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1 Introduction

Since the advent of metal nanoclusters with atomic precision, these novel nanomaterials have garnered significant interest because of their accurate compositions/constructions and intriguing physicochemical properties.¹⁻⁸ Indeed, owing to their quantum size effect and discrete electronic energy levels, metal nanoclusters and cluster-based nanomaterials display atomic structure tunable properties, that is, slight tailoring of structures of nanoclusters can trigger remarkable differences in their performances.⁹–¹³ Besides, metal nanoclusters have been used as ideal platforms for the meticulous investigation of structure– property correlations.¹⁴⁻²⁰ Consequently, metal nanoclusters are an emerging class of programmable nanomaterials for several promising applications, such as catalysis, drug delivery, energy storage, and biological applications.²¹⁻²⁴

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Surface environment complication makes Ag_{29} nanoclusters more robust and leads to their unique packing in the supracrystal lattice†

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Silver nanoclusters have received unprecedented attention in cluster science owing to their promising functionalities and intriguing physical/chemical properties. However, essential instability significantly impedes their extensive applications. We herein propose a strategy termed "surface environment complication" to endow Ag₂₉ nanoclusters with high robustness. The Ag₂₉(S-Adm)₁₈(PPh₃)₄ nanocluster with monodentate PPh₃ ligands was extremely unstable and uncrystallizable. By substituting PPh₃ with bidentate PPh₂py with dual coordination sites (i.e., P and N), the Ag₂₉ cluster framework was twisted because of the generation of N–Ag interactions, and three $NO₃$ ligands were further anchored onto the nanocluster surface, yielding a new Ag₂₉(S-Adm)₁₅(NO₃)₃(PPh₂py)₄ nanocluster with high stability. The metal-control or ligand-control effects on stabilizing the Ag₂₉ nanocluster were further evaluated. Besides, $Ag_{29}(S-Adm)_{15}(NO_3)_{3}(PPh_2py)_{4}$ followed a unique packing mode in the supracrystal lattice with several intercluster channels, which has yet been observed in other M₂₉ cluster crystals. Overall, this work presents a new approach (i.e., surface environment complication) for tailoring the surface environment and improving the stability of metal nanoclusters. **EDGE ARTICLE**
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In the past two decades, silver nanoclusters have received unprecedented attention in cluster science.²⁵⁻⁴⁰ It is widely accepted that silver nanoclusters exhibit promising functionalities and intriguing physical/chemical properties that are obviously different from their gold counterparts.²⁷ Ag-based metal nanoclusters generally display strong photoluminescence that renders them optically active nanomaterials for sensors or biological applications.⁴¹⁻⁴³ However, Ag nanoclusters are essentially unstable relative to Au nanoclusters, which significantly impedes their extensive applications. Developing new approaches to enhance the nanocluster stability remains highly desired.

Recently, our group has developed a $M_{29}(S-Adm)_{18}(PPh_3)_4$ (where S-Adm is 1-adamantanethiol) nanocluster system for mapping the structure–property correlations at the atomic level.⁴⁴⁻⁴⁶ Although several M_{29} nanoclusters, e.g., $Pt_1Ag_{28}(S Adm_{18}(PPh_3)_4$ (Pt₁Ag₂₈-PPh₃ for short), $Au_1Ag_{28}(S-Adm)_{18}$. $PPh_3)_4$, and $Pt_1Ag_{12}Cu_{16}(S-Adm)_{18}(PPh_3)_4$, have been controllably synthesized and structurally determined, the homometal $Ag_{29}(S\text{-Adm})_{18}(PPh_3)_4$ (Ag₂₉-PPh₃ for short) nanocluster was extremely unstable and uncrystallizable.⁴⁶ We remain committed to stabilizing the homo-silver Ag_{29} nanocluster with a new approach.

Herein, a "surface environment complication" strategy has been exploited to endow the Ag_{29} nanocluster with high robustness. By substituting the monodentate $PPh₃$ (with only the P coordination site) in previously reported Ag_{29} -PPh₃ with

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bidentate PPh₂py (with P and N dual coordination sites), the nanocluster surface structure underwent a twist due to the generation of N-Ag interactions. Besides, three $NO₃$ ligands were further anchored onto the nanocluster surface, making the metallic kernel entirely wrapped. The obtained $Ag_{29}(S Adm)_{15}$ (NO₃)₃(PPh₂py)₄ (Ag₂₉-PPh₂py for short) nanocluster was much more robust relative to Ag_{29} -PPh₃, and its structure was successfully determined by single-crystal X-ray diffraction. Furthermore, based on this nanocluster template, the metalcontrol and ligand-control effects on stabilizing the Ag_{29} framework were evaluated. Moreover, at the supramolecular level, Ag_{29} -PPh₂py followed a unique packing mode in the crystal lattice with several intercluster channels, while such an aggregation pattern has yet been discovered in other M_{29} cluster crystals. Edge Article. Chemis of the Aga months of the Aga months were series are controlled to the article in the state of th

2 Experimental methods

Materials

All the following reagents were purchased from Sigma-Aldrich and used without further purification: silver nitrate $(AgNO₃,$ 99.5%, metal basis), hexachloroplatinic($\rm w$) acid ($\rm H_2PtCl_6 \cdot 6H_2O$, 99.9% metals basis), 1-adamantanethiol (Adm-SH, $C_{10}H_{15}SH$, 99%), triphenylphosphine (PPh₃, 99%), diphenyl-2pyridylphosphine (PPh2py, 97%), sodium borohydride (NaBH₄, 99%), methylene chloride (CH₂Cl₂, HPLC grade), methanol (CH₃OH, HPLC grade), ethanol (CH₃CH₂OH, HPLC grade), and *n*-hexane $(C_6H_{12}$, HPLC grade).

Synthesis of $Ag_{29}(S-Adm)_{18}(PPh_3)_4$ (Ag₂₉-PPh₃)

The preparation of Ag_{29} -PPh₃ was based on a reported method.⁴⁶

Synthesis of $Pt_1Ag_{28}(S-Adm)_{18}(PPh_3)_4$ ($Pt_1Ag_{28}-PPh_3$)

The preparation of Pt_1Ag_{28} -PPh₃ was based on a reported method.⁴⁶

Preparation of $Ag_{29}(S-Adm)_{15}(NO_3)_3(PPh_2py)_4 (Ag_{29}-PPh_2py)$

In a 50 mL round-bottom flask, 94 mg of AgNO₃ was dissolved in 5 mL of MeOH and 10 mL of EtOH, and 50 mg of Adm-SH was added under vigorous stirring. After 20 min, 100 mg of PPh_2py was added. Shortly after this, 10 mg of NaBH₄ (dissolved in 1 mL of EtOH) was poured in, and the reaction was continued for 12 hours. The obtained solution was centrifuged at 10 000 rpm for 5 minutes, and then the supernatant was collected and evaporated to get the dry product, which was then washed several times with *n*-hexane to get the final product, *i.e.*, Ag_{29} -PPh₂py. The yield was about 30% based on the Ag element (calculated from $AgNO₃$).

Preparation of Pt₁Ag₂₈(S-Adm)₁₈(PPh₂py)₄ (Pt₁Ag₂₈-PPh₂py)

94 mg of AgNO₃ used to synthesize Ag_{29} -PPh₂py was substituted by 94 mg of AgNO₃ and 10 mg of $H_2PtCl_6.6H_2O$. Other conditions remained unchanged. The yield for the synthesis of Pt₁Ag₂₈-PPh₂py was about 45% based on the Ag element (calculated from $AgNO₃$).

Crystallization of the Ag_{29} nanocluster series

Single crystals of Ag_{29} -PPh₂py or Pt₁Ag₂₈-PPh₂py were cultivated at -4 °C by liquid-diffusing *n*-hexane into the CH₂Cl₂ solution of each nanocluster. After a week, red crystals were collected, and the structures of these nanoclusters were determined. Of note, in order to accelerate the crystallization process and improve the crystal quality, the counterions $(i.e., Cl^-)$ in these nanoclusters were replaced by $\mathrm{SbF_6}^-$ or $\mathrm{BPh_4}^{-1.47}$ The reaction equation was $[Ag_{29}(S-Adm)_{15}(NO_3)_3(PPh_2py)_4]Cl_3 + 3SbF_6^- \rightarrow [Ag_{29}(S-Adm)_{15}(-S_3)^2]$ $NO₃$ ₃(PPh₂py)₄](SbF₆)₃ + 3Cl⁻ or [Pt₁Ag₂₈(S-Adm)₁₈(PPh₂py)₄]Cl₂ + $2BPh_4^- \to [Pt_1Ag_{28}(S-Adm)_{18}(PPh_2py)_4](BPh_4)_2 + 2Cl^-.$

Characterization

The optical absorption spectra of nanoclusters were recorded using an Agilent 8453 diode array spectrometer.

Electrospray ionization mass spectrometry (ESI-MS) measurements were performed by using a Waters XEVO G2-XS QTof mass spectrometer. The sample was directly infused into the chamber at 5 μ L min⁻¹. For preparing the ESI samples, nanoclusters were dissolved in $\mathrm{CH_2Cl_2}\,(\mathrm{1\ mg\ mL^{-1}})$ and diluted $(v/v = 1 : 1)$ with CH₃OH.

Infrared (IR) measurements were recorded on a Bruker Vertex 80sv Fourier transform IR spectrometer.

X-ray crystallography

The data collection for single-crystal X-ray diffraction (SC-XRD) of Ag₂₉-PPh₂py was carried out on a Bruker Smart APEX II CCD diffractometer under a nitrogen flow, using graphitemonochromatized Mo K α radiation ($\lambda = 0.71073$ Å). The data collection for single-crystal X-ray diffraction (SC-XRD) of $Pt₁Ag₂₈$ -PPh₂py was carried out on a Stoe Stadivari diffractometer under a nitrogen flow, using graphite-monochromatized Cu K α radiation ($\lambda = 1.54186$ A). Data reductions and absorption corrections were performed using the SAINT and SADABS programs, respectively. The structure was solved by direct methods and refined with full-matrix least squares on F^2 using the SHELXTL software package. All non-hydrogen atoms were refined anisotropically, and all the hydrogen atoms were set in geometrically calculated positions and refined isotropically using a riding model. All crystal structures were treated with PLATON SQUEEZE. The diffuse electron densities from these residual solvent molecules were removed. The CCDC number of the Ag_{29} -PPh₂py nanocluster is 2115749. The CCDC number of the Pt_1Ag_{28} -PPh₂py nanocluster is 2117814.

3 Results and discussion

 Ag_{29} -PPh₃ was prepared by a literature method.⁴⁶ Although the Ag₂₉-PPh₃ nanocluster was uncrystallizable because of its weak stability, several of its alloyed derivatives have been structurally determined, including Pt_1Ag_{28} -PPh₃, Au₁Ag₂₈(S-Adm)₁₈(PPh₃)₄, and $Pt_1Ag_{12}Cu_{16}(S-Adm)_{18}(PPh_3)_4$.⁴⁴⁻⁴⁶ In this context, alloying has been used as an efficient approach to improve the stability of the M₂₉ framework.⁴⁶ Fig. 1 depicts the proposed structure of Ag_{29} -PPh₃. Of note, the Ag₁₃ kernel in Ag_{29} -PPh₃ might follow a FCC (face-centered cubic) configuration for two reasons: (i) the

Fig. 1 Structural comparison between unstable Ag₂₉-PPh₃ and robust Ag₂₉-PPh₂py. Compared with Ag₂₉-PPh₃, the Ag₂₉-PPh₂py nanocluster contained retained S-Adm ligands, while the surface PPh₃ ligands were altered to PPh₂py, and several NO₃ ligands were arranged on the nanocluster surface. Color legends: light blue sphere, Ag; red sphere, S; magenta sphere, P; blue sphere, N; green sphere, O. For clarity, all C and H atoms are omitted. Of note, the structure of Ag_{29} -PPh₃ is proposed in this figure.

consistent FCC configuration of the M_{13} kernel in PPh₃ and S-Adm co-stabilized M_{29} nanoclusters,⁴⁴⁻⁴⁶ and (ii) the different absorption profiles of Ag_{29} -PPh₃ and Ag_{29} -PPh₂py (discussed below). However, such a verification calls for more experimental efforts.

At the same time, we unremittingly made efforts to stabilize the homo-silver Ag_{29} and determine its atomically precise structure. Considering that (i) the unchanging S-Adm ligand could retain the basic framework of the Ag_{29} nanocluster^{47,48} and (ii) the introduction of N-coordination sites in original ligands would generate new N–metal interactions that might enhance the structural robustness,⁴⁹⁻⁵² we were motivated to substitute the PPh₃ ligand with PPh₂py while retaining the S-Adm ligand in the nanocluster synthesis. A new Ag_{29} nanocluster, formulated as $Ag_{29}(S-Adm)_{15}(NO_3)_3(PPh_2py)_4$ $(Ag_{29}$ PPh₂py), was synthesized and further structurally determined owing to its high stability (Fig. 1 and S1†).

Compared with Ag_{29} -PPh₃, Ag_{29} -PPh₂py contained three fewer S-Adm ligands and three more $NO₃$ ligands, and the number of the phosphine ligands retained was four (Fig. 1). Because of the interactions between N (in PPh₂py) and Ag (in the cluster), the surface structure of Ag₂₉-PPh₂py displayed more obvious distortion relative to Ag_{29} -PPh₃ (Fig. 1 and S2 \dagger). Besides, three $NO₃$ ligands were observed on the nanocluster surface via Ag–O interactions. For the three O atoms in each NO3, the two inward O linked to two Ag atoms or one Ag atom, while the outward O was naked (Fig. 1 and S2†). The presence of $NO₃$ in the cluster system has been verified by IR measurement (Fig. S3†). ESI-MS measurement was performed to validate the molecular composition and determine the valence state of the nanocluster. As shown in Fig. S4,† the experimental mass signals at 2292.30 and 2271.64 Da matched well with the theoretical results of $[Ag_{29}(S\text{-Adm})_{15}(\text{NO}_3)_3(\text{PPh}_2\text{py})_4]^{3+}$ and $[Ag_{29}(S\text{-Adm})_{15}(\text{NO}_3)_3(\text{PPh}_2\text{py})_4]^{3+}$ Adm) $_{15}$ [NO $_{3})_{2}$ [PPh $_{2}$ py) $_{4}$]³⁺, respectively. In this context, the NO $_{3}$ ligand on the nanocluster surface was more prone to be

dissociated relative to S-Adm and PPh₂py ligands. Besides, the "+3" valence state of Ag_{29} -PPh₂py was tallied with the presence of $3SbF_6^-$ counterions with an Ag₂₉ cluster molecule in the crystal lattice (Fig. S1†). According to the valence state of the Ag₂₉-PPh₂py nanocluster, its nominal electron count was determined to be 8^{53} *i.e.*, $29(Ag) - 15(SR) - 3(NO_3) - 3(charge)$ $=$ 8e, the same as that of Ag_{29} -PPh₃.

Structurally, the Ag_{29} -PPh₂py nanocluster contains an icosahedral Ag₁₃ kernel (Fig. 2A). Of note, for other structurally determined $M_{29}(S-Adm)_{18}(PR_3)_4$ nanoclusters, their Ag₁₃ kernels follow a FCC configuration.⁴⁶ The difference between these two kernel configurations originates from their distinguishable surface environments via a "surface-kernel structure transfer effect". The Ag_{13} kernel of Ag_{29} -PPh₂py is first wrapped by three same $Ag_4(S-Adm)_2(PPh_2py)_1$ motif structures that are further fixed by three S-Adm bridges (Fig. 2B and C), giving rise to an $Ag_{25}(S-Adm)_{9}(PPh_{2}py)_{3}$ structure (Fig. 2D). Such three Ag₄(S-Adm)₂(PPh₂py)₁ motifs or three S-Adm bridges are in C_3 axial symmetry. Besides, an $Ag_4(S-Adm)_6(PPh_2py)_1$ surface unit caps the $Ag_{25}(S-Adm)_{9}(PPh_{2}py)_{3}$ structure to present an $Ag_{29}(S Adm)_{15}(PPh_3py)_4$ structure (Fig. 2E and F). In this context, the four PPh_2py ligands follow different bonding modes in the nanocluster framework: three PPh₂py are dually bonded onto the nanocluster via both Ag–P and Ag–N interactions, while the remaining one is singly bonded onto the nanocluster vertex via the Ag–P interaction (Fig. S2†). Of note, the $Ag_{29}(S-Adm)_{15}$. $PPh_3py)_4$ structure is still bare to a certain extent, and three NO_3 ligands, which originated from the $AgNO₃$ reactant, are further anchored onto the nanocluster surface (Fig. 2G), making the Ag₂₉ kernel fully protected and yielding the overall structure of Ag_{29} -PPh₂py (Fig. 2H). The complete structure of Ag_{29} -PPh₂py follows a C_3 axial symmetry, and the axis of the symmetry passes through the vertex P and the innermost Ag atoms (Fig. S5†).

In the crystal lattice of Ag_{29} -PPh₂py, two nanocluster enantiomers were observed, labeled as the R-nanocluster enantiomer

Fig. 2 Structural anatomy of the Ag₂₉-PPh₂py nanocluster. (A) The icosahedral Ag₁₃ kernel. (B) and (C) The Ag₄(S-Adm)₂(PPh₂py)₁ surface and S-Adm bridge-like units. (D) The Ag₂₅(S-Adm)₉(PPh₂py)₃ structure. (E) The Ag₄(S-Adm)₆(PPh₂py)₁ surface unit. (F) Ag₂₉(S-Adm)₁₅(PPh₃py)₄ structure. (G) The surface NO₃ ligand. (H) Overall structure of the Ag₂₉-PPh₂py nanocluster. (J) and (I) The Ag₂₉-PPh₂py nanocluster enantiomers. Color legends: orange sphere, kernel Ag; light blue sphere, surface Ag; red/yellow sphere, S; magenta sphere, P; blue sphere, N; green sphere, O. For clarity, all H atoms and several C atoms are omitted.

and S-nanocluster enantiomer in Fig. 2I and J. Each type of enantiomer displayed a bilayer rotation: (i) for the S-nanocluster enantiomer, the inner-layer (*i.e.*, the Ag₄(S-Adm)₆(PPh₂py)₁) was counterclockwise while the outer-layer (i.e., assembly of three surface $Ag_1(S-Adm)$ ₁(PPh₂py)₁) was clockwise (Fig. 2I); (ii) for the R-nanocluster enantiomer, the rotations of the inner-layer and outer-layer were opposite to those of the S-nanocluster enantiomer (Fig. 2J). Since the quantities of R- and S-nanocluster enantiomers are the same in the crystal lattice, the nanocluster samples were racemic.

The Ag_{29} -PPh₃ and Ag_{29} -PPh₂py nanoclusters with distinguishable kernel structures and surface environments exhibited different optical absorptions. The CH_2Cl_2 solution of Ag_{29} -PPh₃ showed an intense absorption at 413 nm and a shoulder band at 506 nm (Fig. S6,† black line). By comparison, the CH_2Cl_2 solution of Ag₂₉-PPh₃ showed several apparent UV-vis signals at 401, 438, and 530 nm (Fig. S6,† red line). The difference in optical absorptions of these two Ag₂₉ nanoclusters suggested their distinct electronic structures.54,55 The photoluminescence properties of Ag_{29} -PPh₃ and Ag_{29} -PPh₂py nanoclusters were further compared. As shown in Fig. $S7$,† the CH_2Cl_2 solution of Ag_{29} -PPh₃ was red emissive with an intense signal at 622 nm. By comparison, the Ag_{29} -PPh₂py was non-emissive in the solution state. The different photophysical properties originated from their distinct electronic structures.^{54,55}

The thermal stability of these two Ag_{29} nanoclusters was then compared in air. As shown in Fig. 3A, the characteristic optical peaks of Ag_{29} -PPh₃ continuously decreased in the first three hours and completely disappeared within six hours, demonstrating the decomposition of the nanoclusters. In this context, the Ag_{29} -PPh₃ nanocluster was unstable. In vivid contrast, the optical absorptions of Ag_{29} -PPh₂py remained unchanged for 24 hours (Fig. 3B), which suggested the high robustness of this nanocluster. Besides, the difference in stability was primarily responsible for the crystallographic discrepancy of these two Ag₂₉ nanoclusters: the Ag₂₉-PPh₃ nanocluster was uncrystallizable, whereas the crystal structure of Ag_{29} -PPh₂py was successfully determined.

Collectively, as depicted in Fig. 4A, two approaches have been presented to endow the unstable Ag_{29} -PP h_3 nanocluster with enhanced stability: (i) the metal control approach $(e.g.,)$ from unstable Ag_{29} -PPh₃ to stable Pt_1Ag_{28} -PPh₃),⁴⁶ and (ii) the ligand control approach (i.e., from unstable Ag_{29} -PPh₃ to stable Ag_{29} -PPh₂py). These two disparately stabilizing approaches raised an interesting question: which type of the $Pt₁Ag₂₈$ nanocluster would be generated when the metal control and the ligand control were performed simultaneously in the synthesis (Fig. 4B)?

As inspired by the aforementioned results, two types of $Pt₁Ag₂₈$ nanoclusters with different surface environments might be generated (Fig. 4B): $Pt_1Ag_{28}(S-Adm)_{18}(PPh_2py)_4$ with a maintained framework or $Pt_1Ag_{28}(S-Adm)_{15}(NO_3)_3(PPh_2py)_4$ with a twisted framework. After the crystallographic analysis, we determined its structure as the framework-retained $Pt_1Ag_{28}(S-$

Fig. 3 Stability of different Ag₂₉ nanoclusters. (A) Time-dependent optical absorptions of Ag₂₉-PPh₃ in CH₂Cl₂ in air. (B) Time-dependent optical absorptions of Ag_{29} -PPh₂py in CH₂Cl₂ in air.

Fig. 4 Metal control versus ligand control on the Ag_{29} nanocluster template. (A) From unstable Ag_{29} -PPh₃ to stable Pt_1Ag_{28} -PPh₃ via metal control, or from unstable Ag₂₉-PPh₃ to stable Ag₂₉-PPh₂py via ligand control. (B) From unstable Ag₂₉-PPh₃ to stable Pt₁Ag₂₈-PPh₂py via both metal control and ligand control. Color legends: light blue sphere, Ag; dark green sphere, Pt; red sphere, S; magenta sphere, P; blue sphere, N; green sphere, O. For clarity, all C and H atoms are omitted.

 $Adm)_{18}(PPh_2py)_4$ (Pt₁Ag₂₈-PPh₂py for short). The structure of $Pt₁Ag₂₈-PPh₂pv$ was almost the same as that of $Pt₁Ag₂₈-PPh₃$ (Fig. S8†).^{44,46} Although the four PPh₂py ligands in Pt₁Ag₂₈- PPh_2 py exposed N coordination sites, these N sites remained uncoordinated in the nanocluster formation (Fig. S8†). Consequently, in the competition between metal control and ligand control in this nanocluster system, the metal control seized a dominant position (Fig. 4B). In other words, when the Pt heteroatom was introduced into the innermost region of the nanocluster, the M_{29} structure was robust enough to hinder the formation of surface Ag–N interactions, which resulted in

a retained cluster framework without any distortion. Besides, in the previously reported intercluster transformation from $Pt₁Ag₂₈-PPh₃$ into $Pt₁Ag₂₈(BDT)₁₂(PPh₃)₄$ (BDT = 1,3-benzenedithiolate), the presence of BDT afforded the kernel transformation from FCC into icosahedron.⁵⁶ In this context, for the Pt₁Ag₂₈ cluster template, the bidentate thiolate ligand (*i.e.*, BDT) showed enhanced ability for directing the nanocluster configuration relative to the bidentate phosphine ligand (i.e., PPh_2py).

The Ag₂₉-PPh₂py nanocluster molecules followed a crystallographic pattern of "lamellar eutectic" between R-nanocluster and S-nanocluster enantiomers, viewed from both x and y axes (Fig. S9A–C†). The interlayer distance along the z axis was determined to be 34.064 \AA (from cluster kernel to cluster kernel, as shown in Fig. S9B†). Significantly, the supracrystal lattice of Ag_{29} -PPh₂py showed several intercluster channels with the same diameter of 18.875 Å from the (001) crystalline plane (Fig. 5A and S9D†), which was reminiscent of the behavior of MOFs (metal– organic frameworks).57,58 However, the channel diameter should be remarkably less than 18.875 \AA due to the presence of carbon tails from peripheral ligands of nanoclusters (Fig. S10†). The intercluster channel was constructed by symmetrically assembling six cluster molecules into a hexagon, where three molecules were R-nanocluster enantiomers (marked in orange in Fig. 5B), while the other three were S-nanocluster enantiomers (marked in blue in Fig. 5B). Specifically, the intercluster hexagon was composed of two cluster-based triangles in parallel planes in opposite directions, and each triangle contained three cluster molecules in the same enantiomeric configuration (Fig. 5B and C). The intermolecular distance of the cluster-based triangle was 22.224 Å, and the interlayer distance between two adjacent triangles was 18.816 Å (Fig. 5B and C). Furthermore, the arrangement of ${\rm SbF_6}^-$ counterions in the supracrystal lattice was analyzed. As shown in Fig. $S11$, \dagger $2/3$ of SbF_6^- counterions were uniformly organized in the intercluster channels while the others were packed along the C_3 axis of symmetry of Ag_{29} -PPh₂py nanoclusters. Of note, such a hexagon-like crystallographic packing of Ag₂₉-PPh₂py cluster molecules in the supracrystal lattice was unique, which has yet been detected in other M_{29} nanocluster crystals.⁴⁴–46,48,59,60 For example, for the crystal lattice of Pt_1Ag_{28} -PPh₂py, the nanocluster molecules were packed in

Fig. 5 Packing of Ag₂₉-PPh₂py in the supracrystal lattice. (A) Crystalline packing of Ag₂₉-PPh₂py, viewed from the (001) plane. (B) Vertical and (C) lateral views of the aggregation pattern of Ag₂₉-PPh₂py molecules in the supracrystal lattice. Color legends: light blue sphere, Ag in the Snanocluster enantiomer; orange sphere, Ag in the R-nanocluster enantiomer; red sphere, S; magenta sphere, P; blue sphere, N; green sphere, O. For clarity, all C and H atoms are omitted.

a layered assembly mode from the x axis, y axis, or z axis, and no intercluster channel was detected (Fig. S12†). In this context, such unique intercluster channels may render the Pt_1Ag_{28} -PPh₂py crystals potential nanomaterials for gas adsorptionrelated applications.⁶¹–⁶⁵

4 Conclusions

In summary, a strategy termed "surface environment complication" has been exploited to render unstable Ag₂₉ highly robust. The surface structure of unstable $Ag_{29}(S-Adm)_{18}(PPh_3)_4$ underwent directional distortion due to the generation of Ag–N interactions by substituting the monodentate $PPh₃$ ligand with bidentate PPh₂py. Besides, three $NO₃$ ligands were anchored onto the nanocluster surface to entirely protect the Ag₂₉ kernel, yielding a new $Ag_{29}(S-Adm)_{15}(NO_3)_3(PPh_2py)_4$ nanocluster with high robustness. Owing to its enhanced stability, the $Ag_{29}(S-$ Adm)₁₅(NO₃)₃(PPh₂py)₄ nanocluster was crystallizable, and its atomically precise structure was successfully determined. On the supramolecular level, the $Ag_{29}(S\text{-Adm})_{15}(NO_3)_3(PPh_2py)_4$ nanocluster molecules followed a unique crystallographic packing mode and displayed several intercluster channels. This study thus presented a novel strategy for tailoring the surface environment of metal nanoclusters, and also provided fundamental insights into the controllable synthesis of highly robust silver nanoclusters. Future work will focus on promoting this strategy to other ligand-protected metal nanoclusters.

Data availability

All the data supporting this article have been included in the main text and the ESI.†

Author contributions

C. X. and Q. Y. carried out the experiments and analyzed the data. X. W., H. L. and H. S. assisted in the analysis. X. K. and M.

Z. designed the project, analyzed the data, and wrote the manuscript.

Conflicts of interest

There are no conflicts to declare.

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