Regulation of Surface Structure of \([\text{Au}_{9}\text{Ag}_{12}(\text{SAdm})_{4}(\text{Dppm})_{6}\text{Cl}_{6}]\)(\text{SbF}_{6})_{3}\) Nanocluster via Alloying

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Tailoring of specific sites on the nanocluster surface can tailor the properties of nanoclusters at the atomic level, for the in-depth understanding of structure and property relationship. In this work, we explore the regulation of surface structure of \([\text{Au}_{9}\text{Ag}_{12}(\text{SAdm})_{4}(\text{Dppm})_{6}\text{Cl}_{6}]\)(\text{SbF}_{6})_{3}\) nanocluster via alloying. We successfully obtained the well-determined tri-metal \([\text{Au}_{9}\text{Ag}_{8}@\text{Cu}_{4}(\text{SAdm})_{4}(\text{Dppm})_{6}\text{Cl}_{6}]\)(\text{SbF}_{6})_{3}\) by the reaction of \([\text{Au}_{9}\text{Ag}_{12}(\text{SAdm})_{4}(\text{Dppm})_{6}\text{Cl}_{6}]\)(\text{SbF}_{6})_{3}\) with the Cu(SAdm) complex precursor. X-ray crystallography identifies that the Cu dopants preferentially replace the position of the silver capped by Dppm ligand in the motif. The Cu doping has affected the optical properties of Au\(_9\)Ag\(_{12}\) alloy nanocluster. DPV spectra, CD spectra and stability tests suggest that the regulation of surface structure via Cu alloying changes the electronic structure, thereby affecting the electrochemical properties, which provides insight into the regulation of surface structure of \([\text{Au}_{9}\text{Ag}_{12}(\text{SAdm})_{4}(\text{Dppm})_{6}\text{Cl}_{6}]\)(\text{SbF}_{6})_{3}\) via alloying.

Keywords: regulation of surface structure, alloy engineering, optical properties, electrochemical properties, intercluster reactions

INTRODUCTION

Atomically precise core-shell nanoclusters have become a promising material in catalysis, biomedicine, and chemical sensing due to the unique quantum confinement effect resulting in optical properties (Jin et al., 2016; Yao et al., 2018; Xu et al., 2019; Jin R. et al., 2021; Sun et al., 2021; Zheng et al., 2021). The studies on correlation between the properties and structures of cluster compounds based on the determined crystal structures show that the core and shell structures have different effects on the performance of the cluster compounds, and modifications on the core and shell structures may induce variations on clusters properties (AbdulHalim et al., 2014; Chakraborty and Pradeep, 2017; Khatun et al., 2018; Yan et al., 2018; Jin Y. et al., 2021). The Pt core-doped nanocluster PtAu\(_{24}(\text{SC}_{6}\text{H}_{13})_{18}\) exhibits higher hydrogen production than that of Au\(_{25}\) (Kwak et al., 2017), and the dopant AuAg\(_{24}\) shows stronger fluorescence performance (Bootharaju et al., 2016). Surface shell dopant Au\(_{24}\)Cu\(_{6}\) exhibited superior catalytic activity compared to other homometallic and Au-Cu alloy nanoclusters (Chai et al., 2019). Therefore, alloying could serve as an efficient approach to tailor the properties of nanoclusters for more applications (Ghosh et al., 2018; Jin et al., 2018a; Wang et al., 2018; Dias and Leite, 2019).

Current alloy research mainly focuses on bimetallic clusters, and there are few studies on trimetallic clusters due to factors such as synthesis, characterization, and crystallization, etc. (Kang et al., 2016; Sharma et al., 2016; Yan et al., 2016; Hossain, et al., 2018; Kang et al., 2019a; Kang et al., 2019b; Kang et al., 2020; Chai et al., 2019; Khatun et al., 2018; Jin et al., 2018a; Wang et al., 2018; Dias and Leite, 2019).
Optical absorption spectra and the XPS spectra of \([\text{Au}_9\text{Ag}_{12}(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6](\text{SbF}_6)_3\) (black line) and \([\text{Au}_9\text{Ag}_8\text{Cu}_4(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6](\text{SbF}_6)_3\) (red line); ESI-MS spectra of (C) \([\text{Au}_9\text{Ag}_{12}(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6]\) and (D) \([\text{Au}_9\text{Ag}_8\text{Cu}_4(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6]\). The peaks labeled by asterisks in Panels (C, D) correspond to \([\text{Au}_9\text{Ag}_{12}(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6]^{2+}\) and \([\text{Au}_9\text{Ag}_8\text{Cu}_4(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6]^{2+}\), respectively.

The metal exchange from \([\text{Au}_9\text{Ag}_{12}(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6]\) to \([\text{Au}_9\text{Ag}_8\text{Cu}_4(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6]\) (Color labels: Golden = Au; Sky blue = Ag; red = S; purple = P; Gray = C; light green = Cl; Turquoise = Copper).
2019b; Kang et al., 2020) When the third metal is doped into the bimetallic alloy clusters, what site will it occupy and what effect will it have on the overall performance? Recently, for the active metal Cu doping, several surface Cu-doped nanoclusters such as Au13Cu_x (x = 2, 4, 8) (Yang et al., 2013), Cu_xAu_{25-x} (Yang et al., 2017), Cu_3Au_{34} (Yang et al., 2017), Ag_{28}Cu_{12} (Yan et al., 2016), Ag_{30}Cu_{14} (Li et al., 2020) and Cu-internal-doped nanoclusters like Ag_{61}Cu_{30} have been observed and well-determined by x-ray crystallography (Zou et al., 2020). Specifically, the outer Au shells always are partially alloyed by the incorporated Cu heteroatoms for Au-based nanoclusters, while core-shell alloy nanoclusters with a shell-by-shell configuration could be generated for Ag-based nanoclusters. However, for the Au-Ag alloy nanocluster, how will the copper atoms choose the sites? Herein, we use position-determined alloy clusters [Au_9Ag_{12}(SAdm)_4(Dppm)_6Cl_6](SbF_6)_3 as templates for the doping of the third metal copper (Jin et al., 2018b). The crystallography analysis suggested that the four Cu atoms priority replace the position of the silver capped by Dppm ligand in the motif for [Au_9Ag_{8}Cu_4(SAdm)_4(Dppm)_6Cl_6](SbF_6)_3 (Scheme 1). And the Cu doping affected the electronic structure, resulting in the difference of optical properties in CD spectra, DPV spectra and so on. This provides a good observation method for understanding the doping position.

MATERIALS AND METHODS

Materials
Tetrachloroauric(III) acid (HAuCl_4·3H_2O, 99.99%), silver nitrate (AgNO_3, 98%), tetrabutyl ammonium chloride (TBAC, 98%), sodium borohydride (NaBH_4, 99.99%), bis-(diphenylphosphino)methane (Dppm, 98%), 1-Adamantanethiol (C_10H_{16}S, 99%), sodium hexafluoroantimonate (NaSbF_6, 98%), toluene (Tol, HPLC grade, Aldrich), methanol (CH_3OH, HPLC, Aldrich), n-hexane (Hex, HPLC grade, Aldrich), dichloromethane (CH_2Cl_2, HPLC grade, Aldrich), Pure water was purchased from Wahaha Co. Ltd. All reagents were used as received without further purification.

Synthesis of [Au_9Ag_{12}(SAdm)_4(Dppm)_6Cl_6](SbF_6)_3 nanocluster
The synthesis of [Au_9Ag_{12}(SAdm)_4(Dppm)_6Cl_6](SbF_6)_3 was obtained by the method reported (Jin et al., 2018a). Typically, HAuCl_4·3H_2O (40 mg) and AgNO_3 (60 mg) was mixed in 15 ml toluene with TBAC (200 mg). Stirring for 5 min, 50 mg bis-(diphenylphosphino)methane and 50 mg 1-Adamantanethiol were added together. 15 min later, a solution of 20 mg NaBH_4 (1 ml H_2O) was added. The reaction sustained for 12 h at room temperature. The crude product was spied dry and washed by hexane. 30 mg NaSbF_6 in 3 ml CH_3OH was added to replace the anion of the cluster for easy crystallization. The yield of [Au_9Ag_{12}(SAdm)_4(Dppm)_6Cl_6](SbF_6)_3 is as high as 70% based on the Ag element, which was determined by ESI-MS and X-ray crystallography. The CCDC number is 2114779.

Synthesis of CuISR Complex Precursor
CuCl (0.05 g, 0.5 mmol) was dissolved in 5 ml CH_3CN, and AdmSH (0.09 g, 0.55 mmol) was dissolus in 5 ml CH_3CN and added drop-wise to the solution under vigorously stirred. The resulting solution mixture was then washed several times with hexane. Then the final product was used directly.

Synthesis of [Au_9Ag_{8}Cu_4(SAdm)_4(Dppm)_6Cl_6](SbF_6)_3 nanocluster:
The 20 mg [Au_9Ag_{12}(SAdm)_4(Dppm)_6Cl_6](SbF_6)_3 dissolved in 7 ml methylene chloride, CuSR (1 mg) was added to the solution.
The reaction lasted for 10 min at room temperature. After that, the reaction mixture was centrifuged at 8,000 rpm. The organic layer was separated from the precipitate and evaporated to dryness. [Au9Ag8Cu4(SAdm)4(Dppm)6Cl6](SbF6)3 was obtained. The yield of [Au9Ag8Cu4(SAdm)4(Dppm)6Cl6](SbF6)3 is as high as 60% based on the [Au9Ag12(SAdm)4(Dppm)6Cl6](SbF6)3. Orange crystals were crystallized from CH2Cl2/hexane at room temperature after 7 days. The CCDC number is 2114780.

**Characterization**

All UV/Vis absorption spectra of nanoclusters are recorded on a Techcomp UV1000 spectrophotometer. Electrospray ionization time-of-flight mass spectrometry (ESI-TOF-MS) measurement was performed using a UPLC H-class/XEV0G2-XS QTOF high-resolution mass spectrometer. The sample was directly infused into the chamber at 5 μL/min. Photoluminescence spectra were measured using an FL-7000 spectrofluorometer with the same

![FIGURE 3](image-url) (A) the overall structure of [Au9Ag12(SAdm)4(Dppm)6Cl6](SbF6)3; (B) the overall structure of [Au9Ag8Cu4(SAdm)4(Dppm)6Cl6](SbF6)3; Packing models of (C) [Au9Ag12(SAdm)4(Dppm)6Cl6]3+ and (D) [Au9Ag8Cu4(SAdm)4(Dppm)6Cl6]3+ from default view a, b, c. Color labels: Golden = Au; Sky blue = Ag; red = S; purple = P; Gray = C; light green = Cl; Turquoise = Copper.

![FIGURE 4](image-url) The circular dichroism (CD) spectra of the enantiomer in (A) [Au9Ag12(SAdm)4(Dppm)6Cl6](SbF6)3 and (B) [Au9Ag8Cu4(SAdm)4(Dppm)6Cl6](SbF6)3 nanoclusters.

The reaction lasted for 10 min at room temperature. After that, the reaction mixture was centrifuged at 8,000 rpm. The organic layer was separated from the precipitate and evaporated to dryness. [Au9Ag8Cu4(SAdm)4(Dppm)6Cl6](SbF6)3 was obtained.

The yield of [Au9Ag8Cu4(SAdm)4(Dppm)6Cl6](SbF6)3 is as high as 60% based on the [Au9Ag12(SAdm)4(Dppm)6Cl6](SbF6)3. Orange crystals were crystallized from CH2Cl2/hexane at room temperature after 7 days. The CCDC number is 2114780.
optical density (OD) of ∼0.2. X-ray photoelectron spectroscopy (XPS) measurements were performed using a Thermo ESCALAB 250 configured with a monochromated Al Kα (1486.8 eV) 150 W X-ray source, 0.5 m mm circular spot size, a flood gun to counter charging effects, and an analysis chamber base pressure lower than 1 × 10⁻⁹ mbar, and the data were collected with FAT/equals 20 eV. CD spectra are recorded with a BioLogic MOS-500 CD-spectropolarimeter in a 0.1-cm path length quartz cell. The spectra are recorded in diluted solutions of dichloromethane and the signal of the blank solvent is subtracted. The enantiomers of chiral \([\text{Au}\text{9Ag}\text{Cu}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) were separated by HPLC on an Agilent 1260 system equipped with a Chiralcel OD-H column (5 µm, 4.6 mm ø × 250 mm). A diode array detector (DAD) in situ monitors the entire optical absorption spectrum (190–950 nm range) of the eluted solution, and the 427, 482 and 710 nm wavelength were used for the chromatogram. The nanoclusters were pre-dissolved in solvent which has the same composition of the mobile phase (methanol/isopropanol = 35/65). The flow rate was at 0.4 ml min⁻¹ and the temperature set at 20°C.

RESULTS AND DISCUSSION

The synthesized \([\text{Au}\text{9Ag}\text{12}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) based on the reported method was determined by ESI-MS and X-ray crystallography. The next is the regulation of surface structure of \([\text{Au}\text{9Ag}\text{12}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) with Cu\text{I}(\text{SAdm}) complex precursor. As shown in Figure 1A, the \([\text{Au}\text{9Ag}\text{8Cu}\text{4}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) shows main peaks at 322, 365, 427, 480 and 670 nm, respectively, and \([\text{Au}\text{9Ag}\text{8Cu}\text{4}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) shows 322, 366, 427, 482 and 710 nm, respectively. In contrast, most of the peaks for both nanoclusters have not changed significantly, except for the red shift of the 670 nm peak to 710 nm. The binding energy of Cu\text{2p} from XPS data confirmed the Cu doping in the \([\text{Au}\text{9Ag}\text{8Cu}\text{4}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) (Figure 1B), and energy level positions of other elements have basically not changed (Supplementary Figure S1). The peak at m/z 2084.85 corresponds to the 3 + charge of \([\text{Au}\text{9Ag}\text{12}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) and can be perfectly assigned by the calculated result (m/z 2084.81) (Figure 1C). And peak at m/z 2026.20 corresponds to the 3 + charge of \([\text{Au}\text{9Ag}\text{8Cu}\text{4}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) (Cal. 2026.18) (Figure 1D).

Furthermore, in order to have a deep understanding of the regulation process, the time-dependent UV-Vis spectra and ESI mass spectra of \([\text{Au}\text{9Ag}\text{12}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) in CH₂Cl₂ after adding Cu\text{I}(\text{SAdm}) complex precursor were performed. As shown in Figure 2A, with the increase of time of CuI(SAdm) complex precursor adding, the peak centered at 427 nm always maintained, and the peak centered at 480 nm only redshifts. While the 670 nm peak gradually red shift to 710 nm, with a redshift value of 40 nm. ESI mass spectra suggested the copper atoms are gradually replacing silver atoms, which leads to red shift (Figure 2B). The successful determination of \([\text{Au}\text{9Ag}\text{8Cu}\text{4}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) structure allowed us to know the site of doping clearly. As shown in Figure 3, the overall structure of \([\text{Au}\text{9Ag}\text{12}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) and \([\text{Au}\text{9Ag}\text{8Cu}\text{4}(\text{SAdm})\text{4}(\text{Dppm})\text{6Cl}\text{6}]\text{(SbF}\text{6})\text{3}\) are basically the same: firstly, five gold atoms and eight silver atoms constitute the icosahedron, then the Au₅Ag₈ icosahedron and four gold atoms constitute the Au₄@Ag₈Au₅ metallic kernel. The Au₄@Ag₈Au₅ is first capped by four Dppm ligands and two Cl ligands, forming Au₄@Ag₈Au₅(Dppm)₂Cl₂ framework. After the Au₄@Ag₈Au₅(Dppm)₂Cl₂ is further protected by two peripheral

FIGURE 5 | (A) Photon energy spectra of Au₉Ag₁₂ (black line) and Au₉Ag₈Cu₄ (blue line). (B) DPV of Au₉Ag₁₂ (black line) and Au₉Ag₈Cu₄ (blue line).
structures DppmAg$_2$Cl$_2$(SR)$_2$, the Au$_9$Ag$_{12}$ was obtained. By contrast, Au$_5$Ag$_8$@Au$_4$@Cu$_4$ is obtained when four copper atoms doped the position of the silver of peripheral structures DppmAg$_2$Cl$_2$(SR)$_2$. Meanwhile, the copper doping has little effect on the bond length and angle of the icosahedron metal core (Supplementary Figure S4). Based on the doping sites of copper atoms, we realize that the Au$_4$@Ag$_8$Au$_5$ will be a stable metal core. In the packing model, the difference of arrangement can be observed clearly, and it is worth mentioning that the doping can affect the arrangement of clusters in the unit cell (Figures 3C,D) from a crystal engineering point of view.

As reported, the chirality of metal clusters mainly come from chiral metalcore, the arrangement of chiral ligands and local chiral patterns on an achiral surface (Zeng and Jin, 2017). The chirality of [Au$_9$Ag$_{12}$(SAdm)$_4$(Dppm)$_6$Cl$_6$]$^3+$ comes from the chiral Au$_4$@Ag$_8$Au$_5$ metallic kernel. After doping, the cluster will have a different CD spectrum compared to the parent compound. Importantly, herein, the Cu dopants also have some impacts on the chiral properties. As shown in Figure 4, the CD spectra of [Au$_9$Ag$_{12}$(SAdm)$_4$(Dppm)$_6$Cl$_6$]$^3+$ reveal multiple CD-active peaks at 325, 363, 428 and 483 nm, respectively, and some weak peaks. While the CD spectra of [Au$_9$Ag$_8$Cu$_4$(SAdm)$_4$(Dppm)$_6$Cl$_6$]$^3+$ show peaks at 340, 373, 442, and 493 nm, respectively. The Au$_{38}$ cluster with Pd atoms leads to core-doped Pd$_2$Au$_{36}$(SC$_2$H$_4$Ph)$_{24}$. Comparison between the CD spectra of Au$_{38}$(SC$_2$H$_4$Ph)$_{24}$ and Pd$_2$Au$_{36}$(SC$_2$H$_4$Ph)$_{24}$ shows significant differences, revealing core-doping has strong impacts on the electronic structure of the cluster (Barrabés et al., 2014). The comparison between the CD spectra of [Au$_9$Ag$_{12}$(SAdm)$_4$(Dppm)$_6$Cl$_6$]$^3+$ and [Au$_9$Ag$_8$Cu$_4$(SAdm)$_4$(Dppm)$_6$Cl$_6$]$^3+$ shows that all the peaks from [Au$_9$Ag$_8$Cu$_4$(SAdm)$_4$(Dppm)$_6$Cl$_6$]$^3+$ have redshift, different from the differences between Au$_{38}$(SC$_2$H$_4$Ph)$_{24}$ and Pd$_2$Au$_{36}$(SC$_2$H$_4$Ph)$_{24}$. The doping location may have different impacts on the CD spectra.

In addition to the CD spectra, the electronic structures of [Au$_9$Ag$_{12}$(SAdm)$_4$(Dppm)$_6$Cl$_6$](SbF$_6$)$_3$ and [Au$_9$Ag$_8$Cu$_4$(SAdm)$_4$(Dppm)$_6$Cl$_6$](SbF$_6$)$_3$ are investigated by optical and electrochemical spectroscopies. Differential pulse voltammetry (DPV) of Au$_9$Ag$_{12}$ and Au$_9$Ag$_8$Cu$_4$ are carried out. The scan direction was detected from $+1.6$ to $-1.6$ V and then back from $-1.6$ to $+1.6$ V. As shown in Figure 5, the HOMO-LUMO gaps of Au$_9$Ag$_{12}$ and Au$_9$Ag$_8$Cu$_4$ are determined as 1.54 and 1.44 eV, respectively. For the differential pulse voltammetry (DPV) curves, there is a reduction peak at $-1.32$ V (R1) and two oxidation peaks at 0.40 V (O1) and 0.58 V (O2) for Au$_9$Ag$_{12}$, while there are two reduction peaks at $-0.82$ V (R1) and $-1.01$ (R2) and one oxidation peaks at 0.83 V (O1) for Au$_9$Ag$_8$Cu$_4$. So, the electrochemical energy gap is 1.72 eV for Au$_9$Ag$_{12}$ and 1.65 eV for Au$_9$Ag$_8$Cu$_4$. The HOMO-LUMO gaps calculated from DPV are consistent with those derived from the optical absorption spectra. So, the regulation of surface structure via Cu alloying changes the electronic structure, thereby affecting the electrochemical properties. Besides, the [Au$_9$Ag$_{12}$(SAdm)$_4$(Dppm)$_6$Cl$_6$](SbF$_6$)$_3$ in CH$_2$Cl$_2$ solution...
shows non-fluorescence, while the Cu dopant 
\[\text{[Au}_9\text{Ag}_{12}(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6](\text{SbF}_6)_3\] in CH$_2$Cl$_2$ solution shows weak fluorescence at 638 nm, once again verifying the changes in the electronic structure. (Supplementary Figure S5).

The 
\[\text{[Au}_9\text{Ag}_{12}(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6](\text{SbF}_6)_3\] and 
\[\text{[Au}_9\text{Ag}_{12}(\text{SAdm})_4(\text{Dppm})_6\text{Cl}_6](\text{SbF}_6)_3\] show good stability in an ambient environment (Figures 6A,D) and the stability tests (i.e., under oxidizing/reducing environments) for Au$_9$Ag$_{12}$ and Au$_9$Ag$_8$Cu$_4$ are also performed to explore the effects of copper dopants on the stability of nanoclusters. Under the oxidizing environment (by mixing 200 µl of H$_2$O$_2$ (50%) with 6 mg of cluster in 10 ml of CH$_2$Cl$_2$), the Au$_9$Ag$_{12}$ can stabilize for several hours (Figures 6B,E), and the peaks of the UV-vis spectra are obvious. However, the Au$_9$Ag$_8$Cu$_4$ decompose quickly to form complexes within several mins (Figures 6C,F). This difference may be because the peripheral copper atom is easier to be oxidized. Meanwhile, the copper doping has an impact on the properties of clusters on reducing environment (by mixing the 10 ml CH$_2$Cl$_2$ solvent of 6 mg of cluster with 200 µl of EtOH solvent of 1 mg of NaBH$_4$). The UV-vis of Au$_9$Ag$_{12}$ changes quickly until there are no obvious peaks within 30s. And the UV-vis of Au$_9$Ag$_8$Cu$_4$ also changes quickly, but still some peaks can be observed within 60 min. These indicate the regulation of surface structure affects the stability of nanoclusters.

Intercluster reactions between Au$_9$Ag$_{12}$ and Au$_9$Ag$_8$Cu$_4$ (Abs$_{671\text{nm}} = 0.3$ for Au$_9$Ag$_{12}$ and Abs$_{712\text{nm}} = 0.3$ for Au$_9$Ag$_8$Cu$_4$, respectively) are performed (Zhang et al., 2016; Khatun et al., 2020; Neumaier at al., 2021). As shown in Figure 7, the reaction was completed quickly (1 min), similar to the UV-vis spectrum that prolongs the reaction for 3 h. As shown in the Figures 7A,B, intercluster reactions produce a spectrum with 428, 482 and 702 nm, respectively. Learned from the Figures 7C,D, the products are Au$_9$Ag$_8$Cu$_4$, Au$_9$Ag$_9$Cu$_3$, Au$_9$Ag$_{10}$Cu$_2$, Au$_9$Ag$_{11}$Cu$_1$, respectively. Theoretical and experimental isotopic distributions of them matched perfectly as shown in Supplementary Figures S6, 7. This indicates the copper migration between Au$_9$Ag$_{12}$ and Au$_9$Ag$_8$Cu$_4$ upon mixing in solution, similar to silver migration between Au$_{38}$(SC$_2$H$_4$Ph)$_{24}$ and doped Ag$_x$Au$_{38-x}$(SC$_2$H$_4$Ph)$_{24}$ nanoclusters (Zhang et al., 2016).
CONCLUSIONS

In conclusion, the regulation of surface structure of \([\text{Au}_9\text{Ag}_{12} (\text{SAdm})_4 (\text{Dppm})_6 \text{Cl}_6]_3^+\) nanocluster via alloying produced a trimetallic nanocluster formulated as \([\text{Au}_9\text{Ag}_{12} \text{Cu}_4 (\text{SAdm})_4 (\text{Dppm})_6 \text{Cl}_6]_3^+\). X-ray crystallography identifies that the Cu dopants prioritely replace the position of the silver of peripheral structures Dppm\(\text{Ag}_2\text{Cl}_2\text{(SR)}_2\). This controlled target metal exchange method may be extendable to other sized nanoclusters capped by multiple-ligands. Meanwhile the regulation of surface structure affected the CD spectra, DPV spectra, and stability. The \([\text{Au}_9\text{Ag}_{12} (\text{SAdm})_4 (\text{Dppm})_6 \text{Cl}_6]_3^+\) and \([\text{Au}_9\text{Ag}_{12} \text{Cu}_4 (\text{SAdm})_4 (\text{Dppm})_6 \text{Cl}_6]_3^+\) contribute to understanding of the structure-optical property relationship deeply.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

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AUTHOR CONTRIBUTIONS

HD and XL: performed the experiment and wrote the manuscript. XY: assisted the synthesis. SJ and MZ: analyzed the date and revised the manuscript. HD and XL equally contribute to this work.

FUNDING

National Natural Science Foundation of China (21901001, 21631001), Natural Science Foundation of Education Department of Anhui Province KJ2019A0008, Innovation and entrepreneurship project of Returning Overseas Chinese Scholars in Anhui Province 2019LCX021, and the Doctoral Scientific Research Foundation of Anhui University to XL.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2021.793339/full?supplementary-material
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