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1 Introduction

1.1 Construction of multiple active sites – optimization of the reaction pathway

Single-atom catalysts (SACs) have exhibited unique characteristics (e.g., unusual configuration and high atomic utilization) in heterogeneous catalysis.¹ However, it is extremely difficult to synchronously achieve the high-efficiency catalysis of two-step or multi-step reactions for SACs. Dinuclear-site catalysts with two active centers can independently interact with different intermediates and catalyze successive reaction steps, significantly improving the activities and selectivity of multiple-step catalytic reactions. Hu *et al.* (2021) fabricated a Fe-based catalyst including duet Fe₃C and FeN NPs for the hetero-e-Fenton reaction. The Fe₃C and FeN NPs can mediate the electroreduction of O₂ to H₂O₂ and the H₂O₂ activation towards 'OH, respectively (Fig. 1a). However, nanoparticle (NP) or atomic cluster (AC) catalysts usually suffer from inferior catalytic activity, due to the low atomic exposure and thus ineffective utilization of active sites.² To solve this issue, a dual-atomic-site catalyst (CoFe DAC) was fabricated, which can synergistically catalyze 'OH electro-generation. Specifically, Co SACs can promote O₂ reduction to H₂O₂ intermediates, and Fe

SACs were responsible for H₂O₂ activation to 'OH³ (Fig. 2a). In addition, there are reactions focusing on simultaneous reactant enrichment and catalytic elimination near the active center, also highlighting the importance of constructing dual-function sites. For example, surface oxygen vacancies and Co sites can capture H₂O₂/peroxymonosulfate (PMS) (*i.e.*, reduction reaction) and contaminants, respectively, decreasing the migration distance and thus increasing the degradation performance.^{4,5} Besides, dual-function sites can be used to drive simultaneous PMS oxidation and reduction reactions, for example, PMS can be oxidized at Co sites (*e.g.*, generating ¹O₂) and reduced at Bi sites (*i.e.*, generating SO₄²⁻ and 'OH) in Co-doped Bi₂O₂CO₃.⁶ Therefore, the incorporation of the second atomic dispersed active center can be a promising method to improve the efficiency of Fenton or Fenton-like reactions.

1.2 Incorporation of other atomic sites – strengthened reaction dynamics over the intrinsic active center

In addition, the incorporation of the second active center can effectively regulate the reaction activity when the interaction between the original catalytic site and intermediates is too weak or too strong. For example, Mo *et al.* (2023) reported that the incorporation of Fe ACs around the Fe single atoms (SAs) can significantly regulate the electronic structure of Fe SAs (*i.e.*, regulated d-band center, Fig. 1b) and thus the interaction between Fe SAs and PMS (*i.e.*, decreased adsorption energy, Fig. 1c), enabling rapid PMS adsorption and SO₅²⁻ desorption,¹ however, suffering from the drawbacks of low catalytic activity of ACs. Fe₄N₆ failed to function as active sites due to the high

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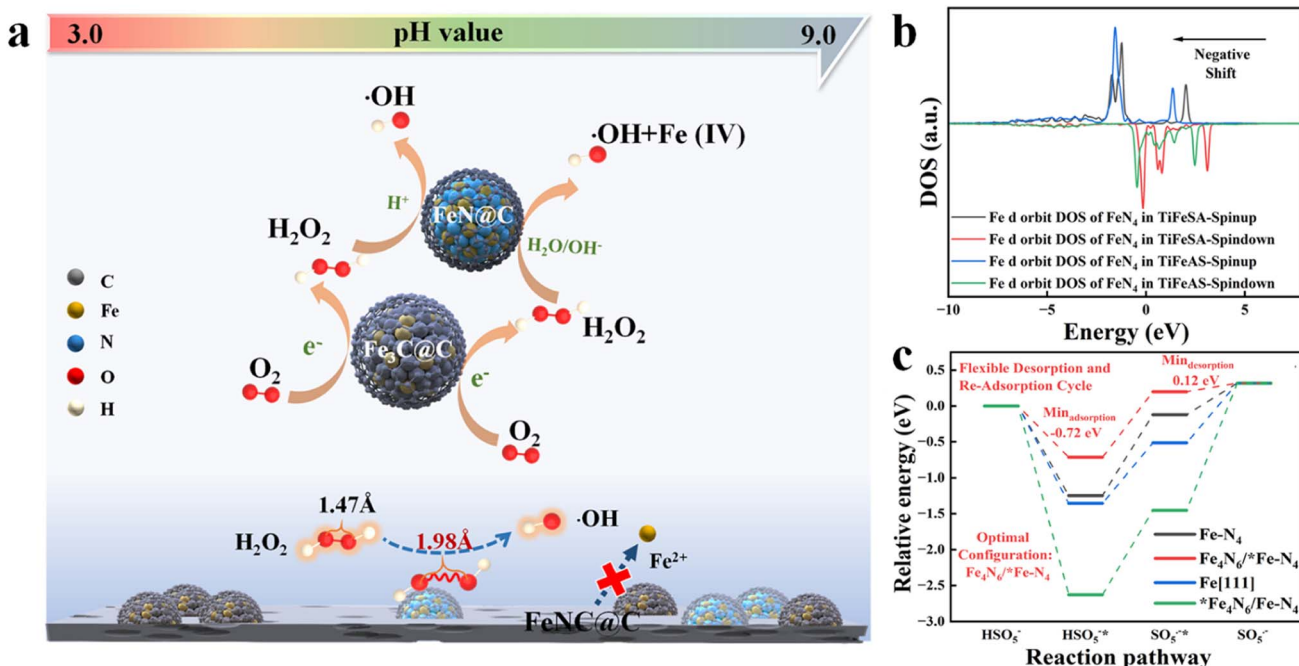


Fig. 1 Probable role of Fe_3C and FeN nanoparticles in the $\text{FeNC}@\text{C}$ catalyst for enhanced heterogeneous electro-Fenton performance, reprinted (adapted) with permission from ref. 2 copyright 2021 ACS. (b) The Fe d orbit DOS of FeN_4 in TiFeSA and TiFeAS. (c) Reaction energy diagrams for PMS oxidation on Fe-N_4 , $\text{Fe}_4\text{N}_6^*/\text{FeN}_4$, $\text{Fe}[111]$, and ${}^*\text{Fe}_4\text{N}_6/\text{Fe-N}_4$, reprinted (adapted) with permission from ref. 1. Copyright 2023 PNAS.

binding energy (Fig. 1c). Under these situations, the preparation cost may be reduced when substituting the ACs or NPs with SAs. For example, the coupling of Fe and Co sites (*i.e.*, FeCoN_6) optimized the adsorption strength between PMS and Fe sites (*i.e.*, 2.30 eV), compared to the sole Fe (*i.e.*, 2.14 eV) or Co SACs (*i.e.*, 2.43 eV), enabling a moderate interaction intensity and thus efficient reactant activation and product desorption⁷ (Fig. 2f). Therefore, the incorporation of heteroatom sites may effectively regulate the electronic configuration of active centers. To sum up, DACs have rich reaction centers, and would show better catalytic activity and customized selectivity toward Fenton or Fenton-like reactions than SACs. However, the applications of DACs in Fenton or Fenton-like reactions are less focused. Therefore, this perspective aims at summarizing the most recent advances in DAC-mediated Fenton or Fenton-like reactions with the underlying mechanisms being revealed. Most importantly, promising research directions are proposed, expecting further development in this field.

2 Most recent advances of DACs in Fenton or Fenton-like reactions

Several researchers have investigated the applications and mechanisms of DACs in Fenton or Fenton-like reactions, but their exploration is still in the initial stage.

2.1 Heterogeneous electro-Fenton reactions

Qin *et al.* (2023) investigated the application of Co-Fe DACs in heterogeneous electro-Fenton reactions.³ Co-Fe DACs were easily

fabricated through high-temperature annealing of Co, Fe-doped ZIF-8 precursors, followed by HNO_3 leaching at 60 °C. The DFT calculation showed a lowest energy barrier for ${}^*\text{H}_2\text{O}_2$ formation on $\text{Co-N}_4(\text{CoFe})$ (0.31 eV) and highest adsorption energy of H_2O_2 on $\text{Fe-N}_4(\text{CoFe})$ with a binding energy of −0.48 eV. Therefore, the synergistic effect can be established as follows: the atomically dispersed Co sites can enhance the O_2 reduction to H_2O_2 intermediates. Subsequently, the single-atom Fe sites can catalyze the transformation of H_2O_2 to $\cdot\text{OH}$ (Fig. 2a). This synergism enabled a higher $\cdot\text{OH}$ production rate of $2.4 \text{ mM L}^{-1} \text{ min}^{-1} \text{ g}_{\text{cat}}^{-1}$ compared to Fe-NC ($1.0 \text{ mM L}^{-1} \text{ min}^{-1} \text{ g}_{\text{cat}}^{-1}$) and Co-NC ($0.8 \text{ mM L}^{-1} \text{ min}^{-1} \text{ g}_{\text{cat}}^{-1}$). Simultaneously, the energy consumption was only $19.0 \text{ kWh kg}^{-1} \text{ COD}^{-1}$ compared to other electrochemical technologies ($29.7\text{--}68.0 \text{ kWh kg}^{-1} \text{ COD}^{-1}$).

2.2 PMS/peroxydisulfate (PDS) activation system

2.2.1 Homo-diatomc catalysts. $\text{Fe}_1\text{-N-C}$, $\text{Fe}_2\text{-N-C}$, and $\text{Fe}_3\text{-N-C}$ were synthesized by Li *et al.* (2022).⁸ Specifically, $\text{Fe}_2\text{-N-C}$ was synthesized using $\text{Fe}_2(\text{CO})_9@\text{ZIF-8}$ as the precursor, followed by annealing at 800 °C for 2 h under N_2 and 0.5 M HCl etching. $\text{Fe}_2(\text{CO})_9$ was replaced with $\text{Fe}(\text{C}_5\text{H}_7\text{O}_2)_3$ and $\text{Fe}_3(\text{CO})_{12}$ when fabricating $\text{Fe}_1\text{-N-C}$ and $\text{Fe}_3\text{-N-C}$. Subsequently, their degradation performance with monoatomic Fe ($\text{Fe}_1\text{-N-C}$), diatomic Fe ($\text{Fe}_2\text{-N-C}$), and triatomic Fe active centers ($\text{Fe}_3\text{-N-C}$) was explored, showing the highest efficiency for $\text{Fe}_2\text{-N-C}$, however, without in-depth mechanistic interpretation. The ${}^1\text{O}_2$ and $\text{Fe}(\text{IV})$ dominated non-radical pathway was verified to play a leading role in the catalytic degradation. Similar catalysts were investigated by Li *et al.* (2022) as well.⁹ Fe_1CN can be obtained

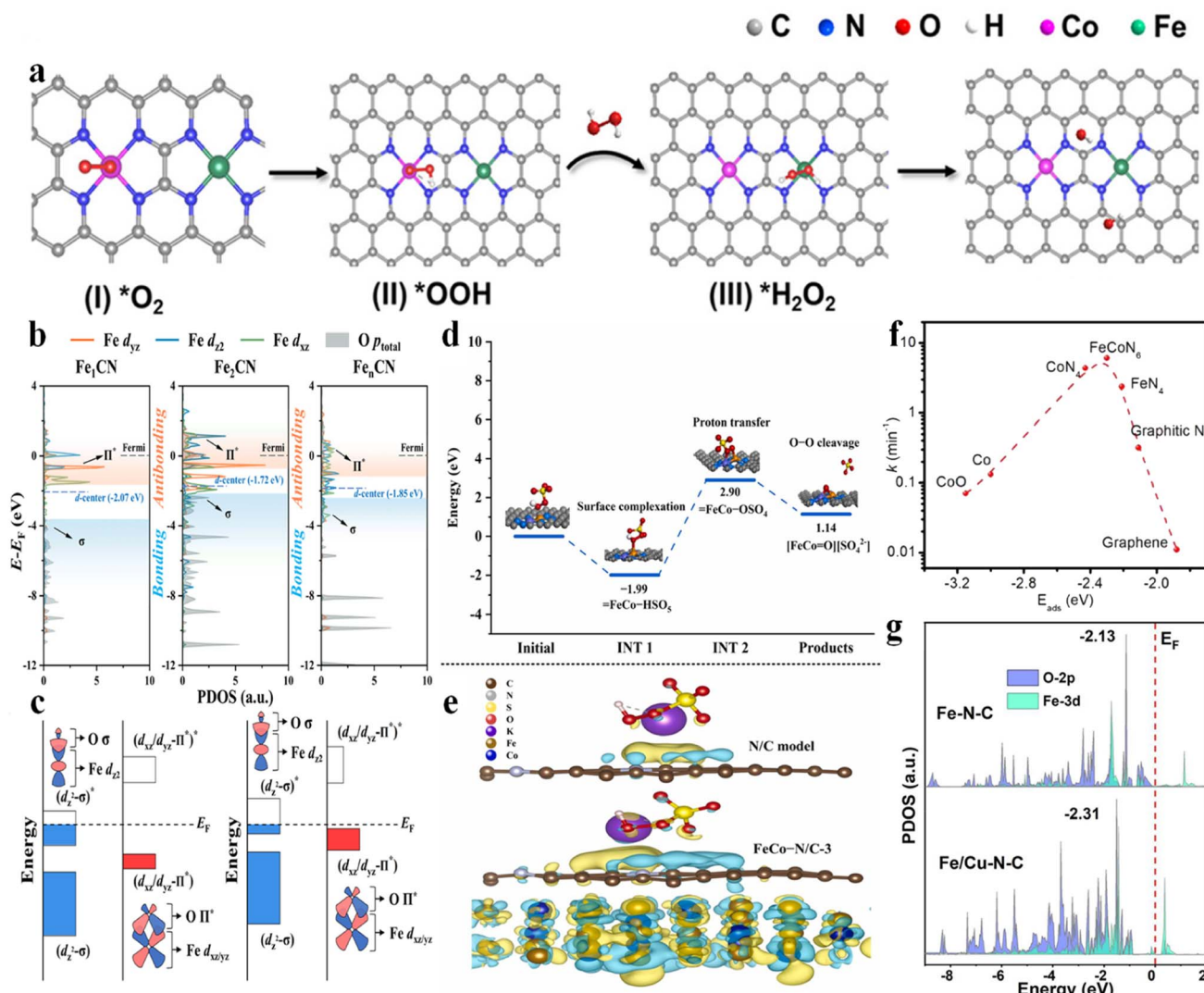


Fig. 2 (a) (I–IV) Proposed mechanism of ·OH electro-generation from the ORR on Co–N₄/Fe–N₄(CoFe), with permission from ref. 3 copyright 2023 ACS. (b) Projected density of states (PDOS) for Fe-3d_{xy}/d_{z²}/d_{xz} orbitals and O-p orbitals of adsorbed PMS, with permission from ref. 9 copyright 2022 ACS. (c) Schematic illustration of the orbital interaction between Fe-3d (d_{z²} and d_{xz/yz}) and O of adsorbed PMS (II* and σ) of Fe sites in Fe₁CN and Fe₂CN, with permission from ref. 9 copyright 2022 ACS. (d) Potential energy profiles of the pathway for the generation of high-valent metal-oxo species, with permission from ref. 10 copyright 2022 Elsevier. (e) The calculated charge density difference diagrams of PMS adsorbed on the N/C model and FeCo-N/C-3. Isosurface contour is 0.0003e per bohr³. The light yellow and light blue correspond to the electron accumulation and electron depletion, respectively, with permission from ref. 10 copyright 2022 Elsevier. (f) Volcano plot of the reaction rate as a function of E_{ads} of PMS at various active sites, with permission from ref. 7 copyright 2021 Elsevier. (g) PDOS values calculated for the O-2p and Fe-3d orbitals in Fe-N-C and Fe/Cu-N-C, with permission from ref. 13 copyright 2022 Elsevier.

through a two-step synthesis strategy (*i.e.*, Fe³⁺ (FeNO₃) adsorption on g-C₃N₄ and heat treatment at 550 °C (NH₃ atmosphere)). Pyrolysis of supramolecular precursors was used to fabricate Fe₂CN (hydrothermal reaction of urea and Fe³⁺ (FeNO₃) in an acidic environment, followed by heat treatment at 550 °C (NH₃ atmosphere)) (Fig. S1†). For Fe_nCN, melamine and FeSO₄·7H₂O were used as precursors. Subsequently, the mixture was stirred at 80 °C for 2 h, aged, filtered, and dried, followed by heat treatment at 550 °C (NH₃ atmosphere). Fe loadings were 3.16, 5.26, and 18.56 wt% for Fe₁CN, Fe₂CN, and Fe_nCN, respectively (the geometric structure is exhibited in Fig. S2†). The results indicated that the homo-diatom Fe sites

modulated the d-band center (*i.e.*, upshift of the d-band center compared to Fe₁CN and Fe_nCN, Fig. 2b) and increased the binding with oxygen-related intermediates (highest adsorption energy between Fe sites and PMS, Fig. S2†), thereby improving the PMS activation kinetics (3.58 times *vs.* single-atom Fe, accompanied by the transformation of the non-radical pathway to the radical pathway). Specifically, the Fe–Fe coordination (*i.e.*, the coupling between Fe-3d orbitals) can decrease the contribution of the antibonding state in the Fe–O bond (Fig. 2c), thereby facilitating the O–O bond cleavage of the Fe₂–HOO–SO₃ complex with a reduced thermodynamic energy barrier of only –0.29 eV.⁹

2.2.2 Hetero-diatomc catalysts. The $\text{FeN}_3\text{-CoN}_3$ moiety and FeCo alloy were constructed on the obtained FeCo-N/C, showing independent functions of dual active sites. The DFT calculation showed that the $\text{Fe}(\text{N}_3)\text{-Co}(\text{N}_3)$ would interact with PMS to generate FeCo=O through $=\text{FeCo-HSO}_5 \rightarrow =\text{FeCo-OSO}_4$ (proton transfer) $\rightarrow \text{FeCo=O}$ (O-O heterolytic cleavage) (Fig. 2d), with an energy barrier of 2.05 eV, which was significantly lower than those of high-valent Fe-O (3.28 eV) and high-valent Co-O (4.54 eV) species. Simultaneously, the FeCo alloy can provide electrons to the carbon lattice (Fig. 2e), reducing the E_{ads} of PMS on the N/C and facilitating self-decomposition of PMS to generate $^1\text{O}_2$. Therefore, the dual active sites enhanced the Fenton-like activity.¹⁰ A similar catalyst was reported by Yang *et al.* (2020), showing synergistic effects between the optimal FeCoN_6 configuration and pyrrolic N as the adsorption site. A volcano-shaped relationship between the E_{ads} of PMS and the rate constant was established, showing optimum adsorption energy at 2.31 eV, which was close to the E_{ads} of PMS on the FeCoN_6 site (Fig. 2f), highlighting the pivotal role of the $\text{Fe}(\text{N}_3)\text{-Co}(\text{N}_3)$ coordination.⁷ Zhu *et al.* (2023) fabricated Fe-Co DACs using $\text{NH}_2\text{-UIO-66}$ as a porous carbon support, showing favorable synergism as well. Specifically, the higher adsorption energy of PMS at FeCoN_6 (-0.958 eV ($-\text{SO}_3$ moiety near Fe atom) or -0.918 ($-\text{SO}_3$ moiety near Co atom) eV) was verified compared to FeN_4 (-0.568 eV) and CoN_4 (-0.471 eV), enabling high PMS concentration at the catalyst surface. In addition, the Fe-Co DACs possess higher DOS near the Fermi level compared to the Fe or Co SAC counterparts, inducing superior reactivity for PMS activation (*i.e.*, favorable electron exchange from the lone electron pair of O to the metal 3d vacant orbital and *vice versa*).¹¹ Besides, a spin-state reconstruction mechanism on Fe-Co DACs was used to elucidate the performance enhancement. The isolated Fe-Co DACs significantly increased the spin state of the Co center, while lowering the spin state of the Fe center, compared to the Co SAC and Fe SAC counterparts. These variations favorably resulted in the moderate adsorption energy of PMS on Fe-Co DACs (-1.71 eV on Co and -1.91 eV on Fe, moderate) compared to Fe SACs (-2.08 eV, too high) and Co SACs (-1.64 eV, too low).¹²

The Fe-Cu DACs were fabricated through hydrothermal synthesis ($\text{Fe}(\text{acac})_3\text{/Cu}(\text{acac})_3@ZIF-8$) and pyrolysis (Fe/Cu-N-C),¹³ showing an enhanced chloramphenicol removal rate from 0.073 (Fe-N-C) to 0.093 min^{-1} (Fe/Cu-N-C). The synergistic effects can be illustrated as follows: on the one hand, the electron would transfer from Cu to Fe, enabling the generation of low-valence Fe for peroxydisulfate (PDS) activation. On the other hand, the d-band center of Fe-3d downshifted from -2.13 eV (Fe-N-C) to -2.31 eV (Fe/Cu-N-C) when adding Cu-N-C, thus promoting the adsorption (-2.97 eV *vs.* -2.35 eV on Fe-N-C) and activation of PDS (Fig. 2g). An *et al.* (2023) reported the application of Fe-Cu DACs in PMS activation as well. Theoretical simulations illustrated that the electron transfer from Cu to Fe optimized the Fe 3d orbital distribution, showing the decreased d-band center position from -0.456 eV to -0.693 eV, promoting the adsorption and activation of PMS. In addition, the SMX was easily adsorbed on pyrrolic N nearby the Fe-Cu

diatomic sites, decreasing the migration distance of active species (*i.e.*, SO_4^{2-} , $\cdot\text{OH}$, O_2^{2-} , high-valent metal-oxo species, and metal-peroxy intermediates in this system).¹⁴

Studies dealing with the application of DACs in Fenton or Fenton-like reactions are still in its infancy (*i.e.*, only 9 published papers), requiring further validation and exploitation. Due to the characteristics of dual active centers of DACs, these catalysts can efficiently mediate multiple-step reactions (*e.g.*, H_2O_2 generation and activation),³ achieve dual-function catalysis (*e.g.*, simultaneous PMS oxidation and reduction),⁶ or strengthen intrinsic activity (*e.g.*, optimizing the realistic active center by incorporating co-catalytic sites). The innovative viewpoints for DAC optimization are listed in the following part.

3 Future outlook

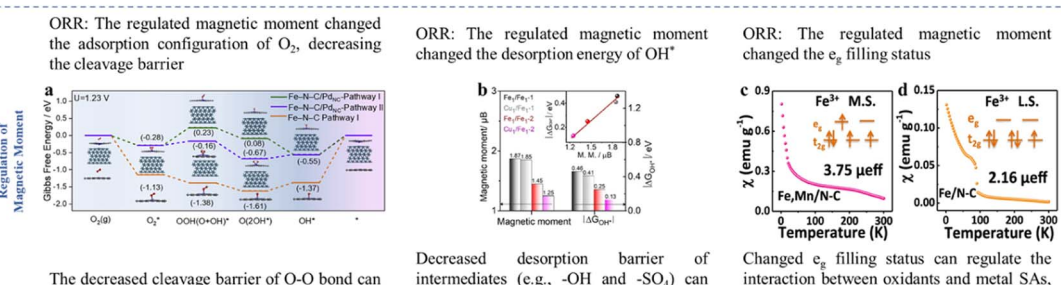
Several theories and methods aiming at improving SAC activity have been put forward, *e.g.*, regulation of the magnetic moment, interatomic distance effect, strain engineering, heteroatomic engineering, coordination regulation, ligand design, *etc.*, which may lay the theoretical foundation for fabricating innovative DACs applied in Fenton or Fenton-like reactions.¹⁵

3.1 Regulation of the magnetic moment

Magnetic moment variation can lead to a significant alteration in reaction activity. The introduction of Pd ACs around Fe SAs achieved the spin transition of low spin (LS) to intermediate spin (MS) for Fe SAs. To be specific, the electron would transfer from Fe SAs to Pd ACs due to the higher electronegativity of Pd compared to Fe^{2+} , leading to the redistribution of Fe 3d-orbital electrons and an increasing spin state of the Fe d_{z^2} orbital. In terms of the d_{z^2} orbital perpendicular to the Fe-N-C plane, the originally empty orbital of LS Fe(n) would change into a partially occupied one in the MS configuration, thereby effectively tuning the orbital overlapping with oxygen-containing intermediates. Interestingly, the O_2 molecule was adsorbed on Fe SAs in an end-on fashion (Pauling model) with the O-O bond elongated by 0.08 Å. Whereas, the side-on adsorption configuration (Griffiths model) was preferred on Fe SAs modified with Pd ACs with the O-O bond being elongated by 0.136 Å, which lowered the cleavage barrier of the O-O bond. In addition, the removal of OH^* was accelerated upon the incorporation of Pd ACs (Fig. 3a).¹⁶ The cleavage barrier of the O-O bond is also important for the Fenton or Fenton-like reactions as the H_2O_2 , PMS, or PDS activation required the peroxide bond dissociation as well. Specifically, the adsorption between the O-containing group and active center (*e.g.*, O-H bond dissociation, O-O bond formation, and O-O bond dissociation) is an important process for the OER (*e.g.*, O-O bond formation (ESI† eqn (3) and (8)) and O-H bond dissociation (ESI eqn (2), (4), (6), and (9)†)) and ORR (*e.g.*, O-O bond dissociation (ESI eqn (3) and (8)†)). Similarly, these processes are key reactions for PMS activation (*e.g.*, O-O bond dissociation in PMS reduction and O-H bond dissociation in PMS oxidation), PDS activation (*i.e.*, O-O bond dissociation), and H_2O_2 activation (*e.g.*, O-O bond dissociation). Therefore, energy-related reactions, especially the OER and

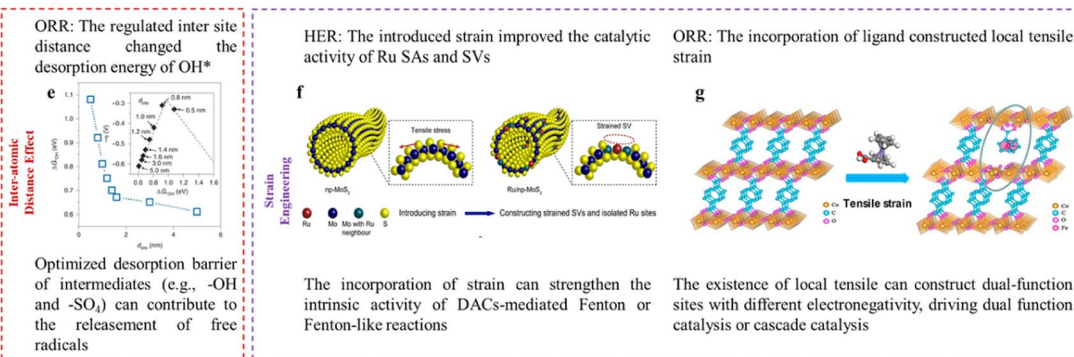


Examples



Potential application in Fenton or Fenton-like reactions

Examples



Potential application in Fenton or Fenton-like reactions

Examples

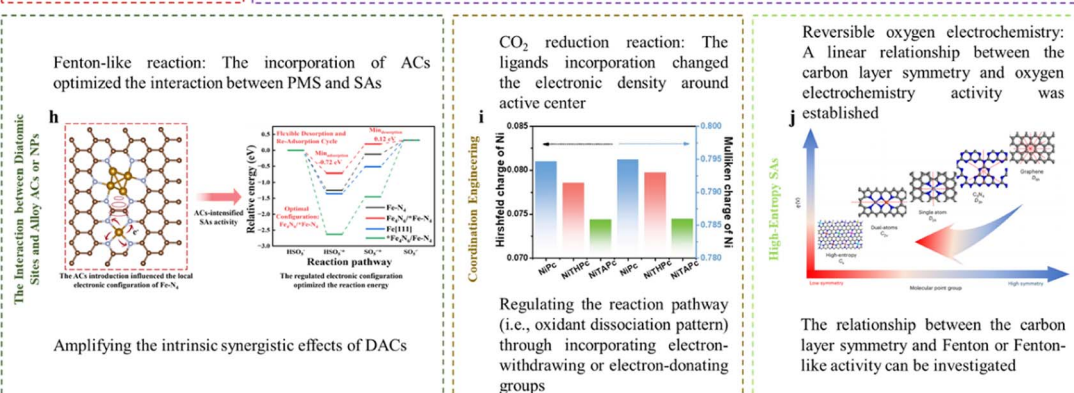


Fig. 3 (a) Gibbs free energy diagram of the ORR after the consideration of the solvent effect. Reprinted (adapted) with permission from ref. 16. Copyright 2022 Cell Press. (b) Comparison of the magnetic moment and ΔG_{OH^*} . Reprinted (adapted) with permission from ref. 17. Copyright 2022 Wiley. (c and d) Magnetic susceptibility of (c) Fe–Mn/NC, (d) Fe/NC (M. S. represents medium-spin, L. S. represents low-spin), with permission from ref. 18 copyright 2021 Springer Nature. (e) The d_{site} -dependent ΔG_{OH^*} obtained by DFT calculations. (Inset) Volcano plot of calculated overpotentials for the ORR against ΔG_{OH^*} . Reprinted (adapted) with permission from ref. 20. Copyright 2021 Springer Nature. (f) Illustration of the construction of Ru/np-MoS₂. Reprinted (adapted) with permission from ref. 21. Copyright 2021 Springer Nature. (g) Crystal structures of CoBDC and CoBDC FcCA models obtained from DFT simulations. The microreactor is composed of directly coordinated carboxyl oxygen atoms of FcCA (Co1) and unsaturated coordinative Co2, together with the FcCA linker labeled by the circle. Reprinted (adapted) with permission from ref. 23. Copyright 2021 ACS. (h) The regulatory role of Fe ACs in the Fe SA-mediated PMS oxidation reaction. Reprinted (adapted) with permission from ref. 1. Copyright 2023 PNAS. (i) Hirshfeld and Mulliken charges of the Ni atom in NiPc, NiTHPc, and NiTAPc. Reprinted (adapted) with permission from ref. 30. Copyright 2021 Wiley. (j) The relationship between symmetry and electrocatalytic performance, with permission from ref. 32 copyright 2023 Springer Nature. Notably, high-resolution figures can be viewed in ESI (Fig. 3a–j corresponding to Fig. S4–S11†).

ORR, can be used as important references in designing DACs. Thus, optimizing the magnetic moment can potentially regulate the Fenton or Fenton-like reaction activity through influencing the interaction between active sites and O-containing intermediates.

In addition, the magnetic moment is closely associated with the intermediate desorption. For example, the desorption energy of $-OH$ (ΔG_{OH^*}) was proved to be roughly linearly correlated with the Fe magnetic moment (Fig. 3b). The

incorporation of Cu atoms adjacent to Fe centers reduced the Fe magnetic moment, accompanied by an improved ORR activity (*i.e.*, decreased energy barrier for OH^* desorption).¹⁷ Specifically, the Fe SACs with Cu incorporation showed increased d_{z^2} electron filling near the Fermi energy level (marked by the red circle, appearing as a decreased magnetic moment) (Fig. S3a and b†), indicating more active electrons for reduction reaction. Thus, the desorption energy of the OH^* (*i.e.*, potential decomposition product of H_2O_2 or PMS) can also be optimized



through regulation of the magnetic moment, enabling rapid reaction dynamics.

The adsorption configuration of PMS, H_2O_2 , PDS, *etc.* on SAs was proved to be metal–O interaction. For example, Mi *et al.* (2021) proposed that the Co SAs would interact with PMS through the interaction of O on the $-\text{SO}_4$ side, therefore, highlighting the pivotal role of metal–O interaction in Fenton or Fenton-like reactions. Also, radical and nonradical pathways are closely associated with the binding energy between metal SAs and O sites in oxidants. Promisingly, this interaction can be regulated through the magnetic moment as well. Furthermore, the adjacent atomically dispersed Mn–N moieties can activate the Fe^{III} sites through modulation of the spin-state and electronic configuration, showing that the Fe^{III} with one e_g electron ($t_{2g}^4 e_g^1$) can readily penetrate into the antibonding π -orbital of O.¹⁸ In detail, the zero-field cooling (ZFC) temperature-dependent magnetic susceptibility was measured. The calculated effective magnetic moment of Fe–Mn/NC and Fe/NC was $3.75 \mu_{\text{eff}}$ and $2.16 \mu_{\text{eff}}$, respectively (Fig. 3c and d). Therefore, the number of unpaired d electrons (n) of Fe^{III} ions was determined using the following equation:

$$\mu_{\text{eff}} = \sqrt{n(n+2)} \quad (1)$$

The n value of Fe/N–C was 1.3, suggesting the low-spin state of Fe^{III} ions without e_g filling, enabling a strong interaction between the $\text{Fe}^{\text{III}}/\text{O}_2$ interaction due to the fact that no electron occupied the σ^* antibonding orbital of FeN_4 . Whereas, an increased n value³ was determined for Fe–Mn/NC, indicating single e_g filling. These results indicated that the unusual low-spin state of neighboring Mn^{III} moieties permitted Fe^{III} in FeN_4 to achieve the ideal e_g filling, endowing the catalysts with optimal binding energy with O_2 and thus favorable ORR activity. Hence, this strategy can be potentially used to regulate the metal–O interaction, thus regulating the radical and non-radical pathways.

Therefore, the incorporation of second metal sites (*i.e.*, constructing DACs) can optimize the magnetic moment (*i.e.*, spin state) of the intrinsic metal active site through the 3d electron interaction, especially the orbital filling of d_{z^2} . In further studies, the exact correlation between the magnetic moment of the active center and Fenton or Fenton-like reactions should be established, especially focusing on the adsorption configuration, energy barrier for the peroxide bond dissociation, and $-\text{OH}$ desorption. Particularly, the trade-off between the energy barrier for O–O dissociation and intermediate desorption (*e.g.*, $-\text{SO}_4$ or $-\text{OH}$) should be deeply investigated.

3.2 Inter-atomic distance effect

DACs with direct M–M interaction (*e.g.*, $\text{M}_1\text{--M}_2\text{--N}_6$)⁷ and indirect M–M interaction (*e.g.*, $\text{M}\text{--N}_4/\text{M}\text{--N}_4$)³ were both prepared for Fenton or Fenton-like reactions. Simultaneously, the advanced SAC synthesis field is devoted to improve the mass loading of metal sites (*i.e.*, exceeding 20% through a two-step annealing method).¹⁹ In addition to the metal loading changes, the inter-site distance would change accordingly, which is not deeply investigated to date. This scientific problem would be extremely interesting in the diatomic configuration.

As discussed in the “Regulation of the Magnetic Moment”, O–O bond cleavage and OH^* desorption are important in both the ORR and Fenton or Fenton-like reactions. Therefore, the study focusing on the ORR can be used for reference. For example, Jin *et al.* (2021) reported the effect of Fe inter-atomic distance in regulating the ORR activity.²⁰ The results showed that the inter-site distance would induce valence changes and distorted D_{4h} symmetry (changed pre-edge peaks and pre-edge absorption density, as confirmed by XANES analysis), spin state changes (D_1 sites, assigned to low-spin $\text{Fe}^{\text{II}}\text{--N}_4$ or high-spin $\text{Fe}^{\text{III}}\text{--N}_4$ accounted for 56 and 63% when the Fe inter-site distance was 1.9 and 0.5 nm, respectively, as confirmed by Mössbauer spectroscopy), spin moment (marked decrease in the on-site magnetic moment when the Fe inter-site distance was lower than *ca.* 1.6 nm, as verified by the density-functional theory (DFT)), and changed e_g electron filling (a higher degree of filling for the e_g electron would weaken the interaction intensity of OH^* , as verified by the DFT). The experimental results showed improved activity when decreasing the distance from 1.2 to 0.7 nm (optimum), below which the intrinsic activity would be slightly diminished, which can be ascribed to the optimized ΔG_{OH^*} (Fig. 3e).

Specifically, the XANES analysis identified that the decreased inter-atomic distance led to the decreased valence state and more distorted D_{4h} symmetry. It was demonstrated that the electronic activity would increase with decreased basal-plane symmetry. Therefore, the more distorted D_{4h} symmetry regulated by the inter-atomic distance may enhance the ORR performance, originating from the lowered aromaticity and more localized electronic structure. The inter-atomic distance-induced spin state changes verified by Mössbauer spectroscopy can potentially influence the d_{z^2} orbital filling, thus influencing the ORR efficiency. Similar underlying mechanisms can be established for the magnetic moment, which is important for the overlap between the metal d_{z^2} orbital and p_z orbital of oxygen-related adsorbates. The downshift of the energy of the d_{z^2} orbital when two Fe atoms get closer allows a higher orbital filling for the e_g electron, thereby resulting in a lower on-site spin moment. Consequently, the increased filling degree of the partially occupied d orbitals would weaken the bonding strength between the metal site and adsorbed hydroxyl intermediates (*e.g.*, OH^*). Therefore, the correlation between the inter-atomic distance and G_{OH^*} can be established through the changed basal symmetry, spin state, magnetic moment, *etc.* Similarly, the inter-atomic distance in the DAC configuration can possibly influence the Fenton or Fenton-like reaction activity, originating from the varied valence state, basal symmetry, d_{z^2} orbital filling (*i.e.*, magnetic moment and spin state), *etc.* To solve this issue in the Fenton or Fenton-like reaction field, the following aspects may be focused or considered:

(i) The inter-site distance can be easily regulated through changing the metal precursor concentration, *i.e.*, the site density-mediated active center distance, which should be examined by inductively coupled plasma mass spectrometry (ICP-MS). Particularly, acid etching should be performed to remove metal ACs or NPs. For example, Jin *et al.* (2021) placed the catalysts in 0.5 M H_2SO_4 solution for 8 h to remove the aggregated metal ACs and NPs.²⁰



(ii) To precisely assess the atomic dispersion of DACs, ion-beam thinning should be conducted to reduce the thickness of entities (where all metal atoms were able to be counted), followed by high-angle annular dark-field image (HAADF-STEM) analysis and X-ray absorption spectroscopy (XAS).

(iii) Most importantly, the inter-site distance can be estimated by statistical distribution (*i.e.*, HAADF-STEM images taken in a thin area) and *in situ* surface interrogation scanning electrochemical microscopy (SI-SECM).

(iv) Subsequently, the correlational analysis between site density and reaction activity can be established, mainly focusing on the valence using XAS analysis, spin state using Mössbauer spectroscopy analysis, and spin moment and intermediate binding energy using DFT.

3.3 Strain engineering

The synergistic effects of DACs are widely verified, however, it is still desirable to amplify their synergism and thus promote the practical application. Promisingly, the incorporation of strain (*i.e.*, the construction of a high-curvature surface) into DACs can optimize the electronic configuration.²¹ To the best of our knowledge, limited studies have paid attention to this strategy, in order to strengthen the Fenton or Fenton-like reaction efficiency, thus requiring the investigation and validation of strain engineering in this field.

In this part, several examples dealing with the hydrogen evolution reaction (HER) and ORR were referenced to provide insights for catalyst-mediated Fenton or Fenton-like reactions. In Fenton or Fenton-like reactions, the strain construction can amplify the known synergistic effects of DACs. When introducing the curvature, the mass transport of H_2O_2 from Co SAs to Fe SAs and their respective catalytic efficiency (*i.e.*, Co SA-mediated H_2O_2 generation and Fe SA-mediated H_2O_2 activation) may be facilitated in Co-Fe DAC-catalyzed electro-Fenton reaction.³ For example, the incorporation of tensile stress on Ru SAs and S vacancies (SVs) strengthened their intrinsic activity. The reaction density around SVs and H_2O dissociation/ H_2 evolution on Ru SAs was promoted, respectively (Fig. 3f). Similar results were reported for the high-curvature surface-loaded FeN_4 (*i.e.*, introducing compressive stress), showing the down-shifted d-band center and thus optimized bonding strength towards oxygenated intermediates.²² This strategy can be useful in regulating the DAC-mediated Fenton or Fenton-like reactions, as the optimized d-band position is a critical factor in influencing the activation efficiency of oxidants (*e.g.*, H_2O_2 , PMS, or PDS).¹

Promising substrates that can be applied to exert pressure are listed as follows:

(i) Nanoporous substrates that can introduce curvature, for example, the nanoporous Au with the Au ligament, followed by the coverage or loading of DAC materials.

(ii) Helical carbon structure with abundant high-curvature surface, which can be obtained by carbonization of helical polypyrrole that was templated from self-assembled chiral surfactants, followed by metal loading.

(iii) MOFs with the incorporation of other ligands, for example, when substituting partial terephthalic acid (BDC) in Co-BDC with

ferrocene carboxylic acid (FcCA), the crystal lattice CoBDC FcCA with tensile strain was obtained, generating Co coordinated with the carboxyl oxygen atom of FcCA (Co1) and unsaturated coordinative Co2 (Fig. 3g). Under this situation, the intermediate spin state ($t_{2g}^6 e_g^1$) Co can be constructed, showing the optimal OH^* adsorption energy and OOH^* formation energy.²³

3.4 Interaction between diatomic sites and alloy ACs or NPs

Recently, several reports have highlighted the role of *in situ* formation of ACs or NPs, and proposed the viewpoints as follows:

(i) The role of ACs/NPs in SAC-mediated catalytic reaction was ignored.²⁴ Specifically, the transformation of Cu SAs to ACs/NPs was observed during electrochemical reduction of nitrate to ammonia. Interestingly, the aggregated Cu NPs would reversibly disintegrate into SAs and then be restored to the $\text{Cu}-\text{N}_4$ structure upon being exposed to an ambient atmosphere.²⁴

(ii) The contribution of ensembled multiple atom sites in chemical reactions should be highly regarded.²⁵ Specifically, the contribution of various surface Pd species, ranging from SAs to ACs and NPs, in dehydrogenation of dodecahydro-*N*-ethylcarbazole (DNEC, for H_2 transportation and utilization) was determined.²⁵ Interestingly, the results showed that the Pd SAs were inactive to this reaction. The fully exposed Pd ACs with an average Pd-Pd coordination number of 4.4 can actively activate the reactant and promote product desorption, due to the fact that this reaction required multiple metal sites to initiate the DNEC adsorption. Whereas, the strong interaction between the reactant and crystalline surface (*i.e.*, Pd(111)) enabled a sluggish desorption process.

Theoretically, the incorporation of ACs or NPs can amplify the synergistic effects of DACs, increasing the charge asymmetry, thus influencing the reaction pathways. For example, the AC-induced charge redistribution of Fe-N-C was verified, showing a down-shifted d-band center and decreased Bader charge, thus strengthening the interaction between Fe-N-C and PMS. In detail, the incorporation of ACs optimized the HSO_5^- oxidation and SO_5^{2-} desorption steps, accelerating the reaction progress¹ (Fig. 3h). Therefore, the reinforcement of the catalytic efficiency of DACs by incorporating ACs or NPs can be expected. Definite catalyst types are listed as follows, especially in dual-metal forms.

(i) Metal 1 and metal 2 ensembled DACs, accompanied by the M1/M2 alloy NPs or ACs. For example, the coupled $\text{FeN}_3-\text{CoN}_3$ moiety and FeCo alloy moiety loaded on N-doped C were reported, working separately to generate high-valence metal (*i.e.*, $\text{FeCo}-\text{O}$) and $^1\text{O}_2$.¹⁰

(ii) The M1/M2 alloy NP or AC supported M1 SAs or M2 SAs. For example, the CoPt alloy-strengthened Pt SAs. The Pt SA aided by the CoPt alloy showed high d-band occupation for promoting the HER reaction kinetics.²⁶

(iii) The M1 NP or AC supported M1 SAs coupled with M2 NP or AC supported M2 ACs. This strategy is extremely less explored, which presents wide prospects. For example, the constructed $\text{Fe}_4@\text{FeN}_4$ and $\text{Ni}_4@\text{NiN}_4$ showed decreased adsorption energy of OH^* for Fe SAs modified with Fe ACs and

increased adsorption energy of OH* for Ni SAs modified with Ni ACs, promoting the ORR and OER activities, respectively.²⁷

3.5 Coordination engineering

Several in-plane structure engineering strategies were widely studied and previously reported, such as the regulation of coordination atoms, coordination number, *etc.* Analogous to the above discussion, these strategies can also effectively regulate the electronic configuration, for example, the d-band center, charge status (*i.e.*, Bader charge, valence number, *etc.*), *etc.* Undoubtedly, the bonding energy between intermediates and active centers can be optimized, showing optimal adsorption energy and a decreased energy barrier. These types of fabrication methods can be summarized as follows, which can be used in DAC performance optimization.

(i) The doping of heteroatoms with different electronegativities. For example, the Fenton-like activity (*i.e.*, PMS activation activity) can be enhanced or reduced by integrating electron-deficient boron (B) or electron-rich phosphorus (P) into the carbon substrate, respectively. Specifically, the long-range interaction with B atoms decreased the electronic density of single-atom Cu sites and down-shifted the d-band center, leading to the optimized adsorption energy for PMS activation.²⁸

(ii) The optimization of the coordination number. For example, the coordination number of single-atom Ni can be easily regulated through changing the annealing temperature for the ZIF precursor, generating Zn–N₃ and Zn–N₄ at 900 and 800 °C, respectively.²⁹ Theoretically, the different coordination numbers would significantly regulate the DAC performance, through influencing the intermediate formation.

(iii) The incorporation of ligands. Two types of reaction pathways can be expected for the PMS activation reaction (*e.g.*, PMS oxidation and PMS reduction), depending on the electronic density of the active center. Typically, PMS would be oxidized at electron-deficient sites and reduced at electron-rich sites.⁶ For example, PMS was reduced at Bi sites and oxidized at Co sites in Co-doped in Bi₂O₂CO₃, generating SO₄²⁻/OH and O₂, respectively.⁶ Promisingly, the electron status of the active center can be easily regulated by incorporating an organic ligand. Polymer modification (poly-4-vinylpyridine (P4VP)) on Co sites in cobalt phthalocyanine (CoPc) through Co-pyridine coordination was observed, enabling the transformation of Co(II) to more nucleophilic Co(I). In addition, the incorporation of electron-donating groups (*e.g.*, hydroxyl or amino) can induce electronic localization at the Ni site, increasing its electronic density (Fig. 3i). In contrast, electron-withdrawing groups would lead to the electron delocalization of the Ni site.³⁰ Therefore, this strategy can undoubtedly be used to design the PMS reaction pathway.

3.6 High-entropy SAs

DACs should be highly regarded due to the intrinsic drawbacks of SACs, *i.e.*, single-function active centers. This strategy can be extended to multiple-atom systems, due to the fact that there typically existed limited active centers and other heteroatoms would participate in the catalytic system as cocatalysts. For example, triple-atom sites (Co–O–Bi) were fabricated through

Co doping in Bi₂O₂CO₃. The O sites would function as electron mediators, whereas, the Co and Bi would initiate reduction function (PMS → SO₄²⁻ and OH) and oxidation function (PMS → O₂), respectively.⁶ In addition, the incorporation of Co optimized the binding between Fe sites and PMS. In this situation, the Fe sites and Co sites would function as catalytic sites and co-catalytic sites, respectively.⁷ However, the excessive complex system (*i.e.*, multiple-atom system) may be avoided as the optimal interaction strength can be obtained especially when only introducing two metals.³¹ Otherwise, the mechanism interpretation and high cost would be the significant problems.

The recent study proposed an innovative and interesting concept (*i.e.*, high-entropy single atom). Researchers embedded five single atoms (*i.e.*, Fe, Mn, Co, Ni, and Cu) with two sources of N, effectively minimizing the local symmetry to destabilize the π -electron network of graphitic carbons, thus endowing the catalysts with ideal binding energies (*e.g.*, neither too strong nor too weak) (Fig. 3j).³² Specifically, graphitic carbons with sp² hybridization showed unique aromaticity with delocalized π electrons, derived from the high symmetry (D_{6h}) of single graphene layers. However, π -conjugated electron network-protected stable graphitic carbons are extremely stable and inert towards chemical reactions. Several strategies such as heteroatom doping, defects, edge, and morphology modification have been developed to break this symmetry, so as to increase reactivity. The electrocatalytic activity enhancement has been verified when decreasing the basal-plane symmetry from D_{6h} to D_{2h} and C_{2v}. Therefore, incorporating several metal atoms can ideally decrease the symmetry to a higher level. Hence, an interesting theory or principle can be established, we should focus on the local structure when fabricating DACs, however, may pay attention to the holistic characteristics when incorporating several metal heteroatoms (*i.e.*, carrier engineering). Intriguingly, the symmetry of carbon layers and Fenton or Fenton-like reactions may be focused, aiming at establishing a correlation.

3.7 Trade-off between activity and stability: the pivotal role of *in situ* characterization technology

It is widely acknowledged that SAs would be the active sites during numerous reactions.^{33–35} However, the transformation of Cu SAs to ACs/NPs was observed during electrochemical reduction of nitrate to ammonia. Interestingly, the aggregated Cu NPs would reversibly disintegrate into SAs and then be restored to the Cu–N₄ structure upon being exposed to an ambient atmosphere,²⁴ thus highlighting the pivotal role of *in situ* formation of ACs/NPs in catalytic reactions. In another study dealing with single-atom Pt on CeO₂ for CO oxidation, interesting results were reported as well. The dynamic evolution of two types of single-atom Pt on CeO₂, *i.e.*, adsorbed Pt₁ in Pt/CeO₂ and square planar Pt₁ in Pt_{AT}CeO₂, fabricated at 500 °C and 800 °C (atom-trapping method), respectively, was investigated. Adsorbed Pt₁ in Pt/CeO₂ was mobile with the *in situ* formation of Pt ACs during CO oxidation, however, showing high reactivity with near-zero reaction order in CO. In contrast, Pt_{AT}CeO₂ with strongly anchored Pt₁ sites led to relatively low reactivity with a positive reaction order in CO. It's worth noting that both Pt/CeO₂ and



$\text{Pt}_{\text{AT}}\text{CeO}_2$ in CO transformed Pt_1 to Pt NPs, but retaining their initial Pt_1 state after re-oxidative treatments.³⁶ Therefore, the changed atomic sites during the reaction can be considered as realistic active centers to some extent. Atomic catalysts with high stability may face inferior reactivity. These issues should be further focused with advanced *in situ* characterization technology, such as *in situ* XAS, *in situ* X-ray photoelectron spectroscopy (XPS), and *in situ* electron microscopy.

4 Conclusion

DACs are promising candidates in Fenton or Fenton-like reactions. On the one hand, the diatomic configuration is conducive to the achievement of cascade reaction, bifunctional reaction, *etc.* On the other hand, the incorporation of the second active center can strengthen the intrinsic active sites through influencing the electronic configuration. In consideration of the limited focus on DAC-mediated Fenton or Fenton-like reactions (*i.e.*, only 9 published papers), several innovative developmental directions were put forward. Specifically, the in-depth mechanisms in designing DACs included regulation of the magnetic moment (*i.e.*, spin state), inter-atomic distance, and strain and incorporation of NPs/ACs and heteroatoms/ligands with specific electronic characteristics, through influencing the valence electron state, basal plane symmetry, orbital filling (*e.g.*, d_{z^2} electron), *etc.*, eventually regulating the reaction pathways (*e.g.*, cascade reaction, bifunctional reaction, oxidation reaction, reduction reaction, *etc.*) due to the optimized interaction (*e.g.*, adsorption configuration) between metal sites and O-containing intermediates. It is suggested that DACs can effectively design and regulate the Fenton or Fenton-like reactions, especially for multi-step reactions, thus representing a new avenue in Fenton-based water treatment technologies.

Author contributions

All authors contributed to the writing – review & editing of the manuscript. Particularly, Qixing Zhou was responsible for the funding acquisition.

Conflicts of interest

There are no conflicts of interest to declare.

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