Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2024, 15, 19513

d All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 27th July 2024 Accepted 18th October 2024

DOI: 10.1039/d4sc05014b

rsc.li/chemical-science

Steering N/S coordination number to accelerate catecholase-like catalysis over low-coordinated Cu site†

Meng Yuan,^a Nannan Xia,^b Ziheng Huang,^a Chaofeng Huang,^c Xun Hu^b*^a and Fei He^b*^a

Modulation of coordination configuration is crucial for boosting the biomimetic catalytic activity of nanozymes, but remains challenging. Here, we found that the non-first-shell amino group in the ligand was capable of steering the N/S coordination number through remote induction to enable the formation of a low-coordinated CuN_2S_1 configuration. This endowed the resulting nanozyme (ATT-Cu) with an upshifted d-band center compared with a control nanozyme (TT-Cu) with CuN_1S_3 configuration, enhancing the adsorption capabilities of ATT-Cu for O_2 and H_2O_2 intermediates as well as its affinity for catechol. Additionally, the low-coordinated CuN_2S_1 configuration caused more charges to accumulate at the atomic Cu site, which improved the capabilities of ATT-Cu for both donating electrons to oxygen-related species and accepting electrons from catechol. As a result, this ATT-Cu nanozyme with a low-coordinated CuN_2S_1 moiety presented a faster initial oxygen reduction step, which in turn accelerated catechol oxidation, thus greatly boosting the catecholase-like activity of ATT-Cu that exceeded those of many catecholase-mimicking artificial enzymes/nanozymes with $Cu-N_xO_y$ sites as well as those of Cebased, Zr-based and Pt-based nanozymes.

Introduction

Nanozymes with enzyme-like catalytic functions are potential candidates for replacing natural enzymes due to their tunable activities, excellent recyclability and low cost. ¹⁻⁹ Despite these advantages, the broad application prospects of nanozymes are usually hindered by their poor catalytic activities. To address this challenge, various strategies have been employed to improve the biomimetic catalytic activities of nanozymes. Taking nanozyme-mimicking catechol oxidase (CO) as an example, regulating multinuclearity, ² chiral microenvironment, ¹⁰ coordination saturation ¹¹ and the types of coordination center atoms ¹²⁻¹⁴ have generally been used to manipulate the catalytic activities of CO-like nanozymes. Despite great advances, many CO-mimicking nanozymes or artificial enzymes, especially those carrying M-N_xO_y moieties (where M represents a metal) as catalytic sites, still exhibit unsatisfactory

activities. ^{10,15} One possible reason is the strong electronegativity of coordinated N and/or O atoms with small atomic radius, which results in the metal sites either have unsuitable free energy for adsorption/desorption of reaction-related species (including reactants, intermediates and/or products) ¹⁶ or inappropriate electron transfer capability for donating and/or accepting electrons.

To overcome these obstacles, a potential solution is to choose a coordinating atom with comparatively large atomic size and weak electronegativity¹⁷ to regulate the interface configuration of the central metal atoms, ¹⁶ thus boosting the catalytic activity of CO-mimicking nanozymes by adjusting the electronic state of the metal site to optimize the potential barriers to adsorption/desorption as well as electron transfer capability during biomimetic catalysis. In this regard, using S to partially replace N or O as the coordinating atom may be a promising choice due to its large atomic size and weak electronegativity compared to N and O atoms. However, few CO-mimicking nanozymes with M-N_xS_y or M-O_xS_y moieties have been reported.

With this goal in mind, we herein used the non-first-shell amino group to perturb the N/S coordination number around an atomic Cu center to construct two nanozymes with CuN_2S_1 and CuN_1S_3 moieties for CO-mimicking catalysis. We found that the low-coordinated CuN_2S_1 configuration endowed the resulting nanozyme with an upshifted d-band center, which enhanced the adsorption strength of the Cu site for O_2 and

[&]quot;School of Material Science and Engineering, University of Jinan, Jinan 250024, China. E-mail: xun.hu@outlook.com; mse_hef@ujn.edu.cn

bState Key Laboratory of Biobased Material and Green Papermaking, Key Laboratory of Pulp & Paper Science and Technology of Shandong Province/Ministry of Education, Qilu University of Technology (Shandong Academy of Sciences), Jinan 250353, China School of Chemistry and Chemical Engineering/State Key Laboratory Incubation Base for Green Processing of Chemical Engineering, Shihezi University, Shihezi 832000, China

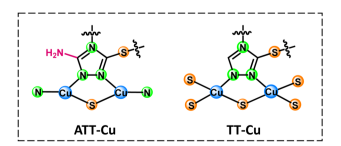
[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d4sc05014b

 ${
m H_2O_2}$ intermediates as well as the affinity for catechol. Moreover, this low-coordinated ${
m CuN_2S_1}$ moiety promoted charge accumulation over the atomic Cu center, enabling the resulting nanozyme to more easily donate electrons to oxygen-related species and accept electrons from catechol. Accordingly, when applied to CO-mimicking catalysis, the unsaturated ${
m CuN_2S_1}$ site accelerated the initial oxygen reduction process, which in turn promoted catechol oxidation, thereby endowing ATT-Cu with a superior CO-like activity that surpassed that of a control nanozyme with a ${
m CuN_1S_3}$ moiety and those of many CO-mimicking artificial enzymes/nanozymes with ${
m Cu-N_xO_y}$ moieties as well as those of other Zr-based, Ce-based or Pt-based nanozymes.

Results and discussion

Considering that the active site of natural CO involved Cu, ¹⁸ we chose Cu as the coordination center atom. To construct a nanozyme with a CuN_xS_y moiety, two ligands, *i.e.*, 3-thiol-1,2,4-triazole (TT) and 3-amino-5-thiol-1,2,4-triazole (ATT), were used to synthesize nanozymes (TT-Cu and ATT-Cu, Scheme 1) through a simple one-step coordination reaction between the ligands and CuSO_4 in water at room temperature. (See the Experimental section in ESI.†) The only difference between TT and ATT was whether there existed an amino group, which was introduced to perturb the coordination microenvironment, thus modulating the coordination configuration of the Cu center.

A transmission electron microscope (TEM) was first used to observe the morphologies of TT-Cu and ATT-Cu. Interconnected nanoparticles with average sizes of 15.4 nm and 42.2 nm were observed for TT-Cu and ATT-Cu (Fig. 1A and B), respectively. The larger size of ATT-Cu could probably be attributed to the existence of -NH₂ in the ATT ligand, contributing to forming an extra H-bond network.¹⁹ Such a morphology with the formation of interconnected nanoparticles was also confirmed by scanning electron microscope (SEM) (Fig. 1C and S1†). The survey scan spectra of X-ray photoelectron spectroscopy (XPS) detected the elemental signals of C, N, O, S and Cu (Fig. S2†), which were uniformly distributed in both nanozymes based on energy dispersive spectroscopy (EDS) mapping images (Fig. 1D and S3†). The results of inductively coupled plasma optical emission spectrometry (ICP-OES) indicated that the Cu contents in TT-Cu



Scheme 1 Coordination configurations of atomic Cu sites in ATT-Cu and TT-Cu. Note that only the structures of ligands and local motifs around Cu centers are presented for simplicity.

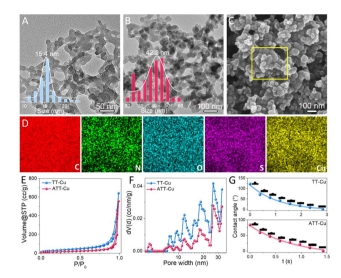


Fig. 1 TEM images of TT-Cu (A) and ATT-Cu (B); SEM image (C) and EDS mapping (D) of ATT-Cu; $\rm N_2$ adsorption—desorption (E) and pore size distribution (F) curves of ATT-Cu and TT-Cu; (G) contact angles of ATT-Cu and TT-Cu.

and ATT-Cu were 43.9% and 31.9% (Fig. S4†), respectively. According to an N_2 adsorption/desorption experiment, we found that introducing the amino group caused a reduced surface area and pore volume in ATT-Cu compared with those of TT-Cu (Fig. 1E), although both showed similar mesopore-dominated porous structures (Fig. 1F). Meanwhile, both ATT-Cu and TT-Cu exhibited hydrophilic features (Fig. 1G), which would endow the active sites with an easy accessibility for hydrated O_2 during CO-mimicking catalysis.²⁰

The collected N 1s spectra displayed that most N atoms in the TT ligand coordinated with Cu to form Cu-N bonds, in addition to the presence of a few uncoordinated N (Fig. 2A). Likewise, the formation of Cu-N bonds was also found for ATT-Cu (Fig. 2A). Differing from TT-Cu, another signal in the N 1s spectrum of ATT-Cu was also observed, which probably corresponded to exocyclic -NH2.21 The calculated ratio of Cu-N and C-NH₂ was close to 3, implying that Cu possibly tended to coordinate with heterocyclic N atoms rather than exocyclic -NH2. This result was attributable to the weak Lewis basic feature of -NH2 in the ATT ligand,22 probably resulting in the poorer coordination ability of -NH2.23 The O 1s spectra of ATT-Cu and TT-Cu detected H₂O and SO₄²⁻ originating from CuSO₄ (Fig. S5†). In the S 2p spectra, the important Cu-S signals were observed for both nanozymes apart from C-S and SO₄²⁻ (Fig. 2B),²⁴ confirming the occurrence of coordination between Cu and the S atom of the ligand.

To further confirm the existence of Cu–N and Cu–S coordination, time-of-flight secondary ion mass spectrometry (ToF-SIMS) was conducted to disclose coordination information about the Cu center by identifying the mass fragments ejected from ATT-Cu and TT-Cu. In Fig. 2C, the characteristic fragment ions with mass-to-charge ratios (m/z) of 255.8 (Cu₃C₂N₃H₁) and 287.7 (Cu₃C₂N₃S₁H₁) were detected for ATT-Cu, which may be assigned to Cu₃-coordinated ATT fragments involving Cu–N and

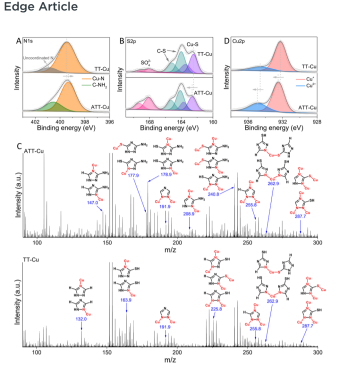


Fig. 2 N 1s (A) and S 2p (B) XPS spectra of ATT-Cu and TT-Cu; (C) ToF-SIMS analysis of ATT-Cu and TT-Cu and possible fragment assignments; (D) Cu 2p XPS spectra of ATT-Cu and TT-Cu.

Cu–S coordination, indicating that the S atom and three heterocyclic N atoms of ATT could bond with Cu. The possible existence of Cu–N and/or Cu–S coordination in the Cu₂-coordinated ATT fragment ions, featuring m/z of 191.9 (Cu₂C₂N₃), 208.9 (Cu₂C₂N₄H₃) and 240.8 (Cu₂C₂N₄S₁H₃), as well as the Cu₁-coordinated ATT fragments with m/z of 147.0 (Cu₁C₂N₄H₄), 177.9 (Cu₁C₂N₄S₁H₃), 178.9 (Cu₁C₂N₄S₁H₄) and 262.9 (Cu₁C₄-N₆S₂H₄) supported coordination mainly occurring between the heterocyclic N and S atoms of ATT and Cu. Similar to ATT-Cu, the Cu₁/Cu₂/Cu₃-coordinated TT fragments involving Cu–N and/or Cu–S coordination were also found for TT-Cu. These ToF-SIMS data, together with the XPS results, physically demonstrated that the three heterocyclic N atoms and S atoms of ATT/TT tended to coordinate with Cu to form CuN_xS_y configurations.

To explore how the coordinated N/S atoms in ATT and TT affected the electronic state of Cu, we collected the Cu 2p XPS spectra of ATT-Cu and TT-Cu (Fig. 2D). The Cu⁺ and Cu²⁺ species coexisted in both ATT-Cu and TT-Cu. Importantly, the binding energy of Cu in ATT-Cu was shifted positively compared with that in TT-Cu, verifying the accumulation of more positive charges at the Cu center of ATT-Cu. In Cu K-edge X-ray absorption near-edge structure (XANES) spectroscopy, the absorption edge energies of both ATT-Cu and TT-Cu were higher than that of Cu foil (Fig. 3A), indicating that Cu was in the oxidized state in both nanozymes. Compared with TT-Cu, the absorption edge energy of ATT-Cu was shifted positively, and the intensity of the white line peak corresponding to the 1s \rightarrow 4p_{xy} transition also increased (Fig. 3A). These results reconfirmed the accumulation of more positive charges at the

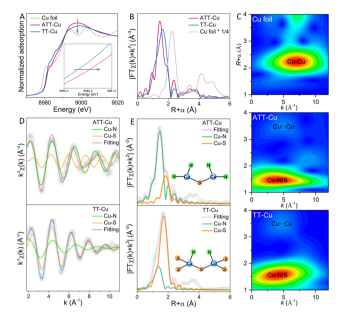


Fig. 3 X-ray absorption spectra of ATT-Cu, TT-Cu and Cu foil. (A) Cu K-edge XANES spectra; (B) Fourier-transform EXAFS curves in *R* space; (C) wavelet-transform EXAFS spectra; Fourier-transform EXAFS fitting curves in *k*-space (D) and *R*-space (E).

Cu center of ATT-Cu, agreeing well with the Cu 2p XPS result. Upon rechecking the N 1s and S 2p XPS spectra, we found that the Cu–N bond in ATT-Cu was shifted to a higher binding energy than that in TT-Cu, while the binding energy of the Cu–S bond in ATT-Cu was lower than that in TT-Cu (Fig. 2A and B). These results together with the Cu 2p and Cu K-edge XANES spectra illustrated that the more electrons of the Cu–N bond in ATT-Cu flowed toward the S atom compared with TT-Cu, causing more positive charges to accumulate at the Cu center of ATT-Cu.

To gain an insight into the origin of charge accumulation at the Cu center, extended X-ray absorption fine structure (EXAFS) spectroscopy was conducted to investigate the atomic-scale coordination configuration of Cu. As shown in Fig. 3B, the characteristic peaks of ATT-Cu and TT-Cu in Fourier-transform EXAFS spectra in R space were located at about 1.44 and 1.66 Å, respectively, and no Cu-Cu scattering peak at ~ 2.24 Å was observed for them, suggesting that the Cu atom of ATT-Cu and TT-Cu existed in the form of atomic dispersion. In wavelet transform (WT)-EXAFS spectra (Fig. 3C), the intensity maximum for ATT-Cu and TT-Cu were situated at about 3.90 $\mbox{\AA}^{-1}$ and 4.05 Å⁻¹, respectively, which were clearly different from that for Cu foil (7.05 Å^{-1}), again supporting the atomic dispersion of Cu in both nanozymes. Additionally, we observed another prominent scattering path at \sim 3.48 Å in WT-EXAFS (Fig. 3C), which could possibly be assigned to the nonbonded Cu···Cu scattering originating from the adjacent geminal copper centers, similar to the reported triazole-Cu complex. 19,21

The Fourier-transform EXAFS signal in *k*-space revealed that both ATT-Cu and TT-Cu presented two different oscillation periods, which corresponded to the Cu-N and Cu-S paths (Fig. 3D). This favored the formation of Cu-N and Cu-S

Chemical Science

coordination, coinciding with the results of the N 1s/S 2p spectra and ToF-SIMS data. We thus performed Fouriertransform EXAFS fitting analysis in R space to determine the first-shell N/S atom numbers around the Cu center (Fig. 3E and Table S1†). For ATT-Cu, the average number of S atoms coordinating with Cu was 0.5, suggesting that each S atom was shared by two Cu atoms. Meanwhile, each Cu atom in ATT-Cu was also connected with two N atoms. Therefore, the Cu center of the ATT-Cu nanozyme formed the coordinationunsaturated CuN2S1 configuration. In contrast to ATT-Cu, each Cu center in TT-Cu boned with one N atom, and the average number of S atoms coordinating with Cu attained 2.5, suggesting that the adjacent two Cu atoms shared five S atoms, thus forming the CuN₁S₃ coordination configuration in the TT-Cu nanozyme. These XANES and EXAFS data, together with the XPS and ToF-SIMS results as well as the different coordination structures of the nanozymes illustrated that introducing an amino group into the ATT ligand could modulate the coordination-unsaturated configuration of atomically dispersed Cu centers by steering the first-shell N/S atom numbers, thus promoting the more notable accumulation of positive charge at the Cu center of ATT-Cu.

The CO-like catalytic activities of ATT-Cu and TT-Cu were evaluated using 3,5-di-tert-butylcatechol (3,5-DBTC) as a model molecule.2 In the absence of nanozyme, 3,5-DBTC in an airsaturated solution showed a characteristic absorption peak at 280 nm (Fig. 4A). Once the ATT-Cu nanozyme was introduced into the reaction solution, a new absorption peak located at 416 nm appeared, which was assigned to the oxidation product (i.e., 3,5-di-tert-butyl-o-benzoquinone, 3,5-DTBQ) (Fig. 4A). When the reaction solution containing ATT-Cu was filtered using a filtration membrane with an average pore size of 100 nm, the reaction rate in this system obviously decreased (Fig. S6†). Meanwhile, increasing the O2 concentration accelerated the CO-mimicking catalysis of ATT-Cu, suggesting that ATT-Cu activated O2 to oxidize 3,5-DBTC (Fig. S7†). These results jointly verified the CO-mimicking activity of the ATT-Cu

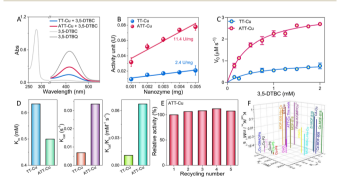


Fig. 4 Evaluation of CO-like activities of ATT-Cu and TT-Cu. (A) UVvis spectra of 3,5-DTBC, 3,5-DTBQ and nanozyme-catalyzed oxidation of 3,5-DTBC in air-saturated PBS (pH = 8)/CH₃CN solution; (B) specific activities; (C) Michaelis-Menten curves; (D) kinetic parameters (K_{m}, K_{cat}) and K_{cat}/K_{m} ; (E) recyclability; (F) comparison of the kinetic parameters (K_m , K_{cat} and K_{cat}/K_m) of ATT-Cu with reported CO-like artificial enzymes.

nanozyme. Moreover, the CO-like activity of ATT-Cu increased with the increasing pH of the solution (Fig. S8†). For comparison with reported CO-like nanozymes, 2,10 a solution of pH = 8 was used. Additionally, we found that sodium metabisulfite (Na₂S₂O₅) acting as an inhibitor could significantly inhibit the catalytic activity of ATT-Cu nanozyme (Fig. S9A†), analogous to reported CO and CO-like nanozymes.21,26

Interestingly, compared with ATT-Cu, TT-Cu displayed poor CO-mimicking activity (Fig. 4A), as demonstrated by its lower specific activity (Fig. 4B). Moreover, both the Cu content and the surface area of ATT-Cu were lower than those of TT-Cu, manifesting that the Cu content and surface area were not the crucial causes dominating the different CO-like activities between ATT-Cu and TT-Cu. To further compare the difference in intrinsic catalytic activities between ATT-Cu and TT-Cu, kinetic parameters such as substrate affinity $(K_{\rm m})$, catalytic rate constant $(K_{\rm cat})$ and catalytic efficiency (K_{cat}/K_m) were assessed at room temperature under the condition of pH = 8 by varying the initial concentration of 3,5-DTBC. The catalytic reaction rates of both ATT-Cu and TT-Cu nanozymes followed Michaelis-Menten kinetics while performing CO-like catalysis (Fig. 4C), analogous to reported CO-mimicking nanozymes.2,10 According to the Michaelis-Menten equation, K_m , K_{cat} and K_{cat}/K_m were acquired through non-linear curve fitting. ATT-Cu exhibited better substrate affinity, as confirmed by its lower Km value compared with TT-Cu (Fig. 4D). The K_{cat} and $K_{\text{cat}}/K_{\text{m}}$ values of ATT-Cu were obviously superior to those of TT-Cu, demonstrating the better CO-mimicking activity of ATT-Cu (Fig. 4D). After five catalytic cycles, the CO-like activity of the ATT-Cu nanozyme did not notably decrease (Fig. 4E), indicating its good recyclability. Moreover, the corresponding XRD pattern of ATT-Cu did not show a notable difference compared with the original ATT-Cu (Fig. S9B†). Meanwhile, adding interfering ions such as K⁺, Na⁺ or Cl⁻ also did not result in a notable decrease in the catalytic activity of ATT-Cu (Fig. S9C†). Importantly, some of these kinetic parameters presented by this ATT-Cu nanozyme with a low-coordinated CuN2S1 configuration surpassed those of many reported CO-mimicking artificial enzymes/nanozymes with CuN_xO_v moieties^{10,15} as well as those of Ce-based,^{2,27} Zrbased²⁷ or Pt-based² nanozymes (Fig. 4F and Table S2†). This demonstrated that using large-sized and weakly electronegative S as a coordinating atom could endow the Cu center with superior CO-like activity when the N/S coordination numbers were controlled to manipulate the coordination unsaturation of the Cu center to modulate its charge accumulation.

To explore why this low-coordinated CuN₂S₁ configuration with accumulation of more positive charges at the Cu center accelerated the CO-mimicking catalysis of ATT-Cu, we first used electrochemical technology to understand this biomimetic oxidation process. According to the cyclic voltammetry (CV) curve acquired in the absence of O2, it could be seen that the potential for oxidizing 3,5-DTBC over ATT-Cu was lower than that over TT-Cu (Fig. 5A), indicating that accumulation of more positive charges at the Cu center endowed ATT-Cu with easier reception of electrons during the oxidation of 3,5-DTBC. However, the ATT-Cu nanozyme showed a lower redox constant (k_s) determined in the absence of O_2 based on Laviron's method

Edge Article

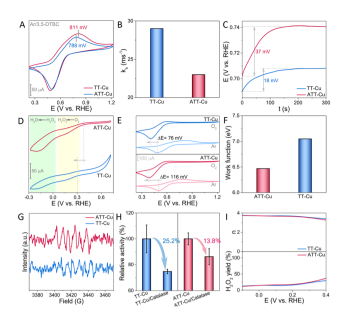


Fig. 5 (A) CV curves of ATT-Cu and TT-Cu in the presence of Ar and 3,5-DTBC; (B) k_s values of ATT-Cu and TT-Cu during oxidation of 3,5-DBTC based on Laviron analysis; (C) changes in OCP over time; (D) CV curves of ATT-Cu and TT-Cu in the presence of O2; (E) CV curves of ATT-Cu and TT-Cu in the presence of 3,5-DTBC as well as O_2 or Ar; (F) work functions of ATT-Cu and TT-Cu; (G) ESR spectra of DMPOtrapped O₂ •-; (H) specific trapping of H₂O₂ during ATT-Cu- and TT-Cu-catalyzed oxidation of 3,5-DTBC using catalase; (I) electron transfer numbers (n) and H₂O₂ yields of ATT-Cu and TT-Cu during

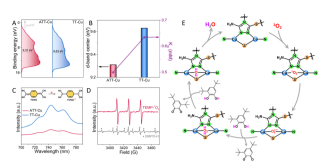
(Fig. 5B and S10†),28 suggesting the slower oxidization rate of 3,5-DBTC over ATT-Cu compared with that over TT-Cu. In fact, the catalytic kinetics experiment disclosed that ATT-Cu exhibited a faster rate for oxidizing 3,5-DBTC compared with TA-Cu while performing CO-like catalysis (Fig. 4C and D). This implied that the process of the oxygen reduction reaction (ORR) should be the key initial step for controlling CO-like catalysis, rather than the oxidation step of 3,5-DBTC.

If the above deduction was reasonable, the preferentially triggered ORR process would make the Cu center of the nanozyme become more electron-deficient, thus elevating the electrochemical potential of the Cu site to accelerate 3,5-DBTC oxidation. Meanwhile, this rising electrochemical potential would also make the nanozyme a weaker reductant, attenuating the ORR rate. Accordingly, the "mixed potential" of reductive and oxidative half-reactions (i.e., the open-circuit potential (OCP))29 would experience a continuous change until the rates of ORR and oxidation of 3,5-DBTC were balanced to attain a steady state, which coincided with Fig. 5C, demonstrating that the ORR process was the key initial step during CO-mimicking catalysis. Furthermore, compared with TT-Cu, ATT-Cu showed a more notable change in OCP when reaching a steady state (Fig. 5C), signifying that the initial ORR step driven by ATT-Cu was faster. Fig. 5D verified this, showing that the onset potential and current of ORR over ATT-Cu exceeded those over TT-Cu. Combining these results and the above structural information indicated that accumulating more positive charges at the lowcoordinated CuN2S1 site contributed to boosting the COmimicking activity of ATT-Cu by accelerating the key initial ORR step.

To clarify the origin of the faster ORR step exhibited by ATT-Cu, we compared the CV curves of the two nanozymes, which were collected in the presence of 3,5-DTBC as well as Ar or O_2 , respectively. We found that introducing O2 into the reaction system resulted in the disappearance of the oxidation peak of 3,5-DTBC, and the reduction peak of 3,5-DTBQ (i.e. the oxidation product of 3,5-DTBC) was negatively shifted (Fig. 5E), which illustrated that the electrochemical oxidation of 3,5-DTBC and the reduction of 3,5-DTBQ became more difficult in the presence of O2, suggesting that O2 may be more easily preferentially adsorbed by both nanozymes to occupy their Cu sites. It is worth noting that after replacing Ar with O2, the 3,5-DTBQ reduction peak presented by ATT-Cu was negatively shifted by 116 mV, which obviously exceeded that shown by TT-Cu (76 mV) (Fig. 5E), reflecting the stronger adsorption of O_2 by ATT-Cu during CO-like catalysis. After O2 was adsorbed, the lower work function of ATT-Cu enabled an easier electron transfer to O₂ (Fig. 5F). These factors ensured the formation of more O₂. species, which were confirmed in the presence of air and 5,5dimethyl-1-pyrroline N-oxide (DMPO) without adding 3,5-DBTC through electron spin resonance (ESR) spectra (Fig. 5G).

As the ORR process progressed, this formed O2. would be further transformed into the product H2O via the formation of H₂O₂ intermediate by further accepting electrons stemming from oxidation of 3,5-DTBC during CO-mimicking catalysis, as shown by the result of a catalase trapping experiment (Fig. 5H) and the evaluated electron transfer number approaching 4 (Fig. 5I). Such a 4e reduction of O₂ to form H₂O via the H₂O₂ intermediate was also confirmed by the CV curve. It displayed two obvious stages corresponding to $O_2 \rightarrow H_2O_2 \rightarrow H_2O$ during the nanozyme-catalyzed ORR process (Fig. 5D). Moreover, according to Fig. 5H, we found that the drop in the relative COlike activity of ATT-Cu was smaller than that of TT-Cu after adding catalase into the reaction system. Given that catalase specifically competed with the nanozyme for H2O2, the smaller decrease in relative CO-like activity demonstrated the stronger adsorption capability of ATT-Cu for H₂O₂. As a result, the stronger capabilities for adsorbing O2 and H2O2 intermediate as well as the faster electron transfer jointly enabled the ATT-Cu nanozyme to exhibit better ORR activity than TT-Cu, as confirmed by its more positive onset potential and larger current for ORR (Fig. 5D). Such an enhanced ORR process contributed to boosting the CO-mimicking activity of ATT-Cu.

To further understand why the ATT-Cu nanozyme with more accumulated positive charges enabled by the low-coordinated CuN₂S₁ configuration enhanced adsorption of O₂ and H₂O₂ intermediates, an ultraviolet photoelectron spectroscopy (UPS) test was conducted to evaluate its d-band center. In general, upshifting the d-band center of a catalyst is beneficial for enhancing the adsorption strength of a catalyst for reactive species. As expected, the d-band center of ATT-Cu was upshifted toward the Fermi level compared to that of TT-Cu (Fig. 6A), contributing to enhanced O2/H2O2 adsorption. Furthermore, this upshifted d-band center prompted the ATT-Cu nanozyme



Chemical Science

Fig. 6 (A) UPS spectra of ATT-Cu and TT-Cu; (B) positive relationship between d-band center and $K_{\rm m}$ of ATT-Cu and TT-Cu; (C) UV-vis spectra of TCNQ solutions in the presence of ATT-Cu and TT-Cu; (D) trapping other reactive oxygen species ('OH and $^{1}O_{2}$) during the biomimetic oxidation of 3,5-DTBC over ATT-Cu; 2,2,6,6-tetramethylpiperidine (TEMP) and 5,5-dimethyl-1-pyrroline N-oxide (DMPO) were used to trap $^{1}O_{2}$ and 'OH, respectively; note: the 'OH signal was inconspicuous, and the DMPO-H adduct and TEMPO were found; (E) proposed catalytic mechanism of ATT-Cu.

to present better affinity for 3,5-DTBC, as shown by its lower $K_{\rm m}$ value (Fig. 6B).

Once 3,5-DTBC was adsorbed, the nanozyme would catalyze the oxidation of 3,5-DTBC by accepting its electron. To compare their capabilities for accepting electrons, 7,7,8,8-tetracyanoquinodimethane (TCNQ) was employed as a probe molecule to mix with the nanozyme in acetonitrile solution. After stirring at 80 °C for 20 min, the centrifuged solution was monitored using UV-vis spectra. A peak at 743 nm was observed (Fig. 6C), suggesting that TCNQ accepted one electron from the nanozyme to form the radical anion product TCNQ*-.30,31 However, this TCNQ* peak in the ATT-Cu system was much weaker than that in the TT-Cu system (Fig. 6C), indicating that the ATT-Cu nanozyme found it harder to donate an electron to TCNQ compared with TT-Cu. In other words, the capability of ATT-Cu to accept an electron was stronger than that of TT-Cu, which agreed well with the lower potential of ATT-Cu for the oxidation of 3,5-DTBC (Fig. 5A). Consequently, the better capability to accept and transfer electrons as well as the stronger adsorption for O2, H2O2 and 3,5-DTBC physically enabled ATT-Cu with a low-coordinated CuN₂S₁ moiety to accelerate CO-like catalysis.

To disclose the possible biomimetic catalytic mechanism of ATT-Cu, we further used electron paramagnetic resonance (EPR) to trap other reactive oxygen species (i.e. 'OH and ${}^{1}O_{2}$) apart from the detected O2. and H2O2. 2,2,6,6-Tetramethylpiperidine (TEMP) and 5,5-dimethyl-1-pyrroline N-oxide (DMPO) were used as trapping reagents for ¹O₂ and ¹OH, respectively. As shown in Fig. 6D, an obvious TEMPO signal was observed while the DMPO-OH signal was inconspicuous, suggesting that ${}^{1}O_{2}$ was involved in the oxygen activation process and 'OH was possibly not formed. On the basis of these detected reactive oxygen species, a possible catalytic mechanism was proposed (Fig. 6E). During catalysis, ATT-Cu preferentially adsorbed and activated O_2 to form 1O_2 , which would be transformed into O_2 . The formed O₂ · would be transformed into H₂O₂ by accepting electrons and hydrogen atoms from 3,5-DTBC, thus generating the oxidation product (i.e., 3,5-DTBQ). The produced H₂O₂ was

further transformed into H_2O by obtaining electrons and hydrogen atoms from 3,5-DTBC to produce H_2O and 3,5-DTBQ. Subsequently, the Cu site of ATT-Cu experienced a regeneration process by desorbing the resulting H_2O .

Conclusions

In summary, we constructed CO-like nanozymes with CuN_2S_1 and CuN_1S_3 moieties by steering the N/S coordination number around Cu centers. We found that the d-band center of ATT-Cu with the low-coordinated CuN_2S_1 configuration was upshifted toward the Fermi level compared with that of TT-Cu with the CuN_1S_3 moiety, leading to stronger adsorption of O_2 and H_2O_2 intermediates as well as better affinity for catechol. Meanwhile, this unsaturated CuN_2S_1 configuration provided ATT-Cu with better capabilities for donating electrons to O_2 and accepting electrons from catechol. These advantages accelerated the initial ORR process, which in turn promoted the oxidation of catechol, thus endowing ATT-Cu with enhanced CO-like activity.

Data availability

The data that supports the findings of this study is available from the corresponding author upon reasonable request.

Author contributions

F. H. conceived and designed the experiments. M. Y. and Z. H. performed the synthesis and/or activity evaluation of nanozyme. N. X. and C. H. provided assistance with the funding. F. H. wrote and revised the manuscript. F. H. and X. H. supervised the project.

Conflicts of interest

The authors declare no competing financial interests.

Acknowledgements

This work was financially supported by Doctoral Fund Project of University of Jinan (XBS2409), the Foundation (No. GZKF202335) of State Key Laboratory of Biobased Material and Green Papermaking (Qilu University of Technology, Shandong Academy of Sciences), National Natural Science Foundation of China (No. 52276195 and No. 22408190), Program for Supporting Innovative Research from Jinan (202228072) and Program of Agricultural Development from Shandong (SD2019NJ015). The authors extend their gratitude to Mr Yanda Du from Shiyanjia Lab (https://www.shiyanjia.com) for providing invaluable assistance with the XPS analysis.

References

1 S. Ji, B. Jiang, H. Hao, Y. Chen, J. Dong, Y. Mao, Z. Zhang, R. Gao, W. Chen and R. Zhang, *Nat. Catal.*, 2021, 4, 407–417.

- Z. Yang, R. Cao, R. Sarangi, T. Sun, X. Yang, X. Zheng, W. Yan, Z. Zhuang, J. Li, W. Chen, D. Wang, J. Zhang and Y. Li, *Nat. Commun.*, 2020, 11, 3049.
- M. Li, J. Chen, W. Wu, Y. Fang and S. Dong, J. Am. Chem. Soc.,
 2020, 142, 15569–15574.
 G. Li, H. Liu, T. Hu, F. Pu, J. Pen and Y. Qu, J. Am. Chem. Soc.
- 3 G. Li, H. Liu, T. Hu, F. Pu, J. Ren and X. Qu, *J. Am. Chem. Soc.*, 2023, **145**, 16835–16842.
- 4 D. Chen, Z. Xia, Z. Guo, W. Gou, J. Zhao, X. Zhou, X. Tan, W. Li, S. Zhao, Z. Tian and Y. Qu, *Nat. Commun.*, 2023, **14**, 7127.
- 5 Y. Xu, Y. Ma, X. Chen, K. Wu, K. Wang, Y. Shen, S. Liu, X. Gao and Y. Zhang, *Angew. Chem., Int. Ed.*, 2024, e202408935.
- 6 K. Wang, Q. Hong, C. Zhu, Y. Xu, W. Li, Y. Wang, W. Chen, X. Gu, X. Chen, Y. Fang, Y. Shen, S. Liu and Y. Zhang, *Nat. Commun.*, 2024, 15, 5705.
- 7 Y. Hu, X. J. Gao, Y. Zhu, F. Muhammad, S. Tan, W. Cao, S. Lin, Z. Jin, X. Gao and H. Wei, *Chem. Mater.*, 2018, 30, 6431–6439.
- 8 F. Cao, H. Feng, J. Yao, X. Hou, T. Jin and J. Hui, *Nano Res.*, 2024, **17**, 6376–6385.
- 9 Y. Chen, B. Jiang, H. Hao, H. Li, C. Qiu, X. Liang, Q. Qu, Z. Zhang, R. Gao, D. Duan, S. Ji, D. Wang and M. Liang, *Angew. Chem., Int. Ed.*, 2023, **62**, e202301879.
- 10 M. Sha, L. Rao, W. Xu, Y. Qin, R. Su, Y. Wu, Q. Fang, H. Wang, X. Cui and L. Zheng, *Nano Lett.*, 2023, 23, 701–709.
- 11 S. Xu, H. Wu, S. Liu, P. Du, H. Wang, H. Yang, W. Xu, S. Chen, L. Song and J. Li, *Nat. Commun.*, 2023, **14**, 4040.
- 12 Y. Lin, F. Wang, J. Yu, X. Zhang and G. P. Lu, *J. Hazard. Mater.*, 2022, **425**, 127763.
- 13 P. Makam, S. S. Yamijala, V. S. Bhadram, L. J. Shimon, B. M. Wong and E. Gazit, *Nat. Commun.*, 2022, **13**, 1505.
- 14 W. Zou, Y. Liu, R. Li and R. Guo, ACS Sustain. Chem. Eng., 2022, 10, 10057–10067.
- 15 S. Thanneeru, N. Milazzo, A. Lopes, Z. Wei, A. M. Angeles-Boza and J. He, J. Am. Chem. Soc., 2019, 141, 4252–4256.
- 16 H. Shang, X. Zhou, J. Dong, A. Li, X. Zhao, Q. Liu, Y. Lin, J. Pei, Z. Li, Z. Jiang, D. Zhou, L. Zheng, Y. Wang, J. Zhou,

- 17 K. Li and D. Xue, J. Phys. Chem. A, 2006, 110, 11332-11337.
- 18 T. Klabunde, C. Eicken, J. C. Sacchettini and B. Krebs, *Nat. Struct. Biol.*, 1998, 5, 1084–1090.
- 19 E. Aznar, S. Ferrer, J. Borrás, F. Lloret, M. Liu-González, H. Rodríguez-Prieto and S. García-Granda, Eur. J. Inorg. Chem., 2006, 5115–5125.
- 20 G.-P. Hao, N. R. Sahraie, Q. Zhang, S. Krause, M. Oschatz, A. Bachmatiuk, P. Strasser and S. Kaskel, *Chem. Commun.*, 2015, 51, 17285–17288.
- 21 M. Yuan, K. Han, H. Yang, L. Mi, C. Huang, X. Hu and F. He, *Small*, 2024, e2401756.
- 22 H. S. Park, R. Sun, E. J. Lee, J. Kim and N. H. Hur, ACS Omega, 2022, 7, 40860–40868.
- 23 Z. Shi, Y. Tao, J. Wu, C. Zhang, H. He, L. Long, Y. Lee, T. Li and Y.-B. Zhang, *J. Am. Chem. Soc.*, 2020, **142**, 2750–2754.
- 24 Q. Wen, Y. Lin, Y. Yang, R. Gao, N. Ouyang, D. Ding, Y. Liu and T. Zhai, *ACS Nano*, 2022, **16**, 9572–9582.
- 25 S. Li, P. Ma, C. Gao, L. Liu, X. Wang, M. Shakouri, R. Chernikov, K. Wang, D. Liu, R. Ma and J. Wang, *Energy Environ. Sci.*, 2022, 15, 3004–3014.
- 26 M. Ü. Ünal, Food Chem., 2007, 100, 909-913.
- 27 S. Liu, Y. He, W. Zhang, T. Fu, L. Wang, Y. Zhang, Y. Xu, H. Sun and H. Zhao, *Small*, 2024, **20**, e2306522.
- 28 P. Ei Phyu Win, J. Yang, S. Ning, X. Huang, G. Fu, Q. Sun, X. H. Xia and J. Wang, *Proc. Natl. Acad. Sci. U. S. A.*, 2024, 121, e2316553121.
- 29 W. C. Howland, J. B. Gerken, S. S. Stahl and Y. Surendranath, J. Am. Chem. Soc., 2022, 144, 11253–11262.
- 30 G. A. Saleh, Talanta, 1998, 46, 111-121.
- 31 J. Song, Z. Ji, Q. Nie and W. Hu, *Nanoscale*, 2014, **6**, 2573–2576.