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Atomically precise alkyne-protected Ag₁₉Cu₂ nanoclusters: synthesis, structure analysis, and electrocatalytic CO₂ reduction application†

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We report the synthesis, structure analysis, and electrocatalytic CO₂ reduction application of Ag₁₉Cu₂(C≡CAr^F)₁₂(PPh₃)₆Cl₆ (abbreviated as Ag₁₉Cu₂, C≡CAr^F: 3,5-bis(trifluoromethyl)phenylacetylene) nanoclusters. Ag₁₉Cu₂ has characteristic absorbance features and is a superatomic cluster with 2 free valence electrons. Single-crystal X-ray diffraction (SC-XRD) revealed that the metal core of Ag₁₉Cu₂ is composed of an Ag₁₁Cu₂ icosahedron connected by two Ag₄ tetrahedra at the two terminals of the Cu–Ag–Cu axis. Notably, Ag₁₉Cu₂ exhibited excellent catalytic performance in the electrochemical CO₂ reduction reaction (eCO₂RR), manifested by a high CO faradaic efficiency of 95.26% and a large CO current density of 257.2 mA cm⁻² at –1.3 V. In addition, Ag₁₉Cu₂ showed robust long-term stability, with no significant drop in current density and FE_{CO} after 14 h of continuous operation. Density functional theory (DFT) calculations disclosed that the high selectivity of Ag₁₉Cu₂ for CO in the eCO₂RR process is due to the shedding of the –C≡CAr^F ligand from the Ag atom at the very center of the Ag₄ unit, exposing the active site. This study enriches the potpourri of alkyne-protected bimetallic nanoclusters and also highlights the great advantages of using atomically precise metal nanoclusters to probe the atomic-level structure–performance relationship in the catalytic field.

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1. Introduction

Thanks to the alloying effect, bimetallic nanoclusters usually display different physicochemical properties and functionalities from their monometallic counterparts.^{1–5} Specifically, in the catalytic field, upon introducing another metal atom into the core, on the one hand, the geometric configuration of the metal nanocluster is changed and new catalytically active sites

can be generated, and on the other hand, it can modulate the electronic structure of the metal nanocluster hence tuning the adsorption capacities to the reaction substrates and intermediates. For instance, the Jiang and Lee group reported that the Au₂₄Pt(SC₆H₁₃)₁₈ cluster possesses much higher hydrogen evolution reaction (HER) activity than Au₂₅(SC₆H₁₃)₁₈, as the central Pt atom can more easily form an M–H chemical bond to lower the energy barrier.⁶ In another study, Deng *et al.* found that Cu doping can significantly boost the electrochemical CO₂ reduction reaction (eCO₂RR) of atomically precise Au nanoclusters. In a gas diffusion electrode-based membrane electrode assembly cell, the Au₁₅Cu₄ nanocluster exhibited a high CO faradaic efficiency (over 90%), markedly larger than that of the Au₁₈ cluster (~60%). Density functional theory (DFT) calculations disclosed that Cu doping induced catalytic synergistic effects, where the exposed pair of AuCu dual sites can accelerate the eCO₂RR process.⁷ As seen above, atomically precise bimetallic nanoclusters have good catalytic activity and selectivity for the eCO₂RR, and more importantly, the atomically precise structure of bimetallic nanoclusters is conducive to the in-depth analysis of the reaction mechanism and the elucidation of the catalyst's structure–performance relationship at the atomic level.

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To prevent the bimetallic core from aggregation, there are various types of small molecules that can be used as capping agents or ligands for preparing bimetallic nanoclusters.⁸ The ligand molecules include but are not limited to thiolate molecules,⁹ N-containing molecules,¹⁰ carbenes,¹¹ hydrides,¹² alkynyl molecules,^{13–15} and so on. The ligand molecules can not only tune the physical and chemical properties of the metal nanoclusters,¹⁶ but also affect the catalytic performance drastically.^{17–19} The most widely employed ligand is thiolate molecules; however, alkynyl molecules as a protecting ligand have been gaining more and more research attention.^{13–15} In a quite recent study, the Wang and Zhou groups reported the photophysics of $\text{Au}_{22}(\text{BuPhC}\equiv\text{C})_{18}$ and $\text{Au}_{16}\text{Cu}_6(\text{BuPhC}\equiv\text{C})_{18}$, where the latter exhibited >99% photoluminescence quantum yield (PLQY) in de-aerated solution at room temperature with an emission maximum at 720 nm tailing to 950 nm and 61% PLQY in an oxygen-saturated solution.²⁰ This is due to the fact that the Cu doping suppressed the non-radiative decay (~60-fold less) and promoted the inter-system crossing rate. Moreover, in the catalytic regime, alkynyl ligands can perturb the electronic structure of metal nanoclusters by forming σ and/or π bonds, thereby promoting the catalytic performance of some specific reactions.²⁰ In 2017, the Wang group fabricated two isostructural $[\text{Au}_{38}\text{L}_{20}(\text{Ph}_3\text{P})_4]^{2+}$ (L = alkynyl or thiolate) clusters with the ligand being the only variable.²¹ The alkynyl-protected Au_{38} cluster is very active in the semi-hydrogenation of alkynes with the substrate conversion ratio over 97% while the thiolate one possesses negligible activity (less than 2%). In another study, our group disclosed that, in the electrochemical CO_2 reduction reaction (e CO_2 RR), alkynyl-protected Ag_{32} clusters can outcompete the phosphine and thiolate co-protected Ag_{32} clusters in terms of FE_{CO} , as the former has a much lower energy barrier for forming the key intermediate *COOH.²² Similar phenomena have also been documented for alkynyl-protected Au_{25} nanoclusters to catalyze the HER,²³ and alkynyl-protected Au_{28} nanoclusters supported by NiFe-LDH for catalyzing the oxygen evolution reaction.²⁴

So far, significant progress has been achieved on the synthesis, electronic properties, luminescent behaviours, and catalytic applications of alkynyl-protected bimetallic nanoclusters, yet the family or potpourri of alkynyl-protected bimetallic nanoclusters is still limited. For AgCu bimetallic nanoclusters, molecules with atomically precise structures and alkynyl protection only include Ag_9Cu_6 ,²⁵ $\text{Ag}_{20}\text{Cu}_{12}$,²⁶ $\text{Ag}_{15}\text{Cu}_6$,²⁷ $\text{Ag}_{22}\text{Cu}_7$,²⁸ $\text{Ag}_{14}\text{Cu}_2$,²⁹ and so on. Inspired by the above studies, herein, we report the synthesis, structural analysis, and e CO_2 RR application of $\text{Ag}_{19}\text{Cu}_2(\text{C}\equiv\text{CAR}^{\text{F}})_{12}(\text{PPh}_3)_6\text{Cl}_6$ nanoclusters (hereafter referred to as $\text{Ag}_{19}\text{Cu}_2$, $\text{C}\equiv\text{CAR}^{\text{F}}$: 3,5-bis(trifluoromethyl)phenylacetylene). $\text{Ag}_{19}\text{Cu}_2$ has characteristic absorbance features, and its overall structure was analyzed by SC-XRD. $\text{Ag}_{19}\text{Cu}_2$ exhibited excellent catalytic performance, evidenced by a high CO faradaic efficiency (FE_{CO}) of 95.26%, a high current density of 257.2 mA cm^{-2} at -1.3 V, and robust stability with no significant current decay during 14 h of continuous operation.

2. Experimental section

See all the experimental details in the ESI.†

3. Results and discussion

The UV-visible absorption spectrum of $\text{Ag}_{19}\text{Cu}_2$ is shown in Fig. 1a, and two characteristic absorbance peaks at 498 nm and 703 nm can be easily identified. The overall structure of $\text{Ag}_{19}\text{Cu}_2$ was then analyzed by single-crystal X-ray diffraction (Fig. 1b). Specifically, the $\text{Ag}_{19}\text{Cu}_2$ nanocluster is positively charged, with BF_4^- as the counter ion. $\text{Ag}_{19}\text{Cu}_2$ crystallizes in the $P\bar{1}(2)$ space group. $\text{Ag}_{19}\text{Cu}_2$ has 21 metal atoms in the core, which is protected by twelve $-\text{C}\equiv\text{C}-\text{Ar}^{\text{F}}$ ligands, six PPh_3 molecules and six chlorine ions. The calculated free valence electron number is 2, indicating that it is a superatomic cluster with a closed-shell electronic structure. In fact, after being stored at room temperature for 3 months, the absorbance features of the $\text{Ag}_{19}\text{Cu}_2$ clusters remained almost unchanged (Fig. S5†), confirming the excellent chemical stability of $\text{Ag}_{19}\text{Cu}_2$ under ambient conditions.

Subsequently, the metal core configuration of $\text{Ag}_{19}\text{Cu}_2$ was detailedly examined. As illustrated in Fig. 2a, $\text{Ag}_{11}\text{Cu}_2$ forms an icosahedron with two Cu atoms on two diagonal points, and two Ag_4 tetrahedra are connected with two faces of the two opposite sides in the $\text{Ag}_{11}\text{Cu}_2$ icosahedron to form the $\text{Ag}_{19}\text{Cu}_2$ core. Apparently, the $\text{Ag}_{19}\text{Cu}_2$ core is centrosymmetric. Interestingly, the average Ag–Ag bonding length in the $\text{Ag}_{11}\text{Cu}_2$ kernel is 2.9300 Å, slightly longer than that in the Ag_4 tetrahedra (see Fig. S2†). Next, the metal–ligand coordination mode between Ag/Cu and the alkynyl molecule was analyzed and is summarized in Fig. 2b. Basically, there are four types of coordination modes: Motif A, B, C, and D. In Motif A, the alkynyl molecule binds with three Ag atoms with σ bonding and one Ag atom with both σ and π bonding; hence Motif A adopts a $\mu_4-\eta_1, \eta_1, \eta_1, \eta_2$ coordination mode. In Motif B, the alkynyl molecule binds with four Ag atoms with σ bonding; hence the coordination mode is $\mu_4-\eta_1, \eta_1, \eta_1, \eta_1$. In Motif C, the alkynyl molecule binds with two Ag atoms with σ bonding and one Cu atom with both σ and π bonding; hence Motif C adopts a $\mu_3-\eta_1, \eta_1, \eta_2$ coordination mode. Motif D has a

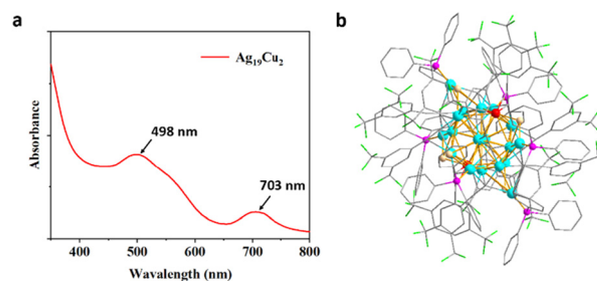


Fig. 1 (a) Absorbance spectra of the $\text{Ag}_{19}\text{Cu}_2$ nanocluster in CH_2Cl_2 . (b) The overall structure of $\text{Ag}_{19}\text{Cu}_2$. Color legend: red, Cu; blue, Ag; green, Cl; purple, P; laurel green, F; white, H; gray, C.

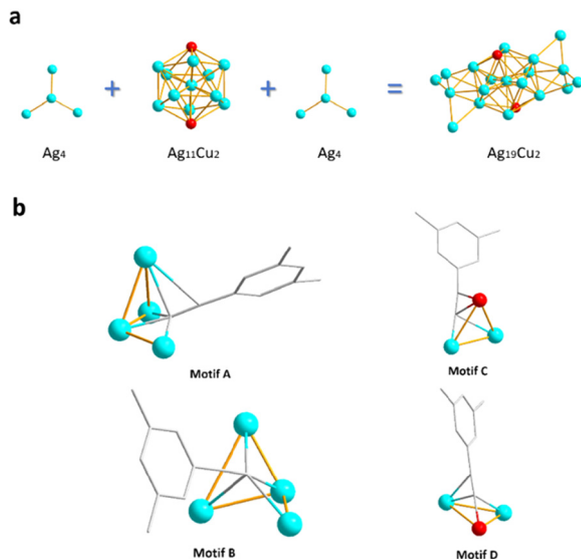


Fig. 2 (a) The structural anatomy of the $\text{Ag}_{19}\text{Cu}_2$ core. (b) Four different coordination modes between Ag/Cu and one alkyne molecule. Color legend: red, Cu; blue, Ag; green, Cl; purple, P; laurel green, F; white, H; gray, C.

similar coordination pattern to Motif C, and the alkyne molecule binding with both σ and π bonding is one Ag atom in Motif D but one Cu atom in Motif C; therefore, Motif D adopts a $\mu\text{-3-}\eta^1, \eta^2, \eta^3$ coordination mode. There are twelve alkyne molecules capping the metal core, and the total number of Motif A, B, C, and D is 4, 4, 3, and 3, respectively. In addition, six PPh_3 ligands are coordinated with six Ag atoms on two sides in the cluster (Fig. S3[†]), and six Cl atoms are also coordinated with $\text{Ag}_{11}\text{Cu}_2$ by σ bonds in the cluster (Fig. S4[†]). Four Cl atoms are connected to three Ag atoms and two Cl atoms are connected to one Cu atom and two Ag atoms. All the detailed structure parameters are summarized in Table S1.[†]

The elemental composition and charge states of $\text{Ag}_{19}\text{Cu}_2$ were subsequently analyzed using X-ray photoelectron spectroscopy (XPS). The survey scan spectra shown in Fig. S6a[†] confirmed the presence of Ag, Cu, C, F, and P atoms. The high resolution XPS spectra of the Ag 3d and Cu 2p electrons in $\text{Ag}_{19}\text{Cu}_2$ can be found in Fig. S6b and S6c,[†] respectively. Notably, the binding energy of the Ag 3d_{5/2} electrons is 368.7 eV, suggesting that the valence state of Ag is between (0) and (I). Such a value is also in good agreement with the previously reported $\text{Ag}_{15}\text{Cu}_6$ ²⁷ and $\text{Ag}_{20}\text{Cu}_{12}$ ²⁶ clusters, whose Ag 3d_{5/2} binding energies are 368.1 eV and 368.84 eV, respectively. Meanwhile, the binding energy of the Cu 2p_{3/2} electrons is 933.34 eV, indicating that the valence state of Cu is (0, 1). The XPS results further validate that the superatomic cluster of $\text{Ag}_{19}\text{Cu}_2$ is successfully obtained.

The $\text{Ag}_{19}\text{Cu}_2$ nanoclusters were then loaded onto carbon nanotubes (CNTs) to investigate the eCO₂RR catalytic performance. As shown in Fig. 3a, in N₂ saturated 1 M KOH, both the CNTs and $\text{Ag}_{19}\text{Cu}_2/\text{CNTs}$ exhibited no activity, while in CO₂ saturated 1 M KOH, both displayed effective activity. At the

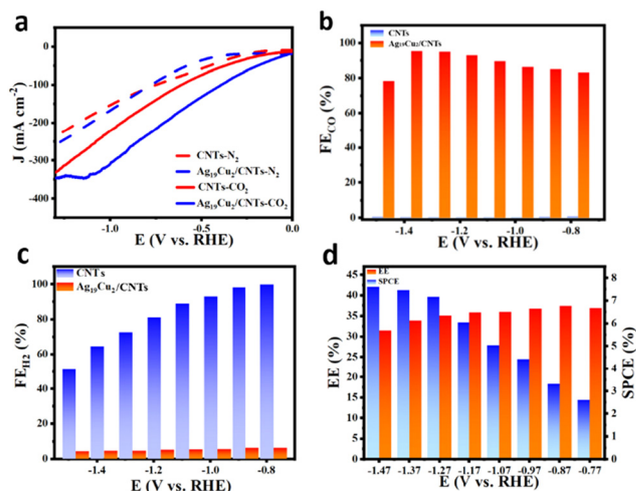


Fig. 3 (a) Polarization curve of CNTs and $\text{Ag}_{19}\text{Cu}_2/\text{CNTs}$ in N₂/CO₂ saturated 1 M KOH. (b) FE_{CO} of CNTs and $\text{Ag}_{19}\text{Cu}_2/\text{CNTs}$. (c) FE_{H₂} of CNTs and $\text{Ag}_{19}\text{Cu}_2/\text{CNTs}$. (d) EE and SPCE of the eCO₂RR.

same applied potential, $\text{Ag}_{19}\text{Cu}_2/\text{CNTs}$ has a much higher current density than CNTs, indicating superior catalytic activity. In the eCO₂RR, the main product is CO, along with H₂ as the side product. Faradaic efficiency (FE) is defined as the amount of product collected relative to the amount that can be produced by the total charge passing through, expressed as a fraction or percentage. In electrochemical reactions, a higher FE means that the actual power of the reaction is closer to the theoretical power and the efficiency of the reaction is higher. The FE_{CO} for $\text{Ag}_{19}\text{Cu}_2/\text{CNTs}$ and CNTs is presented in Fig. 3b. The FE_{CO} for CNTs is nearly zero in the whole tested potential window, but for $\text{Ag}_{19}\text{Cu}_2/\text{CNTs}$, when the applied potential becomes more negative, the FE_{CO} first increases and then decreases, and the highest FE_{CO} is achieved at -1.37 V with a value of 95.26%. When the potential becomes quite negative, the hydrogen evolution reaction starts to dominate, which is the main reason why FE_{CO} decreases. Meanwhile, $\text{Ag}_{19}\text{Cu}_2/\text{CNTs}$ displays a strong capability to suppress hydrogen evolution, as the FE_{H₂} is below 5% in the whole potential window. However, the HER process dominates the reaction for CNTs, as the FE_{H₂} increases from $\sim 51\%$ to nearly 100% (Fig. 3c). To further evaluate the catalytic performance of the $\text{Ag}_{19}\text{Cu}_2/\text{CNT}$ catalyst, we also calculated the energy efficiency (EE), which is defined as the conversion efficiency of the chemical energy from the input electrical energy to the final product, and a higher EE means a higher yield of the reaction product and less energy wasted. From -1.47 V to -0.77 V, the EE first increased and then decreased, reaching a maximal value of 39.76% at an EE of -0.97 V (Fig. 3d). In addition, the single-pass conversion efficiency (SPCE) of the $\text{Ag}_{19}\text{Cu}_2/\text{CNT}$ catalyst increased when the potential changed from -0.77 V to -1.47 V, achieving a maximal value of 7.46% at -1.47 V. Furthermore, long-term stability is another important criterion to further assess the intrinsic catalytic properties of a catalyst. As demonstrated in Fig. S7,[†] after continuous oper-

ation for 14 h, the current density and FE_{CO} of the $Ag_{19}Cu_2/CNT$ catalyst remained almost unchanged, suggesting excellent long-term stability.

The spin-polarized density functional theory (DFT) calculations were subsequently carried out to investigate the electrocatalytic activity and selectivity of $Ag_{19}Cu_2$ nanoclusters by using the Vienna *ab initio* simulation package (VASP5.4.4).³⁰ In order to save the computational cost, we simplified the $-C\equiv CAr^F$ groups to $-C\equiv CR$ ($R = 3,5-C_6H_3F_2$) and the $-PPh_3$ groups to $-P(CH_3)_3$, respectively. The nanoclusters were placed in a cubic box ($26 \text{ \AA} \times 26 \text{ \AA} \times 26 \text{ \AA}$), and their structures were optimized. The interactions of electron exchange–correlation were represented by the Perdew–Burke–Ernzerhof (PBE) functional form of the generalized gradient approximation (GGA).³¹ The projector augmented-wave (PAW) method was utilized to describe the ion–electron interactions,³² and the wave functions of all the computations were extended by a plane-wave cutoff energy of 400 eV. Only the Gamma point was used to sample the Brillouin zone, and the convergence criteria for energy and force were set as 10^{-4} eV and $-0.05 \text{ eV \AA}^{-1}$, respectively. In addition, considering the non-negligible van der Waals interactions between ligands, we used the empirical density functional dispersion (DFT-D3) method.³³

Based on the computational hydrogen electrode (CHE) model,³⁴ the change in Gibbs free energy (ΔG) for each elementary step of the CO_2 reduction reaction and hydrogen evolution reaction can be calculated as follows:

$$\Delta G = \Delta E + \Delta ZPE - T\Delta S$$

where ΔE , ΔZPE , and ΔS represent the changes in the total energy, zero-point energy, and entropy at 298.15 K. The entropy of gas molecules is obtained from the NIST database. The entropy correction at 298.15 K and zero-point energies for CO_2 , CO , and H_2 are presented in Table S2.† For the adsorbed intermediates, only the vibrational entropy calculated from the vibrational frequencies is considered. Furthermore, to address the inaccuracies in the description of CO_2 and CO gas-phase molecules by the PBE functional,^{35,36} we added corrections of -0.34 eV and $+0.10$ eV for CO and CO_2 , respectively.

To further understand the high eCO_2RR activity and selectivity of $Ag_{19}Cu_2$ nanoclusters observed in the experiment, DFT calculations were carried out to identify the catalytically active sites for the CO_2RR . To simplify the calculations, we replaced the $-C\equiv CAr^F$ groups with $-C\equiv CR$ ($R = 3,5-C_6H_3F_2$) groups and the $-P(Ph)_3$ groups with $-P(CH_3)_3$ groups, using the $[Ag_{19}Cu_2(C\equiv CR)_{12}(P(CH_3)_3)_6Cl_6]^+$ cluster as the computational model. Here, we consider all cases where the removal of a single ligand exposes under-coordinated metal active sites. We found that after the removal of the $-Cl$ ligand, the spatial accessibility around the under-coordinated metal site (highlighted by red circles, Fig. S8†) is too small to act as a catalytic site; thus its catalytic activity is negligible. We therefore focused on the catalytic performance of $Ag_{19}Cu_2$ clusters after the removal of a single $-C\equiv CR$ or $-P(CH_3)_3$ group. For the intact $Ag_{19}Cu_2$ cluster, $-P(CH_3)_3$ ligands and part of the $-C\equiv CR$ ligands are attached to the Ag_4 unit of the $Ag_{19}Cu_2$

cluster, and the remaining $-C\equiv CR$ ligands are connected with the $Ag_{11}Cu_2$ unit of the $Ag_{19}Cu_2$ cluster. We first consider the case of the removal of a single $-P(CH_3)_3$ ligand from the Ag_4 unit, as predicted by the free energy profile shown in Fig. 4a, where the formation of $*COOH$ is the potential-limiting step (PDS) of the CO_2RR with a high reaction energy up to 1.82 eV. With the removal of a $-C\equiv CR$ ligand from the $Ag_{11}Cu_2$ unit to form $Ag_{19}Cu_2(C\equiv CR)_{11}(P(CH_3)_3)_6Cl_6^{+(-1)}$ clusters, the formation of the $*COOH$ intermediate remains the most uphill step with a relatively high energy barrier of 1.53 eV. It is noteworthy that when a $-C\equiv CR$ ligand is removed from the Ag_4 unit, the reaction energy for $COOH^*$ formation (PDS) on the $Ag_{19}Cu_2(C\equiv CR)_{11}(P(CH_3)_3)_6Cl_6^{+(-2)}$ cluster is lower (1.09 eV). In comparison, the $Ag_{19}Cu_2(C\equiv CR)_{11}(P(CH_3)_3)_6Cl_6^{+(-2)}$ cluster has a better catalytic activity for the CO_2RR . The optimized structures of all substrates and intermediates are depicted in Fig. S9 and S10† and Fig. 4c. We found that for the $Ag_{19}Cu_2(C\equiv CR)_{11}(P(CH_3)_3)_6Cl_6^{+(-2)}$ cluster, the under-coordinated central Ag atom in the Ag_4 unit prefers to act as a catalytically active site for the CO_2RR , with both $COOH^*$ and CO^* adsorbed to the central Ag atom.

Furthermore, we analyzed the competitive HER process on the surface of $[Ag_{19}Cu_2(C\equiv CR)_{12}(P(CH_3)_3)_5Cl_6]^+$, $[Ag_{19}Cu_2(C\equiv CR)_{11}(P(CH_3)_3)_6Cl_6]^{+(-1)}$ and (2), and the free energy diagrams are shown in Fig. 4b. Our calculations show that the hydrogen-adsorption step is the potential-determining step (PDS) for the HER on the three NCs, with limiting potentials calculated to be 1.02, 0.54 and 1.24 eV respectively. All schematic presentations of the adsorption structures of intermediates are shown in Fig. S11.† Compared with the result of the electroreduction of CO_2 to CO , we found a contrasting product selectivity of $Ag_{19}Cu_2$ clusters with the removal of different ligands. $Ag_{19}Cu_2$ clusters after the removal of a single $-P(CH_3)_3$ group from the Ag_4 unit or a single $-C\equiv CR$ group from the $Ag_{11}Cu_2$ unit are more favourable to undergo the HER process. However, the limiting potential of the HER (1.24

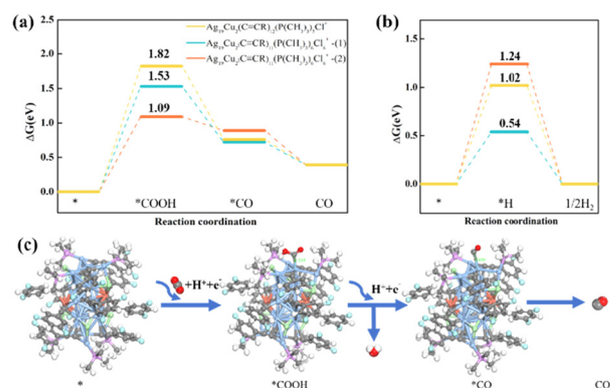


Fig. 4 Free energy diagrams (ΔG) for CO_2 reduction (a) and hydrogen evolution (b) on $Ag_{19}Cu_2$ clusters with different ligands removed at zero potential. (c) Schematic presentation of the electrocatalytic CO_2 reduction process on the $[Ag_{19}Cu_2(C\equiv CR)_{11}(P(CH_3)_3)_6Cl_6]^{+(-2)}$ cluster. Colour legend: Ag, baby blue; Cu, dark orange; C, grey; Cl, light green; F, cyan; P, purple; H, white; O, red.

eV) significantly exceeds that of the CO₂RR (1.09 eV) on the [Ag₁₉Cu₂(C≡CR)₁₁(P(CH₃)₃)₆Cl₆]⁺(-2) cluster, fully supporting the experimentally observed high CO selectivity. Overall, the DFT results suggest that the high CO selectivity of Ag₁₉Cu₂ clusters is probably due to the removal of the -C≡CR group from the Ag₄ unit, exposing the under-coordinated Ag site as a catalytically active centre to promote electrochemical CO₂ reduction.

4. Conclusions

In summary, we successfully synthesized an atomically precise nanocluster of Ag₁₉Cu₂(C≡CAr^F)₁₂(PPh₃)₆Cl₆, which possesses two valence electrons and characteristic absorbance features. The overall structure of Ag₁₉Cu₂ was analyzed by SC-XRD, and it has a novel Ag₁₉Cu₂ core composed of a Ag₁₁Cu₂ kernel connected by two Ag₄ units and various metal-ligand binding coordination modes. When loading into carbon nanotubes, it exhibited excellent catalytic performance, manifested by a high FE_{CO} value of 95.26%, a large CO current density of 257.2 mA cm⁻² at -1.3 V, and robust long-term stability for a continuous 14 h test. DFT calculations revealed that the high CO selectivity is due to the shedding of the -C≡CR group from the Ag₄ unit during the catalytic process, which exposes the undercoordinated Ag atom as the catalytically active site. This study enriches the family of alkynyl-protected bimetallic nanoclusters, and also provides atomic-level mechanistic insights into employing bimetallic nanoclusters as catalysts for complex electrochemical reactions.

Data availability

All data supporting the findings of this study are available within the paper and its ESI.†

Conflicts of interest

There are no conflicts to declare.

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References

- X. Kang, Y. Li, M. Zhu and R. Jin, *Chem. Soc. Rev.*, 2020, **49**, 6443–6514.
- I. Chakraborty and T. Pradeep, *Chem. Rev.*, 2017, **117**, 8208–8271.
- X.-M. Luo, Y.-K. Li, X.-Y. Dong and S.-Q. Zang, *Chem. Soc. Rev.*, 2023, **52**, 383–444.
- A. Ghosh, O. F. Mohammed and O. M. Bakr, *Acc. Chem. Res.*, 2018, **51**, 3094–3103.
- C. Sun, B. K. Teo, C. Deng, J. Lin, G.-G. Luo, C.-H. Tung and D. Sun, *Coord. Chem. Rev.*, 2021, **427**, 213576.
- K. Kwak, W. Choi, Q. Tang, M. Kim, Y. Lee, D.-e. Jiang and D. Lee, *Nat. Commun.*, 2017, **8**, 14723.
- G. Deng, H. Yun, M. S. Bootharaju, F. Sun, K. Lee, X. Liu, S. Yoo, Q. Tang, Y. J. Hwang and T. Hyeon, *J. Am. Chem. Soc.*, 2023, **145**, 27407–27414.
- W. Jing, H. Shen, R. Qin, Q. Wu, K. Liu and N. Zheng, *Chem. Rev.*, 2022, **123**, 5948–6002.
- Y. Li, M. Zhou and R. Jin, *Adv. Mater.*, 2021, **33**, 2006591.
- S. F. Yuan, W. D. Liu, C. Y. Liu, Z. J. Guan and Q. M. Wang, *Chem. – Eur. J.*, 2022, **28**, e202104445.
- H. Shen, G. Tian, Z. Xu, L. Wang, Q. Wu, Y. Zhang, B. K. Teo and N. Zheng, *Coord. Chem. Rev.*, 2022, **458**, 214425.
- Y. Lv, T. Jiang, Q. Zhang, H. Yu and M. Zhu, *Polvometalates*, 2024, **3**, 9140050.
- Z. Lei, X.-K. Wan, S.-F. Yuan, Z.-J. Guan and Q.-M. Wang, *Acc. Chem. Res.*, 2018, **51**, 2465–2474.
- X. Ma, Y. Tang, G. Ma, L. Qin and Z. Tang, *Nanoscale*, 2021, **13**, 602–614.
- L. Chen, L. Wang, Q. Shen, Y. Liu and Z. Tang, *Mater. Chem. Front.*, 2023, **7**, 1482–1495.
- Y. Wang, Z. Liu, A. Mazumder, C. G. Gianopoulos, K. Kirschbaum, L. A. Peteanu and R. Jin, *J. Am. Chem. Soc.*, 2023, **145**, 26328–26338.
- Q. J. Wu, D. H. Si, P. P. Sun, Y. L. Dong, S. Zheng, Q. Chen, S. H. Ye, D. Sun, R. Cao and Y. B. Huang, *Angew. Chem., Int. Ed.*, 2023, **62**, e202306822.
- H. Shan, J. Shi, T. Chen, Y. Cao, Q. Yao, H. An, Z. Yang, Z. Wu, Z. Jiang and J. Xie, *ACS Nano*, 2023, **17**, 2368–2377.
- Z. Liu, H. Tan, B. Li, Z. Hu, D.-e. Jiang, Q. Yao, L. Wang and J. Xie, *Nat. Commun.*, 2023, **14**, 3374.
- W.-Q. Shi, L. Zeng, R.-L. He, X.-S. Han, Z.-J. Guan, M. Zhou and Q.-M. Wang, *Science*, 2024, **383**, 326–330.
- X.-K. Wan, J.-Q. Wang, Z.-A. Nan and Q.-M. Wang, *Sci. Adv.*, 2017, **3**, e1701823.
- L. Chen, F. Sun, Q. Shen, L. Qin, Y. Liu, L. Qiao, Q. Tang, L. Wang and Z. Tang, *Nano Res.*, 2022, **15**, 8908–8913.
- X. Li, S. Takano and T. Tsukuda, *J. Phys. Chem. C*, 2021, **125**, 23226–23230.
- Q.-L. Shen, L.-Y. Shen, L.-Y. Chen, L.-B. Qin, Y.-G. Liu, N. M. Bedford, F. Ciucci and Z.-H. Tang, *Rare Met.*, 2023, **42**, 4029–4038.
- X. Ma, F. Sun, L. Qin, Y. Liu, X. Kang, L. Wang, D.-e. Jiang, Q. Tang and Z. Tang, *Chem. Sci.*, 2022, **13**, 10149–10158.
- G. Ma, F. Sun, L. Qiao, Q. Shen, L. Wang, Q. Tang and Z. Tang, *Nano Res.*, 2023, **16**, 10867–10872.

- 27 G. Deng, J. Kim, M. S. Bootharaju, F. Sun, K. Lee, Q. Tang, Y. J. Hwang and T. Hyeon, *J. Am. Chem. Soc.*, 2022, **145**, 3401–3407.
- 28 G. Deng, K. Lee, H. Deng, S. Malola, M. S. Bootharaju, H. Häkkinen, N. Zheng and T. Hyeon, *Angew. Chem., Int. Ed.*, 2023, **62**, e202217483.
- 29 Q. Shen, X. Cong, L. Chen, L. Wang, Y. Liu, L. Wang and Z. Tang, *Dalton Trans.*, 2023, **52**, 16812–16818.
- 30 G. Kresse and J. Furthmüller, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1996, **54**, 11169–11186.
- 31 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1997, **78**, 1396–1396.
- 32 I. L. Garzón and A. Posada-Amarillas, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1996, **54**, 11796–11802.
- 33 J. K. Nørskov, J. Rossmeisl, A. Logadottir, L. Lindqvist, J. R. Kitchin, T. Bligaard and H. Jónsson, *J. Phys. Chem. B*, 2004, **108**, 17886–17892.
- 34 A. A. Peterson, F. Abild-Pedersen, F. Studt, J. Rossmeisl and J. K. Nørskov, *Energy Environ. Sci.*, 2010, **3**, 1311–1315.
- 35 P. E. Blöchl, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1994, **50**, 17953–17979.
- 36 Q. Tang, Y. Lee, D.-Y. Li, W. Choi, C. W. Liu, D. Lee and D.-e. Jiang, *J. Am. Chem. Soc.*, 2017, **139**, 9728–9736.