Designing narcissistic self-sorting terpyridine moieties with high coordination selectivity for complex metallo-supramolecules

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Coordination-driven self-assembly is a powerful approach for the construction of metallo-supramolecules, but designing coordination moieties that can drive the self-assembly with high selectivity and specificity remains a challenge. Here we report two ortho-modified terpyridine ligands that form head-to-tail coordination complexes with Zn(II). Both complexes show narcissistic self-sorting behaviour. In addition, starting from these ligands, we obtain two sterically congested multitopic ligands and use them to construct more complex metallo-supramolecules hexagons. Because of the non-coaxial structural restrictions in the rotation of terpyridine moieties, these hexagonal macrocycles can hierarchically self-assemble into giant cyclic nanostructures via edge-to-edge stacking, rather than face-to-face stacking. Our design of dissymmetrical coordination moieties from congested coordination pairs show remarkable self-assembly selectivity and specificity.
Coordination-driven self-assembly has been proven to be an effective approach for constructing supramolecular structures, due to its high predictability. As such, a variety of discrete metallo-supramolecular architectures have been fabricated through different types of self-assembly strategies, including the coordinations of pyridine, bipyridine, terpyridine, or heteroligand complexes with the loss of different numbers of PF6− ions, with the formation of self-sorting complexes that are highly selective in precise and controllable supermolecular assembly.

2,2′,6′,2″-tpy and its derivatives have been widely employed in coordination-driven self-assembly on account of their variable coordination ability with transition metal ions, as well as the unique properties and wide applications in optical devices, catalysis, self-healing, and drug delivery after complexation. Among the design of tpy-based metallo-supramolecules, the most common strategy is connecting tpy units and directing units at the central pyridine for subsequent coordination, which usually causes the poor selectivity of coordination units. To further enhance geometrical diversity and complexity of assemblies, a hierarchical stepwise assembly strategy has been developed to promote the controllability of the self-assembly process by using metals that can form strong coordinative bonds with tpy, such as <tpy-Ru(II)-tpy> or <tpy-Fe(II)-tpy>. However, the tedious synthesis and column separation procedures limit its extensive application in supramolecular construction. Currently, it still remains a challenge to precisely construct complex supramolecular structures through the one-step method.

Nature has given us the enlightenment in the exploration of hierarchical assembly of metallacycles by introducing orthogonal interactions. In the previous reports, it is found that the significantly steric congestion and non-coaxial structure of Zn6(LA)6 caused the restriction in the edging rotation of the hexagons, which further causes the hexagons to hierarchically assemble into giant cyclic nanostructures and metallogels via edge-to-edge stacking rather than face-to-face stacking.

Results and discussion

Synthesis and characterization of model system MA and complex Zn9(MA)2. In our design, the modification at terpyridyl 6-position played a central role in the self-sorting assembly. The simple and efficient synthesis process of MA only included a three-step reaction of the starting material, followed by the one-step purification of the product using column chromatography. Especially, the key compound was synthesized by a typical Körhrieks reaction with pyridinium salt 5 (Supplementary Fig. 1b). The final motif MA was obtained via Suzuki coupling reaction in a good yield (77%) and characterized by NMR and mass spectrometry (Supplementary Figs. 2–8). After that, MA and Zn(NO3)2·6H2O (with a precise stoichiometric ratio of 1:1, Fig. 2a) were mixed in CHCl3 and MeOH (1:3, v/v) at 50 °C for 12 h, followed by the addition of excessive NH4PF6 (to exchange NO3− with PF6−) in methanol to give a white precipitate Zn9(MA)2 in a yield of 91%.

1H NMR spectra of MA and complex Zn9(MA)2 were shown in Fig. 2b. Given the dissymmetrical nature of MA, five kinds of expected pyridines signals attributed to the terpyridine moieties were confirmed by 2D-COSY results (Supplementary Fig. 6). In the spectrum of complex Zn9(MA)2, the peaks were sharp and well-resolved. The 2D-COSY and NOESY for Zn9(MA)2 (Supplementary Figs. 9–11) also showed five sets of pyridines signals as MA, suggesting a highly symmetrical structure of complex Zn9(MA)2. Compared with MA, 6-position and 4-position signals of tpy were shifted upfield due to the electron shielding effect. At the same time, the proton signals of e-tpy, f-Ph, and g-Ph were also shifted to upfield, which should be attributed to the existence of π-π interaction between tpy moiety and phenyl in the other ligand. In ESI-MS (Supplementary Fig. 15A), one prominent set of peaks with charge states from 2+ to 4+ was observed (due to the loss of different numbers of PF6−). After deconvolution, the molecular weight of complex Zn9(MA)2 was 1940 Da, matching well with its expected chemical composition of Zn9(MA)2. Complex Zn9(MA)2 showed a series of charge states with narrow drift time distribution ranging from 2+ to 4+, indicating the formation of a discrete product but without any isomers and conformers (Supplementary Fig. 15B).

To further confirm the structure, the single crystal of Zn9(MA)2 was obtained by slowly diffusing the vapor of ethyl acetate into Zn9(MA)2 in acetonitrile for over 2 weeks. As expected, X-ray crystallographic analysis (Fig. 2c, d and Supplementary Table 1) showed that two Zn(II) are sandwiched between two MA to form a dimer with the head-to-tail structure...
The phenyl groups in the middle are parallel to each other (yellow part), and the distance between the two interlayers is 3.57 Å, indicating the existence of π–π interactions\(^{41,62}\).

Synthesis and characterization of model system MB and complex Zn\(_3\)(MB)\(_2\). By referring to the synthetic and characterization procedures of MA, we further designed and obtained another model system MB (Fig. 3a and Supplementary Fig. 17), which includes one additional tpy unit compared to MA and thereby exhibits an enhanced sterically congested effect. An extra tert-butyl was introduced to improve the solubility. Tpy-based molecules have been extensively studied since the 1930s, however, it is worth noting that the unique structure of MB, has never been reported\(^{41,63,64}\). The synthesis of MB (yield 75%) and self-assembly of complex Zn\(_3\)(MB)\(_2\) (yield 87%) (MB:Zn(II) = 2:3) followed the same procedure as that of MA and Zn\(_2\)(MA)\(_2\).

The \(^1\)H NMR data of MB and Zn\(_3\)(MB)\(_2\) were shown in Fig. 3b. In the spectrum of MB with three tpy units, eight kinds of pyridine signals could be observed (Supplementary Figs. 18–24). Moreover, the complex Zn\(_3\)(MB)\(_2\) exhibited a more complicated structure and spectrum than Zn\(_2\)(MA)\(_2\) (Supplementary Figs. 25–30), due to the enhanced steric hindrance effect caused by the introduction of the additional tpy unit. For each tpy, the differences in chemical shift (\(\Delta\delta\)) were 1.48 ppm and 0.8 ppm for 6 position proton and a6 proton, respectively, suggesting that the 6-position of a-tpy shows the strong shielding effect. The signals of tpy-3 and tpy-5 displayed upfield or downfield shifts after...
Fig. 2 The self-assembly and characterization of head-to-tail complex Zn$_2$(MA)$_2$. a The self-assembly of model system MA with Zn(II). b The $^1$H NMR spectra (600 MHz, 300 K) of MA in CDCl$_3$ and complex Zn$_2$(MA)$_2$ in CD$_3$CN (3 mg/mL). c X-ray crystal structure in a side view of complex Zn$_2$(MA)$_2$. d X-ray crystal structure in a front view of complex Zn$_2$(MA)$_2$. Non-coordinated anions and solvent are omitted for clarity (C, gray or yellow; H, white; N, blue; Zn, orange).

Fig. 3 The self-assembly and characterization of head-to-tail complex Zn$_3$(MB)$_2$. a The self-assembly of model system MB with Zn(II). b The $^1$H NMR spectra (600 MHz, 300 K) of MB in CDCl$_3$ and Zn$_3$(MB)$_2$ in CD$_3$CN (3 mg/mL). c X-ray crystal structure in a front view of Zn$_3$(MB)$_2$. Non-coordinated anions and solvent are omitted for clarity (C, gray or yellow; H, white; N, blue; Zn, orange).
coordination, which were consistent with the previous studies. In addition, the restrictive structure induced the split of NMR signals. Both the signals of the middle phenyls and tpy-b6 split into two sets of peaks, which indicate their rotation restriction caused by the altered chemical environment. ESI-MS and TWIM-MS (Supplementary Figs. 31, 32) spectra of Zn3(MB)2 also exhibited a similar prominent set of peaks with different charge states and narrow drift time distribution ranging from 2+ to 4+, suggesting the formation of a single product without any overlapping isomers or conformers. The molecular weight of complex Zn3(MB)2 (3020 Da) agrees well with its expected chemical composition of two MB ligands, three Zn(II) ions, and six PF6−.

The single-crystal data of Zn3(MB)2 was obtained by slowly diffusing the vapor of carbon tetrachloride into acetonitrile solution for over 3 weeks. Complex Zn3(MB)2 also showed a sandwich-like structure with a certain degree of distortion, suggesting higher steric congestion caused the helical shape (Fig. 3c). The different distances between the two middle phenyls (yellow part) were 3.6 and 3.7 Å, respectively, which confirmed the helical structure (Supplementary Data 2, Supplementary Table 2, and Supplementary Movie 2). More importantly, under the sterically congested environment, the different tpy units coordinated with the same Zn(II) cannot maintain the vertical alignments and are forced to exhibit twisted structures. Furthermore, Zn3(MB)2 has two chiral conformations due to the unique head-to-tail coordination mode, and both of the conformations could be found in a single crystal (Supplementary Fig. 33).

Self-sorting behavior of MA, MB, and tpy. To evaluate the selectivity of these model systems, we investigated the self-sorting behavior of MA, MB, and conventional tpy. In the self-sorting study, MA, MB, and conventional tpy were mixed together in an equimolar ratio with the corresponding amount of Zn(II) for the overnight self-assembly at 50 °C. ESI-MS clearly illustrated three sets of signals for the corresponding complexes but without statistical complexes (Fig. 4a, b), indicating that these model complexes exhibit ideal self-sorting properties in a pluralistic system. Both ESI-MS and NMR of any binary mixture (Supplementary Figs. 34–36) showed independent signals for three complexes Zn2(MA)2, Zn3(MB)2, and Zn(tpy)2, suggesting a characteristic narcissistic self-sorting. Moreover, we also monitored the kinetic process of narcissistic self-sorting, and the system basically reached equilibrium after 24 h (Supplementary Fig. 37). As expectation, in accord to the maximum occupancy of coordination sites proposed by Lehn, MA, MB, and tpy tend to narcissistically coordinate to form the most energy favorable structure. In this context, we are inspired to use MA and MB as self-recognition sites to further construct complex supramolecular structures.

Synthesis and self-assembly of ligands LA and LB with multiple tpy moieties. It is reported that the ditopic tpy ligands with 120° angle tend to self-assemble into a mixture of macrocycles with uncontrollable size and structure. However, we expect that the dissymmetrical tritopic (LA) and tetratopic (LB) ligands (Fig. 1c) ligands with 120° angle that contain narcissistic self-sorting moieties could be used to precisely construct complex supramolecular macrocycles Zn9(LA)6 and Zn12(LB)6 (Supplementary Movie 3 and Supplementary Movie 4). In order to improve the solubility of ligands, alkoxy chains (R−OC6H13) were introduced. Compared with LA (yield, 76%), the synthesis of LB with one more tpy units turns to be significantly challenging (Supplementary Fig. 38 and Supplementary Fig. 39). Fortunately, we successfully obtained LB (yield, 37%) by adding CH3COOH/DMF as an auxiliary agent. The obtained ligands LA and LB were assembled into complexes Zn9(LA)6 (yield, 86%) and Zn12(LB)6 (yield, 84%) and were characterized by NMR, MALDI-TOF, ESI-MS, and TWIM-MS (Fig. 5 and Supplementary Figs. 40–70).
Compared with LA and LB, $^1$H NMR spectra of the complexes Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$ were significantly broadened (Fig. 5a, d), suggesting the assembly of large complexes with a slower tumbling on the NMR time scale.$^{24}$ Importantly, the peaks of pyridines on the b-tpy and c-tpy turn broader, due to the sterical congestion of the coordination units and the restriction in free rotation of the phenyl caused by the high steric congestion of alkoxy chains. In the spectrum of LA, seven kinds of expected pyridines signals were observed with respect to the tpy moieties. Owing to the continuous dissymmetrical modification of a- and b-tpy, LB displayed more complicated proton signals (i.e., ten distinguished kinds of pyridine peaks) than that of LA. All six positions of pyridines were shifted towards upfield, because of the electron shielding effects after complexation. However, the chemical shifts of different protons differ significantly owing to the variable shielding strength. For instance, the proton at a$_6$ position showed a maximum chemical shift of $\Delta\delta = 1.41$ ppm; while a slightly upfield-shift $\Delta\delta = 0.55$ ppm was observed for b$_6$ position. 2D-COSY and NOESY for two complexes Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$ (Supplementary Figs. 53–56 and Supplementary Figs. 59–62) also showed the same sets of tpy signals as ligands, suggesting the symmetrical structures of these two complexes. The diffusion-ordered NMR spectroscopy (DOSY) provided dimensional information for Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$. As shown in Supplementary Fig. 66, all proton signals appeared at the same band, indicating the formation of discrete assemblies. The diffusion coefficients in CD$_3$CN were $\log D = -9.61$ for Zn$_9$(LA)$_6$ and $\log D = -9.64$ for Zn$_{12}$(LB)$_6$. The experimental hydrodynamic radii ($r_H$) for Zn$_9$(LA)$_6$ (2.4 nm) and Zn$_{12}$(LB)$_6$ (2.6 nm) agreed well with the modeling structures.

In ESI-MS (Fig. 5b, e), one prominent set of peaks with charge states (from 7$^+$ to 14$^+$ for Zn$_9$(LA)$_6$ and from 8$^+$ to 14$^+$ for Zn$_{12}$(LB)$_6$) was observed on account of the loss of different numbers of PF$_6^-$ . After deconvolution, the molecular weights of these two complexes were 8869 and 12974 Da, respectively, matching well with the expected chemical composition of Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$. 

Fig. 5 Characterization of the hexagonal macrocycles Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$. a $^1$H NMR spectra (600 MHz, 300 K) of ligand LA in CDCl$_3$ and hexagon Zn$_9$(LA)$_6$ in CD$_3$CN (4 mg/mL). b ESI-MS, and c TWIM-MS plots (m/z vs drift time) of Zn$_9$(LA)$_6$. d $^1$H NMR spectra (600 MHz, 300 K) of ligand LB in CDCl$_3$ and hexagon Zn$_{12}$(LB)$_6$ in CD$_3$CN (4 mg/mL). e ESI-MS and f TWIM-MS plots (m/z vs drift time) of Zn$_{12}$(LB)$_6$. 

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Zn₉(LA)₆ (6 LA ligands, 9 Zn(II) ions, and 18 PF₆⁻) and Zn₁₂(LB)₆ (6 LB ligands, 12 Zn(II) ions, and 24 PF₆⁻). All the experimental isotope patterns agreed excellently with the corresponding simulated isotope patterns (Supplementary Figs. 69, 70). Moreover, the TWIM-MS spectra of complexes Zn₉(LA)₆ and Zn₁₂(LB)₆ also showed narrow drift time distribution, supporting the formation of single species with rigid structures (Fig. 5c, f). The full characterization of NMR, ESI-MS, and TWIM-MS confirmed that the two types of tpy moieties in ligands LA and LB could precisely self-assemble into hexagonal macrocycles through the narcissistic self-recognition mechanism. Such moieties with high coordination selectivity and specificity may find their potential in the constructions of more complex metallo-supramolecules with precisely controlled shapes and sizes.

Interestingly, when complexes Zn₉(LA)₆ and Zn₁₂(LB)₆ were mixed in an equimolar ratio at 50 °C for overnight, ESI-MS and TWIM-MS clearly illustrated two additional sets of signals besides Zn₉(LA)₆ and Zn₁₂(LB)₆ (Supplementary Fig. 71). The compositions of Zn₁₀(LA)₄(LB)₂ and Zn₁₁(LA)₂(LB)₄ were confirmed by analyzing the ESI-MS and TWIM-MS data (Supplementary Figs. 72, 73). No exchange for odd number ligands was observed in this mixture (Supplementary Figs. 74–77), suggesting that it could be a subcomponent exchange based on the dominant coordination dimer rather than the usual ligand exchange⁶⁹,⁷⁰. It indicated that the dimer structures based on
ditopic and tritopic tpy moieties not only exhibit higher selectivity but also show higher stability than single-tpy unit.

Hierarchical self-assembly. Our study shows that, with increasing the concentration of Zn₉(LA)₆ and Zn₁₂(LB)₆ in acetonitrile at room temperature, both of the solutions could form metallo-gels at a concentration of 45 mg/mL for Zn₉(LA)₆ and 25 mg/mL for Zn₁₂(LB)₆ (Fig. 6l). The gels were also temperature-responsive and could be converted into solutions after heating at 50 °C for 1 h. To explore the mechanism of gelation, we then used TEM to investigate the morphology of Zn₉(LA)₆ and Zn₁₂(LB)₆ at different concentrations (Supplementary Figs. 78, 79). At low concentrations (10⁻⁶ M), the assemblies were almost distributed in a monodispersed manner. Both complexes exhibited unique hollow structures, suggesting high rigidity, stability, and large inner space. The measured sizes of the two complexes were 5.5 nm for Zn₉(LA)₆ and 6.5 nm for Zn₁₂(LB)₆ respectively (Fig. 6b, c, f, g). However, at a high concentration (10⁻⁵ M), some giant cyclic nanostructures in larger size were observed (Fig. 6d). The diameters of these structures were much larger than that of individual Zn₉(LA)₆, implying the clustering of Zn₉(LA)₆. The height of the hierarchically assembled nanostructures measured by the AFM image (Fig. 6d, k and Supplementary Figs. 80–83) was about 1.5 nm, which was very close to the height of the monolayer. When the concentration further increased, these giant macrocycles formed a three-dimensional network via intermolecular interaction (Fig. 6e). For Zn₁₂(LB)₆ with a larger size, although similar monolayer nanostructures were observed (Fig. 6h), the degree of aggregation was significantly increased (Supplementary Figs. 84–86). Further, at a much higher concentration of 10⁻⁴ M, the formed giant macrocycles showed obvious vertical stacking in space (Fig. 6l).

As well-known, the <tpy-M(II)-tpy> unit could rotate freely around the axis without space obstruction. However, due to the special spatially complementary coordination of LA and LB, these two axial dissymmetrical ligands exhibited entirely steric congestions. Therefore, it can be inferred that the hexagons are basically composed of vertical edges, since the rotation of the dimer can result in the disassembly (Fig. 6a). In order to study the edge-to-edge stacking of Zn₉(LA)₆, we tried to grow single crystal of Zn₉(LA)₆. However, all efforts to grow a single crystal of Zn₉(LA)₆ has proven to be unsuccessful. Instead, we synthesized a model ligand MA-OC₆H₁₃ (Supplementary Figs. 87–94) and further got the complex Zn₂(MA-OC₆H₁₃)₂ which has exactly the same structure as the ditopic part of Zn₉(LA)₆ (Supplementary Figs. 95–102). The single-crystal packing of Zn₂(MA-OC₆H₁₃)₂ provided strong evidence for the edge-to-edge stacking of Zn₉(LA)₆ (Supplementary Data 3 and Supplementary Table 3). As shown in Fig. 7, the intermolecular packing of Zn₂(MA-OC₆H₁₃)₂ is more compact than that of Zn₂(MA)₂ (Supplementary Fig. 103), which indicated that the introduction of alkyl chains can enhance the intermolecular interactions. The distance...
between the alkyl chain and central phenyl of another molecule is originating from intra-ligand charge transfer (ILCT). In sharp macrocycles Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$, the restrictive intramolecular conjugation caused the restriction in rotations of the hexagons, which further leads to their hierarchical assembly into giant cyclic nanostructures and metallogels. Moreover, these complexes showed significantly enhanced fluorescence intensity in the solution state than the complexes based on conventional tpy due to the additional ligand conjugation and the restricted chemical environment. Our design and fabrication of dissymmetrical coordination moieties could pave a new avenue for the development of a set of congested coordination pairs with high selectivity and specificity for the assembly of sequence-specific metallo-supramolecular architectures.

### Methods

**General procedures.** All reagents were purchased from Sigma-Aldrich, Matrix Scientific, Alfa Aesar, Jilin Chinese Academy of Sciences—Yanshen Technology Co. Ltd., and used without further purification. Column chromatography was conducted using SiO$_2$ (VWR, 40–60 μm, 60 Å) and the separated products were visualized by UV light.

**Synthesis.** All the new compounds were fully characterized and spectra are given in Supplementary Methods.

**Nuclear magnetic resonance (NMR).** NMR spectra data were recorded on a 400, 500, and 600 MHz Bruker Avance NMR spectrometer in CDCl$_3$ or CD$_3$CN with TMS as reference. For NMR spectra see Supplementary Information.

**ESI-MS and TWIM-MS.** Electrospray ionization (ESI) mass spectra were recorded with a Waters Synapt G2 tandem mass spectrometer, using solutions of 0.5 mg sample in 1 mL of MeCN/MeOH (3:1, v/v) for complexes. The TWIM-MS experiments were performed under the following conditions: ESI capillary voltage, 3 kV; sample cone voltage, 30 V; extraction cone voltage, 3.5 V; source temperature 100 °C; desolvation temperature, 100 °C; cone gas flow, 10 L/h; desolvation gas flow, 700 L/H (N$_2$); source gas control, 0 mL/min; trap gas control, 2 mL/min; helium cell gas control, 100 mL/min; ion mobility (IM) cell gas control, 30 mL/min, sample flow rate, 5 μL/min; IM traveling wave height, 25 V; and IM traveling wave velocity, 1000 m/s.

**Single-crystal X-ray diffractions.** X-ray diffraction data for Zn$_9$(MA)$_6$ and Zn$_9$(MA-OC$_6$H$_{13}$)$_6$ were measured by a Bruker D8 Venture X-ray single-crystal diffractometer using a Cu·Kα radiation (λ = 1.54178 Å) at 100 K. X-ray diffraction data for Zn$_9$(MB)$_6$ was collected using synchrotron radiation and MAR325 CCD detector at Shanghai Synchrotron Radiation BL17B Beamline. Crystallographic data and structural characteristics of Zn$_9$(MA)$_6$, Zn$_9$(MA-OC$_6$H$_{13}$)$_6$, and Zn$_9$(MB)$_6$ are summarized in Supplementary Tables 1–3, respectively. Crystallographic information files for the complexes are provided in Supplementary Data 1–3, respectively.

**Transmission electron microscopy (TEM) analysis.** The sample of Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$ were measured by a JEOL JEM-2100F transmission electron microscope. The TEM images of the drop cast samples were taken with a JEM-2100F transmission electron microscope.

**AFM imaging.** AFM imaging was performed on a Bruker Dimension Icon AFM system with ScanAsyst and the data was processed by NanoScope Analysis version 2.0 (Bruker Software, Inc.). The sample of Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$ were dissolved

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**Photophysical properties.** Considering the enhanced conjugation and restricted rotation of the structures, we further investigated UV/Vis absorption and photoluminescence of all ligands and complexes in the solution state (10$^{-6}$ M). As shown in Fig. 8a, compared with the absorption spectra of ligands (Supplementary Figs. 104, 105), all complexes displayed a significant redshift originating from intra-ligand charge transfer (ILCT). In sharp contrast with Zn(tpy)$_2$, the maximum emission wavelength of Zn$_9$(MA)$_2$ displayed a ca. 75 nm redshift, ascribed to the enhanced conjugation (Fig. 8b). However, complex Zn$_9$(MB)$_2$ with a more complicated structure also showed a similar maximum emission wavelength to Zn$_9$(MA)$_2$, the reason should be discussed in the text. For the introduction of alkoxy chains as electron donors, the maximum emission peaks of Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$ exhibited 20 nm red-shifted than that of Zn$_9$(MA)$_2$ and Zn$_9$(MB)$_2$. The maximum emission peaks of Zn$_9$(LA)$_6$ and Zn$_{12}$(LB)$_6$ exhibited 20 nm red-shifted than that of Zn$_9$(MA)$_2$ and Zn$_9$(MB)$_2$.
in CH$_2$CN at a concentration of 10$^{-5}$ M. The solution was left overnight and then dropped cast onto a silicon wafer after surface cleaning.

**Photophysical measurements.** UV-vis spectra of solutions were recorded on a PerkinElmer LAMBDA-365 spectrophotometer. Fluorescence emission spectra were measured by a Shimadzu spectrofluorimeter RF-5301PC. Solutions were placed in 1 cm path length quartz cells.

**Molecular modeling.** Energy-minimized structures were obtained following the settings in the literature. Calculations were performed with Geometry Optimization and followed by Anneal in Forcite module of Materials Studio version 8.0 program (Accelrys Software, Inc.).

**Data availability**

The authors declare that all data supporting the findings of this study are available within the article and Supplementary Information files, and also from the corresponding author upon reasonable request. The X-ray crystallographic coordinates for Zn$_2$(MA)$_2$ (Supplementary Data 1), Zn$_2$(MB)$_2$ (Supplementary Data 2), and Zn$_2$(MA-OC$_2$H$_5$)$_2$ (Supplementary Data 3) have been deposited at the Cambridge Crystallographic Data Centre (CCDC), under deposition numbers CCDC 2050233, 2050080, and 2106067, respectively. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre at www.ccdc.cam.ac.uk/data_request/cif.

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Author contributions

M.W. conceived and designed the experiments. J.M. and K.L. completed the synthesis. J.M., T.L., Y.X., Z.L., J.S., Q.B., Z.Z., X.-Q.H., Z.C. and P.W. conducted the characterization, J.M., T.L., X.D. and M.W. analysed the data and wrote the manuscript. All the authors discussed the results and commented on and proofread the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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