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Ligand-passivated Au/Cu nanoclusters with uncoordinated sites give reaction turnover numbers of up to $4 \times 10^4 \, \dagger$

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Ligand-passivated metal nanoclusters with both high stability and catalytic activity have been pursued for a long time to shed light on the underlying rationale of ligand-functionalized metal nanocatalysts in chemical transformations. Nevertheless, these atomically precise nanocatalysts often exhibit low catalytic activity due to the extensive coverage of active sites. In this work, we report a robust gold-copper alloy nanocluster with 74 nuclei with high catalytic performance. The nanocluster with molecular composition of $Au_{41}Cu_{33}(RS)_{40}(MeCN)$ (RSH is 4-fluorothiophenol) has been obtained in a simple way. The cluster contains 34 free electrons, making it a superatom that exhibits high thermal stability as envisioned. Surprisingly, rich uncoordinated copper sites are present on the surface of the cluster, leading to its extremely high catalytic activity (up to 39 269 TON) in C-O coupling reactions with a broad substrate scope.

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Introduction

Ligands have been routinely used in the synthesis of metal nanocatalysts for controlling their shape, structure, and catalytic performance.¹⁻⁴ The role of ligands when synthesizing metal nanocatalysts, however, has remained a topic of debate for a long time: although several recent studies have demonstrated their promotional effects on catalytic activity, most consider them as poisons.⁴⁻⁶ It is the shortage of molecular characterization techniques for nanocatalysts that makes it more challenging to decipher how they work in the presence

of extensive ligand coverage.⁷⁻⁹ As a bridge between molecules and nanoparticles, atomically precise metal nanoclusters have attracted increasing interest in fundamental research in the field of catalysis due to their high degree of monodispersity, enabling the determination of their crystal structure.^{10–23} In this regard, they are proposed to be model systems to rationalize the underlying reaction mechanism of metal nanocatalysts in terms of activity, selectivity, and stability.^{24–28}

The past decades have witnessed an explosive increase in employing ligand-stabilized atomically precise metal nanoclusters as catalysts to drive a wide range of chemical reactions.²⁹⁻⁴¹ In catalysis, the surface ligands of metal nanoclusters are often removed by suitable thermal treatment or in some cases by automatic migration to the supports to activate the substrates. 42,43 This, however, raises another interesting doubt whether the integrity or structure of metal nanoclusters remains intact after their treatment. 44,45 To our delight, recent reports have claimed that clusters protected by bulky and rigid ligands would exhibit catalytic ability without any pre-treatment. For example, the group of Zheng demonstrated that the surface coordination of bulky N-heterocyclic carbene endowed a family of gold nanoclusters including Au13, Au25 and Au44 with high catalytic activity in the activation of alkyne compounds. 46-48 Moreover, Wang and co-workers revealed that the Au₂₃ cluster protected by rigid dipyridylamido and phosphine ligands exhibited high catalytic activity in the aerobic oxidation of benzyl alcohol. 49 The MAu₈ (M = Pd or Pt) series of clusters also displayed excellent catalytic hydrogenation

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ability owing to the introduction of bulky ligands, such as triphenylphosphine. 50,51 In general cases, the high activity of those clusters protected by bulky or rigid ligands relies on the presence of uncoordinated metal sites created by the repulsion between the surface ligands. Therefore, two more questions arise when employing metal nanoclusters as catalysts: 1. whether or not clusters featuring uncoordinated metal sites can be gained in the absence of bulky or rigid ligands? 2. How active can ligand-passivated metal nanoclusters be with limited numbers of exposed sites?

Herein, we report a rare example of a ligand-protected metal nanocluster with both robustness and surprisingly high catalytic activity. The new gold-copper nanocluster with 74 nuclei, with the molecular formula of Au₄₁Cu₃₃(RS)₄₀(MeCN) (labeled as Au₄₁Cu₃₃ hereafter, RSH is 4-fluorothiophenol), has been selectively obtained by a two-step reduction strategy. The flexible thiolate ligands stabilize the metal core in a tight and diverse manner, endowing it with superatomic characteristics of 34 free electrons and high stability. What is interesting to note is that some surface-uncoordinated copper sites are available within the cluster. As a result, it exhibits an extremely high catalytic activity (up to 39269 TON) in catalyzing Ullmann coupling reactions with a broad substrate scope without any pre-treatment.

Results and discussion

The Au₄₁Cu₃₃ cluster was prepared by a two-step strategy (Fig. S1, see details in the ESI†). The first step involved the NaBH₄-mediated reduction of Cu(MeCN)₄BF₄ in the presence of an appropriate amount of 4-fluorobenzenethiol, affording a brown solution (Fig. S2†). It is proposed that copper hydride species are produced in this step. 35,52 To the solution, RSAu was then added in solid form. During the aging period, the solid dissolved gradually, causing the color of the solution to change from brown to black (Fig. S2†). Layering hexane upon the black solution afforded black crystals as final products with a yield of 26.6% (based on Au, Fig. S3†). We note that the product obtained by the two-step synthetic strategy differs from that obtained by the one-step reduction of the mixture containing Cu and Au precursors, as the two strategies represent distinct reduction kinetics during the synthesis.

We first performed X-ray diffraction to determine the structure of the crystalline products (Fig. S4†). The analysis revealed that Au₄₁Cu₃₃ crystallizes in the triclinic system of the Pa3 space group (Table S1†), with two independent clusters in each unit cell (Fig. S5†). Each cluster comprises 41 Au atoms, 33 Cu atoms, 40 thiolate ligands and one acetonitrile ligand, giving rise to a large cluster of $Au_{41}Cu_{33}(RS)_{40}(MeCN) \sim 2.4$ nm in size (Fig. S6†). The absence of counterions in the lattice suggests the neutral state of the cluster. 53,54 The overall structure of Au₄₁Cu₃₃, shown in Fig. 1, resembles a three-necked flask in which the C_2 axis is present across the center (Fig. 1 and S7†).

The cluster adopts a core-shell architecture. As shown in Fig. 2a, the metal core of the cluster can be anatomized as a

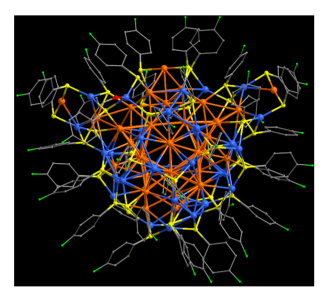


Fig. 1 Total structure of Au₄₁Cu₃₃(RS)₄₀(MeCN). Color legend: orange spheres, Au; blue spheres, Cu; yellow spheres, S; turquoise spheres, F; and gray spheres, C. All hydrogen atoms are omitted for clarity.

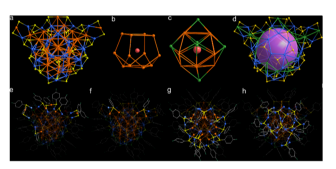


Fig. 2 Detailed structure analysis of the metal framework and coordination modes of 4-fluorothiophenol on Au₄₁Cu₃₃(RS)₄₀(MeCN). (a) Layers of nested Russian nesting structures, (b) icosahedral Au₁₃ coreshell structure, (c) core-shell-shell structure of the Au@Au₁₂@Au₆ skeleton, and (d) outermost hybrid Au–Cu–RS cage. (e) $\mu_2 - \eta^1$ (Cu), η^1 (Au) coordination mode, (f) $\mu_3 - \eta^1(Cu)$, $\eta^1(Au)$, $\eta^1(Au)$ coordination mode, (g) $\mu_3 - \eta^1(Cu)$, $\eta^1(Cu)$, $\eta^1(Au)$ coordination mode, and (h) $\mu_3 - \eta^1(Cu)$, $\eta^1(Cu)$, $\eta^{1}(Cu)$ coordination mode. Color legend: orange spheres, Au; blue spheres, Cu; yellow spheres, S; turquoise spheres, F; and gray spheres, C. All hydrogen atoms are omitted for clarity. For a clearer visual representation, atoms of different shells in the diagram are shown in different colours (magenta Au in the core, orange Au in shell 1, green Au in shell 2, and blue Au and green Cu in shell 3, respectively).

concentric Russian structure consisting of Au@Au₁₂@Au₆@ Au₁₁Cu₃₃. The central part of the cluster core consists of an Au atom, which is surrounded by a distorted Au₁₂ cage that forms a core-shell structure of Au₁₃ (Fig. 2b). Interestingly, the Au₁₂ cage is composed of interlaced pentagonal and triangular windows on the sides and quadrilateral ones on the top and bottom. The top window size in the Au_{12} cage $(4.29 \times 4.39 \text{ Å})$ is slightly larger than the bottom window size $(4.20 \times 4.37 \text{ Å})$. The Au₆ shell is further wrapped around the periphery of the Au₁₂ cage. Of note, each Au atom of the Au₆ shell is located

outside the pentagonal and quadrilateral windows, adopting an octahedral structure for the Au₆ shell (Fig. 2c). The Au@Au₁₂@Au₆ skeleton is further encapsulated by a hybrid Au-Cu-RS cage that forms the Russian nesting structure of Au₄₁Cu₃₃ (Fig. 2d). Careful dissection of the structure reveals that, in the Au-centered icosahedral core, the average distance between the Au center and the surrounding Au₁₂ shell is 3.1093 Å and in the Au₁₂ shell is 2.9150 Å. These values are slightly larger than the distances in the Au@Au12 units of other nanoclusters, e.g. $Au_{25}(SCH_2CH_2Ph)_{18}$, $[Au_{13}(NHC^{Bn})_9Cl_3]^{2+}$ and $[Au_{19}(PR_3)_3(C \equiv CR)_9]^{2+}$, indicating weak interactions.^{55–57} The Au₆ octahedron caps the Au₀Au₁₂ core tightly, as suggested by their short Au-Au bond lengths (average 2.8065 Å). The internal cavity of the Au-Cu-RS cage can be approximated as a sphere with a cavity size of about 4.6 Å, which firmly confines the gold cluster skeleton of Au@Au12@Au6 and thus maintains the stability of its overall structure.

The forty 4-fluorothiophenol ligands are attached to the Au₁₁Cu₃₃ shell to construct the complete structure of the cluster. As for the thiolate ligands on the surface, they can be divided into four types: nine thiolates are coordinated in the $\mu_2 - \eta^1(Cu)$, $\eta^1(Au)$ mode (Fig. 2e), two in $\mu_3 - \eta^1(Cu)$, $\eta^1(Au)$, $\eta^1(Au)$ (Fig. 2f), sixteen in $\mu_3-\eta^1(Cu)$, $\eta^1(Cu)$, $\eta^1(Au)$ (Fig. 2g) and thirteen in μ_3 - $\eta^1(Cu)$, $\eta^1(Cu)$, $\eta^1(Cu)$ (Fig. 2h). The diverse coordination patterns of thiolates with the metal atoms of the cluster lead to its rich surface local structures: either in staple motifs or three-dimensional organometallic oligomers. The binding between the thiolates and the metal atoms is strong, as suggested by their short bond distances (2.277-2.564 Å for Au-S and 2.210-2.426 Å for Cu-S). The diverse coordination patterns of thiolates on the cluster indicate that the organic ligands not only play a key role in the stabilization of metal clusters down to the nanometer scale, but also in the creation of special local structures that facilitate related applications.

It is interesting to note that although the flexible thiolates protect the metal core of the Au₄₁Cu₃₃ cluster tightly, some uncoordinated copper sites are present on the surface. This can be clearly visualized by the coordination of acetonitrile molecules on the surface of the cluster. As shown in Fig. 3a,

one of the surface Cu sites is covered by the ligand of acetonitrile, with a strong interaction (Cu-N bond length of 1.960 Å). On the other hand, some Cu sites are fully exposed, which may facilitate chemical transformations (see Fig. 3b).

Following the structure determination of the cluster, its electronic structure has been then studied both theoretically and experimentally. The count of free electrons of Au₁₁Cu₃₃ observed from its chemical formula corresponds to 34 free electrons, a number that satisfies electron shell closures. The 34 free electrons of the cluster are thus proposed to fill the "superatomic orbitals" of 1S²|1P⁶|1D¹⁰|2S²1F¹⁴|, wherein S-P-D-F-G-H are defined as the angular momentum characters.⁵⁸ To gain insight into the electronic structure of the Au₄₁Cu₃₃ cluster, we further performed density functional theory (DFT) calculations using the experimental structure of the cluster as the model (see technical details in the ESI, Table S2†). 59-62 As shown in Fig. 4a and b, the Au₄₁Cu₃₃ cluster has a gap of 0.52 eV between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO), and both the HOMO and the LUMO distribute on metal and S atoms. The clear HOMO-LUMO gap strongly suggests that the cluster is electronically stable as a neutral species, which is also supported by high-resolution electrospray ionization mass spectrometry and energy-dispersive X-ray spectrometry (EDS) elemental mapping (Fig. S8 and S9†). The partial density of states (PDOS) also suggests that the Au/Cu-d and S-p states dominate both frontier orbitals (see the red and orange lines in Fig. 4c). The measured ultraviolet-visible spectrum (UV-Vis) of the cluster in dichloromethane exhibits nearly no characteristic peaks (Fig. 4d). Containing 34 superatomic free electrons and rigid surface protective shells, the cluster also features high stability. As exhibited in Fig. 4e, the time-dependent UV-Vis profiles suggest that the cluster would maintain its structure in solution form upon heating at 65 °C for up to 5 hours.

Thus, with such a nanocluster of Au₄₁Cu₃₃, which is atomically precise, highly stable and has well-exposed metal sites, we set out to investigate in the following section its applicability

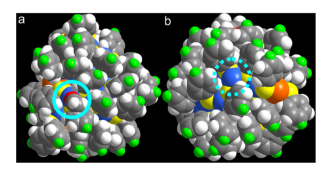


Fig. 3 Au₄₁Cu₃₃(RS)₄₀(MeCN) cluster in space-filling mode showing the presence of uncoordinated atoms on the surface. Color codes for atoms: orange spheres, Au; blue spheres, Cu; magenta spheres, P; purple spheres, Sb; bright green spheres, F; and grey spheres, C. All hydrogen atoms are omitted for clarity.

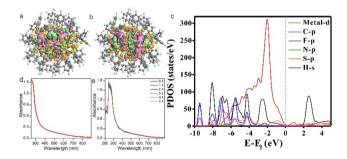


Fig. 4 (a and b) Distribution of the HOMO and the LUMO of the Au₄₁Cu₃₃ cluster. Color codes for atoms: Au, blue; Cu, orange; P, pink; C, gray; and H, white. (c) The projected density of states (PDOS) of the Au₄₁Cu₃₃ cluster. (d) UV-Vis spectra of the Au₄₁Cu₃₃ cluster in dichloromethane. (e) Time-dependent UV-Vis spectra of Au₄₁Cu₃₃ clusters dissolved in chloroform upon heating to 65 °C.

in organic synthesis (Fig. S10-19†). To our delight, this nanocluster allows preeminent catalytic performance in the C-O bond coupling reaction using aryl iodides and phenols and the results are presented in Table 1. Functional groups on the aryl rings such as trifluoromethyl (1k and 1r), methoxy (1d, 2f, 3g), and nitrile (1c) proved to be compatible with this reaction. Furthermore, other aryl rings, such as naphthyl (2e) and quinolinyl (1b), had little influence on the reaction efficiency to give diaryl ether products in good yields. It is worth mentioning that structurally complicated substrates derived from natural products, such as eugenol and δ -tocopherol, were also well accommodated, affording the desired products (3f and **3h**) in moderate yields. As anticipated, the Au₄₁Cu₃₃ cluster, as a recyclable catalyst, presented extremely high TONs for the C-O bond coupling reaction, significantly outperforming its homogeneous counterpart, owing to its decomposition under catalytic conditions. For example, 4-phenoxybenzonitrile (3c) could be prepared using the catalytic system with a TON value as high as 39 269. The comparison of the TON between the Au₄₁Cu₃₃ cluster and several representative catalysts, including copper precursors, gold-thiolate complexes, and other goldcopper alloy nanoclusters without surface-uncoordinated metal sites, gives direct evidence of its high catalytic activity. As shown in Table S3,† the TON of the Au₄₁Cu₃₃ cluster is much higher than that of other reference catalysts under the same conditions, verifying its high surface reactivity.

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To rationalize the high catalytic activity of the Au₄₁Cu₃₃ cluster and to help understand the catalytic mechanism, we performed DFT calculations to probe the reactive sites. The DFT calculations (see Table S4 and Fig. S20†) reveal that the adjacent uncoordinated copper sites have a stronger interaction with C₆H₅-I and C₆H₅-OH than the other copper sites

Table 1 Au₄₁Cu₃₃-catalyzed C-O bond coupling reaction with high TONs^a

do by about 0.2 and 0.5 eV, respectively. This suggests that the uncoordinated copper sites facilitate the formation of HI and C₆H₅-O-C₆H₅. The Au₄₁Cu₃₃ catalyst exhibits not only high catalytic activity, but also high stability. As shown in Fig. S21,† the catalytic activity of Au₄₁Cu₃₃/XC-72 remains nearly unchanged after three cycles. The identical UV-Vis profiles of Au₄₁Cu₃₃/TiO₂ before and after catalysis further suggest its high stability during the catalysis (Fig. S22†). These results indicate that the Au₄₁Cu₃₃ catalyst is practically useful to prepare a variety of diaryl ether compounds.

Conclusion

In conclusion, a new atomically precise Au/Cu nanocluster with 74 nuclei stabilized by flexible thiolates has been obtained. Owing to both electronic and steric closures, the cluster exhibits moderate stability. The most important feature of the cluster relies on the presence of uncoordinated metal sites on its surface, making it a good candidate for catalyzing Ullmann reactions with high activity and a broad reactant scope. This work demonstrates that the stability and catalytic activity of ligand-stabilized metal nanoclusters can be well balanced by delicately tailoring the local structure of the surface. Also, it provides, at the molecular level, the rationale for the catalytic performance of ligand-functionalized metal nanocatalysts.

Author contributions

Hui Shen supervised the project. Lu Dong carried out most of the characterization and catalysis under the guidance of Hui Shen and Shuo Guo. Xueli Sun and Xiongkai Tang prepared the cluster samples and conducted part of the characterization studies. Jiaqi Tang, Zi-Ang Nan, Dongxu Cao, Yanyuan Jia and Simin Li conducted part of the characterization studies. Linke Yu and Xuexin You conducted DFT calculations under the guidance of Fengyu Li. All authors were responsible for analyzing the data and composing the manuscript. Lu Dong, Linke Yu, Xueli Sun, and Xiongkai Tang contributed equally to this work.

Conflicts of interest

There are no conflicts to declare.

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^a Standard conditions: Au₄₁Cu₃₃ (0.00219 mol%), 1 (1.8 mmol, 1.2 equiv.), 2 (1.2 mmol, 1 equiv.), MS 4 Å (20 mg), in CH₃CN (1.2 mL), stirred at 85 °C for 24 h. Yield of isolated products.

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