Observation of a new type of aggregation-induced emission in nanoclusters†

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The strategy of aggregation-induced emission (AIE) has been widely used to enhance the photoluminescence (PL) in the nanocluster (NC) research field. Most of the previous reports on aggregation-induced enhancement of fluorescence in NCs are induced by the restriction of intramolecular motion (RIM). In this work, a novel mechanism involving the restriction of the “dissociation—aggregation pattern” of ligands is presented using a Ag_{29}(BDT)_{12}(TPP)_{4} NC (BDT: 1,3-benzenedithiol; TPP: triphenylphosphine) as a model. By the addition of TPP into an N,N-dimethylformamide solution of Ag_{29}(BDT)_{12}(TPP)_{4}, the PL intensity of the Ag_{29}(BDT)_{12}(TPP)_{4} NC could be significantly enhanced (13 times, quantum yield from 0.9% to 11.7%) due to the restricted TPP dissociation–aggregation process. This novel mechanism is further validated by a low-temperature PL study. Different from the significant PL enhancement of the Ag_{29}(BDT)_{12}(TPP)_{4} NC, the non-dissociative PtAg_{29}(S-Adm)_{12}(TPP)_{4} NC (S-Adm: 1-adamantanethiol) exhibits a maintained PL intensity under the same TPP-addition conditions. Overall, this work presents a new mechanism for largely enhancing the PL of NCs via modulating the dissociation of ligands on the NC surface, which is totally different from the previously reported AIE phenomena in the NC field.

1 Introduction

Noble metal nanoclusters (NCs) have attracted widespread attention due to their advantages of atomically precise structures, distinct physical/chemical properties, and extensive catalytic/biomedical applications.1–18 Among these interesting physical/chemical properties, photo-luminescence (PL) represents one of the most fascinating features owing to the low toxicity, good photostability, and high biocompatibility of NCs.1,2,8,17–24 Although several luminescent NCs have been reported,4,18,21–31 most of them exhibit lower quantum yields (QYs) compared with other fluorescent nanomaterials (such as lanthanide nanoparticles,12 organic dyes32 and quantum dots33), which severely impedes extensive application of fluorescent NCs in biological and sensing fields. Several strategies have been developed to enhance the PL QY of NCs, and they can be mainly classified into the following three categories: (1) tailoring the capping ligands of NCs (in terms of controlling the ligand to metal charge transfer (LMCT) process);8,15–17 (2) controlling the metal composition in the NC kernel;21,22,26,28,29,38 and (3) aggregation-induced emission (AIE) from non- or weakly luminescent metal NCs (or complexes).19,20,25,39,40 Nowadays, the AIE strategy is being expanded to the hydrocarbon, metal complex, metal NC, and macromolecular research fields.4 The corresponding AIEgens have been widely applied in biological probing, chemical sensing, and organic light-emitting diodes, to name a few.41 In the NC field, the AIE strategy could boost the PL QY through facile approaches (e.g., solvent- or cation-inducing approaches).19,25 Furthermore, the previously reported enhancements of fluorescence in NCs have mostly been achieved by the restriction of intramolecular motion (RIM, a general mechanism in AIE materials).19,25 This raises some interesting questions: are there any other patterns (other than RIM) of AIE that exist in the NC field? If so, how can we enhance the QY of fluorescent NCs with new patterns? Addressing these issues will not only develop a powerful and practical strategy for synthesizing more fluorescent NCs with enhanced PL QYs, but also promote wide-range application of AIE in the NC field.

Herein, a novel AIE pattern (the restriction of the ligand dissociation-aggregation process) is discovered using a Ag_{29}(BDT)_{12}(TPP)_{4} NC (BDT, 1,3-benzenedithiol; TPP, triphenylphosphine) as a model. The PL intensity of the Ag_{29}(BDT)_{12}(TPP)_{4} NC in the solid or crystal state is significantly higher than that in the solution state. Considering the particularly close packing of Ag_{29}(BDT)_{12}(TPP)_{4} as well as the existence of π···π and C–H···π interactions between the ligands, the AIE process is unlikely caused by the RIM completely in the present case of the Ag_{29}(BDT)_{12}(TPP)_{4} NC. Furthermore, electrospray ionization mass spectrometry (ESI-MS) and 31P nuclear magnetic resonance (31P NMR) analyses are performed on the Ag_{29}(BDT)_{12}(TPP)_{4} NC, and they reveal several dissociated
ligands from Ag$_{29}$(BDT)$_{12}$(TPP)$_4$; that is, the mechanism of AIE in Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ could be the restriction of the TPP dissociation–aggregation process. To prove the mechanism, different concentrations of TPP molecules are added to the dissociative Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ and non-dissociative Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ (S-Adm, 1-adamantanethiol) NC solution. Resultantly, the PL intensity of Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ is significantly enhanced (up to 13 times, with a PL QY from 0.9% to 11.7%) by the addition of TPP owing to the restriction of the TPP dissociation–aggregation process (or the chemical equilibrium shifting to TPP aggregation on the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NCs, summarized in Scheme 1). By contrast, the non-dissociative Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ shows nearly the same intensity of PL after the addition of TPP. In addition, the temperature-dependent PL of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC shows two segments of rising curves, which represent both the restriction of the TPP dissociation–aggregation process (the QY varies from 0.9% to 22.5%) and the quenched thermal vibration process (the QY varies from 22.5% to about 100%). However, the non-dissociative Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ only exhibits a quenched thermal vibration process (the QY varies from 9.3% to ~100%).

2 Experimental methods

Materials

All chemicals including silver nitrate (AgNO$_3$, 99%, metal basis), hexachloroplatinic(IV) acid (H$_2$PtCl$_6$·6H$_2$O, 99.99%, metals basis), triphenylphosphine (TPP, 99%), benzene-1,3-diethanol (BDE, 99%), 2,4-dimethylbenzenethiol (HSPhMe$_2$, 99%), 1-adamantanethiol (HSC$_{10}$H$_{15}$, 99%), sodium borohydride (NaBH$_4$, 99.9%), methylene chloride (CH$_2$Cl$_2$, HPLC grade, Aldrich), acetic ether (CH$_3$COOC$_2$H$_5$, HPLC, Aldrich), methanol (CH$_3$OH, HPLC, Aldrich), ethanol (CH$_3$CH$_2$OH, HPLC, Aldrich) and N,N-dimethylformamide (DMF, HPLC, Aldrich) were purchased from Sigma-Aldrich and used without further purification. Pure water was purchased from Wahaha Co. Ltd. All glassware was thoroughly cleaned with aqua regia (HCl : HNO$_3$ = 3 : 1 v/v), rinsed with copious amounts of pure water, and then dried in an oven prior to use.

Synthesis of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC

The synthesis of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC was performed following a method reported by Bakr and coworkers.²⁷

Synthesis of the Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NC

The synthesis of the Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NC was performed according to our previous work.⁴³

Synthesis of the Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$ NC

The synthesis of the Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$ NC was performed following a method previously reported by Bakr and coworkers.⁴⁴

Test of the TPP ligand concentration–PL intensity correlation

10 mg of Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ (or Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ or Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$) NC was dissolved in 10 mL of DMF. Then the TPP ligand was added to the DMF solution with the molar ratio of TPP/NC ranging from 0.1 to 10. The PL spectra were then measured for these mixed solutions.

Test of the temperature–PL intensity correlation

10 mg of Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ (or Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$) NC was dissolved in 10 mL of DMF. Then the solutions were cooled to different temperatures and the PL spectra were measured.

Characterization

All UV-vis absorption spectra of NCs dissolved in DMF were recorded using an Agilent 8453 diode array spectrometer, with the background corrected by using a DMF blank. Solid samples were dissolved in DMF to make a dilute solution, which was transferred to a 1 cm path length quartz cuvette for spectral measurements.

PL spectra were measured on an FL-4500 spectrofluorometer with the same optical density (OD) of ~0.05. In these experiments, the NC solutions were prepared in DMF at a concentration of less than 1 mg mL$^{-1}$.

Absolute quantum yields (QYs) were measured with dilute solutions of NCs (0.05 OD absorption at 445 nm) on a HORIBA FluoroMax-4P.

$^{31}$P NMR spectra were acquired using a Bruker 600 Avance III spectrometer equipped with a Bruker BBO multinuclear probe (BrukerBioSpin, Rheinstetten, Germany). To achieve a sufficient signal-to-noise ratio, the $^{31}$P NMR spectra were recorded by collecting 1k scans with a recycle delay time of 5 s.

Electrospray ionization time-of-flight mass spectrometry (ESI-TOF-MS) measurements were performed on a MicrOTOF-QIII high-resolution mass spectrometer.

3 Results and discussion

The Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC was prepared following a method previously published by Bakr and coworkers.²⁷ They also reported the crystal structure of Ag$_{29}$(BDT)$_{12}$(TPP)$_4$.²⁷ The structure is shown in Fig. 1A and B. Regarding the structural anatomy, Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ is composed of a three-layer
configuration: the icosahedral Ag13 kernel is enclosed by four Ag3S6 motifs to form a Ag29(BDT)12 architecture (it should be noted that the four Ag3S6 motifs are united by the BDT ligands); the Ag29(BDT)12 architecture is further capped by four Ag–TPP units. Fig. 1C shows the UV–vis spectrum of Ag29(BDT)12(TPP)4 NC (dissolved in DMF), which exhibits broad, multiband optical absorption bands centred at 345, 360, 445, and 510 nm. Furthermore, Ag29(BDT)12(TPP)4 NC possesses an emission band centred at 642 nm (Fig. 1D). Additionally, the PL excitation spectrum is almost the same as the absorption spectrum (Fig. 1C and D), which is reminiscent of some previously reported fluorescent NCs as well as the quantum-dot behavior.22–25

It is noteworthy that the PL QY of Ag29(BDT)12(TPP)4 is only 0.9%, which is too weak to be perceived by the naked eye (Fig. 1E).27,28 Interestingly, the Ag29(BDT)12(TPP)4 NC exhibits enhanced fluorescence in the solid or crystal state (Fig. 1F and G). These observations suggest that the aggregation process significantly boosts the emission of Ag29(BDT)12(TPP)4. Previously, the RIM pattern was believed to be the primary cause either in the solvent- or ion-induced AIE process.19,25 In our current work, the AIE observed during the drying or crystallization process is obviously not relevant to any ionic effect, because our process only involves a solvent evaporation process.

The crystal structure of the Ag29(BDT)12(TPP)4 NC is analyzed in order to gain insight into the aforementioned solvent-evaporation-induced AIE. In the solution state, the energy dissipation of photo-excited Ag29(BDT)12(TPP)4 includes two pathways: (i) non-radiative transitions (mainly affected by intramolecular vibrations), and (ii) radiative transitions (through luminescence). Fig. 2A shows an illustration of intra-molecular vibrations of the Ag29(BDT)12(TPP)4 NC, which comprise rotative vibrations and swing-vibrations. However, the considerable close-packing of the structural configuration (see the space-fill pattern in Fig. 2B) suggests that such vibrations are difficult. Furthermore, as mentioned above, each Ag3S6 motif is linked by the BDT ligand, which restricts the vibrations of Ag–S–Ag motifs. Moreover, many π⋯π and C–H⋯π interactions among BDT or TPP ligands are observed. Specifically, every two neighboring benzene rings of BDT ligands interact through π⋯π interaction, which forms 6 pairs of π⋯π interactions in total (Fig. 2C). In addition, the vibration of each of the ortho-position C–H on TPP is rendered difficult owing to the C–H⋯π interaction with the nearby benzene ring of BDT (altogether 12 pairs of C–H⋯π interactions, shown in Fig. 2C). To summarize, the main cause of the weak PL of the Ag29(BDT)12(TPP)4 NC in solution is not thermal vibration, because the architecture of the Ag29(BDT)12(TPP)4 NC exhibits strong rigidity. Consequently, the AIE of the Ag29(BDT)12(TPP)4 NC is unlikely induced by the RIM pattern completely.

The above discussion raises an interesting question: what is the essential role of the AIE in the Ag29(BDT)12(TPP)4 NC? From the ESI-MS spectrum of Ag29(BDT)12(TPP)4 (Fig. 3A, see Fig. S1† for the expansion of the spectrum, and Fig. S2† for the comparison of the experimental and simulated isotope patterns), we find that the TPP ligands are capable of dissociation to generate Ag29(BDT)12(TPP)x (x = 0–3) and the dissociated TPP ligands. For instance, the highest signal in the ESI-MS spectrum is the peak of Ag29(BDT)12(TPP)2, which is generated

![Image](https://example.com/image1)

**Fig. 1** (A) Total structure of the Ag29(BDT)12(TPP)4 NC (redrawn from ref. 27). (B) Structural anatomy of the Ag29(BDT)12(TPP)4 NC. (C) UV–vis spectrum of the Ag29(BDT)12(TPP)4 NC (dissolved in DMF). (D) PL excitation spectrum (black) and emission spectra (other colors) of the Ag29(BDT)12(TPP)4 NC in (E) solution state, (F) solid state, and (G) crystal state under visible light. Color codes: cerulean/red spheres, Ag; yellow spheres, S; purple spheres, P; grey spheres, C; white spheres, H.

![Image](https://example.com/image2)

**Fig. 2** (A) Illustration of the vibration patterns in the Ag29(BDT)12(TPP)4 NC. (B) The space-fill pattern of the crystal structure of the Ag29(BDT)12(TPP)4 NC. (C) (i) The π⋯π interaction between neighboring benzene rings on BDT ligands, and (ii) the C–H⋯π interaction between ortho-position C–H on TPP and benzene rings on BDT ligands. Color codes: cerulean spheres, Ag; yellow spheres, S; purple spheres, P; grey spheres, C; white spheres, H.
by dissociating two TPP ligands of the intact NC. To further validate the TPP dissociation process of Ag_{29}(BDT)_{12}(TPP)_{4} in the solution state, ESI-MS measurements were also performed on the mixture of Ag_{29}(BDT)_{12}(TPP)_{4} NC with extra TPP. The spectrum (Fig. 3A, red line) exhibits a stronger signal of the intact Ag_{29}(BDT)_{12}(TPP)_{4} NC compared with the case of Ag_{29}(BDT)_{12}(TPP)_{4} NC only (Fig. 3A, black line), which indicates a suppressed dissociation process with extra TPP in solution (note that because of the dilution process in the ESI-MS measurements, there were still lots of dissociated Ag_{29}(BDT)_{12}(TPP)_{4} signals in the sample mixture). Therefore, based on the ESI-MS results, the reversible reaction of the Ag_{29}(BDT)_{12}(TPP)_{4} NC in solution could be determined to be Ag_{29}(BDT)_{12}(TPP)_{4} ↔ Ag_{29}(BDT)_{12}(TPP)_{3} + Ag_{29}(BDT)_{12}(TPP)_{2} + Ag_{29}(BDT)_{12}(TPP)_{1} + Ag_{29}(BDT)_{12} + TPP (shown in Fig. 3C). Accordingly, the TPP ligands on the Ag_{29}(BDT)_{12}(TPP)_{4} NC are in dynamic dissociation/aggregation. It is suggested that the breaking of coordination bonds would consume energy, and thus the energy loss with non-radiative transitions would influence the energy release with the radiative transition. Specifically, the TPP dissociation process would consume vast amounts of energy, which could be used to explain the result of low PL of the photo-excited Ag_{29}(BDT)_{12}(TPP)_{4} NC in the solution state. Thus, we speculate that the significantly enhanced PL of the Ag_{29}(BDT)_{12}(TPP)_{4} NC in the solid or crystal state should arise from the restriction of the TPP dissociation–aggregation process—which inhibits the non-radiative pathways, and thus enhances the radiative pathway (PL). Therefore, the AIE of the Ag_{29}(BDT)_{12}(TPP)_{4} NC could be induced by the restriction of the ligand dissociation–aggregation process.

Considering that the TPP dissociation–aggregation process is a reversible reaction, we are motivated to control the reversible process to control the intensity of PL. As shown in Fig. 4, an increasing proportion (molar ratio) of TPP added to the DMF solution of Ag_{29}(BDT)_{12}(TPP)_{4} increases the PL intensity rapidly in the beginning, and then it levels off, with the highest PL QY being 11.7%, which is in striking contrast to the weakly luminescent Ag_{29}(BDT)_{12}(TPP)_{4} NC with no TPP addition (QY = 0.9%). A quantitative test was performed to obtain the relationship between the PL intensity and the amount of TPP added (Fig. 4, bottom-left). When adding a 0.1 molar ratio of TPP (versus NC) to the DMF solution of Ag_{29}(BDT)_{12}(TPP)_{4} NC, the PL intensity increases by almost 3 times compared with the initial state. Furthermore, by the further addition of TPP, the QY enhancement gradually becomes steady. Finally, a 13-fold enhancement compared to the initial state is achieved when the amount of added TPP is greater than 2 (molar ratio). The PL intensity at 642 nm is compared (Fig. 4, bottom-right), which also confirms the PL variation trend in the PL QY test (i.e., the increasing trend as well as the 13-fold enhancement). To sum up, the redundant TPP ligands will prevent the TPP dissociation–aggregation process on the nanocluster surface, and then enhance the PL intensity of the Ag_{29}(BDT)_{12}(TPP)_{4} NC.

For comparison, the same TPP-addition experiment was performed on the Pt_{1}Ag_{28}(S-Adm)_{18}(TPP)_{4} NC (see Fig. S3† for the structural anatomy of Pt_{1}Ag_{28}(S-Adm)_{18}(TPP)_{4}). The ESI-MS of the Pt_{1}Ag_{28}(S-Adm)_{18}(TPP)_{4} NC shows a distinct peak at 3637.63 Da (Fig. 5, bottom-left), which clearly illustrates the non-dissociated state of TPP ligands of this NC in the DMF solution. The unchanged ESI-MS and UV-vis spectra after the addition of TPP ligands illustrate that the as-synthesized nanoclusters will not decompose or transform in this operation (Fig. S4†). Importantly, in sharp contrast to the PL variation trend of the Ag_{29}(BDT)_{12}(TPP)_{4} NC, the PL intensity of the Pt_{1}Ag_{28}(S-Adm)_{18}(TPP)_{4} NC maintains a QY of 9.3% no matter how high the TPP amount added to the solution is (Fig. 5, top). Note that the PL QY of Pt_{1}Ag_{28}(S-Adm)_{18}(TPP)_{4} in CH_{2}Cl_{2} is 4.9%, which illustrates the solvent effect on the PL of Pt_{1}Ag_{28}(S-Adm)_{18}(TPP)_{4}. The PL intensity of the Pt_{1}Ag_{28}(S-Adm)_{18}(TPP)_{4} NC monitored at 672 nm by fluorescence spectroscopy also indicates the unchanged fluorescence (Fig. 5, bottom-right). In other words, the non-dissociative
Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NC does not display any equilibrium dissociation–aggregation process, and thus the addition of TPP would not alter the PL intensity in the solution state. The sharp contrast between the PL variation trends of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ and Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NCs indicates that the restriction of the ligand dissociation–aggregation process should be the major underlying mechanism of AIE in the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC.

Additionally, in order to structurally and compositionally match the Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NC, the TPP-addition experiment was performed on a Pt centrally doped Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC (that is, Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$ NC$^\dagger$). As shown in Fig. S5,† the PL intensity of Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$ also exhibits significant enhancement by the addition of TPP (the maximum PL QY was 18.9%), which is similar to the case of Ag$_{29}$(BDT)$_{12}$(TPP)$_4$; hence, Pt doping is not the critical factor for PL enhancement with TPP addition. Specifically, the fluorescence intensity (centered at 720 nm) rises rapidly during the initial TPP addition and then levels off once the molar ratio (TPP to Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$ NC) is greater than 1.5. A 7-fold enhancement is finally obtained by the addition of TPP to the Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$ NC. Considering the same PL enhancement phenomenon of the Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$ and Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NCs with the addition of TPP ligands, the difference in the metallic composition is unlikely to be the primary cause for being dissociative or not of these NCs. The major difference between the dissociative Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ and non-dissociative Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NCs is the outer ligands (BDT vs. S-Adm). The difference of these two ligands largely affects the structure of the metallic kernel and outer complex shell, which is a critical factor in the dissociative/non-dissociative phenomenon (see Fig. S6† for the structural analogies of the Pt$_1$Ag$_{28}$(BDT)$_{12}$(TPP)$_4$ as well as the Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$).

$^{31}$P NMR was performed to validate the TPP dynamic dissociation–aggregation state of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ and Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NCs. As shown in Fig. S7 and S8,† the $^{31}$P NMR spectrum of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC (without the addition of TPP) exhibits a broad peak (half-peak width $\sim$0.85 ppm), which narrows down continuously with the addition of more TPP (the peak width finally reduces to 0.17 ppm with a large excess of TPP in the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ solution). The notable decrease in the peak width illustrates that the state of P is uniformalized; that is, the TPP dynamic dissociation–aggregation extent of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC in solution is remarkably suppressed with the addition of excess TPP. Furthermore, the maintained width of the $^{31}$P NMR peak (half-peak width by 0.02 ppm, shown in Fig. S9†) for the Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NCs with or without TPP addition demonstrates the TPP non-dissociated state of the Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NC.

It is well known that the temperature would significantly influence the dissociation–aggregation dynamic equilibrium process. Thus, to further verify the above-mentioned AIE mechanism, the temperature–PL intensity correlation of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ and Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NCs was monitored. As shown in Fig. 6A and B and S10,† the temperature-dependent fluorescence of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC shows two stages: (1) when the temperature is reduced from 293 K to 251 K, the fluorescence shows a 25-fold enhancement (in this state the UV-vis absorption is maintained, and thus the PL QY increases from 0.9% to 22.5%), and (2) the fluorescence intensity is increased significantly (a 280-fold boost comparing the 107 K data with the 293 K data) when the temperature is reduced to 107 K, and the UV-vis absorption presents a 2.5-fold enhancement (Fig. S11†). Accordingly, the PL QY increases almost to 100%. The fluorescence of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC appeared to be extremely bright at 107 K, which is in striking contrast to the nearly invisible fluorescence at room
temperature (Fig. 6A, insets). In strong contrast, the non-dissociative Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NC exhibits only one stage during the same temperature lowering process (Fig. 6C and D and S12†), and finally exhibits a 20-fold enhancement of PL intensity and 1.9-fold enhancement of UV-vis absorption (Fig. S13†). Thus, the PL QY of the Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NC in the final stage is also nearly 100%. Furthermore, it should be noted that the single stage of Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ is similar to stage 2 of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC. Therefore, the PL boost in stage 2 of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC as well as in Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ should be completely induced by the suppression of thermal energy dissipation (or by the RIM pattern). Additionally, because the TPP dissociation process is much more sensitive to the temperature, the PL enhancement in stage 1 of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC should be mainly caused by the restriction of the TPP dissociation–aggregation process. Because this restriction process is easily influenced by the temperature variation, stage 1 is observed in the relatively high-temperature region (compared with stage 2 in the lower temperature region). To sum up, decreasing the temperature (from r.t. to 251 K) is effective in restricting the TPP dissociation–aggregation process which will lead to a significant enhancement in the fluorescence of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC in the solution state.

It should be noted that the PL enhancement of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC in the TPP addition process and temperature reduction process is different (13 times versus 25 times, with no increase in UV/vis absorption). In the TPP addition process, we find that the half-peak width (in the $^{31}$P NMR spectrum) of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC shows a continual decrease with the addition of TPP; however, the half-peak width in the final state is also wider than that in the Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ NC (0.17 versus 0.02 ppm). This suggests that the TPP dissociation–aggregation dynamic equilibrium in the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC is not completely prohibited, but is just limited to a certain extent. In other words, the non-radiative pathways caused by the TPP dissociation–aggregation process still exist, even with the excess dose of TPP, which makes it difficult to reach the highest PL (with the non-dissociated state) of the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC. By contrast, the non-dissociated state could be easily achieved in the temperature reduction process, and apparently, the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ NC in the non-dissociated state (at 251 K) exhibits a higher PL QY than the final state in the TPP addition process (22.5% versus 11.7%).

4 Conclusions

In summary, a novel mechanism of aggregation-induced emission is discovered in nanoclusters, which involves the restriction of the ligand dissociation–aggregation process. The fluorescence intensity of Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ can be significantly enhanced (about 13-fold, quantum yield from 0.9% to 11.7%) via promoting the aggregation of TPP onto the easy-to-dissociate nanocluster surface. Furthermore, the TPP dissociation–aggregation dynamic equilibrium process of Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ is also restrained by reducing the temperature, which results in enhanced photoluminescence intensity (25-fold) in Ag$_{29}$(BDT)$_{12}$(TPP)$_4$. In contrast, the same experiments performed on the non-dissociative Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ nanocluster do not show any PL enhancement. These different results are not caused by the presence of the Pt dopant, but by the different thiolate ligands (BDT versus S-Adm). The retained PL intensity in the case of Pt$_1$Ag$_{28}$(S-Adm)$_{18}$(TPP)$_4$ validates the aforementioned mechanism for the Ag$_{29}$(BDT)$_{12}$(TPP)$_4$ nanocluster. Overall, this work presents a new mechanism of aggregation-induced emission in nanoclusters. In addition to previous studies on the enhancement of nanocluster photo-luminescence, this work will hopefully draw greater attention of optical and theoretical chemists to fully understand the photo-luminescence properties of metal nanoclusters. Future work will focus on extending this new AIE mechanism to other fluorescent nanoclusters.

Conflicts of interest

There are no conflicts to declare.

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Notes and references


