Hierarchical self-assembly of discrete bis-[2]pseudorotaxane metallacycle with bis-pillar[5]arene via host–guest interactions and their redox-responsive behaviors†

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A discrete rhomboidal metallacycle R functionalized with bis-[2]pseudorotaxane of [Cu(phenanthroline)]2† derivatives was successfully synthesized via coordination-driven self-assembly. Furthermore, the host–guest complexation of such a bis-[2]pseudorotaxane metallacycle with a bis-pillar[5]arene (bisP5) allowed for the formation of a new family of cross-linked supramolecular polymers R⃗(bisP5)2, which displayed interesting redox-responsive properties. By taking advantage of the substantial structural differences between the coordination geometries of [Cu(phenanthroline)]3+ and [Cu(phenanthroline)]2+1, the weight-average diffusion coefficients D of the supramolecular polymer were adjusted through changing the redox state of the Cu(I)/Cu(II) complexes.

Introduction

Hierarchical self-assembly (HSA) is defined as a multilevel spontaneous organizing process in which the precisely controlled self-assembly of components is realized by non-covalent interactions. It has become increasingly clear that HSA offers a powerful bottom-up strategy for constructing essential biomolecules such as DNA, RNA, and proteins, the structural integrities of which are shown to be superior to those of the nonhierarchical self-assembled ones. Inspired by nature, HAS has been widely applied to the fabrication of desirable functional supramolecular architectures with well-defined structures, intriguing properties, and multiple functions.1–3 Among these, coordination-driven self-assembly, as one of the most feasible and efficient methodologies, makes use of directional coordinative bonds between the building blocks and the metal ions to construct well-defined supramolecular coordination complexes (SCCs) such as two-dimensional (2-D) polygons and (3-D) three-dimensional polyhedron.4–9

Based on coordination-driven self-assembly, successful strategies have been developed for designing supramolecular polymers cross-linked by discrete metallacycles or metalacages. The combination of relatively rigid, discrete metal–organic scaffolds with elastic polymer networks allows for the construction of a new class of hybrid soft materials with stimuli-responsive properties. The discrete metallacages or metalacages were used either as metallacycle skeletons or stimulus response sites in these supramolecular polymers as reported by various research groups.10–19 Such metallosupramolecular architectures are of particular interest not only for their fascinating structures but also because of their stimuli-sensitivities to a wide variety of environmental conditions including solvent composition, pH, temperature, light irradiation, and so on.20,21 However, the metallacycle-based redox-responsive supramolecular polymers are still rare.

One way to access such redox-responsive supramolecular materials is to utilize host–guest repeating units which contain one or more redox-active species, for example disulphide, selenium, ferrocene, tetraethylvalene, transition metal ions, and viologen counterparts.22–24 In particular, coordination complexes based on transition metals and bidentate pyridine ligands, notably Cu(i)–phenanthroline complexes, typically present desirable redox properties for diverse applications such as molecular machines, catalysis, ultrasfast spectroscopy, solar energy conversion, optoelectronics, and luminescence.25,26 Moreover, the Cu(i)–phenanthroline complexes feature some promising characteristics including the extra structural rigidity afforded by the phenanthroline unit, and the Cu(i)–phenanthroline complexes being oxidized to Cu(II)–phenanthroline complexes that undergoes significant and rapid Jahn–Teller (J–T) distortion from D3d to D2
symmetry. With this strategy, Sauvage and co-workers have successfully constructed topologically spectacular architectures such as pseudorotaxanes, rotaxanes, catenates and knots with Cu(i)-phenanthroline complexes. It is therefore possible to further utilize the Cu(i)-phenanthroline complexes as the basic building blocks to develop redox-responsive supramolecular polymers with the topology of [2]pseudorotaxane.

Using the HSA method, we previously reported a novel supramolecular polymer which was based on hexagonal metallacycle skeletons containing the above-mentioned Cu(i)-phenanthroline moiety. It was displayed that the redox-responsive of the resulting supramolecular polymers are strongly dependent on the oxidation state of copper. In order to further examine the versatility of this synthesis strategy and investigate the effects of metallacycle skeleton on the redox-responsive behaviours of the supramolecular polymers, we herein report another type of redox supramolecular polymer constructed by the well-defined rhomboidal bis-[2]pseudorotaxanes metallacycle through the hierarchical orthogonal design strategy based on coordination-driven self-assembly and host-guest interactions. First, the rhomboidal metallacycle \( R \) containing two \([2]\) pseudorotaxane moieties was constructed by the coordination between 1,10-phenanthroline(phen) derivatives containing \([2]\) pseudorotaxane and 60° di-Pt(II) acceptors, resulting in a rhomboidal metallacycle \( R \) of which the skeleton size is less than the previously reported hexagonal metallacycle \( M \). Subsequently, hyperbranched supramolecular polymer was successfully constructed in the high-concentration region through host-guest interactions of rhomboidal metallacycle with the bis-pillar[5]arene (bisP5). Notably, the host-guest interactions were similar to covalent bond end-capping process and the Cu(i)N4 \([2]\) pseudorotaxanes could act as the active sites which respond to the redox stimuli in supramolecular polymers. Supramolecular polymerization was shown to help the otherwise unstable tetrahedral complex Cu(i)N4 maintain enough stability during redox reactions. In addition, the tunability of the diffusion constants upon redox stimuli was also compared between the supramolecular polymers made of the \( R \) and \( M \), showing the influence caused by the skeleton size.

**Results and discussion**

**Synthesis and characterization**

With the 120° macrocycle \( 1 \) incorporating a phen moiety and ligand 2 possessing cyano sites in hand, the construction work of \([2]\)pseudorotaxane with bis-pyridine(120°) donor \( D \) was carried out using an approach derived from the transition Cu(i) template strategy. The threading process leading to \( D \), using Cu(i) as templating agent, was shown to be quantitative by \(^1H\) NMR. By simply mixing the donor building block \( D \) with the corresponding 60° acceptor \( A \) in a 1 : 1 ratio in acetone at 50°C for five hours, a novel bis-[2]pseudorotaxanes rhomboidal metallacycle \( R \) was successfully obtained in near-quantitative yield via coordination-driven self-assembly (Scheme 1). It shall be noted that the skeleton size of such a rhomboidal metallacycle is less than that of the previously assembled hexagonal metallacycle \( M \) which used a 120° acceptor.

In the \(^1H\) NMR spectrum (Fig. 1 and S1†), the \( \alpha \)- and \( \beta \)-pyridyl hydrogen signals both exhibited the obvious downfield shifts (\( \Delta \delta \approx 0.48 \) and 0.53 ppm for \( \alpha \)-H; \( \Delta \delta \approx 0.47 \) and 0.54 ppm for \( \beta \)-H) because of the loss of electron density that occurred upon coordination of the pyridine-N atom with the Pt(II) metal center. Notably, two doublets were observed for both the \( \alpha \)- and \( \beta \)-pyridine protons, which might be attributed to hindered rotation about the Pt-N(pyridyl) bond in a small cavity. The \(^{31}P\) NMR spectra of the metallacycle \( R \) displayed a sharp singlet (\( \delta = 14.48 \) ppm), which shifted upfield from the signal of the starting platinum acceptor \( A \) by approximately 5.46 ppm (Fig. 1). This change, as well as the decrease in coupling of the flanking \(^{195}Pt\) satellites, is consistent with back-donation from the platinum.
atoms. Further characterization by 2-D $^1$H NMR spectroscopic techniques ($^1$H–$^1$H COSY and NOESY) were in agreement with the formation of the discrete bis-pseudorotaxanes metallacycles (Fig. S18†). For instance, the presence of cross-peaks between the signal of the pyridine protons (π-H and β-H) and the PEt₃ protons (CH₂ and CH₃) were found in the 2-D NOESY spectrum, which supported the structure of metallacycle R as a result of the formation of N–Pt bonds (Fig. S18b†).

The structure of metallacycle R was also confirmed by ESI-TOF-MS, which allowed the assembly to remain intact to the maximum extent during the ionization process while retaining the high resolution required for resolving the isotopic distribution. For instance, the ESI-TOF-MS spectrum of R revealed three signals that corresponded to different charge states resulting from the loss of PF₆⁻ counterions, [R–3PF₆]³⁺, [R–4PF₆]⁴⁺, and [R–5PF₆]⁵⁺, respectively, in which R represents the intact assembly (Fig. 2). Further investigation revealed that each isotope pattern of these signals was in good agreement with the corresponding simulation result. All evidence above supports the successful formation of discrete bis-[2]pseudorotaxanes rhomboidal metallacycle R by the donor D and accepter A via coordination-driven self-assembly (Scheme 1).

**Cross-linked supramolecular polymers constructed through host–guest interactions**

The supramolecular chemistry of pillar[n]arenes has developed immensely over the past ten years on account of their excellent inclusion properties and many intriguing applications in the constructions of molecular switches/machines, metal–organic frameworks, pH-responsive vesicles, as well as artificial transmembrane channels. Pillararenes are a new family of macrocyclic host molecules that possess symmetrical and columnar geometries with rigid and π-rich cavities. Compared with other macrocyclic hosts, the most peculiar host–guest properties of pillararenes are their strong affinities towards neutral guests by CH/π hydrogen bonds in organic media. For example, Yang’s group previously reported a multiple stimuli-responsive supramolecular polymer through orthogonal coordination-driven self-assembly and host–guest complexation of multi-pillar[5]arene metallacycles and ditopic neutral guests. We have also previously reported the strong binding behavior between D and pillar[5]arene dimer bisP₅ (Fig. S3†), showing that the linear long chain part of ditopic guest D was able to thread through the cavity of bisP₅ to form poly[3]pseudorotaxanes in solution. By taking advantage of the strong affinities of pillar[5]arenes with neutral guests in organic solvents, we envisioned that cross-linked supramolecular polymers could be prepared from the bisP₅ via host–guest interactions when the neutral tetratopic metallacycles R were employed as guests.

By making use of the recognition motif between the ditopic guest D and the host bisP₅ along with their good solubility in acetone, the mixtures of tetratopic metallacycles R with the host bisP₅ (1 : 2 molar ratio of R and bisP₅) at high concentration was expected to form a new family of cross-linked supramolecular polymers R ⊃ bisP₅. The concentration-dependent $^1$H NMR studies of complexes R ⊃ bisP₅ demonstrated the formation of such cross-linked supramolecular polymers. For instance, $^1$H NMR spectra (500 MHz, acetone-$d_6$, 296 K) of the complex of R and bisP₅ in a 1 : 2 molar ratio over a range of metallacycle R unit concentrations from 0.1 up to 3.0 mM were recorded with the purpose of probing the process of the supramolecular polymer formation (Fig. 3a). At low concentration (~0.5 mM), no obvious signal broadening was found, indicating that poly[3]pseudorotaxanes predominated in solution at low concentration. Upon increasing the concentrations, the signal of H₅, H₆ and H₇ gradually became unobvious and

![Fig. 2](image-url) Theoretical (top) and experimental (bottom) ESI-TOF-MS spectra of bis-[2]pseudorotaxane rhomboidal metallacycles R.

![Fig. 3](image-url) (a) Partial $^1$H NMR spectra (500 MHz, acetone-$d_6$, 296 K) of R ⊃ bisP₅ at different concentrations. (b) 2-D DOSY (500 MHz, 296 K) plot of solutions in acetone-$d_6$ of 1 : 2 molar ratio of R and bisP₅ at different concentrations, respectively. (c) SEM image of cross-linked supramolecular polymers R ⊃ bisP₅ at concentration of 4.0 mM in CH₃COCH₃.
finally disappeared. In the meanwhile, the Hm peak became broadened markedly at higher concentrations. These observations gave clean evidence for the formation of cross-linked supramolecular polymers through host-guest interactions (Fig. S4†) in the high concentration region.\textsuperscript{34}

Two-dimensional diffusion-ordered \textsuperscript{1}H NMR spectroscopy (DOSY) experiments were performed to provide further evidence for the formation of cross-linked supramolecular polymers. From Fig. 3b, the measured weight average diffusion coefficient \((D)\) of \(R \supset \text{bisP5}\) decreased remarkably from \(3.55 \times 10^{-9}\) to \(0.2 \times 10^{-9}\) \(\text{m}^2 \text{s}^{-1}\) \((D 1.0 \text{ mM}/D 12.0 \text{ mM} = 17.8)\) as the concentration of \(R \supset \text{bisP5}\) increased from 1.0 to 12.0 mM. In general, a more than 10-fold decrease of the diffusion coefficient is believed to be the important evidence for the formation of a polymerization with high degree.\textsuperscript{41} Therefore, the results from Fig. 3b clearly indicates that the initially small poly[3]pseudorotaxanes grew into larger across-linked supramolecular polymers with the increase in \(R \supset \text{bisP5}\) concentration. These DOSY experimental results also further revealed that the generation of such supramolecular polymers was concentration dependent. Importantly, a higher diffusion coefficient \((D)\) of the generated supramolecular aggregates were observed for \(R \supset \text{bisP5}\) when comparing with the previously reported \(M \supset \text{bisP5}\) at the same concentration of \([\text{Cu(phenanthroline)}]^{+}\) unit. For example, the \(D\) values of \(R \supset \text{bisP5}\) and \(M \supset \text{bisP5}\) were \(1.26 \times 10^{-9}\) \(\text{m}^2 \text{s}^{-1}\) and \(0.35 \times 10^{-9}\) \(\text{m}^2 \text{s}^{-1}\), respectively, when the concentration of \([\text{Cu(phenanthroline)}]^{+}\) unit was 6 mM in acetone (Fig. S5†). Such a difference in the diffusion constants indicates that the rhomboidal metallacycle \(R\) probably yielded in smaller supramolecular polymers than the hexagonal metallacycle \(M\), therefore proving that the skeleton of the metallacycle has a significant effect on supramolecular polymers size.

In order to further demonstrate the role of \text{bisP5} as the “cross-linking agent” in the formation of such supramolecular polymers. The scanning electron microscopy (SEM) studies were performed, which provided the direct evidence for the existence of supramolecular polymerization of \(R \supset \text{bisP5}\) and further insight into the morphological characteristics of cross-linked supramolecular polymers. As shown in Fig. S6,\textsuperscript{†} the typical spherical morphologies for metallacycle \(R\) in acetone solvents were observed at low concentration (\(\leq 1.0\) mM). With the increasing of concentrations, the formation of cube morphologies and the increase of spherical radius for metallacycle \(R\) in acetone solvents were observed, indicating the predominance of the oligomers when the concentration of metallacycle \(R\) is more than 1 mM. On the other hand, the more irregular and extended three-dimensional (3-D) networks for \(R \supset \text{bisP5}\) systems were observed with the presence of \text{bisP5}, which was consistent with the typical morphology of cross-linked supramolecular polymers (Fig. 3c and S7†).

**Redox-responsive supramolecular polymer**

As we know, the reversible redox state of the Cu(1)/Cu(II) complex transition could be tuned by chemical oxidation and reduction processes. Thus, the redox-responsive properties of the resulting supramolecular polymers \(R \supset \text{bisP5}\) were further exploited by the manipulation of the redox reactions between copper(i) and copper(II) (Scheme 2). We have previously investigated the redox-responded behaviors of the poly-pseudorotaxanes \(D \supset \text{bisP5}\) in acetone.\textsuperscript{34} The formation of poly[3]pseudorotaxanes was seen to be very stable between the \text{bisP5} and cyanide chain through host-guest interactions, which was similar to a covalent bond end-capping process and prevented the structure 2 slipping off the macrocycle 1 in the unstable tetrahedral complex Cu(II)N4 when the Cu(i) was oxidized to Cu(II).

In current work, the redox-responded behaviors of supramolecular polymers \(R \supset \text{bisP5}\) were also examined. Likewise, \textsuperscript{1}H NMR, \textsuperscript{31}P NMR, UV-vis, DOSY and SEM experiments showed that the supramolecular polymer \(R \supset \text{bisP5}\) were almost unbroken upon the redox stimulation. In the partial \textsuperscript{1}H NMR measurements, signals from the protons Hm and Hm' on the

![Scheme 2](image_url)  
**Scheme 2** Schematic representation of the formation of redox-responsive supramolecular polymer.
phenanthroline moiety almost disappeared when Cu(i) was oxidized into Cu(ii) by NOBF₄. Interestingly, the proton peaks of R ≻ [bisP₅]₂ were almost completely restored when the Cu(ii) was reduced to Cu(i) by ascorbic acid (ASA) (Fig. 4a and S8†), showing the reversibility of redox manipulations on R ≻ [bisP₅]₂. More importantly, the signal of the proton Hₖ remained absent during the redox process, indicating that the [bisP₅] exhibited very strong binding affinities towards the dinitriles guest which resulted in the stable cross-linked supramolecular polymers. ³¹P NMR and SEM of supramolecular polymers also showed negligible changes during the redox-responded processes (Fig. 4b and S9†). These results demonstrate that the formation of stable supramolecular polymers with rhomboidal metallacycle scaffolds in the extended network through host–guest interactions and degrees of such cross-linked polymerization remain unchanged during the redox reactions.

To verify the actual occurrence of the redox reactions, the UV-vis spectroscopic measurements on reduced and oxidized R ≻ [bisP₅]₂ in acetone (0.1 mM) were carried out (Fig. 4c). A metal-to-ligand charge-transfer (MLCT) absorption band at 437 nm was observed in UV-vis absorption spectra for the original R ≻ [bisP₅]₂ solution, indicated that the entwined arrangement of both phen units around Cu(i). When R ≻ [bisP₅]₂ was oxidized by addition of NOBF₄, the color of the solution changed from red-brown to emerald-green, while a new d–d absorption band appeared at 667 nm and the absorption peak at 550 nm became weaker. The addition of ascorbic acid formed as well during the oxidation and reduction processes. As was reduced to Cu(i) by ascorbic acid (ASA) (Fig. 4a and S8†), ([Cu(phenanthroline)]²⁺ unit = 9 mM) (D_Cu(i)/D_Cu(ii) = 2.54) (Fig. 4d and S10†). We have previously reported that the oxidation of M ≻ [bisP₅]₂ ([Cu(phenanthroline)]²⁺ unit = 9 mM) resulted in a more dramatic increase of the measured weight-average diffusion coefficients D from 0.3 × 10⁻⁹ to 1.55 × 10⁻⁹ m² s⁻¹ (D_Cu(ii)/D_Cu(i) = 4.70). These observations clearly demonstrated that the larger frame size of metallacycle has greater influence on the size changes of supramolecular polymer after oxidation (Table S1†). Moreover, the weight-average diffusion coefficient D of R ≻ [bisP₅]₂ decreased to 0.62 × 10⁻⁹ when 8.0 equiv. ASA was added to the Cu(ii)N₄ on the R ≻ [bisP₅]₂. Further analysis indicated that the above phenomena mainly resulted from the fact that the Cu(i)N₄ complexes with the distorted tetrahedral geometry of approximate C₂ symmetry were transformed into Cu(ii)N₄ with the D₂ symmetry structures and the symmetry of the D₂ structures was higher than that of the C₂ symmetry structures. The change of metal complexes from the more asymmetric ones to highly symmetric structures drove a loose supramolecular polymer structure to form a more compact network, which might caused an increase in diffusion coefficients (Scheme 2).

### Conclusions

In summary, we have constructed a discrete rhomboidal metallacycle R with bis[2]pseudorotaxanes via coordination-driven self-assembly. The structure of rhomboidal metallacycle R has been determined by ¹H NMR, ³¹P NMR, CSI-TOP-MS, DOSY. The supramolecular polymers R ≻ [bisP₅]₂, [3]pseudorotaxanes containing a Cu(i)–phenanthroline complex as a repeating units, was successfully constructed at the high-concentration region through host–guest interactions of bis[2]pseudorotaxanes metallacycles R with the bis-pillar[5]arene (bisP₅). Subsequently, we have investigated interesting redox-responsive properties of the supramolecular polymers variation in R ≻ [bisP₅]₂ system by switching the redox states of Cu(i)/Cu(ii) ions. The R ≻ [bisP₅]₂ system exhibit reversible chromatographic at low concentration and the weight-average diffusion coefficients D at high-concentration controlled by changing the redox state of the Cu(i)/Cu(ii) complexes. It’s interesting that the smaller frame size of metallacycle has smaller influence on the size changes of redox-responsive supramolecular polymer. The redox-responsive supramolecular polymers are also promising as a unique material that can encapsulate and localize bioactive molecules and cells within the material matrix.

### Conflicts of interest

There are no conflicts to declare.

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Notes and references

1 J.-M. Lehn, Chem. Soc. Rev., 2007, 36, 151–160.
2 G. M. Whitesides, J. P. Mathias and C. T. Seto, Science, 1991, 254, 1312–1319.
3 Y. He, T. Ye, M. Su, C. Zhang, A. E. Ribbe, W. Jiang and C. H. Mao, Nature, 2008, 452, 198–201.
4 R. Chakrabarty, P. S. Mukherjee and P. J. Stang, Chem. Rev., 2011, 111, 6810–6918.
5 T. R. Cook, Y.-R. Zheng and P. J. Stang, Chem. Rev., 2013, 113, 734–777.
6 A. J. McConnell, C. S. Wood, P. P. Neelakandan and J. R. Nitschke, Chem. Rev., 2015, 115, 7729–7793.
7 G. H. Clever and P. Punt, Acc. Chem. Res., 2017, 50, 2233–2243.
8 Y. Takezawa, J. Müller and M. Shionoya, Chem. Lett., 2017, 46, 622–633.
9 L.-J. Chen and H.-B. Yang, Acc. Chem. Res., 2018, 51, 2699–2710.
10 W. Zheng, G. Yang, N. Shao, L.-J. Chen, B. Ou, S.-T. Jiang, G. Chen and H.-B. Yang, J. Am. Chem. Soc., 2017, 139, 13811–13820.
11 X. Yan, F. Wang, B. Zheng and F. Huang, Chem. Soc. Rev., 2012, 41, 6042–6065.
12 W. Zheng, W. Wang, S.-T. Jiang, G. Yang, Z. Li, X.-Q. Wang, G.-Q. Yin, Y. Zhang, H. Tan, X. Li, H. Ding, G. Chen and H.-B. Yang, J. Am. Chem. Soc., 2019, 141, 583–591.
13 G.-Y. Wu, L.-J. Chen, L. Xu, X.-L. Zhao and H.-B. Yang, Coord. Chem. Rev., 2018, 369, 39–75.
14 B. Zheng, F. Wang, S. Dong and F. Huang, Chem. Soc. Rev., 2012, 41, 1621–1636.
15 W. Tuo, Y. Sun, S. Lu, X. Li, Y. Sun and P. J. Stang, J. Am. Chem. Soc., 2020, 142, 16930–16934.
16 Q. Zhang, D. Tang, J. Zhang, R. Ni, L. Xu, T. He, X. Lin, X. Li, H. Qiu, S. Yin and P. J. Stang, J. Am. Chem. Soc., 2019, 141, 17909–17917.
17 L. Xu, X. Shen, Z. Zhou, T. He, J. Zhang, H. Qiu, M. L. Saha, S. Yin and P. J. Stang, J. Am. Chem. Soc., 2018, 140, 16920–16924.
18 X. Yan, T. R. Cook, J. B. Pollock, P. Wei, Y. Zhang, Y. Yu, F. Huang and P. J. Stang, J. Am. Chem. Soc., 2014, 136, 4460–4463.
19 Z. Chen, M. H.-Y. Chan and V. W.-W. Yam, J. Am. Chem. Soc., 2020, 142, 16471–16478.
20 P. Wei, X. Yan and F. Huang, Chem. Soc. Rev., 2015, 44, 815–832.
21 J.-F. Chen, B.-B. Han, J.-F. Ma, X. Liu, Q.-Y. Yang, Q. Lin, H. Yao, Y.-M. Zhang and T.-B. Wei, RSC Adv., 2017, 7, 47709–47714.
22 M. Liu, L. Zhang and T. Wang, Chem. Rev., 2015, 115, 7304–7397.
23 H. Jiang, Y. Jiang, J. Han, L. Zhang and M. Liu, Angew. Chem., Int. Ed., 2019, 58, 785–790.
24 C.-B. Huang, L.-J. Chen, J. Huang and L. Xu, RSC Adv., 2014, 4, 19538–19549.
25 F. Michael, D.-R. Gilad, B. Yonatan, Y. Ravit, C. Rémi, A. A.-G. Miguel and W. Itamar, Polym. Chem., 2018, 9, 2905–2912.
26 C.-L. Sun, J.-F. Xu, Y.-Z. Chen, L.-Y. Niu, L.-Z. Wu, C.-H. Tung and Q.-Z. Yang, Polym. Chem., 2016, 7, 2057–2061.
27 C. O. Dietrich-Buchecker, J. P. Sauvage and J. P. Kintzinger, Tetrahedron Lett., 1983, 24, 5095–5098.
28 M. C. Jimenez-Molero, C. Dietrich-Buchecker and J.-P. Sauvage, Chem.–Eur. J., 2002, 8, 1456–1466.
29 A. Livoreil, J.-P. Sauvage, N. Armaroli, V. Balzani, L. Flamigni and B. Ventura, J. Am. Chem. Soc., 1997, 119, 12114–12124.
30 J.-P. Collin, C. Dietrich-Buchecker, P. Gaviña, M. C. Jimenez-Molero and J.-P. Sauvage, Acc. Chem. Res., 2001, 34, 477–487.
31 G. Chelucci and R. P. Thummel, Chem. Rev., 2002, 102, 3129–3170.
32 W. W. Brandt, F. P. Dwyer and E. D. Gyarfas, Chem. Rev., 1954, 54, 959–1017.
33 J.-P. Sauvage, Acc. Chem. Res., 1998, 31, 611–619.
34 G.-Y. Wu, X.-Q. Wang, L.-J. Chen, Y.-X. Hu, G.-Q. Yin, L. Xu, B. Jiang and H.-B. Yang, Inorg. Chem., 2018, 57, 15414–15420.
35 I. A. Bhat, D. Samanta and P. S. Mukherjee, J. Am. Chem. Soc., 2015, 137, 9497–9502.
36 B. Mondal and P. S. Mukherjee, J. Am. Chem. Soc., 2018, 140, 12592–12601.
37 P. S. Mukherjee, T. K. Maji, G. Mostafa, W. Hibbs and N. R. Chaudhuri, New J. Chem., 2001, 25, 760–763.
38 G.-Y. Wu, X. Shi, H. Phan, H. Qu, Y.-X. Hu, G.-Q. Yin, X.-L. Zhao, X. Li, L. Xu, Q. Yu and H.-B. Yang, Nat. Commun., 2020, 11, 3178–3189.
39 G.-Y. Wu, B.-B. Shi, Q. Lin, H. Li, Y.-M. Zhang, H. Yao and T.-B. Wei, RSC Adv., 2015, 5, 4958–4963.
40 J. Murray, K. Kim, T. Ogoshi, W. Yao and B. C. Gibb, Chem. Soc. Rev., 2017, 46, 2479–2496.
41 K. Jie, Y. Zhou, E. Li and F. Huang, Acc. Chem. Res., 2018, 51, 2064–2072.
42 T. Ogoshi, T.-a. Yamagishi and Y. Nakamoto, Chem. Rev., 2016, 116, 7937–8002.
43 T. Ogoshi, K. Maruyama, Y. Sakatsune, T. Kakuta, T.-a. Yamagishi, T. Ichikawa and M. Mizuno, J. Am. Chem. Soc., 2019, 141, 785–789.
44 T. Kakuta, T.-a. Yamagishi and T. Ogoshi, Acc. Chem. Res., 2018, 51, 1656–1666.
45 K. Jie, M. Liu, Y. Zhou, M. A. Little, A. Pulido, S. Y. Chong, A. Stephenson, A. R. Hughes, F. Sakakibara, T. Ogoshi, F. Blanc, G. M. Day, F. Huang and A. I. Cooper, J. Am. Chem. Soc., 2018, 140, 6921–6930.
46 T. Ogoshi, S. Takashima and T.-a. Yamagishi, J. Am. Chem. Soc., 2018, 140, 1544–1548.
47 S. Fa, M. Yamamoto, H. Nishihara, R. Sakamoto, K. Kamitani, Y. Nishina and T. Ogoshi, Chem. Sci., 2020, 11, 5866–5873.
48 K. Wada, T. Kakuta, T.-A. Yamagishi and T. Ogoshi, Chem. Commun., 2020, 56, 4344–4347.
49 T. Ogoshi, S. Kanai, S. Fujinami, T. Yamagishi and Y. Nakamoto, J. Am. Chem. Soc., 2008, 130, 5022–5023.
50 T. Kakuta, R. Nakanishi, T. Ogoshi and T.-A. Yamagishi, RSC Adv., 2020, 10, 12695–12698.
51 J. Fan, Y. Chen, D. Cao, Y.-W. Yang, X. Jia and C. Li, RSC Adv., 2014, 4, 4330–4333.
52 Q. Lin, L. Liu, F. Zheng, P.-P. Mao, J. Liu, Y.-M. Zhang, H. Yao and T.-B. Wei, RSC Adv., 2017, 7, 34411–34414.
53 T.-B. Wei, X.-B. Cheng, H. Li, F. Zheng, Q. Lin, H. Yao and Y.-M. Zhang, RSC Adv., 2016, 6, 20987–20993.
54 Z.-Y. Li, Y. Zhang, C.-W. Zhang, L.-J. Chen, C. Wang, H. Tan, Y. Yu, X. Li and H.-B. Yang, J. Am. Chem. Soc., 2014, 136, 8577–8589.