

Cite this: *Chem. Sci.*, 2019, 10, 4862

All publication charges for this article have been paid for by the Royal Society of Chemistry

Carboxylic acid stimulated silver shell isomerism in a triple core–shell Ag₈₄ nanocluster†

Zhi Wang,^a Hao-Tian Sun,^a Mohamedally Kurmoo,^c Qing-Yun Liu,^d Gui-Lin Zhuang,^{id}*^b Quan-Qin Zhao,^a Xing-Po Wang,^a Chen-Ho Tung^{id}^a and Di Sun^{id}*^{ab}

Isomerization is highly important in all aspects of science, yet it is rarely observed in nanoscience. Here, we synthesized a unique triple core–shell Ag₈₄ nanocluster displaying isomerism, which is controlled by different carboxylic acids and a one-way transformation (SD/Ag84a → SD/Ag84b). The innermost core is a rare Ag₁₀ nanocluster which comprises an Ag₆ octahedral unit as seen in face-centred cubic (fcc) silver metal and four capped Ag atoms. It templates two crescent-shaped polyoxometalate (W₇O₂₆)¹⁰⁻ shells which are then enclosed in a shell of silver shaped as rugby balls. The organic ligands (PrS⁻, ¹⁴⁷PrCOO⁻ and PhCOO⁻) finally shield the metallic clusters. Due to slight differences in structure at two poles and the steric hindrance of ¹⁴⁷PrCOO⁻ and PhCOO⁻, SD/Ag84a and SD/Ag84b adopt the shapes of flat-headed and cuspidal prolate spheres, respectively. Interestingly, PhCOOH is dominant over ¹⁴⁷PrCOOH whereby crystals of SD/Ag84b were isolated if PhCOOH is added during the synthesis of SD/Ag84a. This demonstrates that PhCOOH not only alters the organic coats but also induces metal shell re-organization. This work reveals carboxylate-controlled skeletal isomerism in silver nanoclusters for the first time, thus deepening the understanding of silver nanocluster assembly, flexibility and reactivity.

Received 19th December 2018
Accepted 28th March 2019

DOI: 10.1039/c8sc05666h

rsc.li/chemical-science

Introduction

Isomerization plays a vital role in many fundamental biological processes in nature such as ligand binding, enzymatic catalysis, protein folding and photosynthesis.¹ Diverse isomerizations in inorganic, organic, and supramolecular chemistry have also attracted attention since isomers—compounds with the same atoms arranged in different manners—may deliver substantially different physicochemical properties and chemical reactivities without changing their compositions;² for example, the sweetness of optically active sugars. For polyatomic systems, such as metal nanoclusters, isomerization is not a highly

probable event because (i) structural variation usually involves the re-organization of several dozens of metal atoms and motion of several ligands and (ii) unravelling atomic-precision in nanometric structures or even macromolecules is still a major challenge. Thus, it is difficult to experimentally achieve atomic-level isomerization of specific sites/regions in a cluster without destroying the other parts of the main skeleton. An important breakthrough was made in 2015 by Jin and Wu who reported the first genuine gold nanocluster isomers, Au₃₈T and Au₃₈Q.³ Following this work, their group reported ligand-induced core isomerization in two thiolate-protected Au₂₈ nanoclusters.⁴ Tsukuda also discovered anion-packing induced isomerization between crown- and butterfly-Au₉.⁵ Although there has been sporadic advances in Au nanocluster isomerization recently,⁶ those for silver are yet to be achieved.

The development of silver nanoclusters has expedited the establishment of general synthetic methodologies such as anion-templation and geometric polyhedral principles.⁷ The gradual accumulation of knowledge about silver nanocluster synthesis is also a reminder for us to revisit the ligand strategy⁸ based on the classic hard-soft-acid-base (HSAB) theory,⁹ that is controlling the ‘hardness’ of carboxylic acids in competition with ‘soft’ thiolates, allowing flexibility of the overall silver nanoclusters. The abundant availability of commercial alkyl or aryl carboxylates provides ample options to tune the structure and flexibility of silver nanoclusters through steric hindrance or/and electronic effects. Flexible silver nanoclusters may

^aKey Laboratory of Colloid and Interface Chemistry, Ministry of Education, School of Chemistry and Chemical Engineering, State Key Laboratory of Crystal Materials, Shandong University, Jinan, 250100, People's Republic of China. E-mail: dsun@sdu.edu.cn

^bCollege of Chemical Engineering and Materials Science, Zhejiang University of Technology, Hangzhou, 310032, People's Republic of China. E-mail: glzhuang@zjut.edu.cn

^cInstitut de Chimie de Strasbourg, Université de Strasbourg, CNRS-UMR 7177, 4 rue Blaise Pascal, 67008 Strasbourg Cedex, France

^dCollege of Chemical and Environmental Engineering, Shandong University of Science and Technology, Qingdao, 266590, People's Republic of China

† Electronic supplementary information (ESI) available: IR, ¹³C NMR, UV-Vis, EDS and PXRD data, and details of the data collection and structure refinements, and crystal data. CCDC 1878911 and 1878912 for SD/Ag84a and SD/Ag84b. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c8sc05666h



produce isomers under specific stimuli including acid, base, ligand exchange and so on. Given this situation, using this soft/hard double-ligand strategy in the rational synthesis of isomeric silver nanoclusters is very attractive and challenging.

This study is born out of the successes of the recent aforementioned synthetic strategy achieving two Ag_{84} nanoclusters (**SD/Ag84a** and **SD/Ag84b**). Their impressive structures comprise a Ag_{10} nanocluster core, a pair of novel crescent-shaped $(\text{W}_7\text{O}_{26})^{10-}$ shells and a 74-silver outer shell, thus establishing a novel common rugby-ball shaped three-shell $[\text{Ag}_{10}@\text{(W}_7\text{O}_{26})_2@\text{Ag}_{74}]$ motif that differs in skeletal organization and ligand coverage at the two poles. Their flat-headed and cuspidal prolate spherical structures, respectively, are likely driven by the different steric hindrances between ${}^n\text{PrCOO}^-$ and PhCOO^- . Although the organic shells are different, these two Ag_{84} nanoclusters have identical elemental contents, and hence, they belong to pseudo-isomers. What's more interesting is that we can isolate **SD/Ag84b** in the mother solution of **SD/Ag84a** by adding PhCOOH at the second-step reaction, which demonstrates that PhCOOH not only changes the organic coverage but also induces metal shell distortion or re-organization.

Results and discussion

Synthesis of **SD/Ag84a** and **SD/Ag84b**

The details of the synthesis of **SD/Ag84a** and **SD/Ag84b** are given in the ESI.† Briefly, $({}^i\text{PrSAg})_n$, silver salts, RCOOH , and Na_2WO_4 were mixed in MeOH/DMF ($v/v = 4/1$) and reacted under solvothermal conditions (Scheme 1). Red crystals were collected and characterized by single-crystal and powder X-ray diffraction (SCXRD, PXRD), IR, UV-Vis, elemental analyses, and luminescence.

X-ray structures of **SD/Ag84a** and **SD/Ag84b**

Both **SD/Ag84a** and **SD/Ag84b** crystallize in the triclinic $P\bar{1}$ space group with half of the cluster existing as the asymmetric unit. Apart from the carboxylate group they have similar formulae of $[\text{Ag}_{10}@\text{(W}_7\text{O}_{26})_2@\text{Ag}_{74}\text{S}_2({}^i\text{PrS})_{40}({}^n\text{PrCOO})_{18}] \cdot 2\text{CH}_3\text{OH}$ (**SD/Ag84a**) and $[\text{Ag}_{10}@\text{(W}_7\text{O}_{26})_2@\text{Ag}_{74}\text{S}_2({}^i\text{PrS})_{40}(\text{PhCOO})_{18}]$ (**SD/Ag84b**). Their overall appearance is a prolate sphere with similar dimensions of about 1.1×2.5 nm. Their coverage by organic ligands is also similar to $40 {}^i\text{PrS}^-$ and 18 RCOO^- ligands (Fig. 1a and b). Both Ag_{84} clusters consist of an Ag_{74} outer shell and two crescent-shaped $(\text{W}_7\text{O}_{26})^{10-}$ inner shells clamping a subnanometer Ag_{10} kernel. The Ag_{10} kernel is constructed from an octahedral Ag_6 core by adding four Ag caps at the triangular faces (Fig. 1c). Two $(\text{W}_7\text{O}_{26})^{10-}$ anions symmetrically wrap the Ag_{10}

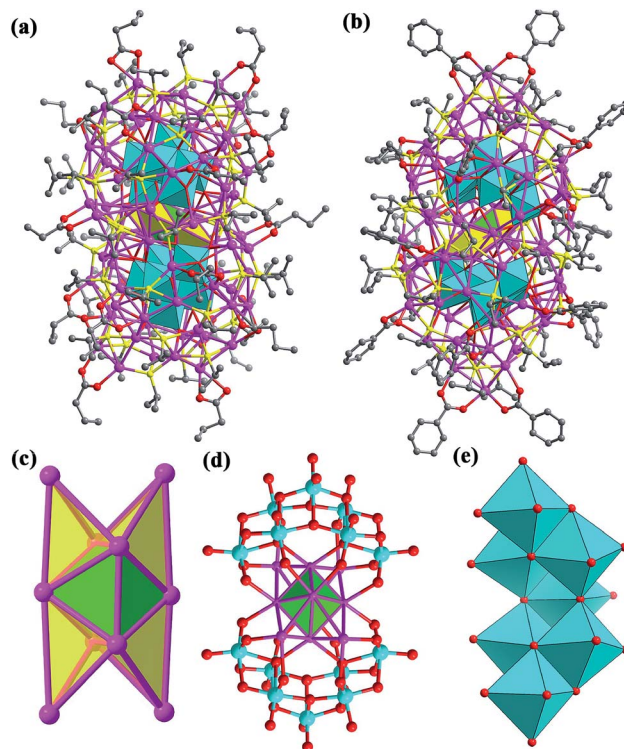
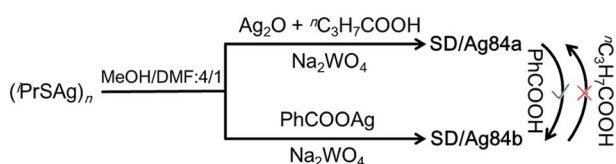


Fig. 1 Structures of the clusters **SD/Ag84a** (a) and **SD/Ag84b** (b). (c) The polyhedral mode showing an Ag_6 octahedron (green) capped by four additional silver tetrahedra (yellow) to form the Ag_{10} kernel; (d) two $(\text{W}_7\text{O}_{26})^{10-}$ anions wrapping an Ag_{10} kernel and (e) polyhedral mode showing the structure of a $(\text{W}_7\text{O}_{26})^{10-}$ anion. Color labels: purple, Ag; cyan, W; yellow, S; gray, C; and red, O.

kernel through Ag–O bonds (Fig. 1d). Each $(\text{W}_7\text{O}_{26})^{10-}$ anion is formed by seven edge-sharing WO_6 octahedra (Fig. 1e) and coordinates to 32 silver atoms in total (Fig. S1†). Such $(\text{W}_7\text{O}_{26})^{10-}$ anions are not only observed in the $\text{POM}@\text{Ag}$ cluster family for the first time but also have never been documented in classical POM chemistry. Two *in situ* generated $\mu_6\text{-S}^{2-}$ ions were inserted into the equatorial waist section, consolidating the linkage between the inner Ag_{10} kernel and the Ag_{74} outer shell (Fig. S2†). We listed several important geometry parameters of **SD/Ag84a** and **SD/Ag84b** such as $\text{Ag}\cdots\text{Ag}$ distances in the Ag_{10} kernel and the Ag_{74} shell, $\text{Ag}\text{-S}_{\text{ligand}}$, $\text{Ag}\text{-S}_{\text{sulfide}}$, $\text{Ag}\text{-O}_{\text{carboxylate}}$, $\text{Ag}\text{-O}_{\text{POM}}$, and $\text{W}\text{-O}$ bond lengths, and coordination modes of ligands and $(\text{W}_7\text{O}_{26})^{10-}$ for comparison (Table S1†).

Notably, such an Ag_{10} kernel is observed for the first time, although its innermost subvalent Ag_6^{4+} octahedron has been reported in silver nanoclusters¹⁰ and some inorganic compounds such as $\text{Ag}_6\text{Ge}_{10}\text{P}_{12}$, Ag_5GeO_4 , Ag_5SiO_4 and Ag_6O_2 .¹¹ The formation of such subvalent Ag nanoclusters is related to the reductive effect of DMF, which has been recognized as a key factor in the controlled synthesis of multiple-twin silver nanocrystals by reducing $\text{Ag}^+ \rightarrow \text{Ag}^0$.¹² Based on the digested ^{13}C NMR spectra of reaction mother solutions of **SD/Ag84a** and **SD/Ag84b** (Fig. S3 and S4†), we observed typical $\text{C}_{\text{carboxyl}}$ resonances at $\delta = 163.16$ and 163.04 ppm corresponding to the oxidation product of DMF, Me_2NCOOH .¹³ In the same region, the



Scheme 1 Schematic representation of the assembly and conversion of **SD/Ag84a** and **SD/Ag84b**.



expected C_{aldehyde} of DMF and C_{carboxyl} of ${}^n\text{PrCOOH}$ and PhCOOH were also observed at $\delta = 164.39$, 176.47, and 168.95 ppm, respectively. These results provided experimental evidence that the redox reaction happened in such a complicated self-assembly process. As the smallest unit in fcc bulk silver metal, the subnanometer Ag_6 octahedron with eight exposed [111] facets can be seen as an embryonic state of any other bigger silver nanocrystals and thus is of particular interest

in the field of nanoparticles.¹⁴ The subnanometer Ag_{10} kernel trapped in **SD/Ag84a** and **SD/Ag84b** can be seen as the Ag_6 “nuclei” grown by adding four tetrahedra on its four [111] facets, manifesting the atomic-level silver nanocrystal growth route, that is, stepwise growth of tetrahedral caps on specific facets. Such a growth route is quite similar to that proposed for larger decahedral and icosahedral silver nanocrystals by the Tsuji group.¹⁵ This result thus sheds light on the atomic details of the growth of silver nanocrystals in the embryonic stage.

Upon further carefully checking and comparing the structural features such as ligand distributions and skeletons of these two silver nanoclusters, we surprisingly found that the distinct differences of silver skeletons between **SD/Ag84a** and **SD/Ag84b** are in the polar sections and have nothing to do with the inner Ag_{10} kernel and $(\text{W}_7\text{O}_{26})^{10-}$, although the silver polygons on the other regions of surfaces are almost identical with slight dislocations and distortions. The Ag_{74} shells in **SD/Ag84a** and **SD/Ag84b** can be described as flat-headed (red skeleton in Fig. 2a) and cuspidal (green skeleton in Fig. 2a) prolate spheres, respectively. The superposed Ag_{74} shells, especially in the polar regions (Ag_{15} caps, Fig. 2b), showed silver polygon migrations and severe distortions, which are caused by the different distributions of ${}^1\text{PrS}^-$ and RCOO^- ligands on this region. There are in total six ${}^1\text{PrS}^-$ and six RCOO^- ligands on the polar regions of **SD/Ag84a** and **SD/Ag84b**. On one pole of **SD/Ag84a**, six ${}^n\text{PrCOO}^-$ ligands are equally distributed at the two sides (Fig. 2c), whereas four PhCOO⁻ ligands are located at one side and the other two at another side of one pole of **SD/Ag84b** (Fig. 2d). Moreover, two $\mu_2\text{-}\kappa^1\text{-}\kappa^1$ PhCOO⁻ ligands are adjacent

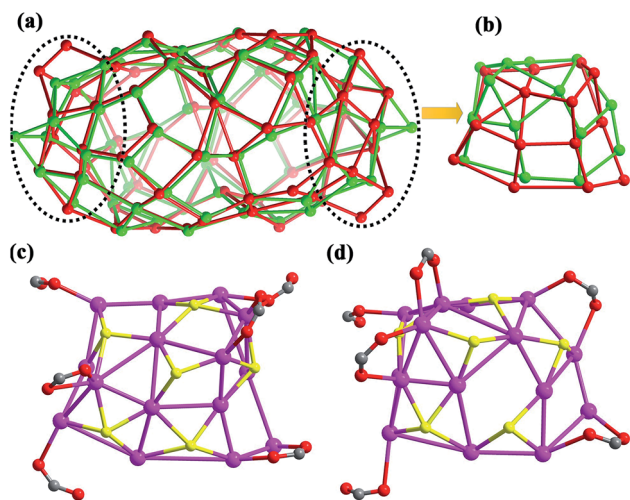


Fig. 2 (a) The superposed Ag_{74} shells of **SD/Ag84a** (red) and **SD/Ag84b** (green). (b) The top views of silver polygons at the pole regions of **SD/Ag84a** (red) and **SD/Ag84b** (green). The top views of ligand distributions at the pole region of **SD/Ag84a** (c) and **SD/Ag84b** (d).

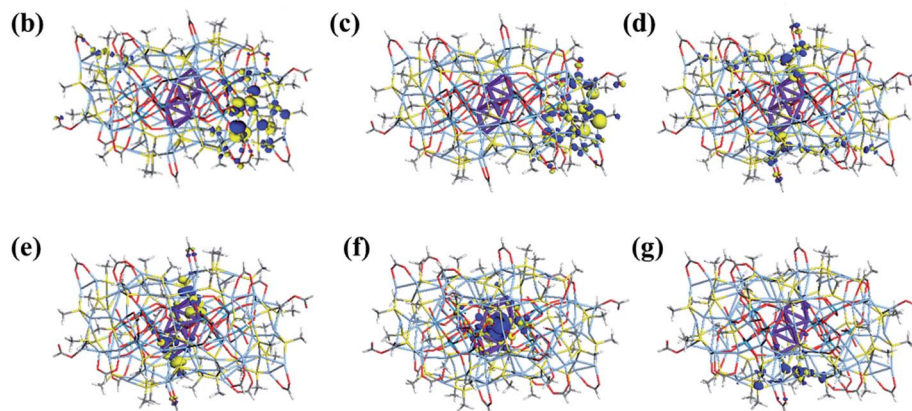
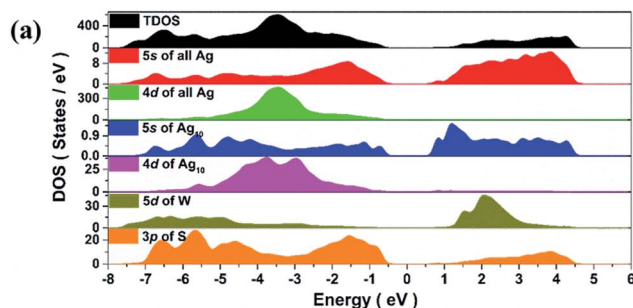


Fig. 3 Total DOS and partial DOS of **SD/Ag84a** (a). Frontier molecular orbitals: HOMO–2 (b), HOMO–1 (c), HOMO (d) and LUMO (e), LUMO+1 (f) and LUMO+2 (g).



and simultaneously coordinate to the same Ag atom (Ag32), creating a single peak (Ag32) on the pole of **SD/Ag84b**. Although the organic coats are different in **SD/Ag84a** and **SD/Ag84b**, the same silver atom counts suggested that they are cage isomers. The driving force of the isomeric silver nanocluster should be most probably generated from the steric hindrance of different carboxylate groups. Thus, we successfully filled in the blanks in isomeric silver nanoclusters using a soft/hard double-ligand strategy.

Inspired by the isomeric silver skeletons of **SD/Ag84a** and **SD/Ag84b**, we also tried to explore the possibility of conversion between them under the carboxylic acid stimulus. The isomerization experiments were performed in respective reaction mother liquors without removing crystals and by adding a portion of another kind of carboxylic acid. Interestingly, we found that **SD/Ag84b** can be isolated after the synthesis of **SD/Ag84a** by adding bulkier PhCOOH for the second-step reaction; however, the transformation from **SD/Ag84b** to **SD/Ag84a** failed by adding smaller ⁿPrCOOH into the system for synthesizing **SD/Ag84b**, which indicated that the bulkier the carboxylic acid, the stronger the inducing effect.

Considering the subvalent characteristics of the Ag₁₀ kernel, we deduced the charge of it to be +4 based on the formula determined by high-quality SCXRD data, which was also further confirmed by the DFT calculations (see details in ESI†). The electronic structure of **SD/Ag84a** (Fig. 3a) demonstrates that both 5s states of the outer Ag₇₄ shell and 3p states of S play a pivotal role in the valence band maximum (VBM), while only the 5s states of Ag (especially from the inner Ag₁₀ kernel) are dominant in the conduction band minimum (CBM). The resultant band gap of 1.12 eV is comparable with the observed value ($E_g = 1.24$ eV) from solid UV-Vis measurement results discussed below. Thus, the corresponding absorption peak can be attributed to electronic transition from the outer Ag₇₄ shell to the inner Ag₁₀ kernel. Moreover, frontier molecular orbitals (Fig. 3b–g) also indicate that the three highest occupied orbitals (HOMO, HOMO–1 and HOMO–2) are concentrated in the 5s orbitals and 4d orbitals of the outer Ag₇₄ shell, while the two lowest unoccupied orbitals (LUMO and LUMO+1) mainly consist of 5s orbitals (derived from octahedron-like Ag₆) and 4d orbitals (involving two wing-like Ag₂ units) of the inner Ag₁₀ kernel. Therefore, it is concluded that the inner Ag₁₀ core features an electron-deficient state, consistent with the +4 valence estimated crystallographically.

The optical properties of **SD/Ag84a**

The solid state UV-Vis absorption spectrum of **SD/Ag84a** was recorded at room temperature in the wavelength range from 300 to 1000 nm. As depicted in Fig. 4, there is one intense absorption band centered at 361 nm and a shoulder peak at 500 nm, respectively. The high energy absorption centered at 361 nm originates from the $n \rightarrow \pi^*$ transition of ⁱPrS⁻, as similarly observed in the absorption spectrum of the (ⁱPrSAg)_n precursor. The low energy absorption centered at 500 nm can be attributed to the charge transfer transition from the S 3p to Ag 5s orbitals. Using a transformed Kubelka–Munk plot (Fig. S5†), the HOMO–

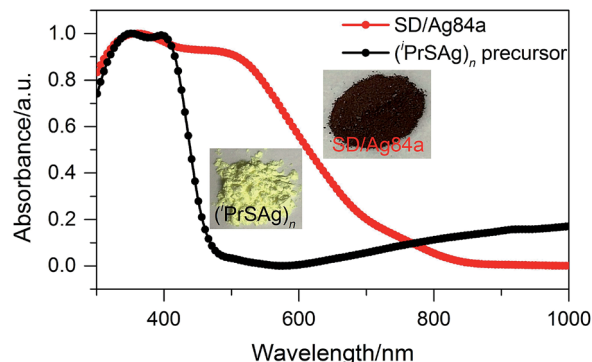


Fig. 4 The UV-Vis absorption spectra of **SD/Ag84a** and the (ⁱPrSAg)_n precursor.

LUMO gap of **SD/Ag84a** was found to be 1.24 eV, indicating that **SD/Ag84a** is a potential narrow-band-gap semiconductor. By contrast, the optical gap of the (ⁱPrSAg)_n precursor was calculated to be 2.52 eV, which matches with the color of the samples: **SD/Ag84a** is red, whereas the (ⁱPrSAg)_n precursor is yellow (see the insets of Fig. 4).

SD/Ag84a does not emit at room temperature in the solid state under a 365 nm hand-held UV light; however, it emits

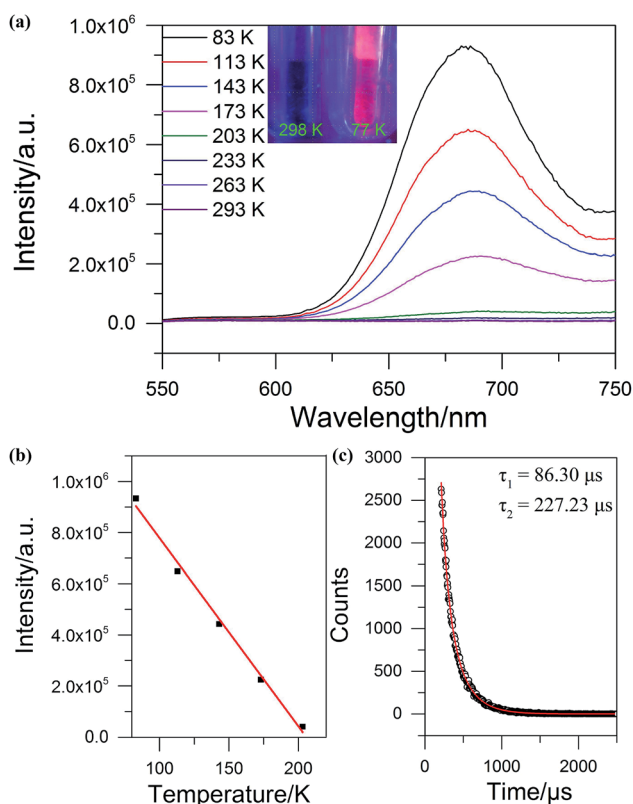


Fig. 5 (a) The temperature-dependent emission spectra of **SD/Ag84a** under 468 nm excitation. Insets: the photographs of **SD/Ag84a** irradiated with 365 nm UV light at 298 and 77 K. (b) The plot of temperature vs. maximum emission intensity (red line is the linear fitting in the range of 83–203 K). (c) Luminescence lifetime of **SD/Ag84a** recorded at 83 K (red line is the fitting curve).



bright-red light at 77 K, which is detectable by the naked eye (see the insets of Fig. 5). The temperature dependent fluorescence spectra were recorded under 468 nm excitation in the temperature range of 293–83 K. From Fig. 5a, we can see that **SD/Ag84a** is almost non-emissive from 293 to 203 K. Upon cooling from 203 to 83 K, **SD/Ag84a** starts emitting with the maximum emission band centered at 689 nm, which is gradually blue-shifted to 685 nm at 83 K along with the increase of luminous intensity. This low-temperature emission should originate from the ligand-to-metal charge transfer (LMCT, charge transfer from S 3p to Ag 5s) perturbed by Ag \cdots Ag interactions.¹⁶ The temperature dependence behavior should be relevant with variable molecular rigidity and Ag \cdots Ag interactions at different temperatures. The emission intensity was found to be sensitive to temperature and showed good linearity with the corresponding temperature ranging from 83 to 203 K (Fig. 5b), with the linear equation $I_{\max} = 1511090 - 7364.75T$. The linear equation correlation coefficient is 0.992, which is suitable for temperature detection in the temperature range of 83–203 K. The fluorescence lifetime of **SD/Ag84a** was measured at 83 K (Fig. 5c), with the lifetime value falling on the scale of microseconds ($\tau_1 = 86.30 \mu\text{s}$ and $\tau_2 = 227.23 \mu\text{s}$), suggesting a triplet state emission.

Conclusions

In conclusion, we successfully achieved the assembly of isomeric Ag₈₄ nanoclusters for the first time by the combination of anion template and soft/hard double-ligand strategies. The comparative structural analysis indicated that the isomerism mainly occurred on the polar regions of the prolate spheres. The differences are found in both the silver skeleton and ¹PrS[−]/RCOO[−] ligand distributions. The driving force for such an isomerism is dominated by the steric hindrance of carboxylates. What's more interesting is that the Ag₁₀ kernel as a grown nanocrystal from the smallest octahedral Ag₆ unit in face-centred cubic (fcc) silver metal was first identified in the innermost region of the Ag₈₄ nanocluster. The results obtained in this study are not only the pioneering results for isomeric silver nanoclusters but also enrich the library of multi-shell silver nanoclusters containing both novel POM templates and ultrasmall reductive silver nanocrystals.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was financially supported by the National Natural Science Foundation of China (Grant No. 21822107, 21571115, 21671172), the Natural Science Foundation of Shandong Province (No. JQ201803 and ZR2017MB061), the Taishan Scholar Project of Shandong Province of China, the Qilu Youth Scholar Funding of Shandong University and the Fundamental Research Funds of Shandong University (104.205.2.5). Dr

Zhuang was financially supported by the Zhejiang Provincial Natural Science Foundation (No. LR19B010001).

References

- (a) D. B. Ritchie and M. T. Woodside, *Curr. Opin. Struct. Biol.*, 2015, **34**, 43–51; (b) S. W. Englander and L. Mayne, *Proc. Natl. Acad. Sci. U. S. A.*, 2017, **114**, 8253–8258; (c) P. Leippe and J. A. Frank, *Curr. Opin. Struct. Biol.*, 2019, **57**, 23–30; (d) N. Ankenbruck, T. Courtney, Y. Naro and A. Deiters, *Angew. Chem., Int. Ed.*, 2018, **57**, 2768–2798; (e) L. A. Acevedo, J. Kwon and L. K. Nicholson, *Proc. Natl. Acad. Sci. U. S. A.*, 2019, **116**, 2589–2594.
- (a) B. Moulton and M. J. Zaworotko, *Chem. Rev.*, 2001, **101**, 1629–1658; (b) J. A. DeVine, M. L. Weichman, B. Laws, J. Chang, M. C. Babin, G. Balerdi, C. J. Xie, C. L. Malbon, W. C. Lineberger, D. R. Yarkony, R. W. Field, S. T. Gibson, J. Y. Ma, H. Guo and D. M. Neumark, *Science*, 2017, **358**, 336–339; (c) Y.-Z. Tan, Z.-J. Liao, Z.-Z. Qian, R.-T. Chen, X. Wu, H. Liang, X. Han, F. Zhu, S.-J. Zhou, Z. Zheng, X. Lu, S.-Y. Xie, R.-B. Huang and L.-S. Zheng, *Nat. Mater.*, 2008, **7**, 790–794; (d) L.-Y. Yao and V. W.-W. Yam, *J. Am. Chem. Soc.*, 2015, **137**, 3506–3509; (e) J.-P. Zhang, X.-C. Huang and X.-M. Chen, *Chem. Soc. Rev.*, 2009, **38**, 2385–2396; (f) C. B. Williamson, D. R. Nevers, A. Nelson, I. Hadar, U. Banin, T. Hanrath and R. D. Robinson, *Science*, 2019, **363**, 731–735; (g) B. Zhang, T. Zhu, M. Ou, N. Rowell, H. Fan, J. Han, L. Tan, M. T. Dove, Y. Ren, X. Zuo, S. Han, J. Zeng and K. Yu, *Nat. Commun.*, 2018, **9**, 2499.
- S. Tian, Y.-Z. Li, M.-B. Li, J. Yuan, J. Yang, Z. Wu and R. Jin, *Nat. Commun.*, 2015, **6**, 8667.
- Y. Chen, C. Liu, Q. Tang, C. Zeng, T. Higaki, A. Das, D.-e. Jiang, N. L. Rosi and R. Jin, *J. Am. Chem. Soc.*, 2016, **138**, 1482–1485.
- S. Yamazoe, S. Matsuo, S. Muramatsu, S. Takano, K. Nitta and T. Tsukuda, *Inorg. Chem.*, 2017, **56**, 8319–8325.
- (a) X.-Y. Liu, Y. Yang, Z. Lei, Z.-J. Guan and Q.-M. Wang, *Chem. Commun.*, 2016, **52**, 8022–8025; (b) L.-Q. Mo, J.-H. Jia, L.-j. Sun and Q.-M. Wang, *Chem. Commun.*, 2012, **48**, 8691–8693; (c) S. Zhuang, L. Liao, J. Yuan, N. Xia, Y. Zhao, C. Wang, Z. Gan, N. Yan, L. He, J. Li, H. Deng, Z. Guan, J. Yang and Z. Wu, *Angew. Chem., Int. Ed.*, 2019, **58**, 4510–4514.
- (a) G.-G. Gao, P.-S. Cheng and T. C. W. Mak, *J. Am. Chem. Soc.*, 2009, **131**, 18257–18259; (b) C. W. Liu, H.-W. Chang, C.-S. Fang, B. Sarkar and J.-C. Wang, *Chem. Commun.*, 2010, **46**, 4571–4573; (c) Y.-P. Xie and T. C. W. Mak, *Angew. Chem., Int. Ed.*, 2012, **51**, 8783–8786; (d) Y.-P. Xie, J.-L. Jin, G.-X. Duan, X. Lu and T. C. W. Mak, *Coord. Chem. Rev.*, 2017, **331**, 54–72; (e) Y.-P. Xie and T. C. W. Mak, *J. Am. Chem. Soc.*, 2011, **133**, 3760–3763; (f) H. Liu, C.-Y. Song, R.-W. Huang, Y. Zhang, H. Xu, M.-J. Li, S.-Q. Zang and G.-G. Gao, *Angew. Chem., Int. Ed.*, 2016, **55**, 3699–3703; (g) R.-W. Huang, Q.-Q. Xu, H.-L. Lu, X.-K. Guo, S.-Q. Zang, G.-G. Gao, M.-S. Tang and T. C. W. Mak, *Nanoscale*, 2015, **7**, 7151–7154; (h) R.-W. Huang, Y.-S. Wei, X.-Y. Dong,



- X.-H. Wu, C.-X. Du, S.-Q. Zang and T. C. W. Mak, *Nat. Chem.*, 2017, **9**, 689–697; (i) C.-Y. Gao, X. He, L. Zhao and M.-X. Wang, *Chem. Commun.*, 2012, **48**, 8368–8370; (j) Q.-M. Wang, Y.-M. Lin and K.-G. Liu, *Acc. Chem. Res.*, 2015, **48**, 1570–1579.
- 8 (a) J.-W. Liu, H.-F. Su, Z. Wang, Y.-A. Li, Q.-Q. Zhao, X.-P. Wang, C.-H. Tung, D. Sun and L.-S. Zheng, *Chem. Commun.*, 2018, **54**, 4461–4464; (b) Y.-M. Su, W. Liu, Z. Wang, S.-A. Wang, Y.-A. Li, F. Yu, Q.-Q. Zhao, X.-P. Wang, C.-H. Tung and D. Sun, *Chem.–Eur. J.*, 2018, **24**, 4967–4972.
- 9 R. G. Pearson, *J. Am. Chem. Soc.*, 1963, **85**, 3533–3539.
- 10 (a) D. Sun, G.-G. Luo, N. Zhang, R.-B. Huang and L.-S. Zheng, *Chem. Commun.*, 2011, **47**, 1461–1463; (b) Y. Kikukawa, Y. Kuroda, K. Suzuki, M. Hibino, K. Yamaguchi and N. Mizuno, *Chem. Commun.*, 2013, **49**, 376–378; (c) H. Yang, J. Lei, B. Wu, Y. Wang, M. Zhou, A. Xia, L. Zheng and N. Zheng, *Chem. Commun.*, 2013, **49**, 300–302.
- 11 (a) H. G. V. Schnering and K. G. Hausler, *Rev. Chim. Miner.*, 1976, **13**, 71–81; (b) W. Beesk, P. G. Jones, H. Rumpel, E. Schwarzmann and G. M. Sheldrick, *J. Chem. Soc., Chem. Commun.*, 1981, 664–665; (c) M. Jansen and C. Linke, *Angew. Chem., Int. Ed.*, 1992, **31**, 653–654; (d) C. Linke and M. Jansen, *Inorg. Chem.*, 1994, **33**, 2614–2616.
- 12 I. Pastoriza-Santos and L. M. Liz-Marzan, *Nano Lett.*, 2002, **2**, 903–905.
- 13 I. Pastoriza-Santos and L. M. Liz-Marzan, *Pure Appl. Chem.*, 2000, **72**, 83–90.
- 14 A. Tao, P. Sinsermsuksakul and P. Yang, *Angew. Chem., Int. Ed.*, 2006, **45**, 4597–4601.
- 15 (a) M. Tsuji, Y. Maeda, S. Hikino, H. Kumagae, M. Matsunaga, X.-L. Tang, R. Matsuo, M. Ogino and P. Jiang, *Cryst. Growth Des.*, 2009, **9**, 4700–4705; (b) M. Tsuji, M. Ogino, R. Matsuo, H. Kumagae, S. Hikino, T. Kim and S.-H. Yoon, *Cryst. Growth Des.*, 2010, **10**, 296–301; (c) M. Tsuji, X. Tang, M. Matsunaga, Y. Maeda and M. Watanabe, *Cryst. Growth Des.*, 2010, **10**, 5238–5243.
- 16 (a) V. W.-W. Yam, V. K.-M. Au and S. Y.-L. Leung, *Chem. Rev.*, 2015, **115**, 7589–7728; (b) A. Barbieri, G. Accorsi and N. Armaroli, *Chem. Commun.*, 2008, 2185–2193.

