

Cite this: *Chem. Sci.*, 2024, 15, 7643

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

Ag¹⁺ incorporation via a Zr⁴⁺-anchored metalloligand: fine-tuning catalytic Ag sites in Zr/Ag bimetallic clusters for enhanced eCO₂RR-to-CO activity†

 Liang-Jun Li,^a Wen-Lei Mu,^a Yi-Qi Tian,^a Wei-Dong Yu,^b Lan-Yan Li,^b Jun Yan^a and Chao Liu^{*a}

Attaining meticulous dominion over the binding milieu of catalytic metal sites remains an indispensable pursuit to tailor product selectivity and elevate catalytic activity. By harnessing the distinctive attributes of a Zr⁴⁺-anchored thiacalix[4]arene (TC4A) metalloligand, we have pioneered a methodology for incorporating catalytic Ag¹⁺ sites, resulting in the first Zr–Ag bimetallic cluster, Zr₂Ag₇, which unveils a dualistic configuration embodying twin {ZrAg₃(TC4A)₂} substructures linked by an {AgSal} moiety. This cluster unveils a trinity of discrete Ag sites: a pair ensconced within {ZrAg₃(TC4A)₂} subunits and one located between two units. Expanding the purview, we have also crafted ZrAg₃ and Zr₂Ag₂ clusters, meticulously mimicking the two Ag site environment inherent in the {ZrAg₃(TC4A)₂} monomer. The distinct structural profiles of Zr₂Ag₇, ZrAg₃, and Zr₂Ag₂ provide an exquisite foundation for a precise comparative appraisal of catalytic prowess across three Ag sites intrinsic to Zr₂Ag₇. Remarkably, Zr₂Ag₇ eclipses its counterparts in the electroreduction of CO₂, culminating in a CO faradaic efficiency (FE_{CO}) of 90.23% at –0.9 V. This achievement markedly surpasses the performance metrics of ZrAg₃ (FE_{CO}: 55.45% at –1.0 V) and Zr₂Ag₂ (FE_{CO}: 13.09% at –1.0 V). Utilizing *in situ* ATR-FTIR, we can observe reaction intermediates on the Ag sites. To unveil underlying mechanisms, we employ density functional theory (DFT) calculations to determine changes in free energy accompanying each elementary step throughout the conversion of CO₂ to CO. Our findings reveal the exceptional proficiency of the bridged-Ag site that interconnects paired {ZrAg₃(TC4A)₂} units, skillfully stabilizing *COOH intermediates, surpassing the stabilization efficacy of the other Ag sites located elsewhere. The invaluable insights gleaned from this pioneering endeavor lay a novel course for the design of exceptionally efficient catalysts tailored for CO₂ reduction reactions, emphatically underscoring novel vistas this research unshrouds.

Received 30th December 2023
Accepted 24th March 2024

DOI: 10.1039/d3sc07005k

rsc.li/chemical-science

Introduction

The electrochemical reduction of CO₂ (eCO₂RR) stands as a highly promising pathway for the conversion of CO₂ into valuable chemical fuels.^{1,2} Among the diverse range of potential electrocatalysts, silver (Ag)-based nanomaterials have garnered significant attention due to their remarkable selectivity in producing CO.^{3–6} Despite notable advancements in the

synthesis of monodisperse Ag nanoparticles, accurately characterizing their structural attributes and identifying catalytic Ag sites remains a formidable challenge.^{7,8} This limitation hampers the attainment of a comprehensive understanding of the intricate relationship between structure and activity, consequently impeding the overall progress in this research field. Hence, a pivotal stride toward advancement involves achieving a controlled synthesis of Ag sites' coordination environments. This endeavor would pave way for the creation of meticulously defined catalytic centers. Embracing such a controlled approach holds the potential to unravel the multifaceted interplay between the structure and activity of Ag nanocatalysts, thereby ushering in new avenues for efficient catalysis.^{9–13}

Metal oxides have gained prominence as substrates for stabilizing Ag nanoparticles while exposing catalytic sites, thereby contributing significantly to the realm of Ag

^aHunan Provincial Key Laboratory of Chemical Power Sources, College of Chemistry and Chemical Engineering, Central South University, Changsha 410083, Hunan, P. R. China. E-mail: chaoliu@csu.edu.cn

^bChina College of Science, Hunan University of Technology and Business, Changsha 410000, P. R. China

† Electronic supplementary information (ESI) available: X-ray crystallographic file in CIF format, full experimental and computational details. CCDC 2290841–2290844. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d3sc07005k>



catalysis.^{14–16} Zirconia (ZrO_2) has demonstrated its potential for effectively stabilizing Ag nanoparticles, offering a diverse range of catalytic applications.^{17,18} Nevertheless, the intricate interface and surface composition of Ag– ZrO_2 nanomaterials remain enigmatic, posing challenges in understanding their complex nature. Extensive research endeavors have been directed towards unraveling the structural and molecular intricacies of Ag– ZrO_2 materials, a pursuit that promises valuable insights into binding patterns and atomic-scale electronic configurations. Integrating active catalytic Ag sites into zirconium-oxygen clusters presents a reliable avenue for crafting an Ag– ZrO_2 molecular model, offering the prospect of precisely modulating the coordination environment of Ag. Remarkably, such bimetallic clusters, however, have yet to be synthesized. The synthesis of Zr/Ag bimetallic clusters hinges on a strategically designed ligand system that facilitates the orchestrated assembly of Zr and Ag. This design imperative is pivotal in enabling their cooperative engagement to shape bimetallic clusters. Thiocalix[4]arene (TC4A), a subclass of calixarenes, has exhibited exceptional coordination traits with metals due to its distinctive –OH and –S– functional groups.^{19–22} As such, TC4A emerges as a promising scaffold for constructing nanosized clusters, rendering it a viable candidate for cluster assembly.^{23–30} By leveraging the principles of soft and hard acid-base theory, Zr^{4+} is classified as a hard Lewis acid with a propensity for phenolic oxygen, while Ag^+ is identified as a soft Lewis acid with an affinity for soft bases such as sulfur atoms.^{31–34} By capitalizing on these principles, the interaction of Zr^{4+} and Ag^+ with TC4A holds the tantalizing potential to yield uncharted bimetallic clusters characterized by distinctive geometric and electronic configurations.³⁵

This article pioneers the sequential assembly of Zr/Ag bimetallic nanoclusters with clear Ag catalytic sites, utilizing Zr^{4+} -anchored TC4A as a metalloligand and further explores the applications of these clusters in eCO_2RR . The groundbreaking synthesis of a dimeric Zr_2Ag_7 , $\{\text{HNET}_3\}_2\{\text{H}_2\text{Zr}_2\text{Ag}_7(\text{TC4A})_4(\text{HSal})_3\}$, is reported. This cluster unveils an innovative triad of distinct catalytic Ag^+ sites, including a pair nested within the $\{\text{ZrAg}_3(\text{TC4A})_2\}$ subunits and one situated between the two subunits, each characterized by unique environments, potentially leading to a diverse range of catalytic activities. We employed mass spectrometry to trace the assembly mechanism of Zr_2Ag_7 , successfully isolating two intermediate structures— ZrAg_3 and Zr_2Ag_2 clusters. These structures contain two or one Ag site(s) from the $\{\text{ZrAg}_3(\text{TC4A})_2\}$ subunits, respectively. By comparing the eCO_2RR performance of three clusters, we accurately compared the activity of those three Ag sites. Remarkably, Zr_2Ag_7 displays remarkable catalytic prowess in eCO_2RR , with a CO faradaic efficiency (FE_{CO}) of 90.23% at -0.9 V, surpassing ZrAg_3 (FE_{CO} : 55.45% at -1.0 V) and ZrAg_2 (FE_{CO} : 13.09% at -1.0 V). To delve deeper, the DFT method was employed to calculate the free energy change during CO_2 to CO conversion in different Ag sites. It was demonstrated that the Ag site located on the $\{\text{Ag}(\text{HSal})\}$ moiety could stabilize $^*\text{COOH}$ in eCO_2RR better than those located on the $\{\text{ZrAg}_3(\text{TC4A})_2\}$ unit.

Results and discussion

Synthesis and characterization

In the initial phase of our investigation, crystals of Zr_2Ag_7 were synthesized through a solvothermal reaction involving salicylic acid (H_2Sal), TC4A, $\text{Zr}(\text{OAc})_4$, $\text{Ag}(\text{CF}_3\text{CO}_2)$ and triethylamine. This reaction took place in MeCN at 80°C for a duration of 3 days. The ensuing single-crystal analysis unveiled that Zr_2Ag_7 crystallizes in the monoclinic crystal system with a $P2_1/n$ space group. The structure comprises a cluster core bearing a total of two negative charges, with counter cations provided by two protonated triethylamines. The molar ratio of Zr to Ag in Zr_2Ag_7 , determined through energy-dispersive X-ray spectroscopy (EDS), is approximately 1:3.5, which is consistent with the crystal structure. The high-resolution Ag 3d spectrum exhibits two peaks at 374.08 and 368.08 eV, corresponding to the binding energies of $\text{Ag } 3d^{5/2}$ and $\text{Ag } 3d^{3/2}$, respectively, indicating an oxidation state of +1 for Ag (Fig. S21†). The Zr_2Ag_7 cluster can be dissected into two distinct $\{\text{ZrAg}_3(\text{TC4A})_2\}$ substructures and an $\{\text{Ag}(\text{HSal})\}$ fragment. Within the $\{\text{ZrAg}_3(\text{TC4A})_2\}$ unit, the configuration of Zr^{4+} and three Ag^+ takes on a parallelogram arrangement, capped by two deprotonated TC4A ligands and sheltered by a single HSal^- ligand, as depicted in Fig. 1A. The Zr^{4+} site adopts an octahedral coordination geometry, facilitated by coordinating with four phenoxide groups and two S atoms originating from two TC4A ligands. Additionally, coordination involves a carboxylic O and a hydroxyl O from a Sal^- ligand. The three Ag^+ sites within the $\{\text{ZrAg}_3(\text{TC4A})_2\}$ unit are categorized into two groups. The two Ag(i) sites in close proximity to Zr^{4+} exhibit a hexacoordinate environment, while the Ag(III) site, situated farther away from Zr^{4+} , adopts a tetracoordinate configuration. Each site is coordinated by the phenoxide groups and S atoms. The Ag(III) site within the $\{\text{Ag}(\text{HSal})\}$ fragment also exhibits a tetrahedral coordination structure, stabilized by two S atoms from the two $\{\text{ZrAg}_3(\text{TC4A})_2\}$ units and two O atoms from one HSal^- ligand. However, these two O atoms originate from the carboxyl group and, compared to the chelation mode in $\{\text{Zr}(\text{HSal})\}$, this coordination mode is relatively unstable. The distances involving Ag–O range from 2.297(2) to 2.518(5) Å, while Ag–S distances span from 2.519(4) to 2.493(7) Å. Additionally, the distance between the benzene ring of Sal^- and Ag(III) site is 3.420 Å, indicating relatively weak π -d interactions, which provide additional stability to the Ag(III) site (Fig. 1B). From this perspective, this particular Ag site can be regarded as a cluster-stabilized catalytic site, potentially leading to unexpected catalytic effects (Fig. 1C).

The PXRD pattern of Zr_2Ag_7 crystals exhibits a favorable correspondence with the simulated pattern, providing confirmation of its phase purity (Fig. S11†). Due to intermolecular interactions, crystalline Zr_2Ag_7 samples are insoluble in most solvents but can dissolve in dichloromethane (CH_2Cl_2). To investigate its solution stability, Zr_2Ag_7 crystals were dissolved in CH_2Cl_2 , and the composition was analyzed using ESI-MS in the positive mode. In the ESI-MS spectrum of Zr_2Ag_7 (Fig. 2), the signal corresponding to the complete cluster was not observed;



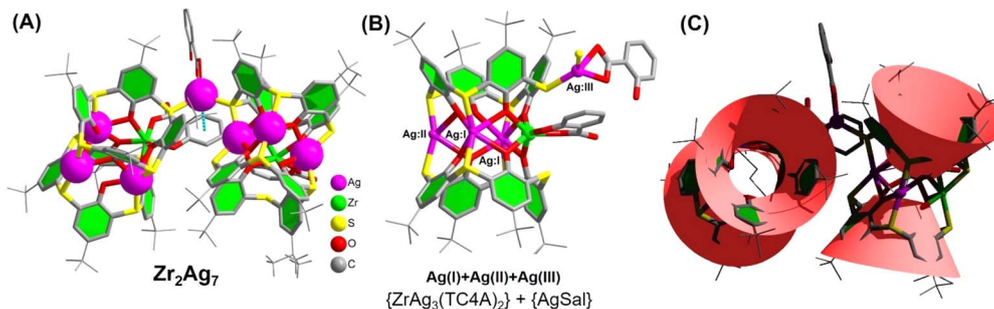


Fig. 1 (A) Molecular structure of the Zr_2Ag_7 cluster; (B) $\{ZrAg_3(TC4A)_2\}$ and $\{AgSal\}$ units; (C) topology structure of the Zr_2Ag_7 cluster.

instead, rich fragmentation information was detected. The signal that appeared at $m/z = 4081.07$ (1a) represents $[H_4Zr_2Ag_7(TC4A)_4(HSal)_2]^+$, resulting from the loss of a Sal^- ligand from the cluster. Zr_2Ag_7 contains three Sal^- ligands, two of which coordinate with Zr^{4+} , while the other coordinates with Ag^+ . The binding strength of this O-containing ligand on the metals follows the order of $Zr^{4+} > Ag^+$, in accordance with the conventional Hard-Soft Acid-Base (HSAB) theory. This suggests that the Sal^{2-} coordinated with Ag^+ in the $Ag(HSal)$ fragment may detach, exposing the $Ag(III)$ active center. Moreover, in the lower m/z range, signals corresponding to the units of $\{ZrAg_3 \pm x(TC4A)_2\}$ (where $x = 0-2$) are observed with a high abundance, indicating an equilibrium between parent clusters and fragments in the solution. The most prominent envelope, centered at $m/z = 1989.57$, can be attributed to the species $[H_3ZrAg_3(TC4A)_2(Sal)]^+$ (calculated $m/z = 1989.50$), corresponding to half of Zr_2Ag_7 . Additionally, signal peaks corresponding to the species $\{H_2ZrAg_4(TC4A)_2(Sal)\}^+$ (1b), $[H_4ZrAg_2(TC4A)_2(Sal)]^+$ (1d), and $[H_4ZrAg(TC4A)_2(Sal)]^+$ (1f) are observed, which are derived from $ZrAg_3$ by either losing a Ag^{1+} or gaining an additional Ag^{1+} ion. The gain and loss of Ag^{1+} from the cluster units indicate that the Zr^{4+} -anchored metalloligand $\{Zr(TC4A)_2\}$ can act as a carrier for Ag^{1+} , with its surface-rich S/O sites providing binding sites.

The Zr_2Ag_7 cluster unveils an innovative triad of distinct Ag^+ sites. To conduct precise comparative assessments of the catalytic activity of those sites, the isolation of structural intermediates assumes paramount importance. Through meticulous ESI-MS analysis, a breakthrough surfaced: the identification of the monomeric $\{ZrAg_3(TC4A)_2\}$ entity and a discovery that

was subsequently subjected to successful crystallization and structural resolution. Synthesized *via* the interaction between TC4A, $Ag(OAc)$, and $Zr(acac)_4$ in CH_3CN , $ZrAg_3$ (Fig. 3B) mirrored the $\{ZrAg_3(TC4A)_2\}$ unit within Zr_2Ag_7 . Notably, this structural analog bore two distinct Ag sites, $Ag(I)$ and $Ag(II)$, a remarkable alignment. Intriguingly, in the absence of $SalH_2$ during the reaction, another monomeric configuration emerged: Zr_2Ag_2 with the composition of $\{Zr_2Ag_2(TC4A)_2(acac)_2\}$ (Fig. 3C). This isomorphous variant, synthesized under solvothermal conditions using $Zr(acac)_4$, TC4A, and $Ag(OAc)$ in $MeCN/DMF$, exhibited a geometric resemblance to $\{ZrAg_3(TC4A)_2\}$. However, it diverged by substituting one Ag^+ and one $Zr(Sal)$ with two $Zr(acac)$, exclusively showcasing the $[AgO_4S_2]$ coordination geometry, solely encapsulating the $Ag(II)$ sites. Moreover, our investigation extended to an intriguing isomorphous entity, Zr_2Na_2 (Fig. 3D), wherein the precision replacement of two Ag^+ with Na^+ showcased an exceptional flexibility within the $\{Zr_2(TC4A)_2\}$ core. Notably, Na^+ and Ag^+ valence similarities rendered this replacement feasible.

From a structural standpoint, the $\{AgSal\}$ site functions as a bridge, connecting two $\{ZrAg_3(TC4A)_2\}$ units and ultimately leading to the formation of the dimeric Zr_2Ag_7 cluster. The pivotal question at this juncture was whether $ZrAg_3$ could undergo further transformation into Zr_2Ag_7 . To validate this hypothesis, we dispersed $ZrAg_3$ crystals in CH_3CN/DCM , introduced $Ag(OAc)$ and H_2Sal to the solution, and conducted the reaction at $80^\circ C$, monitoring the process using ESI-MS. Fig. 4 illustrates the time-dependent ESI-MS analysis of the mother liquor at various intervals during the reaction. In the initial stage, only a peak corresponding to $\{H_2ZrAg_3(TC4A)_2(H_2O)\}^+$

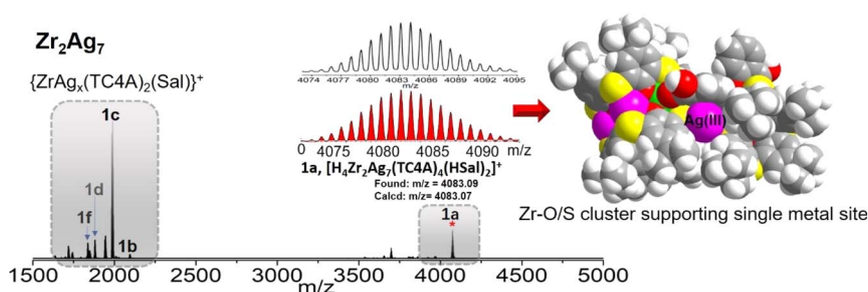


Fig. 2 Positive ion mode ESI-MS of the Zr_2Ag_7 clusters in CH_2Cl_2 .



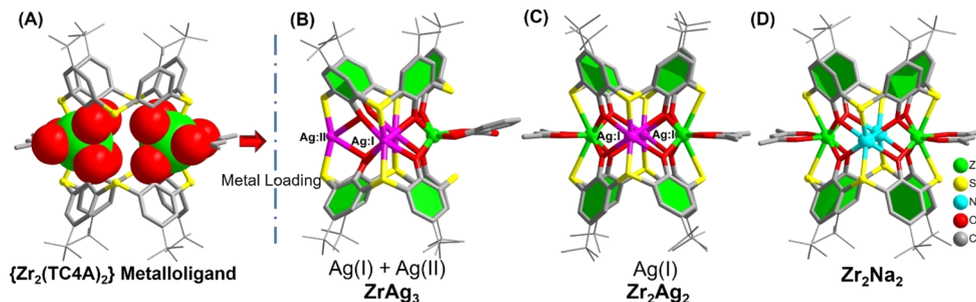


Fig. 3 Molecular structures of the $\{Zr_2(TC4A)_2\}$ metalloligand (A), as well as $ZrAg_3$ (B), Zr_2Ag_2 (C) and Zr_2Na_2 (D) clusters.

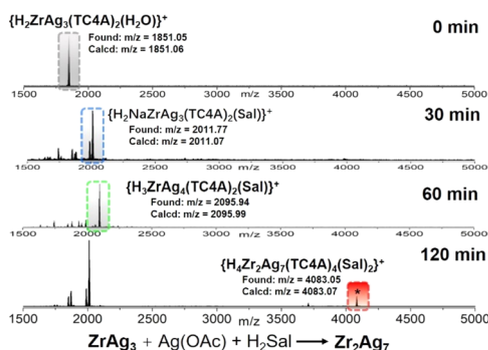


Fig. 4 Time-dependent ESI-MS in the range of m/z 1500–5000 for the reaction of $ZrAg_3$, $AgOAc$ and $SalH_2$ in $MeCN/DCM$ at $100\text{ }^\circ\text{C}$ at 0 min, 30 min, 60 min and 120 min.

(found: $m/z = 1851.05$) was observed in the ESI-MS, indicating the retention of the integrity of the $ZrAg_3$ cluster in solution. However, as the reaction progressed, new signals emerged. Specifically, peaks corresponding to the units of $\{H_4ZrAg_3(TC4A)_2(Sal)\}^+$ ($m/z = 2011.77$) and $\{H_3ZrAg_4(TC4A)_2(Sal)\}^+$ ($m/z = 2095.94$) were detected at 30 min and 60 min, respectively. This signal was formed by the binding of Ag^{1+} to the $\{ZrAg_3(TC4A)_2(Sal)\}$ core, suggesting that $\{ZrAg_3(TC4A)_2(Sal)\}$ could serve as a seed for further growth. At the 120 min mark, the peak corresponding to $\{H_3Zr_2Ag_7(TC4A)_2(Sal)\}^+$ ($m/z = 4083.05$) was observed. These findings clearly demonstrate that Zr_2Ag_7 could be derived from $ZrAg_3$, following a small-to-large assembly pathway. It is worth noting that after two days of reaction, crystals of Zr_2Ag_7 were obtained, as confirmed by single crystal X-ray diffraction.

Electrochemical CO_2 reduction

The role of Ag sites within Zr_2Ag_7 unveils a triadic categorization: one nestled amidst a $\{Ag(HSal)\}$ moiety, and the remaining pair ensconced within two $\{ZrAg_3(TC4A)_2\}$ moieties. Clearly, the catalytic potency of Ag sites is intimately entwined with their coordinating milieu. Undertaking a meticulous comparative analysis of different sites at the molecular level assumes paramount importance to fine-tune the catalyst performance.^{36–47} The structural disparities amongst Zr_2Ag_2 , $ZrAg_3$, and Zr_2Ag_7 offer an opportune platform for discerning CO_2 reduction activity variances across these Ag sites. To ascertain the catalytic

proceeding of Ag, we also introduced Zr_2Na_2 , an isomorph of Zr_2Ag_2 , into our catalytic reactions, aiming to accentuate the superiority of Ag doping. Furthermore, ZrO_2/Ag_2O nanoparticles were synthesized and utilized as electrodes for a comparative analysis.⁴⁸

CO_2 reduction experiments unfolded within a three-electrode H-type electrochemical cell, employing 0.5 M $KHCO_3$ as the electrolyte. Analyzing products entailed a judicious fusion of gas chromatography (GC) and 1H -NMR techniques. Linear sweep voltammetry (LSV) delved into the heart of the matter, unearthing intriguing revelations. Zr_2Ag_7 took the center stage, showcasing a remarkable current density amplification and a more optimistic onset potential within CO_2 -saturated electrolytes, outshining its Ar-purged counterpart—an unmistakable manifestation of its electrocatalytic proficiency (Fig. 5A). $ZrAg_3$ was a close contender, mirroring analogous trends. In stark contrast, the disparity widened significantly upon the entry of Zr_2Ag_2 and Zr_2Na_2 into the competition. In addition, the blank electrode and Ag_2O/ZrO_2 nanoparticles showed no CO_2RR activity (Fig. S29 and S30†). The entire potential spectrum bore witness to the dominance of Zr_2Ag_7 , underscoring its unassailable prominence. It is worth

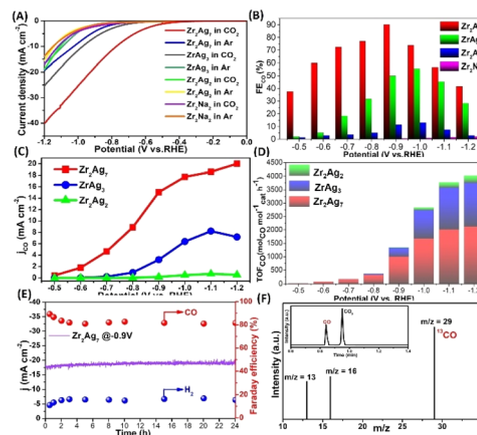


Fig. 5 (A) LSV of samples in Ar or CO_2 saturated 0.5 M $KHCO_3$ solution; (B) FE_{CO} values of Zr_2Ag_7 , $ZrAg_3$, Zr_2Ag_2 and Zr_2Na_2 at different voltages; (C) CO partial current density (j_{CO}); (D) TOF values of Zr_2Ag_7 , $ZrAg_3$ and Zr_2Ag_2 at potentials of -0.5 to -1.2 V vs. RHE; (E) $I-t$ test and FE_{CO} values of Zr_2Ag_7 at -0.9 V in different time periods; (F) GC-MS spectra of ^{13}CO recorded under a $^{13}CO_2$ atmosphere.



nothing that the propensity of Zr_2Ag_7 for more positive onset potentials (-0.49 V vs. RHE) vis-a-vis ZrAg_3 (-0.70 V vs. RHE) and Zr_2Ag_2 (-0.79 V vs. RHE) in CO_2 -saturated electrolytes. While Zr_2Na_2 and Zr_2Ag_2 remained elusive in terms of catalytic activity inference from LSV curves, their participation in electrolysis experiments presented a differential narrative. Analyzing gaseous products solely revealed CO and H_2 for all aforementioned structures, as $^1\text{H-NMR}$ failed to detect any liquid byproducts. Remarkably, Zr_2Ag_7 exhibited an unwavering faradaic efficiency (FE) for CO , culminating in an impressive FE_{CO} of 90.23% at -0.9 V vs. RHE, surpassing the performance of most similar Ag-based CO_2RR catalysts (Table S2[†]). In stark contrast, ZrAg_3 and Zr_2Ag_2 lagged, attaining their peak FE_{CO} of 55.45% and 13.09%, respectively, at -1.0 V vs. RHE, whereas FE_{CO} of Zr_2Na_2 remained insubstantial, almost vanishing across the potential spectrum (Fig. 5B)—an eloquent testament to the preeminence of Ag doping. Fig. 5C paints a revealing picture, juxtaposing the computed CO partial current density (j_{CO}) for Zr_2Ag_7 , ZrAg_3 , and Zr_2Ag_2 . Zr_2Ag_7 emerged triumphant, boasting a j_{CO} of 20.01 mA cm^{-2} at 1.2 V vs. RHE, dwarfing ZrAg_3 (7.19 mA cm^{-2}) and overshadowing Zr_2Ag_2 by a staggering 36-fold margin (0.55 mA cm^{-2}). The CO_2RR turnover frequency calculations further underscored this supremacy, with Zr_2Ag_7 consistently outperforming ZrAg_3 and Zr_2Ag_2 across all potential domains (Fig. 5D). The unwavering consistency across these findings cements the unequivocal excellence of Zr_2Ag_7 in selectively electroreducing CO_2 to CO , eclipsing its catalyst counterparts.

To elucidate factors contributing to the difference in catalytic activity between the catalysts, a Tafel analysis was employed to characterize the reaction kinetics in the electrolyte (Fig. S38[†]). From Zr_2Ag_7 to ZrAg_3 and then to Zr_2Ag_2 , the Tafel slopes of the three species increase sequentially, indicating that the electron transfer energy consumption of the catalyst gradually increases. These findings suggest potential kinetic advantages for Zr_2Ag_7 in eCO_2RR . The electron transfer ability of the catalysts in the electrolyte was further investigated using electrochemical impedance spectroscopy (Fig. S39[†]). The Nyquist curve clearly indicates that Zr_2Ag_7 has a lower charge transfer resistance compared to the other two, indicating good electron transfer ability for Zr_2Ag_7 , which is beneficial for the CO_2RR process. Moreover, electrochemical measurement of the active surface area (ECSA) revealed that Zr_2Ag_7 exhibits a higher density of accessible active sites (Fig. S40[†]). To verify whether Zr_2Ag_7 exhibits a sustained robustness in electrochemical reactions, in a 0.5 hours electrolysis process across the voltage range of -0.6 V to -1.2 V Zr_2Ag_7 demonstrated a stable total current density (Fig. S35[†]). In addition, a consistent current density of about -20 mA and $\text{FE}_{\text{CO}} > 80\%$ were kept for 24 hours when operating in the long-term mode at -0.9 V (Fig. 5E). This excellent performance emphasizes the ultrastable character of Zr_2Ag_7 once again. After the reaction, we recovered the catalyst and conducted ESI-MS measurements. The ESI-MS pattern exhibited a signal corresponding to $[\text{H}_4\text{Zr}_2\text{Ag}_7(\text{TC4A})_4(\text{HSal})_2]^+$, indicating the continued stability of the Zr_2Ag_7 structure (Fig. S41[†]). PXRD and FT-IR revealed that the characteristic signal of the catalyst was preserved after electrolysis (Fig. S42

and S43[†]). Moreover, EDS analysis of the catalyst post-reaction revealed that Zr and Ag elements maintained a consistent 1 : 3.5 ratio (Fig. S47[†]). XPS analysis showed no significant change in the binding energy of Ag in the catalyst after the reaction, indicating the preservation of its coordination environment (Fig. S48[†]). TEM analysis revealed the presence of clusters in the solution as discrete particles, further confirming the stability of the catalyst (Fig. S49[†]). To accurately determine the C source of CO , isotope experiments were conducted using $^{13}\text{CO}_2$ as the C source under similar catalytic conditions. The detection of ^{13}CO ($m/z = 29$) via GC-MS analysis unequivocally confirmed that the generated CO originated from CO_2 (Fig. 5F).

The electrocatalytic reduction of CO_2 using Ag-based materials typically involves three main steps (Fig. 6A).⁴⁹ Initially, CO_2 is adsorbed on the catalyst surface in the form of $^*\text{COOH}$, a process commonly referred to as Proton-Coupled Electron Transfer (PCET) (Step 1). Subsequently, the $^*\text{COOH}$ species acquires H^+ and e^- , converting to $^*\text{CO}$ after releasing H_2O (Step 2). Finally, $^*\text{CO}$ desorbs from the catalyst surface, yielding CO (Step 3). To validate the basic principle of this mechanism, we employed *in situ* electrochemical attenuated total reflection Fourier-transform infrared spectroscopy (ATR-FTIR).^{50–54} This technology monitors the real-time absorption of evanescent waves by catalyst surface substances generated by infrared total reflection. We set the potential range between -0.5 V and -1.5 V (vs. RHE) and compared changes in relevant absorption peaks using ZrAg_3 and Zr_2Ag_7 as electrocatalysts (Fig. 6B and C). Both spectra exhibit similarities, and a distinctive peak near 1698 cm^{-1} is attributed to the stretching of $\text{C}=\text{O}$ in $^*\text{COOH}$, becoming more pronounced with increasing voltage. Additionally, the broad peak in the range of 1346 – 1394 cm^{-1} represents the stretching of $\text{C}-\text{OH}$ in $^*\text{COOH}$, which increases from -0.5 V to -1.5 V. This trend, akin to the intensity change at 1698 cm^{-1} , indicates a significant increase in surface coverage of $^*\text{COOH}$ species with rising voltage. At approximately 2104 cm^{-1} , a weak signal is observed, attributed to the $\text{Ag}-^*\text{CO}$ vibration mode. At lower potentials, the $^*\text{CO}$ band intensity slightly increases, indicating the adsorption of $^*\text{CO}$ on the catalyst surface. However, at higher potentials (above -0.9 V), CO desorption accelerates, leading to a gradual decrease in the 2104 cm^{-1} peak until it becomes undetectable. Notably, the

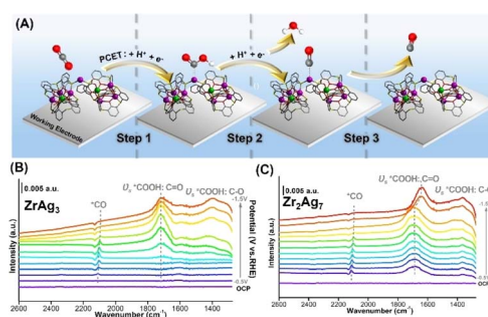


Fig. 6 (A) Schematic depiction of the proposed reaction mechanism of CO_2 reduction to CO on catalysts. (B) and (C) The ATR-FTIR results from 2600 to 1300 cm^{-1} on ZrAg_3 and Zr_2Ag_7 .

intermediate characteristic peak of ZrAg_3 becomes distinctly observable only from an electrode potential of approximately -0.8 V, while the corresponding characteristic peaks of Zr_2Ag_7 are already evident at -0.5 V. This observation indicates that Zr_2Ag_7 demonstrates a higher reaction activity and stronger catalytic proficiency for eCO_2RR , aligning with experimental findings. It should be noted that the position shift of the 1698 cm^{-1} peak in Zr_2Ag_7 may be attributed to changes in the coverage of binding species on the catalyst surface caused by different potentials.

To further comprehend variations in the catalytic performance among the three Ag sites, specific DFT calculations were conducted to investigate the reaction pathways of different Ag sites on the three clusters. Catalyst models were optimized based on the crystal structures of Zr_2Ag_7 , ZrAg_3 , and Zr_2Ag_2 , simplifying $t\text{Bu}$ groups with H atoms to expedite calculation convergence. Additionally, the HSal^- ligand in Zr_2Ag_7 was removed to create the active Ag(III) site. Fig. 7A distinctly illustrates Gibbs free energy changes (ΔG) for each reaction step. For the Ag(I) and Ag(II) sites in $\text{Zr}_2\text{Ag}_7\text{-m}$, CO_2^* captures a proton-electron pair from the electrolyte to form a $^*\text{COOH}$ intermediate with ΔG values of 1.68 and 1.92 eV, respectively. These values are significantly larger than the ΔG of the subsequent step to form $^*\text{CO}$, implying that the formation of $^*\text{COOH}$ is the rate-determining step of the reaction. In contrast, the calculated ΔG value for $^*\text{COOH}$ formation *via* CO_2^* hydrogenation on the Ag(III) site is 0.22 eV, while the ΔG value for the subsequent $^*\text{CO}$ generation step is 0.40 eV. Both values are much lower than those observed on the Ag(I) and Ag(II) sites. Similar conclusions were drawn when calculating the free energy changes for Ag(II)

on $\text{ZrAg}_3\text{-m}$ and Ag(I) on $\text{Zr}_2\text{Ag}_2\text{-m}$, which give larger ΔG values of 1.79 and 2.06 eV for $^*\text{COOH}$ formation, respectively. The calculated free energy of CO_2RR suggests that the Ag(III) sites are more energetically favorable for stabilizing the $^*\text{COOH}$ intermediate compared to the Ag(I) and Ag(II) sites. At the most active Ag(III) site, in addition to the coordination from calixarenes, there is π -d interaction between the benzene ring and Ag site. This unique coordination environment can better disperse the d electron charge of the Ag center, enhancing the ability of the Ag site to stabilize the $^*\text{COOH}$ intermediate and promote the generation of CO.

Conclusions

In summary, our study marks the first systematic comparison of the reactivity of three distinct Ag sites in electrochemical CO_2 reduction, combining experimental and theoretical perspectives through the construction of cluster models. We synthesized an atomically precise bimetallic Zr_2Ag_7 cluster, utilizing a calixarene-protected Zr-O/S core as a substrate for loading Ag^{1+} ions. The Zr_2Ag_7 cluster features three discrete Ag sites: Ag(I) and Ag(II) within the $\{\text{ZrAg}_3(\text{TC4A})_2\}$ subunit and Ag(III) between the two subunits. We meticulously traced the assembly pathway of Zr_2Ag_7 and successfully isolated Ag(II) and Ag(I) sites in ZrAg_3 , as well as Ag(I) sites in Zr_2Ag_2 , providing accurate structural models for a precise comparison of the activity of the three Ag sites. eCO_2RR tests demonstrated that the performance of Zr_2Ag_7 significantly surpassed those of ZrAg_3 and Zr_2Ag_2 . We elucidated the reaction path using *in situ* ATR-FTIR technology and comprehensively calculated the free energy changes for each elementary step of CO_2 conversion to CO at each Ag site using DFT, revealing notable differences in activity among these Ag sites. This study underscores that subtle changes in the coordination geometry of catalytic sites can profoundly influence the catalytic performance. Thus, acquiring atomic structures of nanoclusters is crucial, offering valuable insights for the rational design of cluster structures to achieve efficient catalysis.

Data availability

The data that support the findings of this study are available in the main text and the ESL.†

Author contributions

C. L. and W. D. Yu supervised the project and conceived the idea. L. J. Li carried out synthesis, characterization and catalytic experiment of clusters. L. Y. L. undertook the calculations for this article. C. L. wrote the manuscript. All the authors discussed the experimental results.

Conflicts of interest

There are no conflicts of interest to declare.

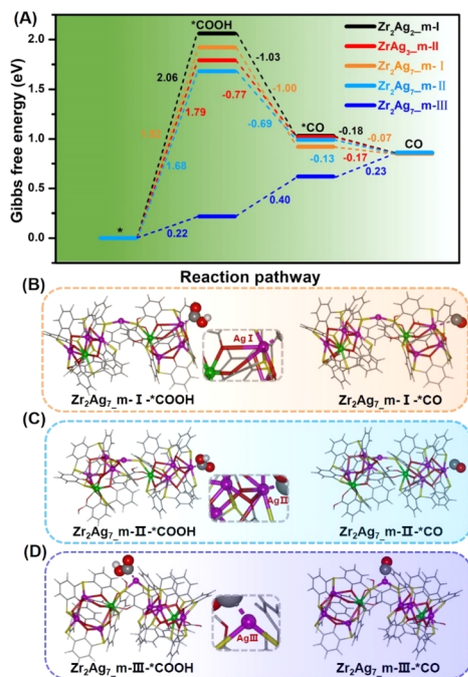


Fig. 7 (A) Free energy diagrams for the CO_2RR pathway of $\text{Zr}_2\text{Ag}_2\text{-m}$, $\text{ZrAg}_3\text{-m}$ and $\text{Zr}_2\text{Ag}_7\text{-m}$. (B), (C) and (D) optimized structural intermediates of $^*\text{COOH}$ and $^*\text{CO}$ for $\text{Zr}_2\text{Ag}_7\text{-m}$.



Acknowledgements

This work was supported by the Natural Science Foundation of Hunan Province (2023JJ30650), Central South University Innovation-Driven Research Programme (2023CXQD061) and Project Foundation of Hunan Provincial Education Department (22B0641).

Notes and references

- 1 P. De Luna, C. Hahn, D. Higgins, S. A. Jaffer, T. F. Jaramillo and E. H. Sargent, *Science*, 2019, **364**, 350–359.
- 2 Y. Z. Xu, C. L. Li, Y. Q. Xiao, C. H. Wu, Y. M. Li, Y. B. Li, J. G. Han, Q. H. Liu and J. F. He, *ACS Appl. Mater. Interfaces*, 2022, **14**, 11567–11574.
- 3 Z. Y. Yang, M. Y. Wan, Z. Y. Gu and F. L. Che, *J. Phys. Chem. C*, 2023, **127**, 17685–17693.
- 4 M. Abdinejad, E. Irtem, A. Farzi, M. Sassenburg, S. Subramanian, H. P. I. V. Montfort, D. Ripepi, M. R. Li, J. Middelkoop, A. Seifitokaldani and T. Burdyny, *ACS Catal.*, 2022, **12**, 7862–7876.
- 5 N. J. Firet and W. A. Smith, *ACS Catal.*, 2017, **7**, 606–612.
- 6 K. Qi, Y. Zhang, J. Li, C. Charmette, M. Ramonda, X. Q. Cui, Y. Wang, Y. P. Zhang, H. I. Wu, W. S. Wang, X. L. Zhang and D. Voiry, *ACS Nano*, 2021, **15**, 7682–7693.
- 7 S. Q. Liu, S. W. Wu, M. R. Gao, M. S. Li, X. Z. Fu and J. L. Luo, *ACS Sustainable Chem. Eng.*, 2019, **7**, 14443–14450.
- 8 S. B. Liu, H. B. Tao, L. Zeng, Q. Liu, Z. H. Xu, Q. X. Liu and J. L. Luo, *J. Am. Chem. Soc.*, 2017, **139**, 2160–2163.
- 9 L. Qin, F. Sun, X. Ma, G. Ma, Y. Tang, L. Wang, Q. Tang, R. Jin and Z. Tang, *Angew. Chem., Int. Ed.*, 2021, **60**, 26136–26141.
- 10 R. Jin, C. Zeng, M. Zhou and Y. Chen, *Chem. Rev.*, 2016, **116**, 10346–10413.
- 11 C. M. Zhao, X. Y. Dai, T. Yao, W. X. Chen, X. Q. Wang, J. Wang, J. Yang, S. Q. Wei, Y. Wu and Y. D. Li, *J. Am. Chem. Soc.*, 2017, **139**(24), 8078–8081.
- 12 J.-D. Yi, X. P. Gao, H. Zhou, W. Chen and Y. Wu, *Angew. Chem., Int. Ed.*, 2022, e202212329.
- 13 X. Q. Wang, Z. Chen, X. Y. Zhao, T. Yao, W. X. Chen, R. You, C. M. Zhao, G. Wu, J. Wang, W. X. Huang, J. L. Yang, X. Hong, S. Q. Wei, Y. Wu and Y. D. Li, *Angew. Chem., Int. Ed.*, 2018, **130**, 1962–1966.
- 14 R. Qi, B. E. Zhu, Z. K. Han and Y. Gao, *ACS Catal.*, 2022, **12**, 8269–8278.
- 15 X. L. Yuan, Y. S. Wu, B. Jiang, Z. S. Wu, Z. X. Tao, X. Lu, J. Liu, T. Qian, H. P. Lin and Q. Zhang, *ACS Appl. Mater. Interfaces*, 2020, **12**, 1664–56649.
- 16 Z. Li, A. A. Haidry, Y. S. Liu, L. C. Sun, L. J. Xie, Q. Fatima and Z. J. Yao, *J. Mater. Sci.: Mater. Electron.*, 2018, **29**, 19219–19227.
- 17 N. Shimoda, S. Umehara, M. Kasahara, T. Hogo, A. Yamazaki and S. Satokawa, *Appl. Catal., A*, 2015, **507**, 56–64.
- 18 X. Wang, Z. Zhao, D. R. Ou, B. F. Tu, D. A. Cui, X. M. Wei and M. J. Cheng, *RSC Adv.*, 2016, **6**, 38153–38158.
- 19 C. Shi, M. Zhang, X. X. Hang, Y. F. Bi, L. L. Huang, K. Zhou, Z. H. Xu and Z. P. Zheng, *Nanoscale*, 2018, **10**, 14448–14454.
- 20 H. T. Han, L. Kan, P. Li, G. S. Zhang, Y. Li, W. P. Liao, Y. L. Liu, W. Chen and C. H. T. Hu, *Sci. China: Chem.*, 2021, **64**, 426–431.
- 21 S. T. Wang, X. H. Gao, X. X. Hang, X. F. Zhu, H. T. Han, W. P. Liao and W. Chen, *J. Am. Chem. Soc.*, 2016, **138**, 16236–16239.
- 22 M. M. Deegan, T. S. Ahmed, G. P. A. Yap and E. D. Bloch, *Chem. Sci.*, 2020, **11**, 5273–5279.
- 23 Z. Wang, L. Li, L. Feng, Z. Y. Gao, C. H. Tung, L. S. Zheng and D. Sun, *Angew. Chem., Int. Ed.*, 2022, **61**, e202200823.
- 24 Z. Wang, H. F. Su, Y. W. Gong, Q. P. Qu, Y. F. Bi, C. H. Tung, D. Sun and L. S. Zheng, *Nat. Commun.*, 2020, **11**, 308.
- 25 Z. Wang, F. Alkan, C. M. Aikens, M. Kurmoo, Z. Y. Zhang, K. P. Song, C. H. Tung and D. Sun, *Angew. Chem., Int. Ed.*, 2022, **61**, e202206742.
- 26 Z. J. Guan, J. L. Zeng, Z. A. Nan, X. K. Wan, Y. M. Lin and Q. M. Wang, *Sci. Adv.*, 2016, **2**, e1600323.
- 27 Z. J. Guan, F. Hu, S. F. Yuan, Z. A. Nan, Y. M. Lin and Q. M. Wang, *Chem. Sci.*, 2019, **10**, 3360–3365.
- 28 C. K. Zhang, Z. Wang, W. D. Si, H. X. Chu, L. Zhou, T. Li, X. Q. Huang, Z. Y. Gao, M. Azam, C. H. Tung, P. Cui and D. Sun, *Nat. Commun.*, 2023, **14**, 6413.
- 29 S. Q. Li, L. F. Dai, Y. Q. Tian, Y. X. Yi, J. Yan and C. Liu, *Chem. Commun.*, 2023, **59**, 575–578.
- 30 L. J. Li, Y. T. Luo, Y. Q. Tian, P. Wang, X. Y. Yi, J. Yan, Y. Pei and C. Liu, *Inorg. Chem.*, 2023, **62**, 14377–14384.
- 31 S. Chen, Z. N. Chen, W. H. Fang, W. Zhuang, L. Zhang and J. Zhang, *Angew. Chem., Int. Ed.*, 2019, **58**, 10932–10935.
- 32 X. Fan, F. Yuan, D. Li, S. Chen, Z. Cheng, Z. Zhang, S. Xiang, S.-Q. Zang, J. Zhang and L. Zhang, *Angew. Chem., Int. Ed.*, 2021, **60**, 12949–12954.
- 33 M. Y. Gao, K. Wang, Y. Y. Sun, D. J. Li, B. Q. Song, Y. H. Andaloussi, M. J. Zaworotko, J. Zhang and L. Zhang, *J. Am. Chem. Soc.*, 2020, **142**, 12784–12790.
- 34 S. Chen, W.-H. Fang, L. Zhang and J. Zhang, *Angew. Chem., Int. Ed.*, 2018, **57**, 11252–11256.
- 35 Y. Q. Tian, W. L. Mu, L. L. Wu, X. Y. Yi, J. Yan and C. Liu, *Chem. Sci.*, 2023, **14**, 10212–10218.
- 36 G. Deng, J. Kim, M. S. Bootharaju, F. Sun, K. Lee, Q. Tang, Y. J. Hwang and T. Hyeon, *J. Am. Chem. Soc.*, 2023, **145**, 3401–3407.
- 37 S. L. Zhuang, D. Chen, L. W. Liao, Y. Zhao, N. Xia, W. Zhang, C. Wang, J. Yang and Z. Wu, *Angew. Chem., Int. Ed.*, 2020, **59**, 3073–3077.
- 38 J. Y. Xu, L. Xiong, X. Cai, S. S. Tang, A. C. Tang, X. Liu, Y. Pei and Y. Zhu, *Chem. Sci.*, 2022, **13**, 2778–2782.
- 39 X. S. Ma, F. Sun, L. B. Qin, Y. G. Liu, X. W. Kang, L. K. Wang, D. E. Jiang, Q. Tang and Z. H. Tang, *Chem. Sci.*, 2022, **13**, 10149–10158.
- 40 J. Wang, F. Xu, Z. Y. Wang, S. Q. Zang and T. C. W. Mak, *Angew. Chem., Int. Ed.*, 2022, **61**, e202207492.
- 41 J. Wang, Z. Y. Wang, S. J. Li, S. Q. Zang and T. C. W. Mak, *Angew. Chem., Int. Ed.*, 2021, **60**, 5959–5964.
- 42 X. Liu, E. D. Wang, M. Zhou, Y. Wan, Y. K. Zhang, H. Q. Liu, Y. Zhao, J. Li, Y. Gao and Y. Zhu, *Angew. Chem., Int. Ed.*, 2022, **61**, e202207685.



- 43 X. Liu, X. Cai and Y. Zhu, *Acc. Chem. Res.*, 2023, **56**, 1528–1538.
- 44 X. Cai, G. J. Li, W. G. Hu and Y. Zhu, *ACS Catal.*, 2022, **12**, 10638–10653.
- 45 Y. Zhang, L. Z. Dong, S. Li, X. Huang, J. N. Chang, J. H. Wang, J. Zhou, S. L. Li and Y. Q. Lan, *Nat. Commun.*, 2021, **12**, 6390.
- 46 R. Wang, J. Liu, L. Z. Dong, J. Zhou, Q. Huang, Y. R. Wang, J. W. Shi and Y. Q. Lan, *CCS Chem.*, 2023, **5**, 2237–2250.
- 47 L. Zhang, X. X. Li, Z. L. Lang, Y. Liu, L. Yuan, W. Y. Lu, Y. S. Xia, L. Z. Dong, D. Q. Yuan and Y. Q. Lan, *J. Am. Chem. Soc.*, 2021, **143**, 3808–3816.
- 48 J. F. Huang and Y. C. Wu, *ACS Sustainable Chem. Eng.*, 2019, **7**, 6352–6359.
- 49 X. Zi, Y. J. Zhou, L. Zhu, Q. Chen, Y. Tan, X. Q. Wang, M. Sayed, E. Pensa, R. A. Geioushy, K. Liu, J. W. Fu, E. Cortes and M. Liu, *Angew. Chem., Int. Ed.*, 2023, **62**, e202309351.
- 50 M. J. Zhuansun, Y. Liu, R. H. Lu, F. Zeng, Z. Y. Xu, Y. Wang, Y. Y. Yang, Z. Y. Wang, G. F. Zheng and Y. H. Wang, *Angew. Chem., Int. Ed.*, 2023, **62**, e202309875.
- 51 A. P. Ayanwale, A. d. J. Ruíz-Baltazar, L. Espinoza-Cristóbal and S. Y. Reyes-López, *Dose-Response*, 2020, **18**, 1–13.
- 52 Z. Z. Liu, X. M. Lv, S. Y. Kong, M. T. Liu, K. H. Liu, J. B. Zhang, B. W. Wu, Q. Zhang, Y. Tang, L. P. Qian, L. J. Zhang and G. F. Zheng, *Angew. Chem., Int. Ed.*, 2023, **62**, e202309319.
- 53 Y. Lin, T. Wang, L. L. Zhang, G. Zhang, L. L. Li, Q. F. Chang, Z. F. Pang, H. Gao, K. Huang, P. Zhang, Z. J. Zhao, C. L. Pei and J. L. Gong, *Nat. Commun.*, 2023, **14**, 3575.
- 54 M. Dunwell, Q. Lu, J. M. Heyes, J. Rosen, J. G. G. Chen, Y. S. Yan, F. Jiao and B. X. Han, *J. Am. Chem. Soc.*, 2017, **139**, 3774–3783.

