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Introduction

In natural systems, the structural transformation of biomolecules plays a crucial role in biological processes like enzymatic catalysis, signal transduction, allosteric regulation, etc.1-3 Inspired by this, chemists have constructed a series of artificial systems to mimic biological transformations based on supramolecular self-assembly,4-8 expecting to prepare advanced materials with sophisticated functions comparable to biomolecules. In this regard, fusion-based structural transformation has received much attention because it can integrate all the building blocks of precursors into the final constructs, forming assemblies with improved structural and functional complexity (Fig. 1).9,10 However, such a process is highly challenging because elegant ligand design and tedious synthesis are generally required to prevent the formation of thermodynamic mixtures and meet the requirements of quantitative supramolecular transformation.

Owing to their moderate bond strength and fixed directionality, metal-coordination bonds have been widely used for the construction of two-dimensional metallacycles and three-



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The exploration of novel structures and structural transformation of supramolecular assemblies is of vital importance for their functions and applications. Herein, based on coordination-driven self-assembly, we prepare a neutral truncated tetrahedron and a heteroleptic truncated octahedron, whose structures are unambiguously confirmed by X-ray diffraction analysis. More importantly, the truncated tetrahedron is quantitatively transformed into the truncated octahedron through its fusion with another cationic truncated tetrahedron, as evidenced by fluorescence, mass and NMR spectroscopy. This study not only deepens our understanding of the process of supramolecular fusion but also opens up possibilities for the subsequent preparation of advanced supramolecular assemblies with complex structures and integrated functions.

dimensional metallacages,^{11–23} which are further employed in molecular recognition, stabilization of active substances, catalysis, *etc*.^{24–29} Moreover, these metal–organic assemblies are promising candidates for the construction of transformable supramolecular systems based on their dynamic and moderate stability. Although some progress has been made in this field,⁹ fusion-based structural transformation, especially the fusion of two geometrically similar metallacages into a new structure, has



Fig. 1 Representations of (a) cell fusion and (b) cage fusion.



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Fig. 2 Illustration of the self-assembly of metallacages 4a-4c.

been rarely reported, which is probably because it is difficult to design and prepare two complementary three-dimensional coordination assemblies for quantitative fusion. Such a study will not only deepen our understanding of the mechanism of biological and abiological fusion, but also offer new opportunities for the direct functional integration of different supramolecular assemblies. Therefore, the development of fusionbased supramolecular systems from three-dimensional selfassembled structures is highly demanding, yet challenging.

The Stang and Mukherjee groups pioneered the integrative self-assembly of pyridyl and carboxylic donors with metal acceptors via charge separation,³⁰⁻³² demonstrating its efficiency in preparing multicomponent metallacages. The heteroleptic self-assembly of platinum nodes with both multiple pyridyl and carboxylic ligands has also proved to be an efficient strategy for the construction of multicomponent metallacages in our group.33-44 Moreover, the homoleptic self-assembly of platinum nodes with multiple pyridyl ligands also provides a series of two-component metallacages.45,46 However, the preparation of metallacages via the self-assembly of platinum nodes with multiple carboxylic ligands has never been reported, although some metallacycles have been reported by this strategy,30 which is due to the decreased solubility and difficult structural characterization of such neutral, three-dimensional supramolecular structures. Herein, based on metal-coordination-driven selfassembly, we prepare a neutral, truncated tetrahedral metallacage (4a) and a heteroleptic truncated octahedral metallacage (4c) (Fig. 2), whose structures are unambiguously confirmed by X-ray diffraction analysis. Interestingly, further assembly of 4a with a previously reported cationic, truncated tetrahedron (4b)⁴⁵ leads to the quantitative formation of 4c, due to the energy of the heteroleptic metallacage significantly being lower than that with two homoleptic metallacages,30 representing a fusionbased supramolecular transformation from geometrically the

same truncated tetrahedron-shaped metallacages. This study not only contributes to the construction of novel coordination structures but also explores their fusion-based structural transformation, which will promote the development of supramolecular assemblies with tunable structures and integrated functions.

Results and discussion

Metallacage 4a or 4b was prepared by the self-assembly of hexatopic carboxylic ligand 1 (L₁) or pyridyl ligand 2 (L₂) with cis-Pt(PEt₃)₂(OTf)₂ 3 (M), while metallacage 4c was synthesized by the heteroleptic self-assembly of 1, 2 and 3 (Fig. 2, see the ESI[†] for synthetic details). All the metallacages were characterized by 1D multinuclear (³¹P{¹H} and ¹H) NMR, 2D diffusionordered NMR spectroscopy (DOSY) and electrospray ionization time-of-flight mass spectroscopy (ESI-TOF-MS), as well as X-ray diffraction analysis. In the ${}^{31}P{}^{1}H$ NMR spectra (Fig. 3a and b), only one single peak was observed for metallacages $4a (M_{12}(L_1)_4)$ at 2.52 ppm and **4b** $(M_{12}(L_2)_4)$ at 0.47 ppm, agreeing well with the single phosphorus environment in their symmetric structures. For metallacage $4c (M_{24}(L_1)_4(L_2)_4)$, two doublet peaks with equal intensities were found at 5.35 and 0.10 ppm (Fig. 3c), which is consistent with the different phosphorus environments after the coordination. In the ¹H NMR spectra, upfield chemical shifts were observed for protons H1, H2, H3 and H4 of 4a compared with its carboxylic precursor 1 (Fig. 3d and e). For metallacages 4b and 4c, downfield chemical shifts were observed for α -pyridyl protons H_a and β -pyridyl protons H_b compared with ligand 2 (Fig. 3f-h). Both H_a and H_b split into two sets of signals, which correspond to the protons inside and outside of the metallacages. In the 2D DOSY spectra, all the protons of metallacages 4b and 4c displayed a single diffusion coefficient (Fig. S13 and S17[†]), respectively, suggesting the formation of single discrete supramolecular structures. ESI-TOF-MS was carried out to afford the coordination stoichiometries of metallacages 4a-4c. For metallacage 4a (Fig. S7⁺), peaks at *m*/*z* = 1035.5970, 1180.2861, 1373.1576 and 1643.2198 were observed, corresponding to $[4a + 8Na]^{8+}$, $[4a + 7Na]^{7+}$, [4a +6Na]⁶⁺ and [4a + 5Na]⁵⁺, respectively. For metallacages 4b and **4c**, multiple prominent peaks of $[M-x(OTf)]^{x+}$ (x = 6-20) were observed, owing to the successive loss of counterions (OTf⁻) (Fig. 3i and j). For instance, peaks at m/z = 1266.9769 and 1454.3884 were found, corresponding to $[4b-9OTf]^{9+}$ and [4c-13OTf¹³⁺, respectively. All the peaks were isotopically resolved and matched well with their theoretical distributions. These results are consistent with previous reports,33-44 suggesting the successful formation of metallacages 4a-4c.

Single crystals of metallacages **4a** and **4c** were obtained through the slow diffusion of dioxane into a DMF (for **4a**) or ethyl acetate into a DMSO (for **4c**) solution of the samples for more than two months. In the crystal structure of **4a**, four hexatopic carboxylic ligands were connected by 12 platinum(II) nodes, forming a truncated tetrahedron (Fig. 4a). The diameter of **4a** was 3.4 nm and the shortest Pt…Pt distance was 0.82 nm. The angle of O–Pt–O was *ca.* 84.1°, while the angle of P–Pt–P was *ca.* 97.8°. The size of **4a** is close to that of **4b** (Fig. 4b), whose



Fig. 3 Partial ³¹P{¹H} NMR spectra (243 MHz, CD₃CN, 295 K) of (a) **4a**, (b) **4b** and (c) **4c**. Partial ¹H-NMR (600 MHz, 295 K) of (d) **1** (CD₃OD), (e) **4a** (CD₃CN), (f) **2** (CDCl₃), (g) **4b** (CD₃CN), and (h) **4c** (CD₃CN). The peak for CHCl₃ is marked with an asterisk. ESI-TOF-MS spectra of metallacages (i) **4b** and (j) **4c**.



Fig. 4 (a)–(c) Crystal structures of metallacages 4a, 4b (ref. 45) and 4c. Hydrogen atoms, counterions, and solvent molecules are omitted for clarity.

diameter reached 3.6 nm,⁴⁵ which would benefit their supramolecular fusion. For metallacage **4c**, four hexacarboxylic ligands (**1**) and four hexapyridyl units (**2**) were connected by 24 Pt nodes, forming a truncated octahedron with a diameter of 4.4 nm (Fig. 4c). Each carboxylic ligand was connected with its neighboring pyridyl unit through two O–Pt–N coordination bonds. The shortest Pt…Pt distance was 1.09 nm. The angle of O–Pt–N was *ca.* 84.4°, while the angle of P–Pt–P was *ca.* 96.0°. In the crystal packing of **4c** (Fig. S21[†]), regular packing was facilitated by non-covalent interactions between neighboring molecules, such as van der Waals force between the peripheral PEt₃ units and C–H···O hydrogen bonds. Due to the propellershaped structure of hexaphenylbenzene (HPB), the metallacages constructed with HPB as the main building block exhibit weak host–guest interactions with the common polycyclic aromatic hydrocarbon guest molecule (Fig. S30 and S31[†]).

Fig. 5 (a) Fluorescence spectra of ligand 1 in CH₃CN/H₂O ($\nu : \nu = 9 : 1$), ligand 2 and metallacages 4a–4c in CH₃CN. (b) Time-dependent fluorescence spectra of an equimolar mixture of 4a and 4b in CH₃CN. $\lambda_{ex} = 265$ nm and $c = 10.0 \ \mu$ M.

To the best of our knowledge, metallacage **4a** represents the first neutral metallacage self-assembled from Pt nodes and multiple carboxylic ligands, while metallacage **4c** is the largest metallacage formed by the heteroleptic assembly of Pt nodes with both pyridyl and carboxylic ligands, which will promote the design and preparation of metallacages with advanced geometric structures.

UV/Vis absorption spectroscopy and fluorescence spectroscopy were conducted to investigate the photophysical properties

of these ligands and metallacages. Ligands 1 and 2 displayed weak absorption bands centered at ca. 247 nm and 256 nm, with molar absorption coefficients (ϵ) of 3.67 \times 10⁴ and 1.50 \times $10^4 \text{ M}^{-1} \text{ cm}^{-1}$, respectively. Metallacages **4a** and **4b** exhibited dense absorption bands centered at ca. 354 nm and 276 nm, with ε of 9.27 \times 10⁴ and 1.83 \times 10⁵ M⁻¹ cm⁻¹, respectively (Fig. S22[†]). Metallacage 4c showed two absorption bands centered at *ca.* 297 and 342 nm (Fig. S22[†]), with ε of 3.07 \times 10⁵ and $1.09 \times 10^5 \,\text{M}^{-1} \,\text{cm}^{-1}$, respectively, which were derived from the absorption of ligands 1 and 2. Metallacages 4a and 4c showed weak emission centered at ca. 433 and 420 nm, respectively, while bright emission centered at ca. 450 nm was observed for metallacage 4b (Fig. 5a). The weak emission of 4a and 4c was due to the photoinduced electron transfer,47 while the bright emission of 4b was because of the restriction of molecular motions by metal-coordination bonds,48,49 which has also been observed in other supramolecular systems.^{50,51}

The distinct fluorescence characteristics of these metallacages provide a visual method to monitor their assembly and fusion processes by using the changes in fluorescence as output signals. Mixing **4a** and **4b** in a 1:1 molar ratio resulted in a gradually decreased emission over time, accompanied by



Fig. 6 (a) Time-dependent ESI-TOF-MS spectra of the mixture of 4a and 4b ($c = 500.0 \mu$ M). Partial ¹H (600 MHz) and ³¹P(¹H) NMR (243 MHz) spectra of (b) 4b, and 4b with (c) 0.2, (d) 0.4, (e) 0.6, (f) 0.8 and (g) 1.0 equivalents of 4a (CD₃CN at 295 K).

Chemical Science

a blueshift of the maximum emission (Fig. 5b). After 120 min, negligible changes were observed from the fluorescence spectra (Fig. S23†), suggesting that the reaction reached an equilibrium state and that the supramolecular fusion of geometrically similar **4a** and **4b** into **4c** was complete in 120 min.

ESI-TOF-MS was also carried out to study the kinetics of the fusion (Fig. 6a). Acetonitrile solutions containing the same amount of **4a** and **4b** were stirred at 60 °C and the ESI-TOF-MS spectra were collected at different time points. After 20 min, the peaks corresponding to **4c** started to appear, suggesting that the fusion process began to take place at this time point. As time progressed, the signals of **4b** gradually decreased, while those of **4c** increased. After 120 min, only the signals of **4c** were observed from the spectra, which agreed well with the fluorescence data, indicating that the fusion of **4a** and **4b** was complete.

The concentration-dependent fusion between 4a and 4b was investigated by NMR spectroscopy. The gradual addition of 4a into the acetonitrile solution of 4b resulted in significant changes in the NMR spectra (Fig. 6b-g). The peaks of protons H_a shifted upfield. while downfield chemical shifts were found for protons H_b, H_c and H_d. Two new peaks, derived from protons H₁ and H₂ of 4a, gradually emerged. After the addition of 1.0 equivalent of 4a into 4b, the ¹H NMR spectrum matched well with that of 4c, suggesting a quantitative transformation from the mixture of 4a and **4b** to **4c**. In the ${}^{31}P{}^{1}H$ NMR spectra, the single peak of **4b** slowly disappeared, while two doublet peaks corresponding to 4c were observed over time. Moreover, the variations in solvent and counterions showed little influence on the formation of 4c (Fig. S26-29[†]). Therefore, the fusion process of these assemblies was fully tracked and supported by the combination of fluorescence, mass and NMR spectroscopy, which provides a good example for the construction of supramolecular assemblies with improved structural complexity via supramolecular fusion. To understand the mechanism of the fusion from two truncated tetrahedron-shaped metallacages, we compared the energies of the sum of the two homoleptic $M_{12}L_4$ metallacages with that of 4cusing the semiempirical GFN1-xTB method. According to the comparison between the energy values of metallacages, the energy of 4c is significantly lower than the sum of the energies of the homoleptic metallacages (Fig. S32[†]), suggesting that the assembly of 4c is enthalpically more favorable.

Conclusions

In summary, a neutral truncated tetrahedron and a giant truncated octahedron were successfully prepared by metalcoordination-driven self-assembly. Interestingly, the neutral truncated tetrahedron could be quantitatively transformed into the truncated octahedron *via* its further assembly with another cationic truncated tetrahedron, representing a supramolecular fusion at the three-dimensional level from geometrically the same truncated tetrahedron-shaped metallacages. The fundamental knowledge obtained from the current research not only shows the process of supramolecular fusion, but also provides a simple and efficient strategy for preparing supramolecular assemblies with enhanced structural complexity and integrated functions.

Data availability

The data supporting this article have been included in the ESI.† Crystallographic data for **4a** (2314346) and **4c** (2314347) have been deposited at the Cambridge Crystallographic Data Centre.

Author contributions

M. Zhang and H. Liu conceived the project. H. Liu and Y. Huang carried out the synthesis and some characterization experiments of the compounds. C. Guo carried out the ESI-TOF-MS tests and analyzed the data. Z. Zhou, S. Jian, Z. Zhang, and Y. Hou assisted in the synthesis and characterization of the compounds. The manuscript was written by H. Liu, C. Mu, and M. Zhang and all authors contributed to the final draft of the paper.

Conflicts of interest

The authors declare no competing financial interests.

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