MATERIALS CHEMISTRY

FRONTIERS



View Article Online

REVIEW



Cite this: *Mater. Chem. Front.*, 2020, **4**, 3176

Received 26th July 2020, Accepted 31st August 2020 DOI: 10.1039/d0qm00540a

rsc.li/frontiers-materials

Naphthalene diimide- and perylene diimide-based supramolecular cages

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During the past few years, the construction of naphthalene diimide (NDI)- and perylene diimide (PDI)based cages has attracted increasing attention due to the unique photo- and electrochemical properties and widespread applications of these materials in catalysis, chemical separation, sensing, *etc.* Thus, it is the right time to summarize the recent development of NDI-/PDI-based cages. In this review, the preparations, properties, and applications of NDI-/PDI-based cages in catalysis, chemical separation, and sensing will be discussed.

1. Introduction

Since the discovery of crown ethers and cryptands by Lehn *et al.*, the construction of new supramolecular hosts, including macrocycles and cages, has attracted increasing interest.¹ Compared to two-dimensional macrocycles such as $pillar[n]arene^2$ and metal-coordinated macrocycles,³ which possess an open cavity and allow guests to fill a hole or bind on the surface of the macrocycle, three-dimensional cages possess enclosed cavities in which guests are entirely encapsulated in the confined space in general. Due to the efficient encapsulation of various guests, cages could thus be applied in many fields, such as molecular recognition, catalysis,

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chemical transfer and separation, and stabilization of reactive intermediates.⁴ During the past few decades, a variety of cages with structural complexity, including Platonic and Archimedes polyhedrons, were reported and exhibited unique properties and functions,⁵ which promoted the development of modern supramolecular chemistry.

Rylene diimides (RDIs, n = 0-7) are a series of aromatic diimides; for example, n = 0 or 1, corresponded to naphthalene diimide (NDI) and perylene diimide (PDI), respectively (Scheme 1). NDI and PDI have attracted extensive attention due to their large planar structure, electron-deficient nature, photochemical and electrochemical properties (strong absorption in the visible region, high quantum yield and redox activity), and easy functionalization.⁶ Therefore, in the past few decades, a large number of NDI and PDI derivatives have been prepared with applications in sensing, light harvesting, photocatalysis and supramolecular self-assembly.⁷ For example, some NDI- and PDI-based macrocycles were successfully



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constructed through covalent synthesis or non-covalent selfassembly.8 Since NDI and PDI derivatives exhibit excellent photochemical and electrochemical properties, integrating NDI or PDI subunits into a cage may result in distinctive properties and functions: (1) the electron-deficient nature of NDI/PDI could endow the cage with a strong affinity for electron-rich guests such as C_{60} ; (2) the switchable redox activity of NDI/PDI endows the cage with catalytic performance; and (3) a cage with strong absorption in the visible region may serve as a visible-light photocatalyst. In recent decades, a series of NDI-/PDI-based cages were constructed, and their properties and applications were investigated in detail. It should be noted that among the NDI/PDI cages, metal-coordinated cages are in the majority since coordinationdriven self-assembly has proven to be a powerful tool to construct various supramolecular architectures, including three-dimensional metal-coordinated cages.9 In addition, due to the linear structure of NDI/PDI units, tetrahedral cages were readily prepared through coordination-driven selfassembly.¹⁰

This review will focus on the preparation strategies, photochemical and electrochemical properties of NDI/PDI-based cages as well as their applications. For clarity, this review will be classified into NDI-based cages and PDI-based cages.

2. Naphthalene diimide-based cages

Naphthalene diimide (NDI) derivatives are planar and electrondeficient as well as π -acidic and can be used in anion– π catalysis.¹¹ Typically, NDI derivatives could undergo two reversible one-electron reduction under chemical or electrochemical conditions with half-wave potentials of -1.10 V and -1.51 V ν s. Fc/Fc⁺ in CH₂Cl₂.^{6a} Based on the outstanding features of NDI units and the confined space of cages, NDI-based cages show unique physical and chemical properties and can be applied in various fields, such as catalysis, sensing, and separation of anions.

Nitschke and co-workers have made significant contributions in the field of NDI-based cages. In 2013, Nitschke, Schalley and Sanders et al. reported the synthesis of a tetrahedral NDI cage and investigated its mechanically interlocked behaviour with bis-1,5-(dinaphtho)-[38]crown-10.¹² As shown in Fig. 1, by treating NDI-containing diamine 1, 2-formylpyridine and $Fe(NTf_2)_2$ in a ratio of 6:12:4 in acetonitrile at 50 °C for 24 h, an M₄L₆-type NDI cage 2 was obtained through subcomponent self-assembly. The NDI cage 2 was well characterized by ¹H NMR, ESI-MS, COSY, and NOESY NMR spectroscopy, and the broadening and overlap of the signals in the ¹H NMR spectra indicated the existence of various diastereomers with T, S_4 , and C_3 symmetry of the cage in solution. Due to their planar structure and electron-deficient properties, NDI derivatives readily form donor-acceptor complexes with aromatic species. Thus, NDI cage 2 was further treated with bis-1,5-(dinaphtho)-[38]crown-10 (10 equiv.) 3, and three-dimensional polycatenanes were formed. The ¹H NMR spectrum of the polycatenanes showed two sets of signals, in which one signal corresponded to the crown etherbound NDI cage (an upfield shift in ¹H NMR relative to the initial NDI cage 2) and the other signal corresponded to the free NDI cage. The above result also suggested the slow exchange process between the bound and free crown ether on the NMR timescale. In addition, the mixtures of polycatenanes cannot be



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Fig. 1 (a) Synthesis of tetrahedral cage 2. (b) Graphical representation of the polycatenanes formed upon the addition of excess crown ether 3 into cage 2.

distinguished in NMR spectra due to the overlapping signals. Furthermore, treatment of NDI cage 2 with a large excess of 3 (38 equiv.) provided saturated [7]catenane. DOSY spectra of the polycatenanes also showed two groups of resonances corresponding to the crown ether-bound NDI cages and the free crown ether, consistent with the ¹H NMR spectra. An infrared multiphoton dissociation (IRMPD) tandem MS experiment was further performed to confirm the specific binding between the NDI cages and crown ethers, and the results indicated that $[M_4L_6C_7]^{8+}$ (C = catenated crown ether 3) showed non-specific binding between NDI cage 2 and crown ether 3, while $[M_4L_6C_6]^{8+}$ ions maintained their tetrahedral structure with six catenated crown ethers. Furthermore, the binding isotherm revealed that the binding of the six crown ethers to the NDI cage was not cooperative due to the relatively large space for threading, and the binding constant was determined to be 794 \pm 34 L mol⁻¹. In conclusion, this work successfully constructed three-dimensional polycatenanes through three reversible interactions (metal-ligand coordination, imine bond, and donor-acceptor interactions) and studied the dynamic systems in detail, laying the foundation for the construction of new molecular machines with various applications.

The exploration of synthetic chemical systems that respond to different stimuli can offer insight into the signalling events and new means to synthesize useful stimuli-responsive systems and materials. Nitschke, Schalley and Sanders et al. reported NDI- and porphyrin-based mixed-ligand tetrahedral cages and studied their stimulus-responsive behaviour upon catenation and guest encapsulation.¹³ As shown in Fig. 2a, two different cages, 5 and 7, based on NDI and porphyrin, respectively, were readily obtained through subcomponent self-assembly. The crystal structures of 5 and 7 confirmed their tetrahedral structure, and the NDI ligand and the porphyrin ligand showed almost the same length (18.4 Å), suggesting the possibility of ligand exchange upon mixing of the two cages. In addition, interactions between the NDI and porphyrin cages with crown ether 3 and C₇₀ were investigated (Fig. 2b). The results indicated that at most two equivalents of crown ethers 3 could thread onto the NDI moiety in cage 5, which was less than its

analogue 2 (at most 6 crown ethers), probably due to the steric strain between the adjacent NDI moieties resulting from the smaller space in cage 5. In addition, 5 showed no host-guest interaction with C₇₀ due to the existence of large pores in its external surface, whereas porphyrin cage 7 could form a hostguest complex with C₇₀. Furthermore, in order to investigate the structural reconstitutions through catenation and guest encapsulation, NDI-based cage 5 and porphyrin-containing cage 7 were mixed in a 1:1 ratio at 25 °C for 24 h and a dynamic combinatorial library (DCL) of mixed-ligand tetrahedral cages was obtained (Fig. 2c). Because the species in the DCL had similar ESI-response factors, ESI-MS could be used to evaluate the content of each species. The results indicated that upon simple mixing, various species were observed, and species 4_46_2 and 4_16_5 (the subscripts indicate the number of ligands in one cage) were in the majority, which deviated from the principle of Pascal's triangle, revealing that minor differences in the ligand will largely affect the final results of self-assembly and that the ligands tend to form homoleptic assemblies through self-sorting. In addition, upon addition of C70 into the DCL, the final composition was almost complete for 5 and $C_{70} \subset 7$ species because the host-guest adduct $C_{70} \subset 7$ was thermodynamically stable. Upon the addition of crown ether 3 into the DCL, 4_46_2 and 4_16_5 appeared with the highest abundance to meet the requirement of the maximum degree of catenation, thus maximizing the number of π -interactions between NDI and naphthalene. This work constructed an elaborate dynamic system that showed a response to catenation and host-guest encapsulation, which was helpful for understanding the stimulus-responsive behaviours of biological systems.

In 2015, Nitschke and co-workers reported the construction of an NDI-based heteroleptic metal–organic cage through the donor–acceptor interaction of pyrene and NDI.¹⁴ As shown in Fig. 3, reaction of the pyrene-containing diamine 8 or 11, 2-formylpyridine, and metal ions (Fe²⁺, Zn²⁺ or Co²⁺) in a ratio of 6:12:4 afforded two $M_4^{II}L_6$ type cages, 9 and 12, respectively. X-ray diffraction analysis indicated that cage 9 showed a C_2 -symmetric rhomboidal structure with *mer* metal coordination



Fig. 2 (a) Synthesis of tetrahedral cages 5 and 7 and their crystal structures. (b) Host-guest behaviour of cages 5 and 7 upon the addition of 3 and C70, respectively. (c) Schematic representation of the different templation effects observed in the 1:1 DCL formed upon mixing of 5 and 7. Reproduced from ref. 13 with permission from Royal Society of Chemistry, copyright 2016

and that cage 12 showed a C_3 -symmetric pseudotetrahedron with fac metal coordination. These structures were obviously different from the common tetrahedral $M_4^{II}L_6$ cages, which was ascribed to the π -stacking of pyrene subunits. In addition, the reaction of pyrene-containing diamine 11, 2-formylpyridine and metal ions $(Zn^{2+} \text{ or } Co^{2+})$ in a ratio of 6:12:6 provided a box-like structure 13 that could encapsulate electron-deficient guests such as NDI, which inspired the authors to construct heteroleptic cages through donor-acceptor interactions. By mixing box 13 with additional NDI-based diamine 4 or the reaction of pyrenecontaining diamine 11, NDI-based diamine 4, 2-formylpyridine and metal ions (Fe^{2+} , Zn^{2+} or Co^{2+}) in a ratio of 4:2:12:4 afforded heteroleptic cage 14. X-ray diffraction analysis indicated that cage 14 adopted an elongated sandwich-like structure due to the internal pyrene-pyrene-NDI stacks. This donor-donor-acceptor arrangement was complementary to the common donor-acceptor interactions. Further self-sorting experiments were performed to

investigate the universality of the construction of heteroleptic cages. As shown in Fig. 4, due to the thermodynamic stability of 14, it was readily obtained via integrative self-sorting. However, a mixture of 8 and 4, 2-formylpyridine and $Zn(NTf_2)_2$ in CD_3CN resulted in homoleptic cages Zn-15 and Zn-10, which revealed that the process was narcissistic self-sorting. In addition, by treatment of 16 (or 17) and 4 under the same conditions, a mixture of seven assemblies was observed, and a dynamic combinatorial library (DCL) was formed (similar to the results by mixing cages 5 and 7 without templation effects), which was ascribed to the absence of π -stacking between pyrene and NDI units. In conclusion, an NDIbased heteroleptic cage was successfully constructed through donor-acceptor interactions, which provided a new method to construct metal-organic cages with more complexity.

Supramolecular catalysts, such as metal coordinated cages, have been widely applied in molecular synthesis.¹⁵ In 2019, Nitschke et al. reported the synthesis of an NDI-based tetrahedral



Fig. 3 Synthesis of cages 9, 10, 12, 13, and 14 and crystal structure of cage Zn-14. Reproduced from ref. 14 with permission from American Chemical Society, copyright 2015.

cage with redox-switchable behaviour and could serve as a catalyst for C-C bond coupling.¹⁶ As shown in Fig. 5, treating NDIcontaining diamine 18 and 2-formylpyridine with $Zn(NTf_2)_2$ in acetonitrile afforded NDI cage 19 with a yield of 77%. NDI cage 19 was well characterized by NMR and ESI-MS spectra. Furthermore, cyclic voltammetry (CV) experiments were conducted to investigate the redox properties of 19, and the results indicated that 19 showed two reduction peaks at -1.1 V and -1.52 V vs. Fc/Fc⁺, corresponding to the NDI moieties. In addition, upon the addition of cobaltocene (Cp_2Co) and tetrachloro-1,4-benzoquinone, the cage could be reversibly reduced and re-oxidized without disturbing the cage's framework. ¹H NMR spectra indicated that the cage could form a host-guest complex with C₆₀ derivative 20. Moreover, tetrakis-(4-chlorophenyl)borate 21 was used in place of Cp₂Co as the reductant, and the ¹H NMR and EPR spectra revealed the generation of radical species. Interestingly, the formation of 4,4'-dichlorobiphenyl was also observed during the reduction, which inspired the authors to investigate the cage as a catalyst for the C-C bond coupling of organoborates. Further experiments indicated that the host-guest complex 20 < 19 exhibited a two-fold increase in yield during the catalytic process relative to cage **19** alone, and the control compound, such as the NDI ligand, showed no catalysis. In addition, complex **20** \subset **19** could catalyse the coupling of organoborates with both electron-withdrawing and electron-donating groups, which was complementary to common coupling catalysis. EPR experiments were performed to investigate the catalytic mechanism, and the results indicated that C₆₀ derivative **20** could serve as a radical-stabilizing agent to stabilize the radical species of the cage. Thus, the catalytic performance occurred on the surface of the cage, and C₆₀ acted as a co-catalyst to enhance the catalytic activity for the oxidative coupling of organoborates, which afforded a new view into supramolecular catalysis.

Molecular sensing plays an important role in supramolecular chemistry. In 2020, Valiyaveettil *et al.* reported the construction of a NDI-based water-soluble macrocycle which contains quaternary ammonium groups.^{8*i*} The macrocycle exhibited high affinity for aromatic carboxylate anions such as perylene tetracarboxylate tetrapotassium salt ($K_{sv} = 1.6 \times 10^5 \text{ M}^{-1}$) and biphenyl-4,4'-dicarboxylic acid dipotassium salt ($K_{sv} = 4.3 \times 10^4 \text{ M}^{-1}$), due to the electrostatic interactions between cationic macrocycle and anionic guests, size matching effect and the interaction between electron-deficient NDI units and aromatic perylene.



Fig. 4 (a-c) Results of subcomponent self-assembly with mixtures of various diamines.

Considering the redox-active properties of NDI, at the same vear, Nitschke and co-workers reported the synthesis of a NDIbased cage with redox-responsive guest uptake and release behaviour.¹⁷ Notably, the addition of excess reductant Cp₂Co into cage 19 led to the formation of precipitation resulting from charge neutralization (the negatively charged NDI panels were compensable to the cationic charges of Zn^{2+} vertices), which hampered the further study in solution. Thus, quaternary ammonium-functionalized NDI ligand 23 in place of ligand 18 was prepared to construct the new NDI cage 24 (Fig. 6). X-ray crystallographic analysis indicated that the cage adopted the typical tetrahedral structure, and three carborate anions were encapsulated in the cage due to the multiple positive charges of 24. CV experiments indicated that cage 24 showed two reduction peaks at -0.81 V and -1.19 V vs. Fc/Fc⁺ and two oxidation peaks at -1.29 V and -0.78 V. It should be noted that the reduction peaks were reversible, while they were irreversible for the NDI ligand 23, indicating the self-assembled cages enhanced the robustness of NDI units under redox process. Thus, the cage could be reversibly reduced and re-oxidized upon the addition of Cp2Co and AgNTf2. In addition, ¹H NMR experiments demonstrated that cage 24 could bind only anion species such as BF₄⁻, PhBF₃⁻ and CB₁₁H₁₂⁻ instead of neutral species due to its multiple positive charges. Furthermore, the redox-responsive guest uptake and release behaviour of the cage was investigated, and the results indicated that the guest PhBF₃⁻ was ejected from the cage upon reduction with Cp₂Co, while with the subsequent addition of AgNTf₂, the guest could be re-taken up (Fig. 6b). It should be noted that cage 24 exhibited a suitable volume for binding with fullerenes, which



Fig. 5 (a) Synthesis of NDI-based tetrahedral cage **19**. (b) Host-guest complexation of cage **19** with **20** and the subsequent reduction with Cp₂Co. (c) Transferred tetrakis(4-chlorophenyl)borate **21** into 4,4'-dichlorobiphenyl **22** with the catalysis of complex **20** \subset **19**.

was similar to the analogous polyaromatic cages.¹ⁿ Thus, interestingly, the reduced form of cage 24 showed obvious affinity for C_{60} , probably due to the partially neutralized charges upon reduction (Fig. 6c). Further addition of AgNTf₂ into the complex led to the release of C_{60} , which required more than 30 days due to the kinetically trapped nature of the complex. Moreover, when the $C_{60} \subset 24$ complex was treated in MeCN/EtOAc (1:1) with aqueous Na₂SO₄, cage 24 was transferred into the water phase, and C_{60} was released, which could be applied in chemical purification due to its simplicity and recyclability. In conclusion, this research realized redox-switchable guest uptake and release behaviour within an NDI cage, which could be helpful for designing supramolecular systems for chemical separations and transport.

In 2018, Chand and co-workers reported the preparation of an NDI-based Pd₂L₄-type metal–organic cage, which can form metallogels with multi-stimulus-responsive properties and anionic dye removal ability.¹⁸ As shown in Fig. 7, NDI-containing



Fig. 6 (a) Synthesis of NDI-based tetrahedral cage **24** and its crystal structure. (b) Redox control of KBF₃Ph binding within **24**. (c) The encapsulation and release of C_{60} within cage **24** upon reduction with Cp₂Co and subsequent re-oxidation with AgNTf₂. Reproduced from ref. 17 with permission from Royal Society of Chemistry, copyright 2020.

ligand 25 reacted with Pd(NO₃)₂ in DMSO or MeCN/H₂O to afford cylindrical-drum-shaped NDI cage 26. Interestingly, adding cage 26 in DMSO or MeCN/H₂O led to the formation of gels, and the concentration, solvent, counteranion and metal ions had a considerable impact on the formation of gels. UV-Vis spectral studies showed a hyperchromic shift with increasing concentrations, probably due to the aggregation of cage 26 in solution. Rheology studies confirmed the formation of gels and revealed that the strength of the gel increased with increasing temperature or concentration. In addition, the gel also exhibited multistimulus-responsive properties including a thixotropic nature and chemical-stimulus-responsive behaviour. For example, upon vigorous shaking or addition of TBABr or DMAP, the metallogel changed into a solution state and could be recovered upon subsequent standing or addition of AgNO₃ or TsOH, indicating that the stimulus-responsive property was reversible (Fig. 7b). Moreover, UV-Vis spectral experiments indicated that the gel could uptake pyrene due to the electron-deficient nature of the NDI ligand and the suitable cavity of the cage.



Fig. 7 (a) Synthesis of NDI-based cage **26** and its optimized structure. (b) Stimulus-responsive behaviour of the metallogel upon shaking/resting and the addition of TBABr/AgNO₃ or DMAP/TsOH pairs. (c) Removal of anion dyes AB93 and MO with the metallogel. Reproduced from ref. 18 with permission from American Chemical Society, copyright 2018.

Furthermore, as shown in Fig. 7c, the gel could be used for dye removal, and the results showed that the gel could efficiently remove anionic dyes instead of cationic dyes due to the positive charge character of the NDI cage.

3. Perylene diimide-based cages

Relative to the NDI derivatives, perylene diimide (PDI) derivatives have a larger planar structure and π -conjugation and intense absorption in the visible region. In general, PDI derivatives exhibit two reversible one-electron reduction with half-wave potentials of -0.96 V and -1.22 V vs. Fc/Fc⁺ in CH₂Cl₂.^{6d} Compared with a large number of reported PDI-based macrocycles,⁸ PDI-based cages have rarely been investigated. Herein, we introduce some PDI-based cages, including metalcoordinated and covalent cages, and the photochemical and electrochemical properties of the cages as well as their applications in molecular sensing and photocatalysis.

Würthner and co-workers reported several PDI-based metalorganic cages. In 2013, they reported the preparation of giant tetrahedral PDI cages and investigated their photo- and electrochemical properties and host–guest behaviour.¹⁹ As shown in Fig. 8, treatment of the bipyridine-functionalized PDI ligand **27** with FeX₂ (X = BF⁴⁻, OTf⁻) in a solution of CH₃CN/CHCl₃ (1:1) provided tetrahedral cages **28–29**. The cages were well characterized by ¹H NMR, 2D DOSY and ESI-MS spectra. Furthermore, UV-vis spectroscopy and cyclic voltammetry (CV) were performed to investigate the photo- and electrochemical properties of the cages. UV-Vis spectral studies indicated that cages **28–29** exhibited two main absorption bands at 542 and 580 nm in the visible region (corresponding to the PDI subunits) with molar



Fig. 8 (a) Synthesis of PDI-based tetrahedral cages 28-29 and graphical representation of the complexation of 28-29 with C₆₀. (b) UV-Vis absorption spectrum of 27-29. (c) Optimized structure of C₆₀ \subset 28 or 29 and $2C_{60} \subset$ 28 or 29. Reproduced from ref. 19 with permission from American Chemical Society, copyright 2013.

absorption coefficients of 212700 and 268500 M⁻¹ cm⁻¹, respectively, almost six times as high as the PDI ligand 27. In addition, the peak of the bipyridine units at 297 nm showed a redshift of approximately 20 nm, and the relative intensities of the peaks at 542 and 580 nm also varied due to the metalto-ligand charge-transfer of $[Fe(bipyridine)_3]^{2+}$ complexes (Fig. 8b). CV experiments indicated that the cages showed two reversible oxidation waves at 0.95 (PBI/PBI⁺) and 0.73 V (Fe^{2+}/Fe^{3+}) vs. Fc/Fc⁺ and five reversible reduction waves at -0.94 (PBI/PBI⁻), -1.11 (PBI⁻/PBI²⁻), -1.65, -1.84 and $-2.08 \text{ V} (\text{bpy/bpy}^-) vs. \text{ Fc/Fc}^+$. The higher oxidation potential of Fe²⁺/Fe³⁺ was probably due to the Fe²⁺ metal centre being close to the electron-deficient PDI moieties, and the easier reduction of cage 28 could be ascribed to the preorganized framework and highly porous nature of the PDI cage. In addition, the counteranions $(BF^{4-} \text{ or } OTf^{-})$ had a negligible influence on the photo- and electrochemical properties of the cages. Moreover, the results of ¹³C NMR and ESI-TOF-MS spectra indicated that the cage could efficiently encapsulate C₆₀, and the UV-Vis absorption spectra suggested that each cage bound 1.9 C_{60} on average. Considering the volume of C_{60} (597 Å³) and the PDI cage (950–2150 Å³ due to the rotatable OPh^tBu substituents), the host-guest behaviour of the cage is consistent with the principle of the empirical 55% rule established by Mecozzi and Rebek.²⁰ Additionally, the results of molecular stimulations indicated that when the cage bound one C_{60} , C_{60} was located in almost the centre of the cavity, and there was no interaction between the cage and C_{60} (Fig. 8c). However, when the cage bound two C_{60} , the guests were located in the corner of the cage to afford enough space for binding both guests. Notably, **28** or **29** displayed the ability of binding two C_{60} in one cage, which was different from the common polyaromatic cages¹ⁿ that could only bind one C_{60} , probably due to its larger internal cavity. This research successfully constructed PDI cages with affinity for C_{60} , which afforded a guideline for designing functional cages for guest encapsulation and reaction within the cavity.

Even though PDI cages **28–29** showed efficient bonding with C_{60} , the cages were non-emissive because the coordinated Fe metal atom facilitated intersystem crossing (ISC) and subsequent non-radiative decay, which hampered their fluorescence sensing.²¹ Hence, Würthner and co-workers reported the synthesis of a highly fluorescent tetrahedral PDI cage and studied its host–guest behaviour in 2015.²² As shown in Fig. 9, the reaction of PDI-containing ligand **30**, $Zn(NTf_2)_2$ and tris-(2-aminoethyl)amine (TREN) under microwave heating at 100 °C provided tetrahedral PDI cage **31** in 92% yield. It should be noted that Zn was vital for the formation of the fluorescent cage. Cage **31**



Fig. 9 (a) Synthesis of PDI-based tetrahedral cage **31**. (b) Optimized structure of cage **31**. (c) Absorption and emission spectra of **30** and **31** in CH₃CN (λ_{ex} = 490 nm). (d) Optimized structure of the host-guest complex $P_3 \subset 31$ (P = perylene). (e) Optimized structure of the host-guest complex $C_2 \subset 31$ (C = coronene). (f) Absorption spectrum of **31**, C and $C_2 \subset 31$. (g) Emission spectra upon the addition of C into **31**. Reproduced from ref. 22 with permission from Wiley-VCH, copyright 2015.

was well characterized by ¹H NMR, 2D NMR and ESI-MS spectra. Variable-temperature ¹H NMR experiments showed a diastereomeric ratio of 1:0.9:0.7 for the *T*, S_4 , and C_3 diastereomers of the cage. 2D DOSY NMR indicated that the hydrodynamic radius of cage **31** was 2.2 nm, in good agreement with the result from theoretical simulations (1.9 nm). In addition, the results of the MMFF geometry optimization model indicated that the length of the edge of cage **31** was 3.8 nm and the internal volume was approximately 1300 Å³ (Fig. 9b). Further investigations on the photophysical properties revealed that **31** showed two main absorption bands at 488 nm and 523 nm corresponding to the absorption of PDI units and two emission peaks at 538 and 578 nm with a quantum yield of 0.67 ± 0.02 (Fig. 9c). The decrease in the quantum yield relative to that of the PDI ligand (almost unity) was probably due to excitonic coupling between the adjacent PDI chromophore. The fluorescence lifetime of cage **31** was also calculated and determined to be 4.8 ns. Furthermore, the hostguest behaviour of the cage was studied, and ¹H NMR titrations indicated that cage **31** could form a 1:3 complex with perylene or a 1:2 complex with coronene (Fig. 9d and e). Furthermore, the hostguest complexation induced a decrease in the absorption and emission intensity of **31**, probably due to the photoinduced charge transfer from the PDI units to the guests (Fig. 9f and g). In addition, the phenomenon of amplified emission quenching



Fig. 10 Synthesis of PDI-based tetrahedral cage 32 and its concentration-dependent supramolecular transformation.

(one guest caused the quenching of 2.1 PBI units) could be ascribed to the close contact of the cage and guests. In conclusion, this work realized fluorescence sensing based on a highly emissive PDI cage, which laid the foundation for constructing fluorescent self-assembly hosts for molecular recognition.

Similarly, in 2015, Würthner and co-workers reported the synthesis of a large PDI-based tetrahedron and investigated its supramolecular transformation at various concentrations.²³ As shown in Fig. 10, PDI cage **32** was obtained *via* subcomponent self-assembly of PDI-containing ligand **30**, *p*-anisidine with Zn(NTf₂)₂ under microwave conditions at 100 °C. The NMR, ESI-MS and atomic force microscopy (AFM) results confirmed the formation of the M₄L₆-type tetrahedral cage. However, the concentration had a great influence on the structure of the self-assembly product. ¹H NMR and UV/Vis spectral studies indicated that the initial M₄L₆ tetrahedron disassembles into an M₂L₃-type helicate with some fragments when the concentration was below 8.7 × 10⁻⁶ mol L⁻¹. In addition, similar to **31**, the cage also showed fluorescence upon Zn coordination.

In 2019, Zhang et al. reported the preparation of chiral imine cages based on PDI and investigated their host-guest and photocatalytic behaviour.²⁴ As shown in Fig. 11, treatment of PDI-containing diamine 34 (R or S) and tetraphenylethene (TPE)-containing tetraaldehyde 35 provided PDI-TPE cage 36 (*R* or *S*). The cages were well characterized by ¹H NMR, 2D NMR and ESI-MS spectra. Circular dichroism (CD) spectra of the two cages displayed signals with complete mirror symmetry, demonstrating that the cages were enantiomeric. UV-Vis absorption of cage R-36 exhibited a maximum absorption peak at 526 nm, consistent with the PDI ligand, revealing no interaction between the PDI units in the cage. The fluorescence spectra of cage R-36 indicated that J-type aggregation occurred at higher concentrations resulting from charge transfer between TPE and PDI units. In addition, CV experiments of the cage showed two reduction peaks at -0.84 V and -1.05 V (vs. SCE), which were assigned to the formation of PDI*⁻ and PDI²⁻, respectively. Furthermore, the host-guest behaviour of cages with polycyclic aromatic hydrocarbons (PAHs), such as perylene and naphthalene, was investigated. The fluorescence of the guests declined dramatically upon the addition of cage **R-36**, probably due to the aromatic π - π stacking interaction between the planar PAHs and PDI subunits (Fig. 11c). Fluorescence titration experiments indicated that the cage could bind 18-22 small PAH molecules, such as naphthalene and pyrene, or 8-10 larger PAH molecules, such as perylene and 9,10-diphenylanthracene. In addition, the fluorescence of the cage itself was enhanced upon guest binding because a large number of guests occupied the space of the cage and decreased the structural relaxation of the PDI units. Since cage R-36 had strong absorption in the visible range and could efficiently encapsulate a series of guest molecules, it could further serve as a visible-light photocatalyst. The results indicated that cage R-36 could efficiently transform 2-aryloxybenzoic acids 37 into aryl salicylates 38 (Smiles rearrangement) under a blue LED (Fig. 11d). The higher catalytic efficiency of the cage relative to



Fig. 11 (a) Synthesis of PDI-based imine cage **36** from PDI-containing diamine **34** and TPE-containing tetraaldehyde **35**. (b) The host–guest behaviour of cage **R-36** with perylene. (c) Emission spectrum of perylene following the titration of **R-36**. (d) Smiles rearrangement under the photolysis of cage **R-36**. Reproduced from ref. 24 with permission from Wiley-VCH, copyright 2019.

the PDI ligand was ascribed to intermolecular photoinduced electron transfer from TPE to PDI units, indicating the importance of the existence of the imine cage. Moreover, different substituents on the substrate had an important effect on the catalytic activity, and an electron-donating group in the *ortho* position in 37 could improve the yield. In conclusion, this research constructed a PDI-based imine cage for visible-light photocatalysis, affording a guideline for designing photoactive supramolecular hosts for photochemical reactions.

4. Conclusions

NDI-/PDI-based cages, including metal coordination and dynamic covalent cages, which possess different properties and functions, have been investigated during the past few years. First, due to the planar structure and electron-deficient nature of NDI/PDI, the cages could bind various electron-rich guests, which were further applied in molecular sensing and constructing a dynamic combinatorial library (DCL). Second, the cages with redox-switchable properties resulting from NDI/PDI units showed unique functions, including controllable uptake and release and catalysis for C–C bond coupling. In addition, the NDI/PDI cages could also be applied in dye removal or photocatalysis, indicating their large potential. Even so, there are still some issues that need to be addressed in this field. For example, the types of catalytic reaction of the NDI/PDI cages are very limited. On the other hand, due to the utilization of Fe²⁺ as vertices for constructing metal coordination cages, most of the NDI/PDI cages were non-fluorescent even though NDI/PDI ligands were highly emissive, which hampered their development of fluorescent materials.

Thus, in our view, some improvement should be taken into consideration in further developing NDI/PDI cages. On one hand, NDI/PDI ligands should be designed reasonably. According to some unpublished results in our group, we could make use of NDI/PDI ligands with larger π -conjugation to construct cages that bind more sophisticated guests, thus further extending their catalytic reactions. On the other hand, more kinds of metals should be utilized to construct cages with various shapes and topological architectures. This may not only prevent the fluorescence quenching of NDI/PDI ligands to some extent but also extend their applications, such as photocatalysis. In conclusion, NDI-/PDI-based cages will be considered to play more important roles in a wide range of fields, including supramolecular chemistry and materials science, in the future.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Nature Science Foundation of China (No. 21922506, 21871092, and 21672070), the Fundamental Research Funds for the Central Universities, the Opening Projects of Shanghai Key Laboratory of Green Chemistry and Chemical Processes, and State Key Laboratory of Fine Chemicals (KF 1801).

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Review

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