Highly Robust but Surface-Active: N-Heterocyclic Carbene-Stabilized Au25 Nanocluster as a Homogeneous Catalyst

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Surface organic ligands play a critical role in stabilizing atomically precise metal nanoclusters in solutions. However, it is still challenging to prepare highly robust ligated metal nanoclusters that are surface-active for liquid-phase catalysis without any pre-treatment. Herein, we report a novel N-heterocyclic carbine-stabilized Au25 nanocluster with high thermal and air stabilities as a homogenous catalyst for cycloisomerization of alkynyl amines to indoles. The nanocluster, characterized as [Au25(iPr2-bimy)10Br7]2+ (iPr2-bimy=diisopropyl-benzilidazolium) (1), was synthesized by direct reduction of AuSMe2Cl and iPr2-bimyAuBr with NaBH4 in one pot. X-ray crystallization analysis revealed that the cluster comprises two centered Au13 icosahedra sharing a vertex. Cluster 1 is highly stable and can survive in solution at 80 oC for 12 h, which is superior to Au25 nanoclusters passivated with phosphines or thiols. DFT computations reveal the origins of both electronic and thermal stability of 1 and point to the probable catalytic sites. This work provides new insights into the bonding capability of N-heterocyclic carbene to gold in a cluster, and offers an opportunity to probe the catalytic mechanism at the atomic level.

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Highly Robust but Surface-Active: N-Heterocyclic Carbene-Stabilized Au$_{25}$ Nanocluster as a Homogeneous Catalyst

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ABSTRACT: Surface organic ligands play a critical role in stabilizing atomically precise metal nanoclusters in solutions. However, it is still challenging to prepare highly robust ligated metal nanoclusters that are surface-active for liquid-phase catalysis without any pre-treatment. Herein, we report a novel N-heterocyclic carbine-stabilized Au$_{25}$ nanocluster with high thermal and air stabilities as a homogenous catalyst for cycloisomerization of alkynyl amines to indoles. The nanocluster, characterized as [Au$_{25}$((Pr$_2$-bimy)$_{10}$Br)$_{2}$]$_{2}^{+}$ (Pr$_2$-bimy=diisopropyl-benzilidazolium) (I), was synthesized by direct reduction of AuSMe$_2$Cl and Pr$_2$-bimyAuBr with NaBH$_4$ in one pot. X-ray crystallization analysis revealed that the cluster comprises two centered Au$_{13}$ icosahedra sharing a vertex. Cluster I is highly stable and can survive in solution at 80 °C for 12 h, which is superior to Au$_{25}$ nanoclusters passivated with phosphines or thiols. DFT computations reveal the origins of both electronic and thermal stability of I and point to the probable catalytic sites. This work provides new insights into the bonding capability of N-heterocyclic carbene to gold in a cluster, and offers an opportunity to probe the catalytic mechanism at the atomic level.

INTRODUCTION

Atomically precise metal nanoclusters (NCs) are expected to combine the advantages of homogeneous and heterogeneous catalysis, and provide an ideal model system for investigating the complicated surface chemistry of metal nanocatalysts. For this purpose, the structurally well-defined NCs should be soluble and stable enough in reaction media while having open surface metal sites accessible by reactants. To stabilize metal nanoclusters in solutions, one of the most common strategies is to passivate their surface with organic ligands. The most common ligands used thus far include phosphines, thiols, and alkynyls, already leading to the successful synthesis and structure solution of a large number of ligated atomically precise metal NCs. While the aforementioned organic ligands are essential in NCs preparation, they often restrict the accessibility of reactants to the catalytic site on the metal cluster surface and, as such, may degrade, or in some cases eliminate the catalytic activity of metal NCs. Therefore, limited success has been achieved to demonstrate the direct use of the ligated atomically precise metal NCs as a truly homogeneous catalyst in liquid-phase reactions without being loaded on supports.

Recently, N-heterocyclic carbenes (NHCs) have been shown excellent attributes in stabilizing surfaces, nanoparticles or clusters of coinage metals. Several groups have demonstrated that NHCs readily form robust self-assembled monolayer on coinage metal surface and function as a performance-enhancing modifier. NHCs stabilized Au nanoparticles with excellent water-solubility, thermal stability, and chemical resistance to thiol exchange have also been reported. Several coinage metal clusters stabilized by NHCs have been prepared as well. Theoretical work also suggested that NHCs may function as new ligands for gold surface modification and NCs protection. Given these reports, it occurs to us that NHCs may mimic tertiary phosphines as a new class of ligands in stabilizing metal clusters, and may be a game-changer in metal cluster chemistry.

Herein, we report the synthesis, structure, optical property, stability investigation, and catalytic performance of a NHCs stabilized Au$_{25}$ nanocluster, [Au$_{25}$((Pr$_2$-bimy)$_{10}$Br)$_{2}$]$_{2}^{+}$(I), where Pr$_2$-bimy is diisopropyl-benzilidazolium. Cluster I shows ultrahigh thermal and air stability in solution, far better than that of thiolated Au$_{25}$ clusters. More impressively, the cluster also exhibits high catalytic activities as modeled by cycloisomerization of alkynyl amine to indoles, DFT analysis gives a clear view of the electronic structure and optical absorption characteristics of I, and points to the probable catalytic sites.

RESULTS AND DISCUSSION

Synthesis and atomic structure. In a typical synthesis of cluster I, 0.05 mmol (14.8 mg) AuSMe$_2$Cl was first dissolved in ethanol, and after ultrasonication, equimolar Pr$_2$-bimyAuBr was added. After stirring for 15 min, freshly-prepared NaBH$_4$ in ethanol was added dropwise. The suspension gradually dissolved and the solution turns from pale brown to finally dark brown in 2 min. The reaction continued for 20 h at 20 °C in the
dark. After that, excess NaNO₃ was added to the reaction mixture. After stirring for 30 min, it was centrifuged to afford a black precipitate. 6 mL of mixed solvent of tetrahydrofuran and dichloromethane was used to dissolve the precipitate and the mixture was centrifuged at 14000 rpm for 4 min. Finally, the supernatant was subjected to the diffusion of toluene. Rod-like brown crystals were obtained after 3 weeks (yield: 16.6% based on Au, Figure S1).

X-ray crystallographic analysis revealed that I crystallized in the P-1 space group (Table S1). Each unit cell contains two independent clusters (Figure S2). Cluster I consists of 25 Au atoms, 10 Pr₂-bimy ligands, and 7 Br atoms, formulated as [Au₂₅(Pr₂-bimy)₁₀Br]⁷⁺. The unit cell contains counter anions like NO₃⁻ and Cl⁻, based on which the +2 charge of I was determined. The structure of I (Figure 1) can be described as two centered Au₁₀₃ icosahedra sharing a vertex, giving rise to a prolate shape with an 1:5:1:5:1:5:1:5:1:5:1 a arrangement of Au atoms (the subscripts designate the positions on the fivefold axis). The four Au₅ pentagons are arranged in a staggered-eclipse-staggered configuration. As such, the cluster conforms to idealized D₅h symmetry. The ten N-heterocyclic carbene ligands coordinate to the two outer Au (5₁) pentagons. Of the seven bromides, five are doubly bridging the two inner Au (5₂) pentagons while the remaining two terminally coordinated to the two outermost Au (1₅) atoms. The rod-like metal core measures about 1.1 nm along the fivefold axis (Figure S3). The Au-Au bond lengths, averaged in accord with D₅h symmetry, are: Au₅-Au₅, 2.935 Å, Au₁₅-Au₁₅, 2.723 Å, Au₁₁-Au₁₁, 2.778 Å, Au₅-Au₅, 2.898 Å, Au₁₅-Au₁₅, 2.857 Å, Au₁₁-Au₁₁, 2.736 Å, Au₁₅-Au₁₅, 2.811 Å. It can be seen that intra-icosahedral Au-Au bonds are significantly shorter than those on the surface as well as those between the two icosahedra (Figure S4).

The average Au-Br bond distance of the doubly-bridging bromides is 2.563 Å which somewhat longer than that of 2.442 Å of the terminal bromides. The ten NHC ligands coordinate to the ten Au₅ atoms forming Au-C bonds. The average Au-C distance is 2.069 Å, suggesting strong bonding interaction. For comparison, this value is slightly larger than that in Pr₂-bimyAuBr (1.978 Å), but somewhat less than that in Pr₂-bimy self-assembled gold monolayer (2.118 Å). Interestingly, structure of I quite resembles that of Au₂₅ or Pr₂Ag₂₅ protected by tertiary phosphines, indicating that in shaping structure of nanoclusters, NHCs can adopt similar binding modes with tertiary phosphines. The similar observation has also been revealed in organometallic complexes. Cluster I is also closely related to the series of vertex-sharing bicosaheiral bimetallic (Au-Ag) and trimetallic (Au-Ag-M where M = Ni, Pd, Pt) clusters stabilized by phosphine and halide ligands reported by Teo and coworkers as early as the 1980s.⁴⁷-⁴⁸

**Figure 1.** Crystal structure of [Au₂₅(Pr₂-bimy)₁₀Br]⁷⁺. (a) Overall structure and (b) Au₂₅ metal core, showing the bonding modes between Au atoms and the ligands (one of each is highlighted). Color legend: orange, Au; cyan, Br; blue, N; gray, C. H atoms are omitted for clarity.

**Figure 2.** (a) ESI-MS spectrum of I crystals dissolved in CH₂Cl₂. Inset: the experimental (black trace) and simulated (red trace) isotopic patterns of [Au₂₅(Pr₂-bimy)₁₀Br]⁷⁺. (b) UV-Vis spectrum of crystals of I dissolved in CH₂Cl₂.

The composition of I was confirmed by electrospray ionization mass spectrometry (ESI-MS) under positive ion mode in CH₂Cl₂. As shown in Figure 2a, the ESI-MS spectrum revealed only a single peak at 3756 m/z, corresponding to [Au₂₅(Pr₂-bimy)₁₀Br]⁷⁺. Perfect agreement was observed between the experimental spectrum and simulated isotopic distribution pattern.

**Figure 3.** Visualization of the Π and Σ symmetries of the six highest occupied molecular orbitals HOMO-5 to HOMO (with 12 electrons), marked in Figure S5. HOMO-5 to HOMO-3 (bottom) have clear bonding characters with respect to the horizontal plane bisecting the Au₂₅ metal core, while HOMO-2 to HOMO (top) have antibonding characters. The center graph shows the schematic energy splitting of the bonding/anti-bonding orbitals formed as combination of Π-type orbitals in each icosahedral half-core.
Electronic structure and optical properties. DFT calculations were employed to investigate the ground-state electronic structure of 1 (see Experimental section for technical details). The calculated HOMO-LUMO energy gap (Figure S5) is significant, 1.52 eV, which implies a good electronic stability of the cluster. However, the chemical composition of 1 implies 16 metallic free electrons, which is not an expected shell-closing electron number of metal clusters. This prompted us to look further to the reasons of the electronic stability. According to Figure S5a, six highest occupied orbitals of the system have clear P-type symmetries when the symmetry analysis is done in one of the icosahedral half-cores of the cluster. Visualization of the orbitals (Figure 3) reveals that these six orbitals (with overall Π and Σ symmetries) are arranged in two groups having a clear bonding and antibonding characters with respect to the plane bisecting the core. This implies a strong interaction of the superatom P-type orbitals between the icosahedral half-cores, indicating that the free-electron count of 16 of the whole cluster can be understood as a strongly interacting dimer of two 8-electron systems.

1 exhibits a distinct optical absorption spectrum in the UV-Vis region (Figure 2b) with five features: three shoulders at 333, 362, and 406 nm as well as two peaks at 469 and 650 nm. The absorption spectrum of 1 is very similar to that of phosphine and thiolate co-protected Au25(PPh3)3(SC2H5PH)3Cl2 (2) (Figure 4b), suggesting that discrete electronic states of Au25 with similar metal framework are most likely dictated by the vertex-sharing bicoshedral metal architecture.

Time-dependent DFT was employed to compute and analyze the optical absorption spectrum (see Experimental section for technical details). Figure S6 shows the comparison of the calculated and measure spectra. We were able to identify the same five features in the computed spectrum as in the measured data, although the two lowest-energy peaks are somewhat red-shifted in the calculations. Analysis of these five features in terms of transition contribution maps (TCMs) (Figure S7) decomposing the state symmetries in the D5h point group (Figure S5b) shows that the two lowest-energy peaks are mostly single-state transition between symmetries A2g to A1u (650 nm in the exp. data) and E2u to E1g (469 nm) while the higher-energy transition have multi-state features with contributions from ligands as well.

Excellent stability of 1 in solution. The stability of 1 was investigated by tracking its UV-vis spectrum under thermal treatments. As shown in Figure 4a, the UV-vis profile of 1 was practically unchanged upon heating in 1,2-dichloroethane at 80 °C for 12 h, indicating ultrahigh thermal stability. The intensity of the peak at 650 nm was plotted as a function of heating time in Figure 4d (black curve). The relative intensity showed almost no degradation in the thermal measurement, suggesting that there is no decomposition. Note that 1 is also highly stable in other solvents such as tetrahydrofuran (Figure S8). For comparison, the stability of thiol and phosphine stabilized Au25 NCs were also investigated under the same conditions. As shown in Figure 4b, UV-vis spectra of 2, Au25(PPh3)3(SC2H5PH)3Cl2, deteriorates progressively in the 12 h heating period, and the relative absorbance intensity of the peak at 650 nm decreases significantly (Figure 4d, red curve). Furthermore, thiol-protected Au25(SC2H5PH)18 (3) also exhibits much worse stability than that of 1 (Figure 4c and pink curve in Figure 4d). Earlier DFT calculations on a mixed-ligand [Au11(PPh3)3(NHC52)Cl2]Cl cluster have shown, that the gold-carbene bond can be stronger than gold-phosphine bond by up to 0.9 eV. This explains also the observed superior thermal stability of 1 over 2 in this work.

Surface reactivity and catalytic performance. Next, we explored the functionality of 1 as a homogeneous catalyst. Here we chose the cycloisomerization of alkynyl amines to form indoles as a model reaction. Indoles are important heterocycles which are broadly used in material science and biology. The catalytic reaction was conducted by adding 1.5 % mmol catalyst (based on Au atom) succeeded in accelerating the cycloisomerization reaction of N-(2-ethynylphenyl)-4-methylbenzenesulfonamide to 1-tosyl-1H-indole at 25 °C in 10 h (Table S2, entry 1). The catalytic conditions were optimized by exploring the effects of solvent, time and temperature. The optimal conditions were identified at 50 °C in 1,2-dichloroethane, providing the desired product with over 99% yield. Of note, the reaction generally worked well in aprotic solvents (Table S2, entry 2-5). As illustrated in Figure 5, the cluster displayed an extremely high activity in cycloisomeriza- tion of alkynyl amines at 50 °C as its conversion close to 100% after 10 h. Moreover, the catalyst is stable during the catalytic process, as practically no apparent changes in the UV-vis spectrum were observed before and after the catalytic reaction (Figure S9). Given the optimal reaction conditions (Table S2, entry 6), the catalytic performance of the other two types of Au25 (2 and 3) were also compared. As shown in Table S3, 1 showed a much higher yield of over 99% in comparison to 2 or 3, as well as Au(I) carbene precursors.

We attribute the observed excellent catalysis of 1 to its sterically open halide sites, which are, on the basis of DFT
Figure 5. Catalytic performances of 1.5% mmol 1 recorded at different reaction time points for the catalytic cycloisomerization of alkynyl amine.

CONCLUSION

In conclusion, a novel gold nanocluster Au25 protected solely by N-heterocyclic carbene ligand was synthesized for the first time via a one-pot reaction protocol. X-ray crystallographic analysis revealed that the cluster consists of two icosahedra sharing a vertex. DFT analysis revealed that the electronic subsystems of the two icosahedra can be understood as 8-electron configurations that are strongly interacting giving a stable 16-electron system with a large HOMO-LUMO energy gap. The cluster displays ultra high thermal and air stabilities in solution, superior to all other reported Au25 nanoclusters. In addition, it exhibits high catalytic activity and stability in the cycloisomerization of alkynyl amines to indoles at 50 °C. This work paves ways to use heterocyclic carbene stabilized metal nanoclusters in homogenous catalysis. Finally, it is hoped that, in analogy to its phosphine stabilized kins, the bicosahedral framework can be extended to polyicosahedral architecture which, given their superior thermal and air stabilities, may find other applications in nanotechnology that are difficult to fill with other metal cluster systems.

ASSOCIATED CONTENT

The Supporting Information is available free of charge on the ACS Publications website including experimental and computational details and additional data (PDF) and the crystallographic data (CIF).

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Clusters in solution as the catalyst

heated at 80°C in solution
