could be attributed to the N atom interaction with metal sites (Pt=Fe=N) in Pt = N2 = Fe ABA (Supplementary Fig. 10a and Supplementary Table 1)23. In addition, the fitting results of Pt 4 f and Fe 2 p XPS spectra for Pt = N2 = Fe ABA show that the metal atoms have a positive charge state with no zero-valence species appearing (Supplementary Fig. 11, Supplementary Table 2–3). The oxidation states of Pt and Fe were further analyzed quantitatively based on the X-ray absorption near-edge structure (XANES) spectra (Supplementary Fig. 12a)24. The normalized Pt L3-edge XANES spectra by subtracting the spectra of Pt = N2 = Fe ABA and PtO2 sample to that of Pt foil reference are shown in Supplementary Fig. 12b. The valence state of Pt in the Pt = N2 = Fe ABA sample can be calculated as +3.36 (Supplementary Fig. 12c) through the area integral of normalized white-line peak, which is consistent with the valence state analysis in XPS characterization25. The valence
state of Fe can be quantitatively evaluated as +2.40 based on the difference value ($\Delta E_0$) of the absorption edge, which validates the XPS results (Supplementary Fig. 13).

The extended XAFS (EXAFS) was detected to further identify the coordination environment of metal sites in Pt = N$_2$ = Fe ABA. For the Fourier transformed (FT) EXAFS spectra of the Pt L$_3$-edge, the Pt = N$_2$ = Fe ABA shows a dominant peak at 1.55 Å (Fig. 1f), similar to that in Pt AMS samples, which is typically assigned to the first shell of Pt–N coordination. Concurrently, in contrast with the reference Pt AMS sample, another characteristic peak is clearly observed at 2.63 Å, suggesting the presence of a metal coordination of the second shell. It is noteworthy that the peak position of metal coordination is slightly longer than that of the Pt–Pt shell in Pt foil and shorter than the second shell of Pt–O–Pt in PtO$_2$. The Fe K-edge FT-EXAFS spectrum of Pt = N$_2$ = Fe ABA demonstrates a predominant peak at 1.42 Å (Fig. 1g), which is similar to that in Fe AMS samples, confirming the presence of Fe–N coordination. Moreover, the Pt = N$_2$ = Fe ABA catalyst presents a characteristic peak at 2.60 Å in the second shell that is obviously longer than the Fe–Fe bond in Fe foil, which may be attributed to the second shell of Fe–N–Pt coordination. Combined with the FT-EXAFS results of the Pt L$_3$-edge, it can be demonstrated that the Pt and Fe atoms in the catalyst are assembled with an appropriate intermetallic distance rather than bonding directly. Thus, we infer that the two nearby Pt and Fe atoms may coordinate with bridged N atoms at the second shell. To obtain the quantitative coordination information in the Pt = N$_2$ = Fe ABA catalyst, FT-EXAFS curve-fitting analysis was performed based on the structures of dual-sites (72.6%) and monometallic sites (25.3%) (Fig. 1h, Supplementary Fig. 14 and Supplementary Table 4–5). The fitting results of both Pt L$_3$-edge and Fe K-edge FT-EXAFS spectra show that the bond length ($R$) of the metal coordination is around 2.86 Å, corresponding to the atomic spacing analysis results in Cs-corrected STEM-HAADF images. Furthermore, in the best fitting results of Pt L$_3$-edge EXAFS, the coordination numbers (CNs) for the Pt–N coordination and the second shell of Pt–N–Fe are 4.1 and 1.1, respectively (Supplementary Table 4). The best fitting result for Fe K-edge EXAFS shows CNs of 4.2 and 1.0 for the Fe–N bond and Fe–N–Pt coordination, respectively (Supplementary Table 5). The above results imply that each of the Pt and Fe atoms coordinates with four N atoms to form an M–N$_6$ configuration, and the adjacent Pt and Fe atoms are connected by two shared bridge N atoms to form a Pt–Fe$_3$N$_6$–N$_6$ geometric configuration.

**Electrochemical oxygen reduction performance**

The electrocatalytic ORR activities of the as-synthesized Pt = N$_2$ = Fe ABA, Pt AMS, Fe AMS catalysts and the commercial Pt/C were evaluated on a rotating disk electrode (RDE) system. The cyclic voltammetry curves were recorded for the synthesized platinum-based catalysts (Pt = N$_2$ = Fe ABA and Pt AMS) in N$_2$-saturated 0.1 M KOH electrolyte, which present no hydrogen adsorption and/or desorption.

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**Fig. 1 | Morphology and structural characterizations. a, b** TEM (a) and HAADF-STEM (b) images of the Pt = N$_2$ = Fe ABA catalyst. c Intensity profile along Line 1 and Line 2 in b. d XRD patterns of Pt=N$_2$=Fe ABA, Fe AMS and Pt AMS. e TEM-EDS mapping images for Pt = N$_2$ = Fe ABA. f–h FT-EXAFS spectra of Pt L$_3$-edge (f) and Fe K-edge (g) for Pt = N$_2$ = Fe ABA catalyst and reference samples, and the corresponding fitting curves for Pt = N$_2$ = Fe ABA (h).
characteristic due to the monoatomic dispersion of platinum in the synthesized catalysts (Supplementary Fig. 15)\(^\text{38}\). The linear sweep voltammery (LSV) curves illustrated in Fig. 2a obviously show that Pt = N\(_2\) = Fe ABA has a superior performance compared with the reference catalyst. As quantified in Fig. 2b, Pt = N\(_2\) = Fe ABA possesses a half-wave potential (\(E_{1/2}\)) of 0.95 V vs. RHE and onset potential (\(E_{\text{onset}}\)) of 1.05 V vs. RHE, which are 90 and 100 mV higher than those of commercial Pt/C (0.86 and 0.95 V), also outperforming those of Pt AMS, Fe AMS and almost all reported ORR catalysts (Supplementary Table 6). As for the reaction kinetics, the Pt = N\(_2\) = Fe ABA presents an ultrahigh kinetic current density (\(i_\text{K}\)) of 5.83 mA cm\(^{-2}\) at 0.95 V (Fig. 2b), nearly two orders of magnitude higher than that of commercial Pt/C (\(i_\text{K}\), 0.09 mA cm\(^{-2}\)). The accelerated current density of Pt = N\(_2\) = Fe ABA at 0.85 V is still dozens of times higher than that of commercial Pt/C (Pt = N\(_2\) = Fe ABA, 106.3 mA cm\(^{-2}\); Pt/C, 9.83 mA cm\(^{-2}\)). The superior ORR kinetics of Pt = N\(_2\) = Fe ABA is further confirmed by the smaller Tafel slope (53 mV dec\(^{-1}\)), which is significantly smaller than those of commercial Pt/C (84 mV dec\(^{-1}\), Fe AMS (75 mV dec\(^{-1}\)) and Pt AMS (90 mV dec\(^{-1}\)) (Fig. 2c). Based on the roughly equal numbers of active sites in the as-synthesized catalysts, the improved catalytic performance of the Pt = N\(_2\) = Fe ABA is mainly attributed to the formed Pt–N–Fe dual-sites. Notably, the criteria of turnover frequency (TOF) and mass activity (MA), which can characterize the intrinsic activity of the catalyst, were calculated and shown in Fig. 2d. Pt = N\(_2\) = Fe ABA displays a TOF of 15.3 e–site\(^{-1}\)s\(^{-1}\) and a large MA of 14.1 A mg\(_{\text{Pt}}\)\(^{-1}\) at 0.90 V vs. RHE, 32 and 61 times higher than that of commercial Pt/C reference (0.48 e–site\(^{-1}\)s\(^{-1}\), 0.23 A mg\(_{\text{Pt}}\)\(^{-1}\)), and the high MA of platinum in Pt = N\(_2\) = Fe ABA exceeds most of the reported platinum-based catalysts (Supplementary Table 7). The cyclic voltammetry curves of Pt = N\(_2\) = Fe ABA catalyst and the reference samples with the scanning rates from 10 to 50 mV s\(^{-1}\) are shown in Supplementary Fig. 16. As a result, the Pt = N\(_2\) = Fe ABA catalyst has the highest electrochemically accessible surface area (ECSA) of 210.5 m\(^2\) g\(^{-1}\), corresponding to the highest ORR activity\(^{39}\).

Catalytic selectivity is an important index used to evaluate the performance of electrocatalysts, especially for electrocatalytic oxygen reduction. The ORR pathway was assessed via polarization curve measurements at different rotation rates (Fig. 2e)\(^{38}\). The electron transfer number (\(n\)) was calculated to be 3.98 according to the Levich equation applied in the diffusion-controlled region (inset of Fig. 2e and Supplementary Fig. 17), proving that the Pt = N\(_2\) = Fe ABA catalyst favours the 4-electron pathway. Moreover, the measurement conducted on the rotating ring-disk electrode (RRDE) also shows that \(n\) for Pt = N\(_2\) = Fe ABA is 3.98 (Fig. 2f), which is better than Pt/C and reference samples, confirming that the side reaction of the 2e\(^{-}\) pathway was effectively suppressed. Importantly, the \(\text{H}_2\text{O}_2\) yield of Pt = N\(_2\) = Fe ABA was measured to be below 1.5%, which represents a high selectivity for the 4-electron ORR\(^{38}\). The rapid O–O bond breaking are realized over Pt = N\(_2\) = Fe ABA for an efficient 4-electron ORR process. In addition, stability during operation is particularly important for the commercial application of catalysts. The Pt = N\(_2\) = Fe ABA catalyst demonstrates reliable stability with better carbon oxidation and methanol resistance in chronamperometry measurements, in which the \(E_{1/2}\) only drops 15 mV and the capacitance increases <4% after continuous operation for 100 h (Supplementary Fig. 18–20). The Pt = N\(_2\) = Fe ABA catalyst also exhibits advanced performance in acidic electrolyte in comparison with the monometallic sites catalysts (Pt AMS and Fe AMS) and the standard commercial Pt/C (Supplementary Fig. 21), which suggests the synergistic effect of the dual-sites in Pt = N\(_2\) = Fe ABA for catalyzing the ORR process. These results demonstrate that the Pt = N\(_2\) = Fe ABA catalyst can serve as a promising electrocatalyst with prominent electroreduction activity and selectivity.

In situ XAFS analysis

The in situ XANES and EXAFS spectra, which can reflect the dynamic evolution of active sites during the actual reaction process, were employed to explain the nature of the high ORR activity\(^{38}\). Pt \(L_\gamma\)-edge XANES in Fig. 3a and the corresponding local magnification of Fig. 3b show that the white-line intensity increases obviously when a potential
of 1.05 V is applied, arising from the higher oxidation state caused by the emergence of Pt sites coordinated with oxygen species. With the application of oxygen reduction potentials, the intensity of the white line decreases, implying an electron transition from nearby atoms to Pt sites for an accelerated catalytic reaction process. Interestingly, for the Fe K-edge XANES (Fig. 3b, c), the white-line intensity changes similarly with potential hysteresis, indicating the coordination with oxygen at the Fe sites are closely adjacent to the Pt sites. This implies that Pt and Fe are synergistic active centres that act as adsorption/desorption sites for reactive species.

To further clarify the structural evolution of the bimetal assembly active sites, FT-EXAFS spectra are presented for both Pt L₂-edge and Fe K-edge. For the FT-EXAFS spectra of Pt L₂-edge under different reaction conditions in Fig. 3e, the dominant peak at ~1.55 Å, which can be attributed to the first-shell of Pt–N/O coordination, obviously increases in intensity, especially from at KOH to at 1.05 V. Analogously, in the FT-EXAFS spectra of Fe K-edges (Fig. 3f), we observe the same phenomenon that the intensity of the dominant peak at ~1.42 Å increases with a voltage of 0.95 V, indicating that both Pt and Fe metal sites are involved in the reaction and have a similar coordination evolution in the ORR process. To quantify the evolution of the local structure at Pt = N₂ = Fe dual-sites, the corresponding FT-EXAFS curve fitting was conducted, as shown in Fig. 3e–g, Supplementary Figs. 22–25, and Supplementary Tables 4–5. The best fitting results of the Pt L₂-edge show that the coordination numbers (CNs) of the Pt–N path is 4.0, which is consistent with the Pt–N₄ configuration obtained from the ex-situ Pt – N₂ = Fe ABA sample. When a potential of 1.05 V is applied, considering the inevitable adsorption of oxygen species during the reaction, one additional Pt–O coordination (2.04 Å) is added. It cannot be ignored that Fe–O coordination is added with a CN of 1.2 at 0.95 V, based on the fitting of Fe K-edge FT-EXAFS curves as Fe–N₄ coordination under KOH conditions. Comprehensive analysis of the fitting results for the dual-sites shows that the oxygen molecules can be adsorbed separately on the appropriately distanced Pt and Fe atoms, likely promoting the intermetallic assembling-sites co-coupling of dioxygen intermediates (~O–O–) in a dual-site adsorption form (Pt–O–O–Fe). It is of particular interest that the intermetallic distance of the second-shell Pt–N–Fe coordination at ~2.6 Å in both the Pt L₂-edge and Fe K-edge FT-EXAFS curves tends to shrink under operating conditions, indicating that the N-bridged Pt = N₂ = Fe assembly can be self-adjustable during the reaction process. This behaviour facilitates the formation of Pt–O–O–Fe active intermediate state at the Pt = N₂ = Fe dual-sites. Two adjacent metal atoms in Pt = N₂ = Fe assembly each binding an O atom of a dioxygen species (~O–O–) can provide a strong driving force for direct O–O bond breakage, regulating the scaling relationship of multiple reaction intermediates for a fast 4-electron ORR process.

**In situ SR-FTIR analysis**

In situ SR-FTIR, which can sensitively capture the reactive species and identify the reaction mechanism, was performed under ORR
operating conditions. As shown in Fig. 4a, an IR absorption band located at 1115 cm$^{-1}$ is observed with a gradually applied potential. It is known that the absorption bands within the range of 1060–1150 cm$^{-1}$ are associated with the IR absorption of O–O$^{15,46}$. Consequently, this voltage-dependent vibration suggests that the key intermediate O–O is generated over the Pt$–$N$_2$–Fe dual-sites to form a Pt–O–O–Fe configuration during the ORR process. Combined with the in situ XAFS results, the dual-sites of Pt and Fe favouring the adsorption of O$_2$ molecules directly evolve into O–O radicals. As a reference, Fig. 4b shows the FTIR measurement results of the Pt AMS sample under the same conditions. A new IR absorption band increases rapidly as the adjacent Pt and Fe atoms is contracted, and the metallic spacing in Pt$–$N$_2$–Fe ABA can greatly reduce the reaction energy barrier of the 4-electron reaction process and achieves high activity and selectivity of catalytic ORR.

Fig. 4 | The reaction pathway identified by in situ SR-FTIR. a, b 800–1300 cm$^{-1}$ range of in situ SR-FTIR characterizations for Pt$–$N$_2$–Fe ABA (a) and Pt AMS (b). c FTIR absorption stretching at 1115 cm$^{-1}$ for Pt$–$N$_2$–Fe ABA and at 965 cm$^{-1}$ for Pt AMS. d, e Schematic of the ORR mechanism for Pt$–$N$_2$–Fe ABA (e) and Pt AMS (d).
Applications in renewable energy devices

We assembled the Pt = N2 = Fe ABA as the cathode of a primary Zn–air battery (ZAB) to evaluate the efficiency of the catalyst in renewable energy devices (Fig. 5a). As shown in Fig. 5b, the Pt = N2 = Fe ABA-incorporated Zn–air cell exhibits an open-circuit voltage (OCV) of 1.50 V, which is higher than that of the counterpart Pt/C-based Zn–air cell (1.42 V). Notably, a maximum power density of 198.4 mW cm⁻² is achieved for the ZAB incorporated with the Pt = N2 = Fe ABA catalyst due to the favourable mass transfer (Fig. 5c), outperforming the Pt/C-based ZAB (172.1 mW cm⁻² at 253 mA cm⁻²) and ZABs made from other catalysts (Supplementary Table 8). Furthermore, the ZAB with the Pt = N2 = Fe ABA air cathode delivers a specific capacity of 787.8 mAh g⁻¹ Zn⁻¹ at a discharge current density of 10 mA cm⁻², accounting for 96.1% of the theoretical capacity of ZAB (~820 mAh g⁻¹ Zn⁻¹) (Fig. 5d). The enhanced performance proves that the Pt = N2 = Fe ABA catalyst can deliver efficiency adequate for justifying incorporation into Zn–air cells. The galvanostatic discharge measurements under different current density platforms are shown in Fig. 5e. The cell voltage responses of the ZAB with Pt = N2 = Fe ABA are obviously higher than those of the Pt/C cell at each current density, and the Pt = N2 = Fe ABA cell voltage returns to its original level as the current density reverts to 2 mA cm⁻². This suggests the considerable rate performance and robust stability of Pt = N2 = Fe ABA as a cathode in a ZAB. In addition, this Pt = N2 = Fe ABA-based ZAB can work stably through mechanical recharging and requires only replenishment of the consumed zinc anode and electrolytes. No apparent decline in the output voltage is observed after three charging cycles (Fig. 5f). These results reveal that a unique Pt = N2 = Fe ABA catalyst which maintains satisfactory catalytic activity in practical operation has great application prospect in industrial ORR.

In summary, a highly efficient and selective N-bridged Pt = N2 = Fe bimetal assembly catalyst with appropriate intermetallic atomic spacing and unique electronic structure was developed by a controllable amino functionalized carbon nanoflakes strategy. In combination with in situ XAFS and SR-FTIR techniques, we identify that the key intermediate state (Pt–O–O–Fe) was successfully formed over self-adjustable N-bridged Pt = N2 = Fe bimetal assembly sites under working conditions, which efficiently catalyses the ORR following a dual-sites mechanism, promoting the generation and fast cleavage of O–O radical intermediates without the formation of the traditional reactive product *OOH. The as-obtained Pt = N2 = Fe ABA catalyst delivers a quite appreciable 4-electron selectivity of ~99% and a high kinetic current density which exceeds the commercial Pt/C by two orders of magnitude. Interestingly, the Pt = N2 = Fe ABA catalyst guarantees high efficiency and robust stability in zinc-air batteries. Our results provide a useful design principle for developing highly active and selective ORR catalysts.

Methods

Synthesis of functionalized CNF–NH₂

The functionalized CNF–NH₂ was first synthesized as a precursor. Typically, 1 g pyrene (Aladdin, 99%) was dissolved in 80 ml nitric acid, refluxed and stirred at 80 °C for 12 h, cooled to room temperature, and centrifuged and washed in deionized water until neutral. The nitrated product 1,3,6-trinitropyrene was obtained by vacuum drying. 1,3,6-Trinitropyrene (300 mg) was dissolved in a mixed solution of 35 ml deionized water and 5 ml concentrated ammonia water (30%, Sigma) and ultrasonicated in cold water for 4 h. After the homogenized suspension was formed, it was poured into a Teflon-lined autoclave and heated at 200 °C for 10 h before cooling to room temperature. The solution was filtered through a 0.22 μm microporous membrane to remove the insoluble carbon products and then concentrated to ~20 mL by rotary evaporation to obtain the CNF–NH₂ concentrated solution.

Pt = N2 = Fe ABA synthesis

Typically, a mixed solution was made by adding 5.75 mg of H₂PtCl₆ and 2.27 mg of FeCl₃ to the glycol solvent. The mixture was ultrasonicated and mechanically agitated for 20 min to promote the effective association between electropositive Fe³⁺ and electronegative [PtCl₆]²⁻.
Then, 20 ml of concentrated CNF–NH₂ solution was added to the resulting mixed solution, sonicated in cold water for 15 min, quickly frozen with liquid nitrogen and dried. The product was mixed with urea at a mass ratio of 1:10, heated to 700 °C at a rate of 5 °C/min in a tubular furnace under a N₂ atmosphere, and held at that temperature for 2 h. Finally, the powder was stirred in 3 M HCl for 2 h, then thoroughly cleaned with deionized water and dried in a vacuum drying oven at 65 °C.

**Pt AMS and Fe AMS synthesis**

The synthesis process for Pt atomically monometallic site (Pt AMS) catalyst and Fe atomically monometallic site (Fe AMS) catalyst is similar to that of the preparation of the Pt–N₂–Fe ABA electrocatalyst. The notable difference is that the separate aqueous solution of H₂PtCl₆ or FeCl₃ is mixed directly with the prefabricated ammonia-rich carrier (CNF-NH₂), and the metal sources are replaced with 11.5 mg of H₂PtCl₆ or 5.94 mg FeCl₃ for the synthesis of Pt AMS and Fe AMS, respectively.

**Characterization**

JSM-6700F and JEM-2100F electron microscopes were employed to conduct SEM and TEM measurements at 5 kV and 200 kV, respectively. HAADF-TEM was conducted on a JEM-ARM200F instrument. XRD and XPS were performed on a Philips X'Pert Pro Super X'ray diffractometer with Cu Kα radiation and an ESCALAB MKII diffractometer with Mg Kα radiation, respectively.

**Electrochemical measurements**

All electrochemical measurements for ORR performance were carried out on a CH Instruments CH760E with a standard three-electrode cell. O₂-saturated 0.1 M KOH was used as the electrolyte, and graphite rods and Ag/AgCl were used as the counter electrode and reference electrode, respectively. The working electrode was an RDE with a disk diameter of 3 mm or an RRDE with a disk diameter of 5 mm. For the preparation of the catalyst ink for the RDE test, 2.5 mg catalyst powder was dispersed in a mixed solution of 750 μl DI water, 250 μl ethanol and 20 μl NaNafion™ (5 wt%) and ultrasonicated for 30 min. The catalyst ink was pipetted onto the GC surface with a catalyst loading of -0.1 mg cm⁻².

The mass activity is calculated by:

\[ MA = \frac{J_f}{M_{metal}} \]  

(1)

(the kinetic current densities \(J_f\), mA cm⁻²; the metal loading density \(M_{metal}\), μg metal cm⁻²), the metal loading on electrode of 2.9 μg cm⁻² in Pt = N₂–Fe ABA, 20 μg cm⁻² in commercial Pt/C, 5.9 μg cm⁻² in Pt AMS and 1.96 μg cm⁻² in Fe AMS.

The turnover frequency (TOF) is calculated by the following formula:

\[ \text{TOF} = \frac{I_f}{W_{metal} \times M_A \times \Delta \text{cat loading}} \]  

(2)

Where \(W_{metal}\) and \(M_{metal}\) are the mass fraction (wt%) and molar mass (g mol⁻¹). \(M_{loading}\) is the loading mass of the catalyst on RDE (g cm⁻²). \(N_{Avogadro}\) is Avogadro’s number (6.02 × 10²³ mol⁻¹). The Pt = N₂–Fe pair was regarded as a unit of synergistic active site, and the dual-sites and single Pt/Fe active sites were taken into account in a statistical proportion (about 3:1). For the tests of zinc-air batteries, custom-fabricated electrochemical cells with a two-electrode system were used. Pt = N₂–Fe ABA or Pt/C catalyst loaded onto Teflon-coated carbon fibre paper (1 cm²) was used as the air cathode (catalyst loading was 1 mg cm⁻²). Zn foil was used as the anode after polishing (1 cm²). All the tests were carried out in O₂-saturated 6.0 M KOH at room temperature with CHI 760E instruments.

**In situ SR-FTIR measurements**

All the in situ SR-FTIR data were collected at BLOIB of the National Synchrotron Radiation Laboratory (NSRL, China). An FTIR spectrometer, a KBr beam splitter, an ACT detector and an external microscope (Bruker Hyperion 3000) were used. The instrument adopts the reflection mode of vertical incident infrared light, the measurement width of the infrared spectrum is 600–4000 cm⁻¹, and the spectral resolution is 0.25 cm⁻¹. The custom-fabricated FTIR electrochemical cell was a three-electrode system, in which Ag/AgCl and platinum wire were used as the reference electrode and counter electrode, respectively. A 1 cm² carbon cloth loaded with catalyst was used as the working electrode and pressed onto the ZnSe crystal window with a micron gap. The electrolyte was 0.1 M KOH saturated with O₂ and circulated through a peristaltic pump.

**In situ XAFS measurements**

All XAFS data were collected at BLOIB station of Shanghai Synchrotron Radiation Facility (SSRF, China). The storage ring of SSRF operates at 3.5 GeV with a maximum current of 210 mA. The electrochemical in situ XAFS measurements were carried out in a custom-fabricated three-electrode system with a 1 × 1 cm² carbon cloth loaded with Pt = N₂–Fe ABA catalyst as the working electrode and 0.1 M continuously O₂-saturated KOH solution as the electrolyte. For the in situ XAFS spectral data acquisition of Pt L₃-edge (11564 eV) and Fe K-edge (7112 eV), we calibrated the positions of absorption edges (E₀) by using Pt foil and Fe foil standard samples, respectively, and all spectra were collected in the same beam time by fluorescence mode to ensure comparability.

**Strategies of XAFS curve Fittings**

The ARTEMIS module implemented in IFEFFIT was used to perform the EXAFS quantitative curve fittings. The fittings for the Pt L₃-edge of Pt = N₂–Fe ABA sample were done for k⁻²-weighted \(\chi(k)\) functions in a k-range of 2.5–11.5 Å⁻¹ and Fourier-transformed to real (R) space by Hanining windows at dk = 1.0 Å⁻¹ within an R-range of 1.0–3.1 Å. The number of independent points is \(N_{p,\text{fit}} = 2\pi k \times \Delta R / \pi = (11.5 - 2.5) \times (3.1 - 1.0) / \pi = 12\). For the ex situ Pt = N₂–Fe ABA sample, the coordination peaks at around 1.35 Å assigned to the Pt–N coordination with N atoms that derived from the N-rich–NH₂ functionalized support. And based on the analysis of the dominant heteronuclear dimer atomic spacing in Cs-corrected STEM-HAADF images and the location of metal coordination peak, the peak at 2.65 Å in Fourier-transformed curves should be assigned to Pt–N coordination. Therefore, two separate Pt–N and Pt–N–Fe scattering paths were included for fitting. For the in situ Pt = N₂–Fe ABA sample under 1.05, 0.95 and 0.90 V, the first coordination peaks show an obviously increased strength and location shift compared to the sample in KOH solution without potential applied condition. Considering that the prominent peak at ~1.6 Å can be assigned to the M–N/O coordination and the possible presence of oxygen-associated species, an additional Pt–O path was included during the fitting. The Debye–Waller factors (\(\sigma\)) and bond length (\(\delta\)) were treated as adjustable parameters for all conditions in all paths, the energy shift (\(\Delta E_{\text{O}}\)) for different paths under the same condition was set equal to reduce the number of adjustable fitting parameters, and the coordination numbers (\(N\)) of Pt–N under in situ conditions were fixed with reference to the ex situ sample. For the in situ samples under 1.05, 0.95 and 0.90 V, the number of adjustable parameters was \(N_{\text{para}} = 9 < N_{\text{fit}}\).

Similarly, the fittings for the Fe K-edge of Pt = N₂–Fe ABA sample were done in the R-space within the R-range of 1.0–3.1 Å for k⁻¹-weighted \(\chi(k)\) functions Fourier-transformed with Hanining windows of dk = 1.0 Å⁻¹ to separate the EXAFS contributions from different coordination shells. A k-range of 2.8–11.3 Å⁻¹ was carried out for curve fitting.
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Author contributions

Q.L. and H.S. conceived the project. W.Z. and J.J. carried out the experiments. Q.L., H.S., W.Z., W.C., Y.L., M.L., F.Y., W.W. and S.W. analyzed the experimental data. The manuscript was written by W.Z., H.S. and Q.L. with contributions from all authors.

Competing interests

The authors declare no competing interests.

Additional information

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