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Pyridine dicarbanion-bonded Ag₁₃ organometallic nanoclusters: synthesis and on-surface oxidative coupling reaction†

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Highly reactive organometallic nanoclusters in situ generated in metal-catalyzed reactions are pivotal in the comprehension of catalytic mechanisms. Herein, we develop a two-step synthetic method to achieve three unprecedented aryl dicarbanion-bonded Ag_{13} nanoclusters by using protective macrocyclic ligands. Firstly, various aryl dicarbanion- Ag_4 cluster intermediates are acquired via a silver-mediated annulation reaction within a macrocyclic ligand. These Ag_4 cluster precursors are released from the surrounding macrocycle by protonation, and further undergo an inter-cluster coupling to generate bipyridine products and low-valence silver atoms. The remaining resurgent diide- Ag_4 clusters assemble with low-valence silver atoms to yield a series of organometallic Ag_{13} nanoclusters. These Ag_{13} nanoclusters feature a unique open-shell electronic structure as well as a chiral cluster architecture due to the asymmetric arrangements of surrounding aryl dianion ligands. Furthermore, the pyridyl diide ligands on the surface of the nanocluster further experience an intra-cluster oxidative coupling to produce bipyridine coupling products and large nanoparticles. The coupling reaction-driven cluster-to-cluster transformation is comprehensively tracked by high resolution mass spectroscopy. This work is not only reminiscent of the detailed evolution of cluster species upon the occurrence of coupling reactions, but also reproduces novel interand intra-cluster coupling steps at different reaction stages.

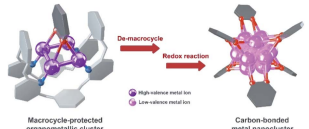
Introduction

The study of metal subnano- or nano-clusters in situ generated in transition metal-catalyzed reactions has become a fascinating and substantial topic in both homogeneous and heterogeneous catalyses.1 These kinds of metal nanoclusters, which are mostly formed through the reduction of metal complexes by reaction substrates, coordinative ligands or additives, 1f2 largely reshape the perception regarding real catalytic species and detailed catalytic mechanisms. They act as either catalytically active species 1d,f,3 or alternative reservoirs of active metal complexes,4 and prevent devitalization of the catalytic species by suppressing the formation of insoluble metal species. Structural investigations on the in situ generated metal nanocluster intermediates, especially carbon-bonded organometallic nanoclusters, enable chemists to gain insight into the structural details of privileged intermediates and promote the comprehension of catalytic mechanisms. However,

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although organometallic clusters have been identified as intermediate species in many metal-catalyzed organic transformations by mass spectrometry, 3d,5 electron microscopy 2a,3a and X-ray absorption spectroscopy, 3b,c their synthesis and structural characterization are still a challenging task due to their transient and highly reactive nature. Accordingly, the transformations of organic ligands attaching on these organometallic nanoclusters are rarely investigated.

The Ag(0/I)-based single-electron redox cycle has been extensively proposed in many silver-catalyzed coupling reactions.⁷ The presence of low valence silver atoms in the Ag(0/1) redox process facilitates the in situ formation of silver nanoclusters, which have been demonstrated to exhibit excellent catalytic activity in organic transformations.8 However, in-depth structural characterization and reactivity studies of these in situ generated silver nanoclusters suffer from the lack of appropriate synthetic methods and stabilizing means as labile silvercarbon bonds prefer a rapid homolysis disproportionation to generate metallic silver and finally cause decomposition.9 In view of the previously reported successful encapsulation and stabilization of reactive organometallic species within supramolecular cages, 10 we envision that the initial encaging of organosilver species within a closed macrocyclic environment followed by a steerable organometallic transformation may provide a viable pathway to acquire organosilver nanoclusters (Scheme 1).



Scheme 1 Synthesis of carbon-bonded metal nanoclusters *via* the transformation of macrocycle-protected organometallic clusters.

In this work, we develop a two-step synthetic method to achieve a series of unprecedented aryl vicinal dicarbanion bonded Ag_{13} nanoclusters ($[Ag_{13}(^{R}Py-H)_{6}](CF_{3}SO_{3})_{6}$, R = Me, n-Pr, Ph). At the initial step, differently substituted pyridine dicarbanion-bonded Ag₄ cluster intermediates are acquired via a silver-mediated cyclization reaction within a macrocyclic ligand. After removing the protective macrocycle by protonation, the dicarbanion-Ag₄ cluster precursors undergo an intercluster oxidative coupling between two pyridine carbanions to in situ generate low-valence silver atoms. The remaining released dicarbanion-Ag₄ clusters assemble with a low-valence silver atom to yield vicinal dicarbanion bonded Ag₁₃ nanoclusters. These organometallic nanoclusters feature a unique open-shell electronic structure as well as a chiral cluster architecture as a result of asymmetric arrangement of surrounding pyridyl dianion ligands. Furthermore, the pyridyl dicarbanion ligands on the surface of a Ag₁₃ nanocluster can undergo an intra-cluster oxidative coupling to generate the bipyridine coupling product and large nanoparticle species, reminiscent of the evolution of cluster species in many metal-catalyzed

reactions. This work not only reveals novel dicarbanion-bonded metal nanoclusters for the first time, but also provides a new perspective for the dynamic evolution of organometallic species in metal catalysis, from isolated single atom species through low-nuclear clusters to nanoclusters along with the occurrence of intra- and inter-cluster coupling reactions.

Results and discussion

Pyridyl dicarbanion bonded Ag₁₃ nanoclusters

In this work, an attempt was made to generate a gem-dimetallic organometallic species as a representative intermediate motif in metal-catalyzed reactions.11 To achieve this goal, a silver(1)mediated intermolecular annulation reaction of ketones and propargylamine¹² was performed in the presence of excessive silver salts and a polydentate macrocyclic ligand octamethylazacalix[8]pyridine (Pv[8]).13 As shown in Fig. 1, various methyl ketones together with propargylamine undergo a continuous multi-step transformation of condensation, imine-enamine isomerization, 6-endo-dig cyclization and oxidative aromatization to yield three differently substituted pyridyl dicarbanion bonded Ag₄ clusters (abbreviated as MePyAg₅, n-PrPyAg₅ and PhPyAg₅) in moderate yields (23-61%). The composition and structural features of these Ag4 clusters were confirmed by highresolution electrospray ionization mass spectrometry (Fig. S1-S3†) and NMR (Fig. S4-S7†).

Crystalline solids of these organosilver cluster compounds are quite stable in the dark even upon exposure to air and moisture for months. The ¹H NMR monitoring on the acetone solution of ^{Me}PyAg₅ shows negligible change for up to seven days (Fig. S8†), suggesting its superior stability in the solution state. Single-crystal X-ray analysis reveals a highly isostructural pyridyl diide-Ag₄ cluster core in ^{Me}PyAg₅ (Fig. 1), ^{n-Pr}PyAg₅

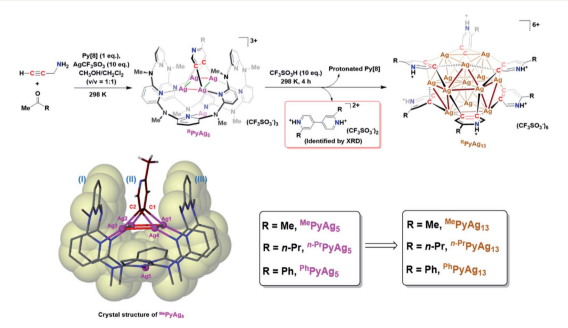


Fig. 1 Synthetic procedures for macrocycle-protected Ag_4 cluster complexes and aryl vicinal dicarbanion bonded Ag_{13} nanoclusters. The crystal structure of $^{Me}PyAg_5$ is shown as an inset with anions and solvent molecules omitted for clarity.

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(Fig. S9†) and PhPyAg₅ (Fig. S10†). Taking MePyAg₅ for instance, two vicinal carbon atoms C1 and C2 of a pyridyl ring are negatively charged and stabilized by a coplanar argentophilic Ag₄ rectangle via four-fold Ag-C σ bonds in the range of 2.148(9)-2.181(9) Å, which are comparable to the CAg₂ species in reported arylsilver(1) complexes. 10e,14 The Ag4 rectangle is composed of two short Ag-Ag edges (Ag1-Ag2: 2.724(1) Å, Ag3-Ag4: 2.683(1) Å) and two long ones (Ag1-Ag4: 3.164(1) Å, Ag2-Ag3: 3.246(2) Å). The whole pyridyl diide-Ag₄ cluster motif is enclosed by a semi-open bowl-shaped macrocyclic ligand Py[8] via four-fold Ag-N bonds. Another silver atom is clamped by two pyridyl nitrogen atoms at the bottom of the bowl. It should be emphasized that the metalated pyridyl diide is sandwiched by two parallel pyridyl rings of Py[8], leading to a fully eclipsed face-to-face π - π stacking with small dihedral angles ($\theta_{I-II} = 5.0^{\circ}$ and $\theta_{\text{II-III}} = 4.4^{\circ}$) and displacement distances ($D_{\text{I-II}} = 0.738 \text{ Å}$ and $D_{\rm II-III} = 0.523$ Å). This unusual eclipsed face-to-face π - π stacking is in sharp contrast to the common displaced π - π stacking between aromatic rings as a result of the requirement of mitigated π – π electron repulsion. ¹⁵ Theoretical studies reveal that the p_{π} electrons of the central metalated pyridine largely participate in the C-Ag₂ multi-centered bonding (Fig. S11†) and thus weaken the π - π repulsion with two bilateral pyridine rings belonging to the peripheral Py[8].

With the macrocycle-protected organosilver cluster intermediates in hand, we subsequently removed the protective Py[8] by protonation to initiate transformations of the diide-Ag₄ clusters. After screening several acids with a wide range of pK_a values (Fig. S12†), the strong trifluoromethanesulfonic acid CF₃SO₃H was selected because of the prompt generation of protonated Py[8] as a white precipitate (Fig. S13†) upon the addition of CF₃SO₃H to the acetone solution of MePyAg₅. This protonation process was simultaneously accompanied by a gradual solution color change from light yellow to brown. Diffusion of diethyl ether into the brown solution deposited two types of brown and colorless crystals in a yield of 52% (Table S1†) and 13%, respectively.

X-ray crystallographic analysis of the MePyAg5-derived brown crystalline sample reveals that its structure includes a twisted cuboctahedral Ag₁₃ core, which is peripherally wrapped by six 2methylpyridyl diides via a μ_4 -C,C- η^2 , η^2 mode and five coordinative triflate anions (Fig. 2a). There is one free CF₃SO₃⁻ serving as an uncoordinated counter anion. This Ag₁₃ nanocluster is different from the reported Ag₁₃ cuboctahedron, each triangle face of which is capped by an Fe(CO)4 group to generate a $[Ag_{13}Fe_8(CO)_{32}]^{4-}$ cluster. ¹⁶ The electrospray ionization mass spectroscopy (ESI-MS) study on the solution of the brown crystals gives rise to two isotopically well-resolved peaks at m/z =2847.7750 and 2865.7704, which can be assigned to the species $\{ [Ag_{13}(^{Me}Py-H)_{6}](CF_{3}SO_{3})_{6} \}^{+} \quad \text{and} \quad \{ [Ag_{13}(^{Me}Py-H)_{6}](CF_{3}SO_{3})_{6} \}^{-} \} = 0$ \cdot H₂O $\}$ ⁺, respectively (Fig. S14†). This result indicates that six 2methylpyridyl diides around the cluster core are all protonated at the pyridyl nitrogen atoms. In the light of the charge balance requirement, we assume that the 13-membered silver cluster core shows twelve positive charges and has an unpaired electron, which is similar to the paramagnetic [Ag₁₃Fe₈(CO)₃₂]⁴⁻ cluster. 16 This conjecture is supported by electron paramagnetic

resonance (EPR) measurement (Fig. S15†)16 and X-ray photoelectron spectroscopy (XPS, (Fig. S16†)). The XPS spectra revealed two intensive peaks in the Ag 3d region at 374.9 and 368.9 eV corresponding to Ag $3d_{3/2}$ and Ag $3d_{5/2}$, respectively (Fig. S16b†). The peak could be deconvoluted into 375.5 eV (3d_{3/} 2, Ag⁰), 374.9 eV (3d_{3/2}, Ag⁺), 369.5 eV (3d_{5/2}, Ag⁰) and 368.9 eV $(3d_{5/2}, Ag^+)$. The area ratios of Ag^+/Ag^0 for Ag $3d_{3/2}$ and Ag $3d_{5/2}$ are 11:1 and 10:1, basically consistent with the presence of two kinds of Ag⁺ and Ag⁰ in the crystal structure. Based on the Auger electron spectroscopy (AES) data under X-ray irradiation of 1486.6 eV, the Auger parameters of Ag⁺ and Ag⁰ are deduced to be 724.5 and 725.1 eV, which are in good agreement with those of Ag₂O (Auger parameter: 724.5 eV)¹⁷ and Ag(0) (Auger parameter: 726.0 eV).18 Consequently, the molecular formula of the brown crystals is determined to be [Ag₁₃(MePy-H)₆](CF₃SO₃)₆ (MePyAg₁₃). The existence of low valence silver atoms in the $[Ag_{13}]^{12+}$ core of $^{Me}PyAg_{13}$ implies that some Ag(i) ions were reduced during the construction of MePyAg13, which is supported by further X-ray crystallographic analysis of the colorless crystals (Fig. 1). The identification of the colorless crystalline product as protonated 2,2'-dimethyl-4,4'-bipyridine (Fig. S17 and S18†) clearly indicates the occurrence of a two-electron oxidative coupling between two C1-Ag2 units from two individual 2-methylpyridine-Ag₄ motifs. This coupling process simultaneously generates low valence silver atoms. In view of the high similarity of pyridyl C2-Ag4 moieties between MePyAg5 and MePyAg13, we deduce that the 13-membered nanocluster core in MePvAg₁₃ arises from the assembly of low valence silver atoms with remaining protonated 2-methylpyridine diide-Ag₄ motifs in situ released from the Py[8] macrocycle of MePyAg₅ (Fig. 1 and S19†).

The twisted cuboctahedral Ag₁₃ core in MePyAg₁₃ is divided into three layers of Ag₃-Ag₇-Ag₃, and the distances between the central silver (Ag1) and the upper and lower Ag3 layers are 2.331 and 2.328 Å, respectively (Fig. 2b). The argentophilic distances between the central silver atom Ag1 and twelve surface silver ions are in the range of 2.863(2)-3.167(2) Å (yellow color in Fig. 2b), which are comparable with the Ag-Ag distances in $[Ag_{13}Fe_8(CO)_{32}]^{4-}$ (2.923 Å). The other 24 surficial argentophilic edges among twelve peripheral silver atoms can be classified into two types. The short edges bridged by the same diide carbon atom (average distances: 2.755 Å, red color in Fig. 2b) and the long ones spanned by two diide carbon atoms of the same pyridine ring (average distances: 3.205 Å, green color in Fig. 2b) are comparable to those in the parent tetranuclear cluster MePyAg₅ (2.703 Å and 3.192 Å, respectively). The maximum value of the argentophilic distance alternations between the long and short edges in MePyAg₁₃ is 0.621 Å, which is significantly larger than that in $[Ag_{13}Fe_8(CO)_{32}]^{4-.16}$ In addition, six protonated 2-methylpyridine diide ligands each is enveloped by a rectangular plane of the Ag₁₃ core through fourfold Ag-C bonds in the range of 2.12(2)-2.29(3) Å, which are slightly longer than the Ag-C bonds in MePyAg₅. Notably, the protonated 2-methylpyridine diide ligands on both upper (I, II and III) and lower layers (IV, V and VI) exhibit different orientations (Fig. 2a). Therein, four diide ligands (I, III, IV and V in pink color) are equatorially bonded to the Ag₁₃ core, while the

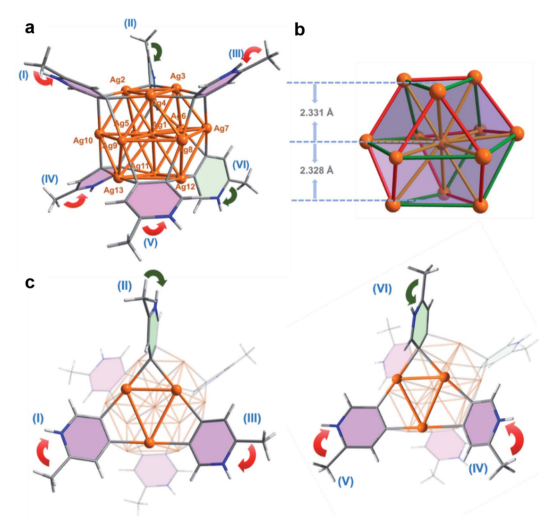


Fig. 2 (a) Crystal structure of $^{Me}PyAg_{13}$ containing equatorial (pink) and axial (light green) 2-methylpyridyl diide ligands. Peripheral $CF_3SO_3^{-1}$ anions are omitted for clarity. (b) The Ag_{13} kernel in $^{Me}PyAg_{13}$. Color coding: Ag, brown; C, gray; H, white; N, blue. (c) Arrangement of the orientated 2-methylpyridine rings attached on the Ag_{13} kernel at the upper and nether sides in $^{Me}PyAg_{13}$. Selected bond lengths (Å) of $^{Me}PyAg_{13}$ highlighted in yellow: $Ag_1^{-1}Ag_2^{-1}2$ 2.885(2); $Ag_1^{-1}Ag_3^{-1}2$ 2.897(2); $Ag_1^{-1}Ag_3^{-1}2$ 3.025(2); $Ag_1^{-1}Ag_3^{-1}2$ 2.863(2); $Ag_1^{-1}Ag_3^{-1}2$ 3.038(2); $Ag_1^{-1}Ag_3^{-1}2$ 2.986(2); $Ag_1^{-1}Ag_3^{-1}2$ 2.993(2); $Ag_1^{-1}Ag_3^{-1}2$ 2.863(2); $Ag_1^{-1}Ag_3^{-1}2$ 3.038(2); $Ag_1^{-1}Ag_3^{-1}2$ 2.780(2); $Ag_2^{-1}Ag_3^{-1}2$ 2.780(2); $Ag_3^{-1}Ag_3^{-1}2$ 2.780(2); Ag_3^{-

other two (II and VI in light green color) are in an axial orientation. The biased spatial arrangement of the protonated 2-methylpyridine ligands can be described by the arrow vector from the methyl substituent toward the pyridyl nitrogen. As to the four equatorial 2-methylpyridine ligands, I and III on the upper layer are related by a C_3 -axis, while IV and V on the nether layer have a mirror symmetry (red arrows in Fig. 2c). On the other hand, the axial 2-methylpyridine ligand II is in an *endo*-form with the methyl group close to the silver nanocluster, while ligand VI shows an *exo*-form with the methyl group pointing away from the cluster core (green arrows in Fig. 2c). In this way, the twelve peripheral silver ions (from Ag2 to Ag13) of the Ag13 core are coordinated by two kinds of pyridyl carbon atoms (C_{para} and C_{meta} refer to the 4- and 5-position carbon atom on the pyridine ring, respectively) in three combinations

of C_{para} –Ag– C_{para} , C_{para} –Ag– C_{meta} and C_{meta} –Ag– C_{meta} (Fig. S20†). The differentiated orientation of the total six 2-methylpyridyl diide ligands breaks the symmetry and thus imparts chirality to the whole cluster structure of Me PyAg₁₃, making it the first {M₁₃}-type chiral metal nanocluster to our knowledge. 19 Herein, both cluster enantiomers are present in the crystal structure of Me PyAg₁₃ (Fig. S21†).

We next carried out density functional theory (DFT) calculations to investigate the bonding and argentophilic interaction in $^{\text{Me}}\text{PyAg}_{13}$. The Wiberg bond order 20 based on the Löwdin orthogonalized basis was calculated and summarized in Table S2.† The sum of Ag–C bond orders for each CAg₂ species is close to one (0.893–0.975) and the multi-center bond order (MCBO, Table S3†) of each CAg₂ species ranges from 0.013 to 0.091, suggesting the formation of a 3c–2e bonding in each CAg₂

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(Fig. S22b and c†).

species. In addition, the Ag–Ag Wiberg bond order of $^{\text{Me}}\text{PyAg}_{13}$ is in the range of 0.189–0.384 as a result of significant argentophilic interaction between two silver atoms. Therein, the average Wiberg bond order of short Ag–Ag edges (red color in Fig. 2b) in a cuboctahedral Ag₁₃ core is 0.359, while those of the long Ag–Ag edges (green color in Fig. 2b) and twelve central ones (yellow color in Fig. 2b) are 0.222 and 0.260, respectively. The independent gradient model based on Hirshfeld partition (IGMH)²¹ was further carried out via sign(λ_2) ρ functions to examine intramolecular interactions in $^{\text{Me}}\text{PyAg}_{13}$. From the IGMH map, the mazarine area between six 2-methylpyridyl diides and the Ag₁₃ core (Fig. S22a†) indicates prominent electron density in these regions and covalent characteristics of the C–Ag bonds. Furthermore, there are blue areas in the isosurface

between two adjacent silver atoms, suggesting the presence of

apparent argentophilic interaction within the Ag₁₃ core

The Atoms in Molecules (AIM) analysis²² also confirms strong argentophilic interaction in $^{\text{Me}}\text{PyAg}_{13}$. As shown in Fig. S23,† the bond critical points exist in each pair of adjacent silver atoms. The negative bond degree parameter (BD = $E(r)/\rho(r)$, E(r) and $\rho(r)$ being the total electron energy density and the electron density value at the Ag–Ag (3, -1) critical point) and the ratio |V(r)|/G(r) > 1 (V(r) and G(r) being the pressures exerted on and by the electrons at the Ag–Ag (3, -1) critical point),²³ indicate apparent argentophilic interactions in the Ag₁₃ core (Table S4†). It is worth noting that some bond critical points lie between H atoms of 2-methylpyridyl diides and F or O atoms of CF₃SO₃ $^-$, suggesting that multiple hydrogen bonding is instrumental in the stabilization of $^{\text{Me}}\text{PyAg}_{13}$.

Our two-step synthetic strategy is also applicable in the synthesis of ^{n-Pr}PyAg₁₃ and ^{Ph}PyAg₁₃ by the reaction of differently substituted ^{n-Pr}PyAg₅ or ^{Ph}PyAg₅ with CF₃SO₃H, respectively. Similarly, during the acidification of ^{n-Pr}PyAg₅ or ^{Ph}PyAg₅, the coupling product 2,2'-dipropyl-4,4'-bipyridine or 2,2'-

diphenyl-4,4'-bipyridine was detected and successfully characterized by mass spectrometry (Fig. S24 and S25†) and single crystal XRD analysis (Fig. S26 and S27†). XRD crystallographic analysis, high-resolution ESI-MS and UV-vis absorption spectra reveal that ^{n-Pr}PyAg₁₃ (Fig. 3a, S28 and S29†) and ^{Ph}PyAg₁₃ (Fig. 3b and S30-S32†) are isostructural with MePvAg₁₃ in the aspects of bond distances and angles. Nevertheless, the dihedral angles between the upper and lower Ag₃ planes in ⁿ-PrPyAg₁₃ and PhPyAg₁₃ are 11.5° and 11.7°, respectively (Table S5†), a bit higher than that of Me PyAg₁₃ (9.7°). In addition, the mean deviation of the central Ag₇ layer in ^{n-Pr}PyAg₁₃ and $^{\text{Ph}}$ PyAg₁₃ is 0.125 and 0.104 Å, respectively, also larger than that of $^{\text{Me}}$ PyAg₁₃ (0.094 Å). The more distorted Ag₁₃ core in $^{n-\text{Pr}}$ PyAg₁₃ and PhPyAg₁₃ should be ascribed to significant steric hindrance among peripheral n-propyl and phenyl substituents. Furthermore, in contrast to the exo and endo spatial orientations of two axial pyridyl diide ligands in ^{n-Pr}PyAg₁₃ and ^{Me}PyAg₁₃, ^{Ph}PyAg₁₃ has two axial 2-phenylpyridine ligands II and VI both in the exoform with the phenyl group pointing away from the cluster core (green arrows in Fig. 3b). This structural difference underlines the oriented diversity of the pyridyl diide ligands around the Ag₁₃ nanocluster core.

The newly synthesized Ag_{13} nanoclusters retain their structures intact in solution based on the $^1\text{H-NMR}$ and ESI-MS spectra. In the ^1H NMR spectrum of $^{\text{Me}}\text{PyAg}_{13}$, three sets of aromatic resonance peaks at 8.94, 8.31–8.19 and 7.98 ppm with a ratio of 1:10:1 were observed (Fig. S33†). This unusual integral ratio corresponding to the pyridyl protons in $^{\text{Me}}\text{PyAg}_{13}$ may be due to different configurational isomers in solution or differentiated ligand orientations in $^{\text{Me}}\text{PyAg}_{13}$. Diffusion ordered spectroscopy (DOSY) of $^{\text{Me}}\text{PyAg}_{13}$ (Fig. S34†) shows only one diffusion coefficient of $D=7.35\times10^{-10}$ m² s⁻¹ for the pyridyl proton peaks (Table S6†). Based on the Stokes–Einstein equation, the calculated diameter was 18.75 Å, which agrees well with the measured distances in the crystal structure of

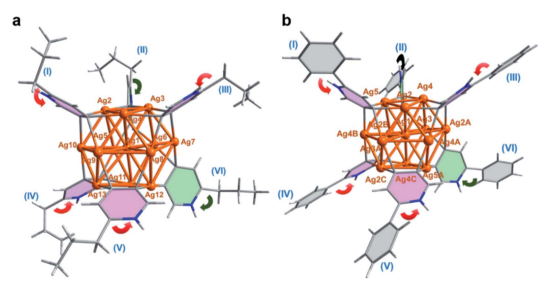


Fig. 3 Crystal structures of the Ag_{13} nanoclusters in (a) $^{n-Pr}PyAg_{13}$ and (b) $^{Ph}PyAg_{13}$ with equatorial (pink) and axial (light green) pyridyl ligands. Peripheral $CF_3SO_3^-$ anions and solvent molecules are omitted for clarity. Color coding: Ag, brown; C, gray and blue gray; H, white; N, blue.

proton NMR peaks.

MePyAg₁₃ (17.55 Å). Besides, variable temperature ¹H NMR spectra (Fig. S35†) and UV-vis spectra (Fig. S36†) display negligible changes with the temperature increase from 298 to 353 K, excluding the possibility of other equilibrium cluster conformations in solution. Therefore, the aromatic integral ratio of 1:10:1 may correspond to 12 inequitable pyridyl protons of six diide ligands. In order to accurately assign the twelve pyridyl proton atoms, the NMR spectrum of MePyAg₁₃ was simulated by the scaling method based on an optimized structure. The simulated results show that the proton signal at 8.94 ppm corresponds to the proton atom attaching on the carbon atom C6 (adjacent to the pyridyl nitrogen) in ligand VI, and the peak at 7.98 ppm is assigned to the C3 hydrogen atom (meta-position to the pyridyl nitrogen) of II (Fig. S37a†). In addition, the chemical shifts of the remaining ten protons are also inequivalent but very close, which are finely influenced by the ligand orientation and the C-Ag₂ units (Fig. S37b†). The ¹H NMR spectrum of $^{n-Pr}$ PyAg₁₃ (Fig. S38†) is very similar to that of MePyAg₁₃ because of their completely consistent ligand orientation scenario. In contrast, the ¹H NMR spectrum of ^{Ph}PyAg₁₃ shows four broad resonance peaks at 8.77, 8.53, 7.90 and 7.66 ppm with an integral ratio of 6:6:12:18 (Fig. S39†). The simulated ¹H NMR spectrum of ^{Ph}PyAg₁₃ (Fig. S40a†) indicates that the resonances at 8.77 and 8.53 ppm are assigned to the protons on the *meta* and *ortho* positions of the pyridine rings in PhPyAg₁₃, and the peaks at 7.90 and 7.66 ppm correspond to the protons on the phenyl rings (Fig. S40b†). The above experimental and theoretical NMR results suggest that distinct spatial arrangements of the pyridyl diide ligands lead to biased charge distribution on carbon atoms and thus account for inequivalent

Inter- and intra-cluster coupling of diide ligands

To gain further insight into the formation process of Ag_{13} nanoclusters, we conducted a cross experiment between the same equivalent $^{\text{Me}}\text{Py}Ag_5$ and $^{n\text{-Pr}}\text{Py}Ag_5$ with CF_3SO_3H in acetone. The ESI-MS monitoring showed a normal distribution of various Ag_{13} nanoclusters containing a ligand combination of protonated $^{\text{Me}}\text{Py}$ and $^{n\text{-Pr}}\text{Py}$ in a ratio from 1:5 to 5:1 (Fig. 4a and Table S7†). In addition, the cross-coupling product 2-methyl-2'-propyl-4,4'-bipyridine was determined as the primary coupling species in ESI-MS spectra, wherein the minor homocoupling products 2,2'-dimethyl-4,4'-bipyridine and 2,2'-dipropyl-4,4'-bipyridine were observed as well. This ESI-MS evidence suggests that the initial formation of the Ag_{13} nanoclusters mostly relies on random combinations of two types of resurgent pyridyl diide- Ag_4 clusters as they are released from the protective macrocycles.

We next studied the transformation of diide ligands on the surface of MePyAg₁₃. The addition of CF₃SO₃H to the methanol solution of MePyAg₁₃ also led to the coupling product 2,2′-dimethyl-4,4′-bipyridine in a high yield of 67%. In order to ensure the occurrence of intra-cluster coupling, the cross experiment by using equivalent MePyAg₁₃ and n-PrPyAg₁₃ was implemented. The homo-coupling compounds 2,2′-dimethyl-4,4′-bipyridine and 2,2′-dipropyl-4,4′-bipyridine were detected as major products in the ESI-MS monitoring (Fig. 4b), while only slight cross-coupling product 2-methyl-2′-propyl-4,4′-bipyridine was observed. In contrast to the prompt inter-cluster coupling among pyridyl diide-Ag₄ clusters at the initial stage, the coupling reaction of diide ligands around a single Ag₁₃ nanocluster at the second stage is relatively sluggish as indicated by

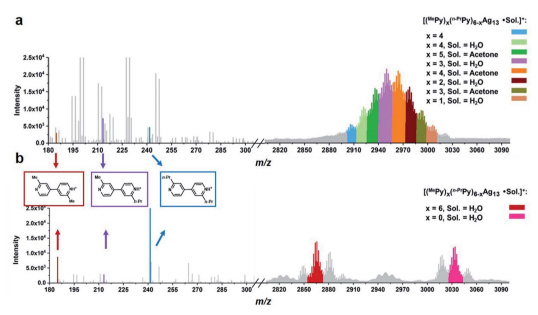


Fig. 4 ESI mass spectra of the acidifying solution samples containing equivalent (a) $^{Me}PyAg_5$ and $^{n-Pr}PyAg_5$ in acetone, and (b) $^{Me}PyAg_{13}$ and $^{n-Pr}PyAg_{13}$ in acetone.

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the detection of unreactive Ag_{13} clusters even after four days (Fig. S41†). Meanwhile, spherical nanoparticles with the sizes in the range of 20–40 nm were identified by transmission electron microscopy (TEM) images (Fig. S42†), suggesting the formation of larger nanocluster species by the association and merging of Ag_{13} nanoclusters after the occurrence of intra-cluster oxidative coupling. Furthermore, infrared spectra of the solid originated from the Ag_{13} –CF $_3$ SO $_3$ H mixture exhibit two stretching vibrations at 1540 cm $^{-1}$ and 1443 cm $^{-1}$ (Fig. S43†), which can be assigned to the characteristic absorption peaks of a pyridine ring. This result indicates that the larger Ag nanocluster species are still coordinated by pyridine ligands. Along with the prolongation of the reaction time, a black precipitate due to the further aggregation of large nanoparticles was observed.

Combining the above experimental results, we propose two phases of inter- and intra-cluster coupling accompanied by the structural evolution of silver cluster species from organic diide-Ag₄ clusters through Ag₁₃ nanoclusters to large nanoparticles. As shown in Fig. 5, the removal of protective Py[8] macrocycles by protonation makes the released organic diide-Ag₄ clusters randomly combine together and undergo a two-electron oxidative coupling to generate low valence silver atoms. The remaining protonated diide-Ag₄ clusters then assemble with the low valence silver atoms to produce the Ag₁₃ nanoclusters. In the second phase, the aryl dicarbanion ligands on the surface of a Ag₁₃ nanocluster experience protolysis at a [C-Ag₂] position and the remaining [C-Ag2] site at the para position undergoes an intra-cluster oxidative coupling to generate homo-coupling bipyridine products and expedite the formation of large nanoparticle species. It is worth noting that the oxidative coupling of organic species around in situ generated metal subnano- or nano-clusters has been proposed in many metal-catalyzed reactions.1a,h However, there has been little knowledge about the rationality of such mechanism and structural details of related intermediates. The present transformation from isolated mononuclear metal complexes through low nuclearity clusters to nanoclusters and nanoparticles is reminiscent of the dynamic structural evolution of cluster species in those metal-

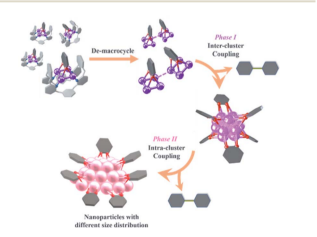


Fig. 5 Structural evolution of organosilver species from organic diide–Ag₄ clusters through nanoclusters to nanoparticles along with the occurrence of coupling reactions at different stages.

catalyzed reactions, and also gives important insights into the coupling process of organic ligands at different reaction stages.

Conclusion

In this work, a series of pyridyl vicinal dicarbanion bonded Ag₁₃ nanoclusters with an open-shell electronic structure are synthesized and characterized for the first time. The newly developed two-step synthetic method involves the occurrence of an intercluster oxidative coupling reaction and the assembly of diidecentered Ag₄ clusters with in situ generated low-valence metal atoms to generate organosilver nanoclusters. Moreover, the aryl dicarbanion ligands on the nanocluster surface further undergo an intra-cluster oxidative coupling to produce homo-coupling products and larger nanocluster species, providing a new perspective for the dynamic evolution of organometallic species in metal catalysis, from isolated metal atoms through low nuclearity organometallic clusters to nanoclusters and nanoparticles. As the in situ generated organometallic nanoclusters have been deemed to be active species in many metal-catalyzed reactions, we believe that this study not only reveals novel inter- and intra-cluster coupling steps in different stages, but also is reminiscent of the evolution of cluster species in metal-catalyzed reactions.

Experimental

Materials and methods

All commercially available chemicals were used without further purification. The solvents used in this study were processed by standard procedures. 1H-NMR and COSY experiments were carried out on a JEOL ECX-400 MHz instrument. Mass spectra were obtained using a Thermo Scientific Exactive Orbitrap instrument and TIMS-TOF mass spectrometer (Bruker Daltonics, Bremen, Germany). DOSY experiments were carried out on a Bruker Avance 600 MHz instrument using a 5 mm TXI H-C/N-D Z-GRD probe. 2D sequence for diffusion measurements were conducted using stimulated echo with 1 spoil gradient. UV-vis spectra were recorded on a Cary 7000 UV-vis-NIR spectrophotometer. EPR experiments were carried out using a JEOL JES-FA200 ESR spectrometer. Transmission electron microscopy (TEM) measurements were performed on a Hitachi H-7650 microscope. The XPS measurement was carried out in a Kratos ULTRA AXIS DLD ultrahigh vacuum photoelectron spectroscopy system. A monochromatic Al Kα X-ray (1486.6 eV) excitation source was used as an excitation source. The details of X-ray crystallographic measurements are summarized in the ESI.† Octamethylazacalix[8]pyridine (Py[8]) was synthesized according to the reported synthetic protocol by Pd-catalyzed fragment coupling reactions between a terminal dibrominated linear pentamer and a terminal diaminated linear trimer.24 Detailed synthesis procedures for aryl dicarbanion bonded Ag4 clusters and hexakis-aryl dicarbanion bonded Ag₁₃ nanoclusters are summarized below.

Synthesis of aryl dicarbanion bonded Ag₄ clusters

Complex Me PyAg₅ ([Ag₅(C₆NH₅)(Py[8])](CF₃SO₃)₃). In a 10 mL round-bottom flask, AgCF₃SO₃ (51.2 mg, 0.2 mmol) and Py[8]

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(17.0 mg, 0.02 mmol) were dissolved in 1.5 mL CH₂Cl₂/CH₃OH $(v/v \ 1:1)$ solution at room temperature. After stirring for five minutes, the propargylamine dissolved in 0.5 mL CH₂Cl₂/ CH₃OH (v/v 1:1) solution was added dropwise. Then acetone (23.2 mg, 0.4 mmol) was added to the system. After stirring for 10 hours under lucifugal conditions, a black precipitate was produced. After centrifugation, the supernatant was processed under vacuum to produce a yellow crude oily product. Yellow block crystals of complex MePyAg₅ were isolated by diffusion of

diethyl ether into a concentrated CH2Cl2/CH3OH (v/v 1:1)

solution of crude product. Yield: 61% (23.2 mg). ¹H NMR (400

MHz, acetone- d_6): δ 8.51–8.23 (m, 2H), 8.08 (m, 4H), 7.67 (s, 2H),

7.61–7.37 (m, 10H), 6.91 (d, J = 7.3 Hz, 1H), 6.85 (d, J = 7.7 Hz, 3H), 6.15 (d, J = 8.2 Hz, 1H), 6.09 (d, J = 8.0 Hz, 3H), 3.87 (s,

12H), 3.14 (s, 12H), 2.42 (s, 3H). HR-MS (ESI) calcd for C₅₆H₅₃- $Ag_5F_6N_{17}O_6S$ ([MePyAg₅-OTf⁻]⁺) 1777.8949, found 1777.9000.

The crystal structure of the complex MePyAg₅ is shown in Fig. 1. Complex $^{n-Pr}$ PyAg₅ ([Ag₅(C₈NH₉)(Py[8])](CF₃SO₃)₃). The synthetic procedure of complex **n-PrPyAg** is similar to that of MePyAg₅, but with 2-pentanone (34.3 mg, 0.4 mmol) in place of acetone. The crystallization yield of *n-PrPyAg5 as yellow block crystals is 56% (21.8 mg). 1 H NMR (400 MHz, acetone- d_{6}) δ 8.42– 8.23 (m, 3H), 8.09-7.95 (m, 6H), 7.81 (s, 1H), 7.77-7.67 (m, 3H), 7.57-7.41 (m, 8H), 7.11 (t, J = 8.1 Hz, 2H), 6.83 (d, J = 7.8 Hz, 5H), 6.08 (d, J = 7.3 Hz, 2H), 5.82 (d, J = 7.8 Hz, 2H), 3.88 (s, 12H), 3.13 (d, I = 17.0 Hz, 12H). HR-MS (ESI) calcd for C_{58} - $H_{57}Ag_5F_6N_{17}O_6S_2$ ([$^{n-Pr}PyAg_5-OTf^-$]⁺) 1805.9262, 1805.9270. The crystal structure of $^{n-Pr}$ PyAg₅ is shown in Fig. S9.†

Complex Ph PyAg₅ ([Ag₅(C₁₁NH₇)(Py[8])](CF₃SO₃)₃). Acetophenone (48.1 mg, 0.4 mmol) is used to replace acetone in the synthetic procedure of PhPyAg₅. The yield of PhPyAg₅ as yellow block crystals is 23% (9.2 mg). ¹H NMR (400 MHz, acetone-d₆) δ 8.42–8.23 (m, 3H), 8.09–7.95 (m, 6H), 7.81 (s, 1H), 7.77–7.67 (m, 3H), 7.57-7.41 (m, 8H), 7.11 (t, J = 8.1 Hz, 2H), 6.83 (d, J = 7.8 Hz, 5H), 6.08 (d, J = 7.3 Hz, 2H), 5.82 (d, J = 7.8 Hz, 2H), 3.88 (s, 12H), 3.13 (d, J = 17.0 Hz, 12H). HR-MS (ESI) calcd for $C_{61}H_{55}Ag_5F_{6}$ $N_{17}O_6S_2$ ([PhPyAg₅-OTf⁻]+) 1839.9106, found 1839.9078. The crystal structure of PhPyAg₅ is shown in Fig. S10.†

Synthesis of hexakis-aryl dicarbanion bonded Ag₁₃ nanoclusters

Complex Me PyAg₁₃ ([Ag₁₃(C₆NH₅)₆H₆](SO₃CF₃)₆). In a 5 mL glass bottle, MePyAg₅ (39.2 mg, 0.02 mmol) was dissolved in acetone at room temperature. Then trifluoromethanesulfonic acid (HOTf) (17.6 µL, 0.2 mmol) was added to the system and the solution was stirred for 5 hours under lucifugal conditions. After centrifugation, the supernatant was diffused by diethyl ether to give brown bulk crystals. Yield: 52% (4.2 mg). ¹H NMR (400 MHz, acetone- d_6): δ 8.94 (s, 1H), 8.31-8.19 (m, 10H), 7.98 (s, 1H), 2.59 (s, 15H), 2.48 (s, 3H). HR-MS (ESI) calcd for $C_{42}H_{36}Ag_{13}F_{18}N_6O_{18}S_6$ ([MePyAg₁₃]⁺) 2847.7919, found 2847.7750. The crystal structure of Me PyAg₁₃ is shown in Fig. 2. The synthetic procedures of other hexakis-aryl dicarbanion bonded Ag₁₃ nanoclusters ^{n-Pr}PyAg13 and ^{Ph}PyAg₁₃

are similar to those for MePyAg₁₃, using the corresponding aryl dicarbanion bonded Ag₄ clusters in place of MePyAg₅.

 $^{n-Pr}$ PyAg₁₃ [Ag₁₃(C₈NH₉)H₆](CF₃SO₃)₆. Yield: 47% (4.1 mg, brown bulk crystals). ¹H NMR (400 MHz, acetone- d_6): δ 8.95 (s, 1H), 8.26 (m, 10H), 7.98 (s, 1H), 2.83 (t, 12H), 1.79 (m, 12H), 0.99 (m, 18H). HR-MS (ESI) calcd for $C_{54}H_{60}Ag_{13}F_{18}N_6O_{18}S_6$ ([** **Pr**PyAg₁₃]⁺) 3017.9637, found 3018.0814. The crystal structure of ^{n-Pr}PyAg₁₃ is shown in Fig. S28.†

 $^{Ph}PyAg_{13} [Ag_{13}(C_{11}NH_7)H_6](CF_3SO_3)_6$. Yield: 23% (2.1 mg, brown bulk crystals). ¹H NMR (400 MHz, acetone- d_6): δ 8.77 (s, 6H), 8.53 (s, 6H), 7.90 (s, 12H), 7.66 (s, 18H). HR-MS (ESI) calcd for $C_{72}H_{48}Ag_{13}F_{18}N_6O_{18}S_6$ ([PhPyAg₁₃]+) 3221.8695, found 3221.8822. The crystal structure of PhPyAg₁₃ is shown in Fig. S30.†

Data availability

The X-ray crystallographic coordinates for structures reported in this work have been deposited at the Cambridge Crystallographic Data Center (CCDC), under deposition number CCDC-1996369 (MePyAg₅), CCDC-2062353 (n-PrPyAg₅), CCDC-2062351 (PhPyAg₅), CCDC-1995296 (MePyAg₁₃), CCDC-2062348 (n-PrPyAg₁₃), CCDC-2062349 (PhPyAg₁₃), CCDC-2175048 (protonated Py[8]), CCDC-1995303 (protonated 2,2'-dimethyl-4,4'bipyridine), CCDC-2060712 (protonated 2,2'-dipropyl-4,4'bipyridine). These data can be obtained free of charge from the Crystallographic Cambridge Data Centre www.ccdc.cam.ac.uk/data_request/cif. For full characterization data including UV-vis spectra, High-resolution ESI-MS, EPR, NMR, XPS, IR, TEM, DFT calculations and experimental details, see the ESI.† Any further relevant data are available from the authors upon reasonable request.

Author contributions

L. Z. conceived and supervised the project. The synthetic experiments and structural characterizations were carried out by C.-C. L. EPR measurements were performed by S. Z. R. J. and Y. X. provided assistance in ESI-MS measurements. C.-C. L., S. Z., J. T. and L. Z. co-wrote the manuscript. All authors discussed the results and commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

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