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An emerging direction for nanozyme design: from single-atom to dual-atomic-site catalysts

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Nanozymes, a new class of functional nanomaterials with enzyme-like characteristics, have recently made great achievements and have become potential substitutes for natural enzymes. In particular, single-atomic nanozymes (Sazymes) have received intense research focus on account of their versatile enzyme-like performances and well-defined spatial configurations of single-atomic sites. More recently, dual-atomic-site catalysts (DACs) containing two neighboring single-atomic sites have been explored as next-generation nanozymes, thanks to the flexibility in tuning active sites by various combinations of two single-atomic sites. This minireview outlines the research progress of DACs in their synthetic approaches and the latest characterization techniques highlighting a series of representative examples of DAC-based nanozymes. In the final remarks, we provide current challenges and perspectives for developing DAC-based nanozymes as a quide for researchers who would be interested in this exciting field.

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1. Introduction

Nanozymes, a class of nanomaterials with intrinsic enzymelike activities, are considered next-generation artificial enzymes. 1-3 Compared with natural enzymes, nanozymes offer the unique advantages of low cost, high stability, tunable catalytic activity, and easy recyclability, features that greatly promote the integration of traditional chemical catalysts and biocatalysts. 4,5 To date, various nanomaterials that show enzyme-like catalytic activities, including noble metals, transition metal oxides, and metal-organic frameworks (MOF), have been discovered.⁶⁻⁹ They have shown great application in biosensing detection, 10,11 biomedicine treatment,^{5,12} food safety,¹³ and environmental protection.¹⁴ However, plenty of challenges remain in employing these nanozymes as enzyme alternatives. Firstly, the heterogeneous nanostructures of nanozymes inevitably yield inhomogeneous composition (size, exposed crystal plane, valence state, and defects), which leads to unsatisfactory enzyme-like selectivity, unclear catalytic active sites, and ambiguous reaction mechanisms. 15 Secondly, the synthetic methods of nanozymes make it rather difficult to fine-tune their structure, and the efficiency of metal utilization is usually insufficient, which leads to unde-

With the rapid development of nanotechnology, 16 singleatom catalysts (SACs) have recently emerged and received great attention in the fields of electrocatalysis, 17 photocatalysis, 18 biocatalysis, 19,20 and thermo-catalysis. 21 SACs integrate the merits of homogeneous and heterogeneous catalysts with maximal atomic usage and unique site structures. 22,23 Multitudinous well-defined SACs have been successfully applied as bioinspired single-atom nanozymes (SAzymes).²⁴ For example, SAzymes containing Fe, 25,26 Co,27 Ni,28 Cu,29 Zn,30 Cr, Mn,31 Mo,32 Pt,33 Ru, Rh, Ir, Pd,34 and Au singleatomic sites anchored on a carbon base,35 N, S, P, B, and F doped carbon-based SAzymes,36 and SAzymes with metal oxide-based supports have been prepared and applied in the fields of biosensing, 10 biomedicine, 37 environmental treatment,38 and food safety analysis.13 The atomically precise configuration of SAzymes also provides ideal models to understand the structure-performance relationship and enzyme-like catalytic mechanisms.39 Thus, the enzyme-like activity and specificity of SAzymes can be precisely adjusted and regulated at the atomic level, which makes them a promising candidate for next-generation nanozymes.40 Despite the success of SAzymes so far, there are still some burning questions to be answered: (1) How can linear scaling relations between different intermediates in enzyme-like reactions be broken for the simple active site of SAzymes? (2) How can the enzyme-like activities of SAzymes be further improved when the loading

sired or limited enzyme-like activities. To overcome these limitations, new design approaches for constructing nanozymes with controllable and homogeneous configurations are crucial.

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amount of a single atom is limited due to metal aggregation? The answers to these questions can be found by exploring advanced strategies for configuring new SAzymes.

In nature, the existence of unique cofactors, together with active centers, plays a synergistic role in the functioning of natural enzymes. For instance, cytochrome c oxidase (CcO) is mainly composed of binuclear sites consisting of a single Cu_B site and a heme a₃ site, which are a cofactor and an active center, respectively. 41 Also, carbon monoxide dehydrogenase and nitrogenase possess active centers configured with dualmetal atomic sites. 42 Inspired by these natural paradigms, dual-atomic catalysts (DACs) have been proposed as an extended and improved version of SACs. In addition to the advantages of atomically defined structures similar to those of SACs, DACs can offer more accessible active sites with the potential to break the linear relationship limitation in enzymatic reactions. 43 Since Warner et al. first reported the characteristics of Fe DACs embedded in graphene vacancies in 2014, 44 DACs have attracted extensive attention, particularly in the catalysis field, and a variety of DACs with unique features, such as adjacent homo-metal sites, 45 adjacent hetero-metal sites, 46 and separated hetero-metal sites, are engaged in heterogeneous catalysis reactions, demonstrating controllable multifunctionalities.47 The development of SAzymes with dualatomic sites has become a promising avenue to enhance their enzyme-like performances and further understand the fundamentals of enzyme-like reaction mechanisms.

The research on nanozymes has rapidly grown in a relatively short timeline (Fig. 1), and it is an appropriate moment to summarize the intriguing new developments in nanozyme design. This mini-review aims to capture the emerging directions in the structural evolution of nanozymes and present a full-scale picture of the nanozymes constructed with dualatomic sites. We will start by presenting a comprehensive and systematic summary of the recent advances made in synthetic approaches and characterization methods of DACs. Then, we will discuss the current status of DAC-based nanozymes in

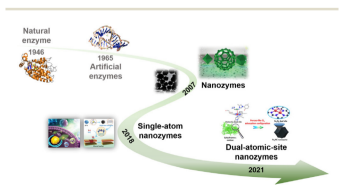


Fig. 1 Timeline of nanozyme development. Natural and artificial enzymes are shown for comparison. Reproduced with permission from ref. 1 Copyright 2007, Springer Nature; ref. 7 Copyright 2019, Royal Society of Chemistry; ref. 26 Copyright 2019, Royal Society of Chemistry; ref. 30 Copyright 2019, Wiley-VCH; ref. 88 Copyright 2019, Springer Nature.

detail highlighting representative examples. We will also show the current challenges and perspectives of DACs in nanozyme fields. It is anticipated that this review will spark more inspiring work on nanozymes and attract more attention from researchers of various backgrounds.

Preparation and characterization methods of DACs

The precise synthesis of DACs involves similar procedures and principles to those for SACs to a certain extent. Determining how to achieve the atomic-level dispersion and overcome the strong surface free energy without forming thermodynamically unstable phases is of great importance. The task of obtaining ideal DACs becomes even more challenging when building two metal atoms in a desired configuration is attempted. For the synthesis of both hetero- and homo-nuclear DACs, the effective manipulation of dual-atomic sites in adjacent or bridged configurations has been proved in some facile synthetic approaches, including atomic layer deposition (ALD), electrochemical deposition (ECD), chemical vapor deposition (CVD) wet chemical methods, and pyrolysis of targeted precursors (Fig. 2). 48,49 In many cases, however, some side products such as single atoms and even clusters are inevitably formed in the entire framework, which complicates the evaluation of the catalytic mechanism and the performance of DACs. Thus, it is of crucial importance to employ modern characterization techniques to confirm the precise atomic structure of the engineered DACs, too.

2.1. Strategies for constructing DACs

Isolated dual-atomic sites in DACs possess high surface energy, leading to challenges in achieving precise control over

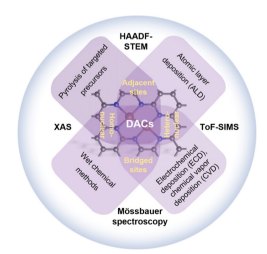


Fig. 2 Illustration of the preparation and characterization methods of DACs. HAADF-STEM: high-angle annular dark-field scanning transmission electron microscopy; XAS: X-ray absorption spectroscopy; ToF-SIMS: time-of-flight secondary ion mass spectrometry.

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the synthesis of metal dimers. In this regard, the adoption of atomic layer deposition (ALD), involving a sequence of self-limiting reactions, can serve as a powerful approach for fabricating DACs with a uniform deposition character. Lu's group reported a precise bottom-up synthesis for Pt2 dimers in a sequential manner using the ALD technique (Fig. 3a). 50 A secondary Pt atom was selectively deposited on the preliminary Pt₁ single-atom sites through the creation of proper nucleation sites in two ALD cycles. The self-limiting nature of ALD combined with orientated surface nucleation sites successfully achieved the synthesis of Pt2 dimers with a high surface area. The ALD method was also extended to heteronuclear DACs. As demonstrated in Fig. 3b, the isolated Pt atoms were first anchored on the surface of N-doped carbon nanotubes under the optimized ALD deposition conditions.⁵¹ By the ALD of Ru precursors on Pt single atoms under precise control, Pt-Ru dimers were successfully constructed. However, the large-scale production of DACs by the ALD technique is challenging due to the special requirement for metal precursor species and expensive equipment. In contrast, electrochemical deposition (ECD) and chemical vapor deposition (CVD) provide more flexible compatibility for expanding the selectable ranges of metal precursors and substrates for preparing DACs. 52-54

Another promising bottom-up strategy for fabricating DACs is the wet-chemistry method. Wet chemistry methods are some of the most widely used techniques for constructing DACs, offering a facile and straightforward approach that can be conducted in conventional chemistry laboratories, without the

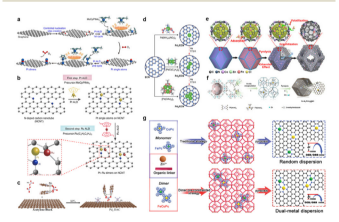


Fig. 3 Schematic diagrams illustrating the synthetic procedure of (a) dimeric $Pt_2/graphene$ (reproduced with permission from ref. 50 Copyright 2017, Springer Nature), (b) Pt-Ru dimers on nitrogen-doped carbon nanotubes (NCNT) (reproduced with permission from ref. 51 Copyright 2019, Springer Nature), (c) supported Pt_2 DAC (reproduced with permission from ref. 57 Copyright 2021, Wiley-VCH), (d) low-nuclearity Pt_3 Cotalysts (reproduced with permission from ref. 58 Copyright 2019, Wiley-VCH), (e) hollow carbon-derived nanostructures with porphyrin-like Pt_3 Fe-Co dual sites ((Pt_4)-Co)/N-C) (reproduced with permission from ref. 63 Copyright 2017, American Chemical Society), (f) dual-atom Pt_4 - Pt_4 -

necessity for specialized techniques or costly equipment. This method leverages dinuclear metal complexes as pre-determined precursors, providing substantial potential for the precise construction of the targeted dual-atomic configurations. Various viable approaches, including co-precipitation procedures, solvothermal/hydrothermal reactions, (electro)chemical reduction processes, are employed to introduce these dual-atom precursors into both organic and inorganic substrates. Notably, Wang and co-workers proposed a "precursor-preselected" wet-chemistry strategy for the synthesis of mesoporous carbon nitride-supported Fe₂ cluster catalysts (Fe₂/mpg-C₃N₄).⁵⁵ This method capitalizes on preselected metal precursors to ensure the formation of specific diatomic Fe clusters, rich in anchoring sites and adept stabilization of mpg-C₃N₄ supports. This strategy was subsequently extended for the synthesis of dual-atom Pt2 catalysts (Pt2/mpg-C₃N₄).⁵⁶ A diatomic Pt precursor, (ethylenediamine)iodoplatinum(II) dimer dinitrate, was anchored on the mpg-C₃N₄ substrates. More recently, this group further engaged an anion replacement deposition-precipitation (ARDP) method in the preparation of a Pd₂ dual-atomic catalyst (Pd₂ DACs, Fig. 3c).⁵⁷ The obtained Pd2 DAC exhibited a PdN2O2 coordination structure akin to its precursor configuration. Noteworthy progress has also been made in introducing single Pd atoms, Pd2 dimers, and Pd₃ trimers into the hetero-macrocycles of carbon nitride (C₃N₄) via microwave-assisted deposition, fine-tuning the metal precursor for precise control (Fig. 3d).⁵⁸ Furthermore, the wet-chemical approach has been expanded to include more accessible synthetic routes, enabling precise control through tunable kinetic and thermodynamic parameters. Liu's group employed a strategy of electrostatically driven self-assembly assisted by phenanthroline (Phen) ligands to fabricate La, Ni bimetallic coordination structures within conjugated boronate-ester-linked covalent organic framework (COFs) substrates.⁵⁹ This work serves as a typical example of the scalable synthesis of well-defined ligand-

In addition to traditional carbon-based substrates, metal oxides can also be used for stabilizing heteronuclear DACs. Huang and co-workers pioneered this avenue by first fabricating a bimetallic carbonyl cluster as organometallic precursors. The subsequent steps involved the anchoring of well-dispersed Ir single atoms (Ir₁) and Mo single atoms (Mo₁) onto TiO₂ supports (Ir₁Mo₁/TiO₂) through adsorption and pyrolysis treatments. Similarly, Su and co-workers synthesized highly active and stable single-atom Sn–Zn pairs within a CuO catalyst via a facile hydrothermal treatment followed by wet impregnation. Soon after, Wang $et\ al.$ employed a photochemical method to create Ir dinuclear heterogeneous catalysts (Ir DHCs). The resulting Ir DHCs catalyst features two active metal centers (Ir–O–Ir) anchored on α -Fe₂O₃ supports.

assisted bimetallic centers.

Typically, the wet chemistry approach was also employed to fabricate specific precursors containing two target metal atoms, which could be subsequently treated under optimized conditions to yield DACs of highly dispersed active sites. For instance, the pyrolysis of these targeted precursors is one of Minireview Nanoscale

the most common synthetic methods for producing DACs. Metal-organic frameworks (MOFs), in which different metal nodes can be coordinated with multifunctional organic linkers, are of particular interest as precursors for this strategy. Moreover, the well-defined porous structures of MOFs can also endow the impregnated precursors with an ideal confinement effect at the molecular level. In an early work reported by Wu et al., 63 the host-guest strategy was used to build Fe-Co dual sites embedded in nitrogen-doped carbon supports ((Fe,Co)/ N-C) (Fig. 3e). A Zn/Co bimetallic MOF (BMOF) was used to host FeCl₃ molecules as a guest in the confined space between Zn and Co metallic nodes. A pyrolytic treatment that decomposed and graphitized the FeCl3-adsorped BMOF successfully removed the vaporized Zn species and produced (Fe,Co)/N-C. Similarly, Hu et al. synthesized atomically dispersed binary Co-Ni sites that were anchored on N-doped hollow carbon supports (CoNi-SAs/NC) via a three-step synthetic strategy.⁶⁴ They first fabricated CoNi-MOFs covered with self-polymerized dopamine by chemical precipitation. The subsequent annealing under an NH3 atmosphere and acid-leaching treatment yielded CoNi-SAs/NC. More recently, Wang and co-workers reported an in situ co-encapsulation method to obtain a more uniform distribution of the Fe-N₄ moiety near the adjacent Pt- N_4 moiety (Fig. 3f). ⁶⁵ The sizes of two precursors, Pt(acac)₂ (d =~10.0 Å) and Fe(acac)₃ (d = ~9.7 Å), were comparable to those of the nanocavities in zeolite imidazole frameworks-8 (ZIF-8, 11.6 Å), which ensured the uniform distribution of isolated Fe and Pt atomic sites in Fe-N₄/Pt-N₄@NC.

Since the sources of metallic sites in DACs are the metal nodes in MOF hosts or guest precursors embedded in MOFs, it is also possible to produce single-atomic and polymetallic species during the pyrolytic process due to random dispersions. Aiming to minimize such random interactions of metal species, Bu et al. recently reported a "pre-constrained metal twins" strategy for fabricating adjacent Fe-N₄ and Co-N₄ DACs embedded in an N-doped graphitic carbon (FeCo-DACs/NC, Fig. 3g).⁶⁶ Different from the conventional method, they first fabricated an FeCo binuclear phthalocyanine dimer and in situ encapsulated it into a ZIF-8 framework. After carbonization without acid treatment, FeCo-DACs/NC containing uniformly dispersed dual-metal centers were obtained. Furthermore, Niu et al. developed a macrocyclic precursor-mediated method to fabricate a series of DACs ranging from homonuclear to heteronuclear bimetal centers (M₁M₂-DAC).⁶⁷ The time-of-flight secondary ion mass spectrometry (ToF-SIMS) provided more definitive evidence for the formation of diatomic fragments in FeCu-DACs, while no typical signals were found in the Fe/Cu-DACs prepared by conventional co-impregnated-pyrolysis. These pioneering studies on pyrolytic methods have opened up a feasible route for constructing DACs with operational flexibility and scale-up possibility.

2.2. Characterization methods for DACs

Despite the extensive reports on the successful synthesis of DACs, some lack direct evidence for the geometric and electronic configuration of DACs due to the limitations of the characterization techniques used. The two most common characterization techniques for DACs are AC HAADF-STEM and XAS, which provide precise information about morphology and local coordination, respectively. Other spectroscopic techniques such as Mössbauer spectroscopy, electron spin resonance (ESR) spectroscopy, Fourier transform infrared (FT-IR) spectroscopy, X-ray photoelectron spectroscopy (XPS), and ultraviolet photoelectron spectroscopy (UPS) can provide complementary identification of DACs. 68

The dimer structure of DACs was first reported by Jamie H. Warner and co-workers in 2014.⁴⁴ They directly captured the atomic structure of Fe atom pairs incorporated within a graphene lattice using an aberration-corrected transmission electron microscope (AC-TEM, Fig. 4a). Various stable structures of Fe dimers were further confirmed by the corresponding multi-slice image simulations using the atomic models based on density-functional theory (DFT) calculations (Fig. 4b and c). In general, AC HAADF-STEM presents a 2D projection along the direction of the incident beam, and thus the subtle features of dual-atomic pairs can be influenced by different projection angles. Wang's group built the spatial models of a homonuclear Fe-Fe catalyst (Fe2-N-C), which were mainly based on the 2D projection along the direction of the incident beam (Fig. 4d).69 The corresponding three-dimen-

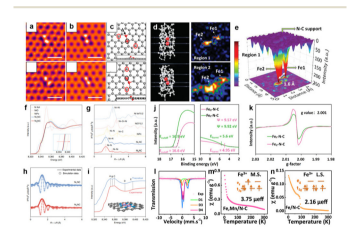


Fig. 4 (a) Smoothed aberration-corrected transmission electron microscope (AC-TEM) images, (b) multi-slice image simulations, and (c) the corresponding DFT optimized atomic models of Fe dimers in graphene vacancies. Reproduced with permission from ref. 44 Copyright 2014. American Chemical Society. (d) 2D projection along the direction of the incident beam of 3D structures and (e) 3D intensity mapping of the homonuclear Fe-Fe catalyst (Fe₂-N-C). Reproduced with permission from ref. 69 Copyright 2023, Wiley-VCH. (f) X-ray adsorption near-edge structure (XANES) spectra at the Ni K edge, (g) k^3 -weighted Fourier transformed extended X-ray absorption fine structure (FT-EXAFS) spectra of Ni single-atom sites (Ni₁NC) and dual-atom sites (Ni₂NC). The corresponding fitted (h) FT-EXAFS and (i) XANES data of Ni₁NC and Ni₂NC. Reproduced with permission from ref. 76 Copyright 2022, Springer Nature. (j) UPS and (k) ESR spectra of Fe₁-N-C and Fe₂-N-C. (I) Room-temperature ⁵⁷Fe Mössbauer spectra and magnetic susceptibility of atomically dispersed Fe,Mn/N-C catalysts. Magnetic susceptibility of (m) Fe,Mn/N-C and (n) Fe/N-C catalysts (M.S.: medium-spin; L. S.: low-spin). Reproduced with permission from ref. 85 Copyright 2021, Springer Nature.

sional (3D) intensity map confirmed the distance of Fe-Fe atom pairs (1.6 Å, Fig. 4e). To gain comprehensive information on the atomic arrangement of DACs, more decisive tools and analysis beyond the AC HAADF-STEM are desired. For instance, electron energy loss spectroscopy (EELS) can provide high-resolution elemental mapping images that can distinguish dual-atomic species. Han and co-workers employed an EELS diagram to demonstrate the co-existence of Ni and Cu atoms in a DAC, demonstrating it to be a powerful tool for differentiating between two atoms of similar atomic numbers.⁷⁰ More recently, scanning tunneling microscopy (STM) and atom probe tomography (APT)⁷¹⁻⁷³ have been shown as important complementary characterization methods for distinguishing dimer adatoms and providing 3D constructions of DACs.

XAS is another key characterization technique to determine the electronic and geometric structure of DACs.74,75 More specifically, X-ray absorption near-edge structure (XANES) spectroscopy can resolve electronic structures of DACs on the atomic scale. In the work by Zhang and co-workers, the absorption edge position of Ni K-edge XANES spectra was used to study the Ni electronic state in a Ni DAC (Ni₂NC),⁷⁶ and a lower Ni oxidation state in the Ni2NC samples than that in Ni₁NC was revealed (Fig. 4f). The information on the coordinate environment of Ni sites in Ni₂NC was further analyzed using the extended X-ray absorption fine structure (EXAFS) technique that revealed two distinctive peaks at ~1.44 and 2.09 Å for the Ni-N and Ni-Ni shell, respectively (Fig. 4g). The complete local coordination structure of Ni₂NC was further refined as N₃-Ni-Ni-N₃ (Ni₂N₆) from the fitted Fourier transform-EXAFS measurements at R space (Fig. 4h). The simulated XANES spectrum based on a DFT model could be used to confirm the experimental data of the Ni₂N₆ configuration (Fig. 4i). In another study, Guo et al. used XAS to characterize the formation of Pt-Fe atomic bonds and fast electron transfer from Fe to Pt.77 Besides these homonuclear and heteronuclear catalysts containing adjacent dual-atom sites, the bridged sites in the DACs can also be probed using XAS techniques. Zhuang et al. confirmed N-bridged Co-N-Ni bimetallic sites based on a series of soft and hard XAS analyses.⁷⁸ Hou and co-workers also identified an O-bridged In-Ni atomic pair in the InNi DACs using EXAFS fitting analysis.⁷⁹ It is worth noting that the XAS characterizations provide only average information regarding the entire structure of DACs. Therefore, other tools are needed to gain a more complete structural view of DACs.

Conventional XPS and UPS characterization studies offer information about the elemental composition, oxidation state, and electronic properties near the catalyst surface. The UPS spectra shown in Fig. 4j compare the surface work function (ψ) values of Fe2-N-C and Fe1-N-C, which have a decreased value in Fe₂-N-C due to its excellent electron-donating capability.⁶⁹ In situ XPS under a synchrotron light source can provide insights into the charge transfer states in DACs during operation.80 Meanwhile, in situ/operando IR spectroscopy and Raman spectroscopy can capture the key intermediates on the DACs. 46 Electron paramagnetic resonance (EPR) is another highly sensitive spectroscopic method, particularly for the electronic structure investigation of paramagnetic species.^{81,82} The enhanced EPR signal at g = 2.001 confirms a higher concentration of unpaired electrons in Fe2-N-C than in Fe1-N-C (Fig. 4k).

Mössbauer spectroscopy is a powerful tool for distinguishing variations in the coordination environments, local oxidation state, and spin states of Fe-, Co-, and Sn-containing catalysts. 83,84 Zhang's group employed Mössbauer spectroscopy to discriminate different Fe species in the atomically dispersed dual-metal Fe,Mn/N-C electrocatalyst (Fig. 4l).85 The quantitative analysis confirmed that the Fe species in Fe,Mn/ N-C was predominantly Fe^{III} with a medium-spin structure. For verification of the electron spin configuration, the temperature-dependent magnetic susceptibilities of Fe,Mn/N-C and Fe/N-C were compared (Fig. 4m and n). The Fe,Mn/N-C displayed a higher magnetic moment value of $3.75\mu_{\rm eff}$ than the Fe/N-C (2.16 $\mu_{\rm eff}$) because of the greater number of unpaired d electrons in the medium-spin state. Besides, inductively coupled plasma mass spectrometry (ICP-MS),86 ToF-SIMS,67 and nuclear magnetic resonance (NMR) spectroscopy are useful characterization techniques for elucidating the structure of DACs.87

Practical applications of DACs in nanozymes

The research of employing DACs as nanozymes is still in its infancy, but its emergence itself is a key stepping stone for the development of nanozyme research. Recently, impressive progress in developing DACs with extraordinary enzyme-like behavior has been made. In the following sections, we will mainly focus on the rational design of DACs with different types of enzyme-like catalytic performance, mechanism understanding, and various biological applications.

In 2022, Zhu et al. designed an Fe₂NC nanozyme with an Fe-Fe dimer similar to natural cytochrome c oxidase (Fig. 5a).88 This as-developed Fe₂NC catalyst exhibited much higher oxidase- (OXD, cytochrome c oxidase, NADH oxidase, and ascorbic acid oxidase) and peroxidase (POD, NADH peroxidase, and ascorbic acid peroxidase)-like activities compared with Fe₁NC counterparts. Liu's group further developed a multi-enzyme cascade antioxidant system by introducing Fe₂NC catalysts into Se-containing MOFs (Fe₂NC@Se, Fig. 5b). 89 The synergistic effect of Fe₂NC and Se significantly promoted the superoxide dismutase (SOD)-, catalase (CAT)-, and OXD-like performances of the Fe₂NC@Se nanozyme.

Considerable research effort has also been directed toward the exploration of heteronuclear DACs with enzyme-like performances. For instance, Zheng et al. successfully fabricated an atomically dispersed Fe, Pt dinuclear nanozyme, featuring Fe-N₃ and Pt-N₄ moieties (Fig. 5c). This nanozyme exhibited a synergistic interplay between Fe-N3 and Pt-N4 sites, effectively catalyzing Fenton-like reactions and demonstrating robust photothermal conversion properties suitable for tumor

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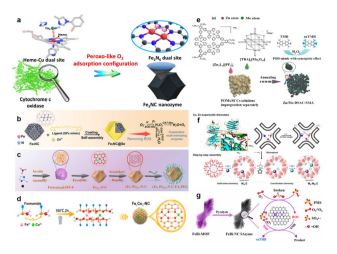


Fig. 5 (a) Schematic illustration of peroxo-like O₂ adsorption configuration of the Fe₂NC nanozyme. (b) The preparation process of the dual-Fe-atom nanozyme in a Se-containing MOF (Fe2NC@Se) with multienzyme cascade antioxidant application. Reproduced with permission from ref. 89 Copyright 2022, Wiley-VCH. (c) Atomically dispersed Fe, Pt dinuclear catalysts ($(Fe,Pt)_{SA}-N-C$) for tumor catalytic therapy. Reproduced with permission from ref. 90 Copyright 2022. American Chemical Society. (d) Schematic illustration of the fabrication process of Fe₁Co₁-NC metal-nitrogen-carbon dual-atom Reproduced with permission from ref. 91 Copyright 2023, American Chemical Society. (e) Schematic illustration of the synthesis of Zn/Mo dual single atom nanozymes supported on a poly(vinyl alcohol) (PVA)based aerogel. Reproduced with permission from ref. 92 Copyright 2022, Wiley-VCH. (f) Schematic diagrams of bio-inspired "step-by-step" assembly approach for the Zn_A-Cu_B samples as native Cu,Zn-containing superoxide dismutase (Cu,Zn-SOD) mimics. Reproduced with permission from ref. 94 Copyright 2022, Elsevier. (g) The synthesis process of Fe-Bi bimetallic MOF-derived carbon supported Fe-N $_4$ and Bi-N $_4$ dual-site FeBi-NC SAzyme. Reproduced with permission from ref. 95 Copyright 2022, Elsevier.

catalytic therapy. Song et al. also reported a general strategy for the development of a library of SAzymes (M1-NC) and DAzymes (M_1/M_2-NC) with similar structures (M = Fe, Co, Ni,Mn, Ru, and Cu) (Fig. 5d). 91 The Fe₁Co₁-NC dual-atom nanozyme with Fe₁-N₄/Co₁-N₄ coordination exhibited remarkable enhancement of POD-like catalytic properties and excellent photothermal conversion efficiency. Wei's group presented a facile approach for crafting Zn/Mo dual atomic-based SAzymes (Zn/Mo DSAC-SMA) via the assembly of supramolecular coordination complexes and polyoxometalates (Fig. 5e). 92 The strong affinity of Mo atoms toward hydroxyl species and their subsequent migration to adjacent Zn sites, facilitated by a low energy barrier of the adsorbed hydroxyl species, resulted in a synergistic effect that substantially enhanced the H2O2 dissociation and POD-like performance of Zn/Mo DSAC-SMA. A Zn-Y dual-atomic catalyst having a ZnY-N₆ configuration (ZnY-DACs/NC) was also developed by Chen's group and showed POD-like performances.⁹³ Inspired by natural enzyme configurations, Lo's group proposed a creative approach for building dual-atom Zn_A-Cu_B bimetallic catalysts (Zn_A-Cu_B-Z) through bio-inspired synthesis procedures (Fig. 5f).⁹⁴ The structural resemblance between the synthesized ZnA-CuB-Z catalysts and natural enzymes (Cu and Zn-containing SODs) makes them appealing SOD mimics. Huang et al. built an FeBi-NC SAzyme featuring Fe-N₄ and Bi-N₄ dual-sites within a cascade enzyme-nanozyme system for excellent peroxymonosulfate activation (Fig. 5g). Furthermore, Chen's research team designed an FeCoZn catalyst containing triple-atom sites coordinated with S and N on the carbon matrix, denoted as FeCoZn-TAC/SNC. Notably, the unique architecture of FeCoZn-TAC/SNC exhibited remarkably enhanced OXD-like catalytic performance compared with those of single/dualatom site catalysts. This distinctive design imparts great potential to FeCoZn-TAC/SNC nanozymes for the colorimetric sensing of ascorbic acid.96

Dual-atomic-site catalysts have demonstrated promising capabilities in emulating the multi-enzyme-like functions observed in natural enzymatic systems. This section aims to comprehend the underlying enzyme-like catalytic mechanisms inherent to dual-atomic-site SAzymes, thereby facilitating the development of next-generation SAzyme designs. Encouraged by theoretical studies based on density functional theory (DFT) calculations, Zhu's group demonstrated that the peroxo-like O₂ adsorption in Fe2NC nanozymes facilitated the activation of O-O bonds with an excellent enzyme-like performance (Fig. 6a). In particular, the corresponding free-energy profiles unveiled that the adsorbed O₂ species (O₂*) tended to evolve

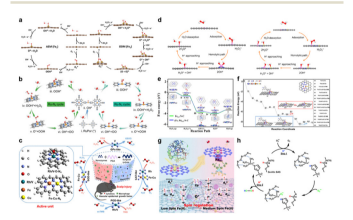


Fig. 6 (a) Schematics of the OXD-like pathway on Fe₁NC and Fe₂NC SAzymes following the adsorption evolution mechanism (AEM, left) and oxygen dissociation mechanism (ODM, right). (b) Proposed CAT-like ROS elimination pathways on Ru-N2 and Ru-N4 catalytic structures. Reproduced with permission from ref. 97 Copyright 2023 Wiley-VCH. (c) Single- and dual-atom nanozymes (RhN_4 , VN_4 , and $Fe-Cu-N_6$) with superior catalytic activities to accelerate scalp healing from brain trauma. Reproduced with permission from ref. 98 Copyright 2022, Springer Nature. Free energy diagrams for (d) the POD-like reaction on the Fe₁-NC and Fe₁Co₁-NC models and (e) the Fenton-like reaction mechanisms of (Fe, Pt)_{SA}-N-C and Fe_{SA}-N-C. (f) OXD-like energy evolution (eV) along the reaction coordinate of FeCoZn-TAC/SNC. Reproduced with permission from ref. 96 Copyright 2022, American Chemical Society. (g) Proposed reaction mechanism for engineering the spin state of Fe single-atom peroxidase-like nanozyme (FeNC) with Pd nanoclusters (PdNC). Reproduced with permission from ref. 99 Copyright 2022, Royal Society of Chemistry. (h) Proposed SOD-like reaction mechanism over Zn_A-Cu_B-Z of the zeolitic micropore.

into OOH*, O*, and OH* intermediates via the adsorption evolution mechanism (AEM) on the Fe₁NC OXD-like nanozyme. In contrast, Fe₂NC SAzymes showed a preference for direct O₂* dissociation, rather than forming OOH* intermediates, following the oxygen dissociation mechanism (ODM). In a parallel pursuit, Han and co-workers constructed a homogenous Ru-N₄/Ru-N₂-based dual-atomic-site SAzyme, endowed with exceptional ROS-scavenging activity (Fig. 6b). 97 They proposed and elucidated a novel nucleophilic attack pathway operating in the Ru-N₄/Ru-N₂-based CAT-like SAzymes.

Beyond the previously discussed homogenous dual-atom sites of SAzymes, Zhang and colleagues recently discovered SAzymes of RhN₄, VN₄, and Fe-Cu-N₆ configurations, each demonstrating efficient multienzyme-mimetic activities (Fig. 6c). 98 Specifically, Rh/VN₄ with an Rh/V-O-N₄ active center displayed CAT-like and glutathione peroxidase-like activities through a unique bilateral reaction mechanism, while Fe-Cu-N₆ showed impressive SOD-like activity and selectivity. Additionally, Rh/VN₄, housing Rh/V-O-N₄ active centers, displayed CAT-like and glutathione peroxidase-like activities propelled by a unique bilateral reaction mechanism. Similarly, Fe-Cu-N₆ showed impressive SOD-like activity and selectivity. A novel bilateral reaction mechanism was also proposed based on the Fe-Cu-N₆ model to explain their efficient multienzyme-mimetic catalysis. Song's group further illuminated the synergistic effect observed in Fe₁Co₁-NC dual-atom POD-like nanozymes. 91 This work revealed that the incorporation of heteronuclear metal sites induced electronic structure modulation and optimized H2O2 substrate adsorption (Fig. 6d). The coexistence of Fe and Co moieties facilitated diverse intermediate stabilization, ultimately reducing the overall reaction barrier. Recent studies by Zheng's group further revealed that the (Fe, Pt)_{SA}-N-C dinuclear nanozyme preferred a proton-mediated H2O2 homolytic pathway (Fig. 6e). 90 Furthermore, Chen's group designed a triple-atom FeCoZn-TAC/SNC nanozyme, showcasing strong interaction between the three metal centers, which facilitates both O2 and TMB intermediate adsorption and activation (Fig. 6f).⁹⁶ Expanding the scope of dual-atomic site SAzymes led to the integration of single-atom sites and clusters. For example, Zhu et al. developed an Fe SAzyme combined with Pd nanoclusters (Pd_{NC}) as a "modulator" (FeNC-Pd_{NC}, Fig. 6g). 99 The electronwithdrawing effect of Pd_{NC} induced the spin transition of Fe active sites in FeNC-Pd_{NC}, culminating an enhanced POD-like performance. This FeNC-Pd_{NC} POD-like mimic could be applied to construct a colorimetric immunosorbent assay for the selective detection of prostate-specific antigen targets. A recent work by Lo and his colleagues unveiled that the suitable molecular distances between Zn_A-Cu_B-Z bimetallic sites would facilitate tandem enzyme-like reactions and the overall enzyme catalytic redox cycle (Fig. 6h).⁹⁴

SAzymes featuring atomically dispersed dual-atomic sites exhibit configurations remarkably akin to those of natural enzymes, thus holding immense potential for bridging the gap between natural enzymes and nanozymes. In this section, the recent advancements in the application of emerging dualatomic site SAzymes, encompassing domains such as biosensing, therapy treatment, and cytoprotection, are highlighted. For instance, Wei's group has established versatile biosensing platforms employing the Zn/Mo DSAC-SMA SAzyme-based system for the precise determination of intracellular H2O2, glucose in serum, cholesterol, and ascorbic acid in commercial beverages (Fig. 7a)⁹² In a similar vein, Chen et al. fabricated ZnCoFe three-atom nanozymes (TAzyme) endowed with PODlike catalytic activities, which were further developed as a nanozyme sensor array for the colorimetric discrimination of phenolic acids (Fig. 7b). 100 Jin and co-workers also expanded the biosensing application of dual-atomic sites SAzyme, venturing into the electrochemiluminescence (ECL) field. 101 Benefiting from the synergistic effect exhibited by Fe, Co dual single-atom nanozymes (Fe, Co D-SACs), they successfully established an ultrasensitive luminol-dissolved O2 (DO) ECL system together with the plasmon enhancement of AgNPs tailored for the detection of prostate-specific antigens (PSAs, Fig. 7c).

SAzymes with dual-atomic sites have received significant recognition for their exceptional OXD-, POD-, SOD-, and SOD-

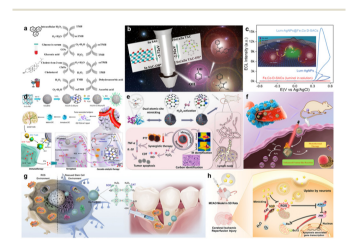


Fig. 7 Schematic illustrations of (a) a colorimetric biosensing platform for intracellular H2O2 detection, glucose detection in serum, cholesterol, and the ascorbic acid in commercial beverages based on the confined Zn/Mo dual single atom nanozyme, (b) visual sensor arrays for the discrimination of phenolic acids based on the ZnCoFe three-atom nanozyme, reproduced with permission from ref. 100 Copyright 2023, American Chemical Society, (c) plasmon-boosted Fe, Co dual singleatom nanozymes for constructing high-performance electrochemiluminescence (ECL) sensing platform of prostate-specific antigen targets, reproduced with permission from ref. 101 Copyright 2023, American Chemical Society, (d) FeCo/Fe-Co dual-metal atom nanozyme (FeCo/Fe-Co DAzyme) with multi-enzyme-like activities that induce cascade immunogenic ferroptosis, reproduced with permission from ref. 102 Copyright 2023, American Chemical Society, (e) biomimetic Fe-Cu dual-atomic-site nanozymes with intrinsic photothermal effect for tumor lymphatic metastasis inhibition, reproduced with permission from ref. 103 Copyright 2023, Elsevier, (f) synergistic application for tumor catalytic therapy based on (Fe, Pt)_{SA}-N-C dinuclear nanozymes, (g) $Ru-N_4/Ru-N_2$ -based antioxidase-like reactive oxygen nanobiocatalysts (ROBCs) to secure stem cells and periodontal tissues, and (h) Fe2NC@Se nanozymes with multi-enzyme cascade properties for reperfusion injury in ischemic stroke.

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like attributes, endowing them with the versatility required for diverse biomedical applications. Wang and colleagues successfully prepared a six-enzyme co-expressed composite nanoplatform (FeCo/Fe-Co DAzyme/PL, Fig. 7d). 102 Leveraging the multi-enzyme-like properties of FeCo/Fe-Co DAzyme/PL, this platform could achieve immunogenic tumor ferroptosis by initiating interferon γ (IFN-γ) activation and targeting arachidonic acid (AA) metabolism. In parallel, Zhu et al. unveiled a biomimetic Fe-Cu dual-atomic-site SAzyme (FeCuNC), designed for efficient activation of H2O2 (Fig. 7e). 103 The intrinsic chemodynamic and photothermal therapies facilitated by FeCuNC SAzymes synergistically induced apoptosis in cancer cells and restrained tumor growth. Similarly, Zheng and colleagues constructed an (Fe, Pt)_{SA}-N-C dinuclear nanozyme, boasting a high photothermal conversion efficiency for efficient tumor catalytic therapy (Fig. 7f). 90 Moreover, reactive oxygen nanobiocatalysts (ROBCs) based on Ru-N2 and Ru-N4 sites exhibited proficient antioxidase-like traits, demonstrating their potential for catalytic ROS elimination through robust CAT-like degradation of H₂O₂ (Fig. 7g).⁹⁷ The unique ultrafast and reversible redox-centers intrinsic to these novel ROBCs offer robust protective effects on stem cells and periodontal tissues, accentuating their superior anti-ROS therapeutic capabilities. Furthermore, the synergistic effect of multifunctional antioxidant Fe2NC@Se nanozymes was demonstrated to counteract oxidative damage and restrain neural apoptosis (Fig. 7h).89

4. Conclusions

SAzymes constructed with dual-atomic sites have attracted extensive attention and provided vast opportunities for developing next-generation nanozymes. In this mini-review, we have summarized the latest research advances made in applying DACs as nanozymes, including their synthesis strategies, characterization methods, and enzyme-like applications. Nevertheless, the development of DACs for nanozyme applications is still in the early stage, and more attention should be paid to some foreseeable challenges:

(1) Compared with SACs, achieving precise control and design in the synthesis of DACs remains a great challenge. Realizing further breakthroughs in preparation methods, precursor construction, and support selection holds great significance for advancing DACs in nanozyme research. On one front, the formation of non-uniform dual-atom sites and even clusters is a prevalent concern during DAC synthesis, hampering the scalable production of DACs with high metal loadings. Addressing this challenge is pivotal for achieving consistent and reproducible catalytic performance across various applications. Equally important, the coexistence of by-products in DACs can complicate the identification of real active sites in enzyme-like reactions. This intricacy underscores the need for precision in both synthesis and characterization to enable a thorough understanding of DACs' catalytic mechanisms. The pursuit of the goal "1 + 1 > 2" in DACs, where the combined

performance surpasses the sum of individual contributions, remains a question demanding further investigation. Sustaining the advantages of DACs over SACs necessitates novel strategies to capitalize on the potential of dual-atom configurations. In this endeavor, traditional synthetic methods might prove insufficient. A promising path forward involves combining multiple approaches rather than relying solely on a single method. Some molecular models featuring hetero-polynuclear or iso-polynuclear coordination compounds (e.g., MOFs, COFs, and hydrogen-bonded organic frameworks (HOFs)) hold the potential to offer the precise dual-atom configuration required for DACs.

- (2) Direct identification of dual-atomic sites with conclusive evidence remains a challenge. Current characterization techniques for DACs primarily rely on HAADF-STEM images and XAS analysis, which may fall short in fully resolving and distinguishing the intricate coordination environment within DACs. Of even greater significance is the accurate identification of the dynamic evolution of dual-atomic sites in SAzymes, a critical aspect that is presently lacking. To address this gap, the development of operando/in situ characterization techniques holds promise. Such methods could offer real-time insights into the dynamic enzyme-like behaviour of dualatomic active sites in SAzymes. Formulating a comprehensive structure-property relationship based on the dual-atom sites in SAzymes is a paramount pursuit. This endeavor is key to unraveling the intricacies of enzyme-like mechanisms at play. Such insights will not only enhance our understanding of catalytic behavior but also guide the rational design of next-generation SAzymes with improved and tailored properties. From the perspective of computational model methods, the integration of machine learning algorithms proves to be invaluable. These algorithms hold the potential to predict the complex relationship between the configuration and properties of SAzymes.
- (3) Bridging the gap between homogeneous biological catalysis and heterogeneous enzyme-like catalysis through SAzymes with dual-atomic sites represents a pivotal question demanding a resolution to usher in the era of next-generation nanozymes. The current landscape of SAzymes with dual-atomic sites has largely been shaped by trial-and-error methods. However, the periodic table offers a wealth of unexplored possibilities for discovering potential SAzymes with dual-atomic sites. The pursuit of groundbreaking strategies, rooted in a deep understanding of the structure–performance relationship inherent to SAzymes with dual-atomic active centers, stands as a pressing need.
- (4) The broad biomedical application of SAzymes with dualatom sites hinges on understanding the intricate structure– biological effect relationship and ensuring long-term biosafety. A thorough biosafety assessment of DAC nanozymes stands as a prerequisite for their potential clinical translation. This concern is not exclusive to DAC-based nanozymes; it is a broader issue encompassing many nanomaterials. The inorganic nature of these heterogeneous nanozymes underscores the initial stage of their interaction with biomedicine. Thus,

substantial efforts are required to comprehensively assess the fundamental interactions between DACs and biological systems. It is noteworthy that biocompatibility is a central point of discussion for various nanomaterials. 2D nanomaterials, in particular, have shown promising initial results in terms of biocompatibility and biodegradability. These materials could serve as suitable substrates for anchoring single atoms or dual atoms in a way that preserves their catalytic activities while maintaining biocompatibility. The evolution of real-time detection techniques, such as optical imaging, holds immense potential in elucidating the relationship between the DAC configuration and biological effects. This advancement enables researchers to closely observe how DACs interact with biological systems, shedding light on critical aspects including immune responses, transport pathways, and cytotoxicity. Such insights are instrumental in understanding the biocompatibility and potential risks associated with DAC-based nanozymes.

(5) The ultimate objective of SAzymes with dual-atomic active sites is to propel the advancement of nanozymes. It is strongly anticipated that a wave of captivating research centered around dual-atomic SAzymes and their practical applications in biology is on the horizon. A comprehensive exploration of the physicochemical and biological attributes of these dual-atomic centers within SAzymes is essential to unlock their potential across diverse biomedical contexts. While the field of SAzymes with dual-atomic sites is still in its formative stages, it holds immense promise. It is believed that dualatomic SAzymes will continually evolve, with new and more promising catalysts emerging to challenge and even replace natural enzymes in various applications.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 L. Gao, J. Zhuang, L. Nie, J. Zhang, Y. Zhang, N. Gu, T. Wang, J. Feng, D. Yang, S. Perrett and X. Yan, Nat. Nanotechnol., 2007, 2, 577-583.
- 2 H. Wei, L. Gao, K. Fan, J. Liu, J. He, X. Qu, S. Dong, E. Wang and X. Yan, *Nano Today*, 2021, **40**, 101269.

- 3 A. Robert and B. Meunier, ACS Nano, 2022, 16, 6956-6959.
- 4 Y. Huang, J. Ren and X. Qu, Chem. Rev., 2019, 119, 4357-
- 5 D. Jiang, D. Ni, Z. T. Rosenkrans, P. Huang, X. Yan and W. Cai, Chem. Soc. Rev., 2019, 48, 3683-3704.
- 6 H. Wei and E. Wang, Chem. Soc. Rev., 2013, 42, 6060-6093.
- 7 J. Wu, X. Wang, Q. Wang, Z. Lou, S. Li, Y. Zhu, L. Qin and H. Wei, Chem. Soc. Rev., 2019, 48, 1004-1076.
- 8 Y. Sun, B. Xu, X. Pan, H. Wang, Q. Wu, S. Li, B. Jiang and H. Liu, Coord. Chem. Rev., 2023, 475, 214896.
- 9 Y. Chong, Q. Liu and C. Ge, *Nano Today*, 2021, 37, 101076.
- 10 Y. Wang, R. Du, L. Y. S. Lee and K.-Y. Wong, Biosens. Bioelectron., 2022, 216, 114662.
- 11 L. Jiao, W. Xu, Y. Wu, H. Yan, W. Gu, D. Du, Y. Lin and C. Zhu, Chem. Soc. Rev., 2021, 50, 750-765.
- 12 X. Cai, L. Jiao, H. Yan, Y. Wu, W. Gu, D. Du, Y. Lin and C. Zhu, Mater. Today, 2021, 44, 211-228.
- 13 Y. Huang, X. Mu, J. Wang, Y. Wang, J. Xie, R. Ying and E. Su, J. Mater. Chem. B, 2022, 10, 1359-1368.
- 14 X. Li, L. Wang, D. Du, L. Ni, J. Pan and X. Niu, TrAC, Trends Anal. Chem., 2019, 120, 115653.
- 15 Z. Wang, R. Zhang, X. Yan and K. Fan, Mater. Today, 2020, 41, 81-119.
- 16 X.-F. Yang, A. Wang, B. Qiao, J. Li, J. Liu and T. Zhang, Acc. Chem. Res., 2013, 46, 1740-1748.
- 17 Y. Wang, H. Su, Y. He, L. Li, S. Zhu, H. Shen, P. Xie, X. Fu, G. Zhou, C. Feng, D. Zhao, F. Xiao, X. Zhu, Y. Zeng, M. Shao, S. Chen, G. Wu, J. Zeng and C. Wang, Chem. Rev., 2020, 120, 12217-12314.
- 18 G. Jia, M. Sun, Y. Wang, X. Cui, B. Huang and J. C. Yu, Adv. Funct. Mater., 2023, 33, 2212051.
- 19 B. Chang, L. Zhang, S. Wu, Z. Sun and Z. Cheng, Chem. Soc. Rev., 2022, 51, 3688-3734.
- 20 D. Wang and Y. Zhao, Chem, 2021, 7, 2635-2671.
- 21 B. Qiao, A. Wang, X. Yang, L. F. Allard, Z. Jiang, Y. Cui, J. Liu, J. Li and T. Zhang, Nat. Chem., 2011, 3, 634-641.
- 22 A. Wang, J. Li and T. Zhang, Nat. Rev. Chem., 2018, 2, 65-
- 23 X. Cui, W. Li, P. Ryabchuk, K. Junge and M. Beller, Nat. Catal., 2018, 1, 385-397.
- 24 L. Jiao, H. Yan, Y. Wu, W. Gu, C. Zhu, D. Du and Y. Lin, Angew. Chem., Int. Ed., 2020, 59, 2565-2576.
- 25 L. Huang, J. Chen, L. Gan, J. Wang and S. Dong, Sci. Adv., 2019, 5, eaav5490.
- 26 W. Ma, J. Mao, X. Yang, C. Pan, W. Chen, M. Wang, P. Yu, L. Mao and Y. Li, Chem. Commun., 2019, 55, 159-162.
- 27 Y. Wang, K. Qi, S. Yu, G. Jia, Z. Cheng, L. Zheng, Q. Wu, Q. Bao, Q. Wang, J. Zhao, X. Cui and W. Zheng, Nano-Micro Lett., 2019, 11, 102.
- 28 M. Zhou, Y. Jiang, G. Wang, W. Wu, W. Chen, P. Yu, Y. Lin, J. Mao and L. Mao, Nat. Commun., 2020, 11, 3188.
- 29 M. Chang, Z. Hou, M. Wang, D. Wen, C. Li, Y. Liu, Y. Zhao and J. Lin, Angew. Chem., Int. Ed., 2022, 61, e202209245.
- 30 B. Xu, H. Wang, W. Wang, L. Gao, S. Li, X. Pan, H. Wang, H. Yang, X. Meng, Q. Wu, L. Zheng, S. Chen, X. Shi,

Minireview

- K. Fan, X. Yan and H. Liu, Angew. Chem., Int. Ed., 2019, 58, 4911-4916.
- 31 Y. Wang, A. Cho, G. Jia, X. Cui, J. Shin, I. Nam, K.-J. Noh, B. J. Park, R. Huang and J. W. Han, Angew. Chem., Int. Ed., 2023, e202300119.
- 32 Y. Wang, G. Jia, X. Cui, X. Zhao, Q. Zhang, L. Gu, L. Zheng, L. H. Li, Q. Wu, D. J. Singh, D. Matsumura, T. Tsuji, Y.-T. Cui, J. Zhao and W. Zheng, Chem, 2021, 7, 436-449.
- 33 Y. Chen, P. Wang, H. Hao, J. Hong, H. Li, S. Ji, A. Li, R. Gao, J. Dong, X. Han, M. Liang, D. Wang and Y. Li, J. Am. Chem. Soc., 2021, 143, 18643-18651.
- 34 M. Chang, Z. Hou, M. Wang, C. Yang, R. Wang, F. Li, D. Liu, T. Peng, C. Li and J. Lin, Angew. Chem., Int. Ed., 2021, 60, 12971-12979.
- 35 G. Liao, L. Zhang, C. Li, S.-Y. Liu, B. Fang and H. Yang, Matter, 2022, 5, 3341-3374.
- 36 S. Ji, B. Jiang, H. Hao, Y. Chen, J. Dong, Y. Mao, Z. Zhang, R. Gao, W. Chen, R. Zhang, Q. Liang, H. Li, S. Liu, Y. Wang, Q. Zhang, L. Gu, D. Duan, M. Liang, D. Wang, X. Yan and Y. Li, Nat. Catal., 2021, 4, 407-417.
- 37 H. Xiang, W. Feng and Y. Chen, Adv. Mater., 2020, 32, 1905994.
- 38 V. Kandathil and S. A. Patil, Adv. Colloid Interface Sci., 2021, 294, 102485.
- 39 Y. Wang, Z. Zhang, G. Jia, L. Zheng, J. Zhao and X. Cui, Chem. Commun., 2019, 55, 5271-5274.
- 40 W. Wu, L. Huang, E. Wang and S. Dong, Chem. Sci., 2020, 11, 9741-9756.
- 41 P. A. Williams, J. Cosme, A. Ward, H. C. Angove, D. Matak Vinković and H. Jhoti, Nature, 2003, 424, 464-468.
- 42 I. G. Denisov, T. M. Makris, S. G. Sligar and I. Schlichting, Chem. Rev., 2005, 105, 2253-2278.
- 43 W. Zhang, Y. Chao, W. Zhang, J. Zhou, F. Lv, K. Wang, F. Lin, H. Luo, J. Li, M. Tong, E. Wang and S. Guo, Adv. Mater., 2021, 33, 2102576.
- 44 Z. He, K. He, A. W. Robertson, A. I. Kirkland, D. Kim, J. Ihm, E. Yoon, G.-D. Lee and J. H. Warner, Nano Lett., 2014, 14, 3766-3772.
- 45 Y. Wang, B. J. Park, V. K. Paidi, R. Huang, Y. Lee, K.-J. Noh, K.-S. Lee and J. W. Han, ACS Energy Lett., 2022, 7,640-649.
- 46 G. Jia, M. Sun, Y. Wang, Y. Shi, L. Zhang, X. Cui, B. Huang and J. C. Yu, Adv. Funct. Mater., 2022, 32, 2206817.
- 47 R. Li and D. Wang, Adv. Energy Mater., 2022, 12, 2103564.
- 48 L. Li, K. Yuan and Y. Chen, Acc. Mater. Res., 2022, 3, 584-
- 49 A. Pedersen, J. Barrio, A. Li, R. Jervis, D. J. L. Brett, M. M. Titirici and I. E. L. Stephens, Adv. Energy Mater., 2022, 12, 2102715.
- 50 H. Yan, Y. Lin, H. Wu, W. Zhang, Z. Sun, H. Cheng, W. Liu, C. Wang, J. Li, X. Huang, T. Yao, J. Yang, S. Wei and J. Lu, Nat. Commun., 2017, 8, 1070.
- 51 L. Zhang, R. Si, H. Liu, N. Chen, Q. Wang, K. Adair, Z. Wang, J. Chen, Z. Song, J. Li, M. N. Banis, R. Li,

- T.-K. Sham, M. Gu, L.-M. Liu, G. A. Botton and X. Sun, Nat. Commun., 2019, 10, 4936.
- 52 L. Bai, C.-S. Hsu, D. T. L. Alexander, H. M. Chen and X. Hu, J. Am. Chem. Soc., 2019, 141, 14190-14199.
- 53 Z. Zhang, C. Feng, C. Liu, M. Zuo, L. Qin, X. Yan, Y. Xing, H. Li, R. Si, S. Zhou and J. Zeng, Nat. Commun., 2020, 11, 1215.
- 54 Y. Yang, Y. Qian, H. Li, Z. Zhang, Y. Mu, D. Do, B. Zhou, J. Dong, W. Yan, Y. Qin, L. Fang, R. Feng, J. Zhou, P. Zhang, J. Dong, G. Yu, Y. Liu, X. Zhang and X. Fan, Sci. Adv., 2020, 6, eaba6586.
- 55 S. Tian, Q. Fu, W. Chen, Q. Feng, Z. Chen, J. Zhang, W.-C. Cheong, R. Yu, L. Gu, J. Dong, J. Luo, C. Chen, Q. Peng, C. Draxl, D. Wang and Y. Li, Nat. Commun., 2018, 9, 2353.
- 56 S. Tian, B. Wang, W. Gong, Z. He, Q. Xu, W. Chen, Q. Zhang, Y. Zhu, J. Yang, Q. Fu, C. Chen, Y. Bu, L. Gu, X. Sun, H. Zhao, D. Wang and Y. Li, Nat. Commun., 2021, 12, 3181.
- 57 N. Zhang, X. Zhang, Y. Kang, C. Ye, R. Jin, H. Yan, R. Lin, J. Yang, Q. Xu, Y. Wang, Q. Zhang, L. Gu, L. Liu, W. Song, J. Liu, D. Wang and Y. Li, Angew. Chem., Int. Ed., 2021, 60, 13388-13393.
- 58 E. Vorobyeva, E. Fako, Z. Chen, S. M. Collins, D. Johnstone, P. A. Midgley, R. Hauert, O. V. Safonova, G. Vilé, N. López, S. Mitchell and J. Pérez-Ramírez, Angew. Chem., Int. Ed., 2019, 58, 8724-8729.
- 59 M. Zhou, Z. Wang, A. Mei, Z. Yang, W. Chen, S. Ou, S. Wang, K. Chen, P. Reiss, K. Qi, J. Ma and Y. Liu, Nat. Commun., 2023, 14, 2473.
- 60 J. Fu, J. Dong, R. Si, K. Sun, J. Zhang, M. Li, N. Yu, B. Zhang, M. G. Humphrey, Q. Fu and J. Huang, ACS Catal., 2021, 11, 1952–1961.
- 61 Q. Shi, Y. Ji, W. Chen, Y. Zhu, J. Li, H. Liu, Z. Li, S. Tian, L. Wang, Z. Zhong, L. Wang, J. Ma, Y. Li and F. Su, Natl. Sci. Rev., 2019, 7, 600-608.
- 62 Y. Zhao, K. R. Yang, Z. Wang, X. Yan, S. Cao, Y. Ye, Q. Dong, X. Zhang, J. E. Thorne, L. Jin, K. L. Materna, A. Trimpalis, H. Bai, S. C. Fakra, X. Zhong, P. Wang, X. Pan, J. Guo, M. Flytzani-Stephanopoulos, G. W. Brudvig, V. S. Batista and D. Wang, Proc. Natl. Acad. Sci. U. S. A., 2018, 115, 2902-2907.
- 63 J. Wang, Z. Huang, W. Liu, C. Chang, H. Tang, Z. Li, W. Chen, C. Jia, T. Yao, S. Wei, Y. Wu and Y. Li, J. Am. Chem. Soc., 2017, 139, 17281-17284.
- 64 X. Han, X. Ling, D. Yu, D. Xie, L. Li, S. Peng, C. Zhong, N. Zhao, Y. Deng and W. Hu, Adv. Mater., 2019, 31, 1905622.
- 65 A. Han, X. Wang, K. Tang, Z. Zhang, C. Ye, K. Kong, H. Hu, L. Zheng, P. Jiang, C. Zhao, Q. Zhang, D. Wang and Y. Li, Angew. Chem., Int. Ed., 2021, 60, 19262-19271.
- 66 M. Liu, N. Li, S. Cao, X. Wang, X. Lu, L. Kong, Y. Xu and X.-H. Bu, Adv. Mater., 2022, 34, 2107421.
- 67 Y.-X. Zhang, S. Zhang, H. Huang, X. Liu, B. Li, Y. Lee, X. Wang, Y. Bai, M. Sun, Y. Wu, S. Gong, X. Liu,

- Z. Zhuang, T. Tan and Z. Niu, J. Am. Chem. Soc., 2023, 145, 4819-4827.
- 68 Y. Ying, X. Luo, J. Qiao and H. Huang, Adv. Funct. Mater., 2021, 31, 2007423.
- 69 Y. Song, F. Kong, X. Sun, Q. Liu, X. Li, L. Ren, Y. Xie, Y. Shen and J. Wang, Adv. Energy Mater., 2023, 13, 2203660.
- 70 L. Zhang, J. Feng, S. Liu, X. Tan, L. Wu, S. Jia, L. Xu, X. Ma, X. Song, J. Ma, X. Sun and B. Han, Adv. Mater., 2023, 2209590, DOI: 10.1002/adma.202209590.
- 71 F. Kraushofer, L. Haager, M. Eder, A. Rafsanjani-Abbasi, Z. Jakub, G. Franceschi, M. Riva, M. Meier, M. Schmid, U. Diebold and G. S. Parkinson, ACS Energy Lett., 2022, 7, 375-380.
- 72 K. Jiang, S. Back, A. J. Akey, C. Xia, Y. Hu, W. Liang, D. Schaak, E. Stavitski, J. K. Nørskov, S. Siahrostami and H. Wang, Nat. Commun., 2019, 10, 3997.
- 73 K. Jiang, S. Siahrostami, A. J. Akey, Y. Li, Z. Lu, J. Lattimer, Y. Hu, C. Stokes, M. Gangishetty, G. Chen, Y. Zhou, W. Hill, W.-B. Cai, D. Bell, K. Chan, J. K. Nørskov, Y. Cui and H. Wang, Chem, 2017, 3, 950-960.
- 74 H. Fei, J. Dong, Y. Feng, C. S. Allen, C. Wan, B. Volosskiy, M. Li, Z. Zhao, Y. Wang, H. Sun, P. An, W. Chen, Z. Guo, C. Lee, D. Chen, I. Shakir, M. Liu, T. Hu, Y. Li, A. I. Kirkland, X. Duan and Y. Huang, Nat. Catal., 2018, 1, 63 - 72.
- 75 Y. Wang, X. Cui, J. Zhao, G. Jia, L. Gu, Q. Zhang, L. Meng, Z. Shi, L. Zheng, C. Wang, Z. Zhang and W. Zheng, ACS Catal., 2019, 9, 336-344.
- 76 Q. Hao, H.-X. Zhong, J.-Z. Wang, K.-H. Liu, J.-M. Yan, Z.-H. Ren, N. Zhou, X. Zhao, H. Zhang, D.-X. Liu, X. Liu, L.-W. Chen, J. Luo and X.-B. Zhang, Nat. Synth., 2022, 1, 719-728.
- 77 X. Miao, W. Chen, S. Lv, A. Li, Y. Li, Q. Zhang, Y. Yue, H. Zhao, L. Liu, S. Guo and L. Guo, Adv. Mater., 2023, 2211790, DOI: 10.1002/adma.202211790.
- 78 J. Pei, T. Wang, R. Sui, X. Zhang, D. Zhou, F. Qin, X. Zhao, Q. Liu, W. Yan, J. Dong, L. Zheng, A. Li, J. Mao, W. Zhu, W. Chen and Z. Zhuang, Energy Environ. Sci., 2021, 14, 3019-3028.
- 79 Z. Fan, R. Luo, Y. Zhang, B. Zhang, P. Zhai, Y. Zhang, C. Wang, J. Gao, W. Zhou, L. Sun and J. Hou, Angew. Chem., Int. Ed., 2023, 62, e202216326.
- 80 X. Li, X. Yang, J. Zhang, Y. Huang and B. Liu, ACS Catal., 2019, 9, 2521-2531.
- 81 S. Baumann, W. Paul, T. Choi, C. P. Lutz, A. Ardavan and A. J. Heinrich, Science, 2015, 350, 417-420.
- 82 C. Ling, X. Liu, H. Li, X. Wang, H. Gu, K. Wei, M. Li, Y. Shi, H. Ben, G. Zhan, C. Liang, W. Shen, Y. Li, J. Zhao and L. Zhang, Angew. Chem., Int. Ed., 2022, 61, e202200670.
- 83 U. I. Kramm, L. Ni and S. Wagner, Adv. Mater., 2019, 31,
- 84 G.-F. Han, F. Li, A. I. Rykov, Y.-K. Im, S.-Y. Yu, J.-P. Jeon, S.-J. Kim, W. Zhou, R. Ge, Z. Ao, T. J. Shin, J. Wang,

- H. Y. Jeong and J.-B. Baek, Nat. Nanotechnol., 2022, 17, 403-407.
- 85 G. Yang, J. Zhu, P. Yuan, Y. Hu, G. Qu, B.-A. Lu, X. Xue, H. Yin, W. Cheng, J. Cheng, W. Xu, J. Li, J. Hu, S. Mu and J.-N. Zhang, Nat. Commun., 2021, 12, 1734.
- 86 A. A. Ammann, J. Mass Spectrom., 2007, 42, 419-427.
- 87 G. Lalande, R. Côté, G. Tamizhmani, D. Guay, J. P. Dodelet, L. Dignard-Bailey, L. T. Weng and P. Bertrand, Electrochim. Acta, 1995, 40, 2635-2646.
- 88 L. Jiao, W. Ye, Y. Kang, Y. Zhang, W. Xu, Y. Wu, W. Gu, W. Song, Y. Xiong and C. Zhu, Nano Res., 2022, 15, 959-964.
- 89 R. Tian, H. Ma, W. Ye, Y. Li, S. Wang, Z. Zhang, S. Liu, M. Zang, J. Hou, J. Xu, Q. Luo, H. Sun, F. Bai, Y. Yang and J. Liu, Adv. Funct. Mater., 2022, 32, 2204025.
- 90 S. Wang, Z. Hu, Q. Wei, P. Cui, H. Zhang, W. Tang, Y. Sun, H. Duan, Z. Dai, Q. Liu and X. Zheng, ACS Appl. Mater. Interfaces, 2022, 14, 20669-20681.
- 91 R. Zeng, Y. Li, X. Hu, W. Wang, Y. Li, H. Gong, J. Xu, L. Huang, L. Lu, Y. Zhang, D. Tang and J. Song, Nano Lett., 2023, 23, 6073-6080.
- 92 C.-B. Ma, Y. Xu, L. Wu, Q. Wang, J.-J. Zheng, G. Ren, X. Wang, X. Gao, M. Zhou, M. Wang and H. Wei, Angew. Chem., Int. Ed., 2022, 61, e202116170.
- 93 X. Jin, X. Feng, G. Wang, R. Tan, Y. Peng, L. Zheng, W. Chen and Z. Chen, J. Mater. Chem. A, 2023, 11, 2326-
- 94 C. K. T. Wun, H. K. Mok, T. Chen, T.-S. Wu, K. Taniya, K. Nakagawa, S. Day, C. C. Tang, Z. Huang, H. Su, W.-Y. Yu, T. K. W. Lee and T. W. B. Lo, Chem. Catal., 2022, 2, 2346-2363.
- 95 Q. Chen, Y. Liu, Y. Lu, Y. Hou, X. Zhang, W. Shi and Y. Huang, J. Hazard. Mater., 2022, 422, 126929.
- 96 R. Wu, M. Sun, X. Liu, F. Qin, X. Zhang, Z. Qian, J. Huang, Y. Li, T. Tan, W. Chen and Z. Chen, Anal. Chem., 2022, 94, 14308-14316.
- 97 J. Guo, Z. Xing, L. Liu, Y. Sun, H. Zhou, M. Bai, X. Liu, M. Adeli, C. Cheng and X. Han, Adv. Funct. Mater., 2023, 33, 2211778.
- 98 S. Zhang, Y. Li, S. Sun, L. Liu, X. Mu, S. Liu, M. Jiao, X. Chen, K. Chen, H. Ma, T. Li, X. Liu, H. Wang, J. Zhang, J. Yang and X.-D. Zhang, Nat. Commun., 2022, 13, 4744.
- 99 X. Wei, S. Song, W. Song, Y. Wen, W. Xu, Y. Chen, Z. Wu, Y. Qin, L. Jiao, Y. Wu, M. Sha, J. Huang, X. Cai, L. Zheng, L. Hu, W. Gu, M. Eguchi, T. Asahi, Y. Yamauchi and C. Zhu, Chem. Sci., 2022, 13, 13574-13581.
- 100 J. Huang, H. Gu, G. Wang, R. Wu, M. Sun and Z. Chen, Anal. Chem., 2023, 95, 9107-9115.
- 101 F. A. Bushira, P. Wang, Y. Wang, S. Hou, X. Diao, H. Li, L. Zheng and Y. Jin, Anal. Chem., 2022, 94, 9758-9765.
- 102 Y. Liu, R. Niu, R. Deng, S. Song, Y. Wang and H. Zhang, J. Am. Chem. Soc., 2023, 145, 8965-8978.
- 103 L. Jiao, N. Tao, Y. Kang, W. Song, Y. Chen, Y. Zhang, W. Xu, Y. Wu, W. Gu, L. Zheng, L. Chen, L. Deng, C. Zhu and Y.-N. Liu, Nano Today, 2023, 50, 101859.