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New aspects in bis and tris(dipyrrinato)metal complexes: bright luminescence, self-assembled nanoarchitectures, and materials applications

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Dipyrrins serve as monovalent bidentate ligand molecules that coordinate to various cations. Their BF_2 complexes, 4,4-difluoro-4-bora-3a,4a-diaza-s-indacene and its derivatives (BODIPYs), exhibit excellent photostability, strong light absorption, and high fluorescence quantum yield, thereby encouraging their application in various fields, e.g., as biological and biomedical fluorescent markers. Dipyrrin may also accept a wide variety of metal ions spontaneously. However, dipyrrin metal complexes have been disregarded from materials science research. This review article summarizes recent progress in bis(dipyrrinato)metal(II) and tris(dipyrrinato)metal(III) complexes from the viewpoint of materials chemistry. Section 2 describes a series of efforts aimed to realize intense luminescence superior to or comparable with that of BODIPYs. The spontaneous coordination of these complexes enables them to construct self-assembled nanoarchitectures, such as supramolecules and coordination polymers that form one-dimensional nanowires, two-dimensional nanosheets, and metal-organic frameworks. Section 3 describes such alluring molecular superstructures. Section 4 discusses potential applications based on these nanoarchitectures, such as thermoelectric and photoelectric conversion.

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1. Introduction

Dipyrrin comprises two pyrroles bridged by a methine bridge, where the two rings are inequivalent to pyrrolic and iminic

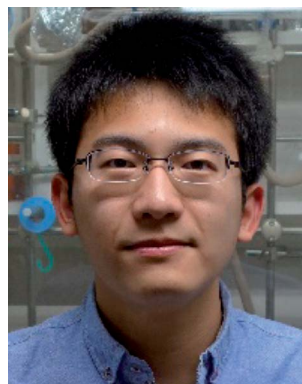
nitrogens (Fig. 1a). Upon deprotonation from the pyrrolic nitrogen, dipyrrin undergoes resonance stabilization, such that the two rings are equivalent to each other (Fig. 1b). In this scheme, dipyrrin serves as a monovalent bidentate ligand molecule that coordinates to various cations to form complexes.^{1–15} Among these complexes, the BF_2 complexes of dipyrrin, 4,4-difluoro-4-bora-3a,4a-diaza-s-indacene (BODIPY,

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Tokyo, joining Prof. Nishihara's group again. His current research interest lies in the construction of molecule-based nanostructures, and photonic and electronic devices using thereof.



Toshiki Iwashima was born in Inazawa City, Aichi, Japan in 1993. He graduated from The University of Tokyo (Japan) in 2015. He is now pursuing his MSc degree with Prof. Hiroshi Nishihara in the Department of Chemistry at The University of Tokyo. His research interest lies in the synthetic design of organometallic complexes, and photonic and electronic devices using them. He is also fascinated by football and a big fan of Arsenal Football Club.



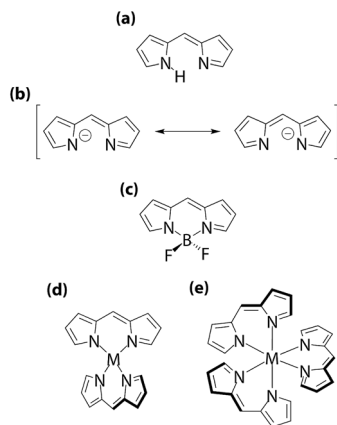


Fig. 1 Dipyrryn family: (a) dipyrryn; (b) dipyrrynate; (c) BODIPY; (d) bis(dipyrrynato)metal(II) complex; (e) tris(dipyrrynato)metal(III) complex.

Fig. 1c) and its derivatives are the most famous and well-investigated ones because of their excellent photostability, strong light absorption in the visible and near infrared regions, and high fluorescence quantum yield (ϕ_F).^{5–16} In fact, BODIPYs are already on the market as biological and biomedical fluorescent markers and are being applied in devices such as solar cells and chemical sensors. However, dipyrryn can also accept a wide variety of metal ions, including nickel(II), copper(II), zinc(II), iron(III), cobalt(III), and rhodium(III), to form bis(dipyrrynato)metal(II) and tris(dipyrrynato)metal(III) complexes (Fig. 1d and e).^{1–4} Since the discovery of bis(dipyrrynato)metal complexes in 1924,¹⁷ dipyrryn metal complexes have long been regarded as trivial substances in materials science.

This review article summarizes recent progress in the development of bis(dipyrrynato)metal(II) and tris(dipyrrynato)metal(III) complexes from the viewpoint of materials chemistry. Section 2 describes a series of efforts aimed to realize intense

luminescence superior to or comparable with that of BODIPYs. Section 3 discusses the self-assembly of bis and tris(dipyrrynato) metal complexes, which results in the formation of alluring nanoarchitectures, such as supramolecules and coordination polymers giving rise to one-dimensional nanowires, two-dimensional nanosheets, and metal-organic frameworks (MOFs) and porous coordination polymers (PCPs). Section 4 describes potential applications based on these nanoarchitectures.

2. Pursuit of bright luminescence

A plain BODIPY shows intense absorption and bright fluorescence at approximately 500 nm, which are derived from the $^1\pi-\pi^*$ transition of the dipyrrynato ligand.^{5–16} The absorption and fluorescence may be redshifted upon the introduction of substituents on the dipyrrynato ligand, covering the 500–900 nm region. BODIPYs are good fluorophores even in polar solvents such as water. In sharp contrast, dipyrryn metal complexes have long been believed to be non-luminescent or weakly luminescent. This drawback seriously reduces the value of dipyrrynato-metal complexes in applications in which they serve, for example, as photosensitizers. In this section, the authors concentrate on efforts to improve the luminescent ability of bis and tris(dipyrrynato)metal complexes. For comprehensive knowledge on the whole types of luminescent dipyrrynato-metal complexes (*i.e.* mono(dipyrrynato)metal complexes with ancillary ligands other than dipyrryns), please see a comprehensive review article contributed by Baudron.²

2.1 Introduction of bulky aryl groups at the *meso*-position

The first essential improvement made to the fluorescence of dipyrryn-metal complexes was addressed by Lindsey, Bocian, and Holten, who focused on the steric effect of the peripheral



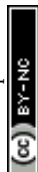
Mizuho Tsuchiya was born in Ueda City, Nagano, Japan in 1989. He received his BSc degree in 2012 from The University of Tokyo and received his MSc in 2014 from the same university under the supervision of Prof. Hiroshi Nishihara. He is now pursuing his PhD degree on the development of molecular photonic devices using fluorescent molecules in the same group. He was selected as a

Research Fellow for Young Scientists of the JSPS in 2014. His research interests include inorganic chemistry, photochemistry, electrochemistry, surface chemistry, and molecular devices.



Hiroshi Nishihara received his BSc degree in 1977, MSc in 1979 and DSc in 1982 from The University of Tokyo. He was appointed research associate of the Department of Chemistry at Keio University in 1982, and he was promoted as lecturer in 1990, and associate professor in 1992. Since 1996, he has been a professor of the Department of Chemistry, School of Science at The University of Tokyo. He also

worked as a visiting research associate of the Department of Chemistry at The University of North Carolina at Chapel Hill (1987–1989), and as a researcher of PRESTO, JST (1992–1996). He received The Chemical Society of Japan Award for Creative Work in 2003, Docteur Honoris Causa from the University of Bordeaux in 2011, and Commendation for Science and Technology by the MEXT in 2014.



aryl group.¹⁸ The researchers demonstrated that the replacement of the phenyl group of bis(dipyrrinato)zinc(II) complex **1** with a 2,6-dimethylphenyl group (*i.e.*, **2**) transformed the molecule from a very weak emitter into highly fluorescent chromophore (Fig. 2). In toluene at room temperature, **1** and **2** exhibit absorption and fluorescence spectra similar to each other (Fig. 3). The spectral features of the compounds are associated with the $^1\pi-\pi^*$ transition of the dipyrrinato ligand, similar to those of BODIPYs. Still, a subtle difference may be found with respect to the compounds' absorption and emission bandwidths, and Stokes shift is slightly reduced in **2**. This series of changes reflects lower conformational freedom in **2**, which is induced by the steric constraint between the 2,6-dimethylphenyl group and dipyrrinato ligand. This steric effect is clearly manifested in the fluorescence quantum yield ($\phi_F = 0.006$ for **1**, 0.36 for **2**) and fluorescence lifetime ($\tau = 90 \pm 10$ ps for **1**, 3 ± 1 ns for **2**) (Table 1). However, the incorporation of a *t*-butyl group into **1** (*i.e.*, complex **3**), which has no associated steric interference, produces a ϕ_F of 0.007; therefore, no enhancement in ϕ_F is observed. This series of experimental findings suggests that hindering the internal rotation of the aryl group at the *meso*-position has a distinctive effect on the fluorescence quantum yield. Similar behavior has been reported for complexes **4** and **5**, which have peripheral 1- and 2-naphthyl groups at the *meso*-position, respectively (Fig. 2 and Table 1).¹⁹ Possessing a greater steric constraint, **4** has shown photoluminescence intensities that are approximately three and ten times greater than those of **5** in THF and polymer media, respectively.

2.2 Heteroleptic bis and tris(dipyrrinato)metal complexes

Although introducing bulky aryl groups at the *meso*-position leads to fluorescence enhancement, quenching in polar solvents remains a serious problem. Sakamoto and Nishihara proposed that thermal transition/equilibrium from/between the emissive $^1\pi-\pi^*$ excited state (D^*-Zn-D or $D-Zn-D^*$) to/and non-emissive charge-separated states ($D'^+-Zn-D'^-$ or $D'^--Zn-D'^+$) would suppress the fluorescence of a bis(dipyrrinato)zinc(II) complex, where **D** and **Zn** denote a dipyrrinato ligand and Zn^{2+} ion, respectively (Fig. 4a).^{20,21} The complex may be regarded as a

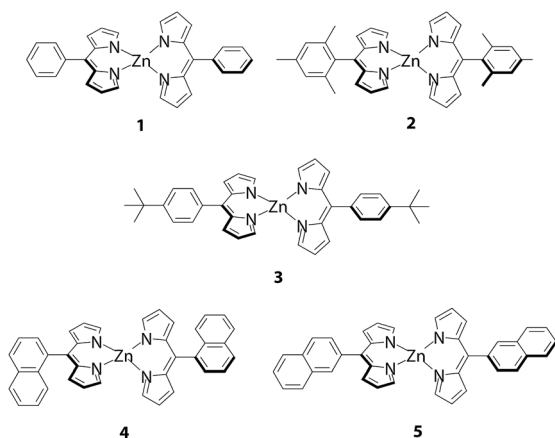


Fig. 2 Luminescent bis(dipyrrinato)zinc(II) complexes **1**–**5**.

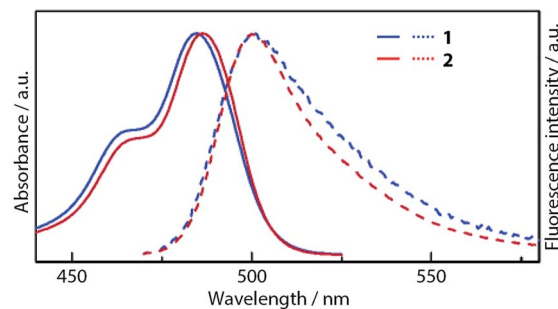


Fig. 3 Absorption and fluorescence spectra of **1** and **2**. Adapted with permission from ref. 18. Copyright 2004 American Chemical Society.

bichromophoric molecule, in which two dye entities (dipyrrinato ligands) are ligated by a zinc(II) ion. Photoexcitation in bichromophoric systems often results in the formation of charge-separated states, which are generated by one-electron transfer.^{22–24} 9,9'-Bianthryl is a representative example of such a system.^{22,23} The charge-separated state is known to be stabilized in polar solvents.

To verify the present hypothesis, and to improve ϕ_F , Sakamoto and Nishihara synthesized a series of zinc(II) complexes **6**–**8** (Fig. 5).²⁰ A small number of heteroleptic bis(dipyrrinato) metal complexes have been reported by Cohen,²⁵ and Radecka,^{26,27} and Dolphin,²⁸ however, **8** is still the first example of a heteroleptic zinc(II) complex that contains two types of dipyrrinato ligands with drastic differences in their electronic structures: one derived from either of the homoleptic complexes **6** and **7**. Fig. 6 shows the absorption and fluorescence spectra of **6**–**8**, and their numerical data are collected in Table 1. Complex **7**, with (4-methylphenyl)ethynyl substituents at the β -position, shows bathochromic shifts in its absorption and fluorescence maxima because of π -extension. Heteroleptic complex **8** features two absorption bands, corresponding to those of **6** and **7** with no obvious shift in wavelength, thereby suggesting no distinctive orbital interaction between the two dipyrrinato ligands both in the ground and excited states. However, **8** exhibits remarkable emission behavior; it exhibits very high ϕ_F in toluene ($\phi_F = 0.76$), and the bright fluorescence is sustained even in polar dichloromethane ($\phi_F = 0.52$). This series of luminescence behaviors contrasts with the behaviors observed for **6** and **7**, which undergo significant quenching from non-polar toluene to polar dichloromethane ($\phi_F = 0.28 \rightarrow 0.00$ for **6**, $\phi_F = 0.72 \rightarrow 0.27$ for **7**). DFT calculation indicates that the configuration of the frontier orbitals (π and π^* orbitals of each dipyrrinato ligand) in heteroleptic complex **8** thwarts thermal transition from the emissive $^1\pi-\pi^*$ state to the non-fluorescent charge-separated states, thereby leading to a greater ϕ_F (Fig. 4b). This positive effect is highlighted in polar solvents. It is also noteworthy for heteroleptic complex **8** that excitation at each ligand results in exclusive emission from the π -extended ligand at the same ϕ_F (Fig. 6 and Table 1). This result demonstrates the presence of quantitative energy transfer between the two dipyrrinato ligands.

Following the publication of the report by Sakamoto and Nishihara, experimental proof was provided by Thompson and

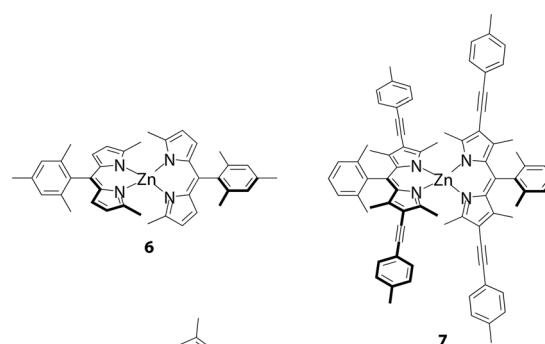
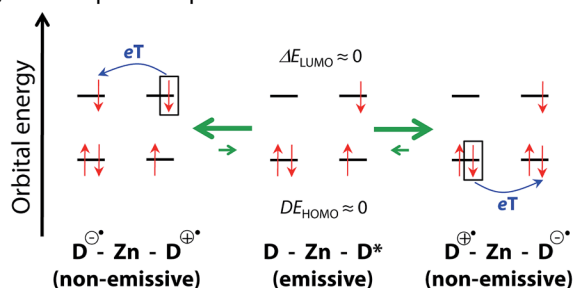


Table 1 Photochemical properties of complexes 1–13 at room temperature in fluid solutions

Compound	Central metal	Solvent	λ_{abs}^a (nm)	λ_{ex} (nm)	λ_{em} (nm)	ϕ_F	τ (ns)	Ref.
1	Zn(II)	Toluene	485	480	501	0.006	0.09 ± 0.01	18
2	Zn(II)	Toluene	487	480	500	0.36	3 ± 1	18
3	Zn(II)	Toluene	—	—	—	0.007	—	18
4	Zn(II)	THF	485	—	516	—	—	19
5	Zn(II)	THF	486	—	518	—	—	19
6	Zn(II)	Toluene	495	495	509	0.28 (0.00 ^b)	—	20
7	Zn(II)	Toluene	553	553	579	0.72 (0.27 ^b)	—	20
8	Zn(II)	Toluene	495, 553	495	578	0.76 (0.52 ^b)	—	20
				553	578	0.75 (0.49 ^b)	—	20
9	In(III)	Hexanes	444, 496	448	522	0.074	1.93 ± 0.02	25
10	In(III)	Toluene	459, 570	459	592	0.41 (0.053 ^b)	2.44	26
11	In(III)	Toluene	478, 564	480	596	0.34 (0.021 ^b)	2.53	26
12	In(III)	Toluene	528, 572	530	600	0.28 (0.015 ^b)	2.49	26
13	Zn(II)	Toluene	485, 646	490, 600	672	0.00043	—	30

^a $^1\pi-\pi^*$ bands are shown. ^b In dichloromethane.

(a) Homoleptic complexes 6 and 7



(b) Heteroleptic complex 8

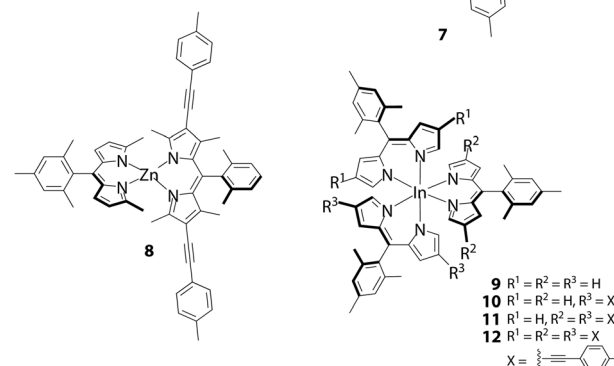
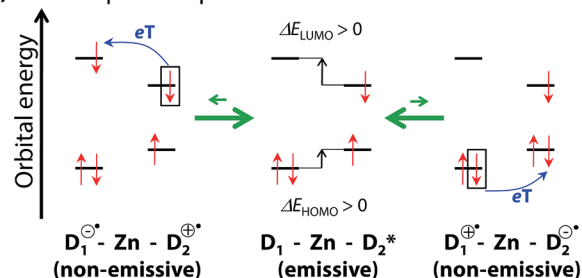


Fig. 4 Schematic illustrations for plausible thermal equilibria among the two non-emissive charge-separated states ($D^{\bullet+}-Zn-D^{\bullet-}$ and $D^{\bullet-}-Zn-D^{\bullet+}$ for homoleptic complexes, $D_1^{\bullet+}-Zn-D_2^{\bullet-}$ and $D_1^{\bullet-}-Zn-D_2^{\bullet+}$ for heteroleptic complexes) and the emissive $^1\pi-\pi^*$ excited state localized on the right-hand dipyrrinato ligand ($D-Zn-D^*$ for homoleptic complexes, $D_1-Zn-D_2^*$ for heteroleptic complexes) in a polar solvent. eT indicates an electron transfer process. Zn and D represent the Zn^{2+} center and dipyrrinato ligand, respectively. Adapted with permission from ref. 20. Copyright 2012 John Wiley & Sons, Inc.

coworkers that indicated that the charge-separated state is the chief non-radiative pathway in homoleptic bis(dipyrrinato)zinc(II) complexes.²⁹ The authors performed transient absorption spectroscopy, thereby disclosing that $^1\pi-\pi^*$ photoexcited homoleptic complexes undergo symmetry-breaking charge transfer to produce the charge-separated state in polar solvents. They also elucidated that in non-polar solvents such as toluene,

Fig. 5 Luminescent heteroleptic bis(dipyrrinato)zinc(II) complex 8, tris(dipyrrinato)indium(III) complexes 10 and 11, and dipyrinato-aza-dipyrinato hybrid complex 13. Corresponding homoleptic complexes 6, 7, 9 and 12 are also shown.

thermal equilibrium occurs between the $^1\pi-\pi^*$ state and the charge-separated state.

To demonstrate the generality and utility of their strategy, Sakamoto and Nishihara pursued extensions to other dipyrrinato-metal complex systems. Cohen and co-workers reported



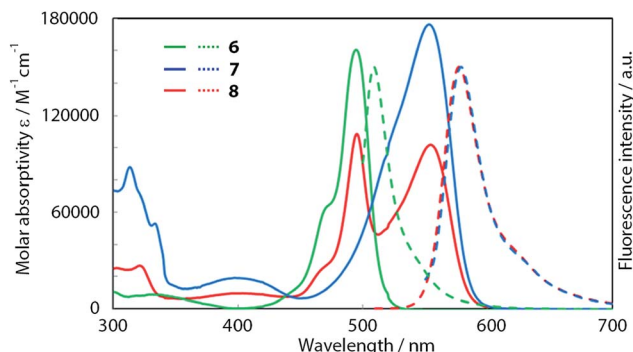


Fig. 6 Absorption (solid lines) and fluorescence spectra (dotted lines) of **6** (green), **7** (blue), and **8** (red) in toluene. Adapted with permission from ref. 20. Copyright 2012 John Wiley & Sons, Inc.

that homoleptic tris(dipyrrinato)gallium(III) and indium(III) complexes were weak emitters;³⁰ however, heteroleptic indium(III) complexes **10** and **11** (Fig. 5) exhibit higher ϕ_F (0.41 and 0.34 in toluene, respectively) than the corresponding homoleptic complexes **9** and **12** ($\phi_F = 0.074$ and 0.28 in hexanes and toluene, respectively) (Table 1).³¹ Surprisingly, the ϕ_F of **9** is greater than that of the corresponding π -expanded BODIPY ($\phi_F = 0.35$). This finding is of great interest because it counters the preconception that BODIPYs emit more brightly than dipyrrinato-metal complexes.

Bis(azadipyrrinato)zinc(II) complexes were believed to be non-fluorescent,^{32–34} in contrast to the behavior observed for BF₂ complexes of azadipyrrins (azaBODIPY). However, Sakamoto and Nishihara revealed that dipyrrinato-azadipyrrinato hybrid zinc(II) complex **13** exhibited detectable fluorescence (Fig. 5 and Table 1).³⁵

2.3 Other molecules

Telfer and coworkers reported that homoleptic tris(dipyrrinato)-rhodium(III) complexes emitted weakly.³⁶ Several supramolecular and coordination polymeric systems show luminescence, as will be discussed in Section 3. Mono(dipyrrinato)metal complexes containing Sn(II),³⁷ Cu(II),^{38,39} Re(II),⁴⁰ Pt(II),⁴¹ Pd(II),⁴¹ Ir(III),^{42–44} Rh(III),⁴⁵ Sn(IV),⁴⁶ Al(III),⁴⁷ Si(IV),⁴⁸ Ca(II),⁴⁹ Zn(II),⁴⁹ and Mg(II)^{50,51} centers also exhibited luminescence, the details of which are discussed in a comprehensive review.²

3. Creation of self-assembled nanoarchitectures

A distinctive advantage of (dipyrrinato)metal complexes over BODIPYs is that they may construct supramolecular and polymer assemblies by taking advantage of facile and spontaneous coordination reactions and multiple coordination sites on the metal centers. In the following section, supramolecules, one-dimensional and two-dimensional coordination polymeric systems, and MOFs and PCPs based on (dipyrrinato)metal complexes are introduced.

3.1 Supramolecules

The development of bis and tris(dipyrrinato)metal complexes is not confined to simple mononuclear ones as described in the previous section. Because many metal ions may accept two or more dipyrrin ligands, supramolecular assemblies linked by bis and tris(dipyrrinato)metal complexes may be designed and obtained. Indeed, the appropriate design of dipyrrin ligands affords supramolecules with intriguing structures.

Numerous multinuclear bis and tris(dipyrrinato)metal complexes featuring oligo(dipyrrin)s, or ligands with multiple dipyrrin subunits joined by spacers, have been developed to date.^{52–72} The length, position, and flexibility of these spacers affect the geometry of resultant metal complexes; their geometries include double-stranded helical,^{52–61,65–72} triple-stranded helical,⁶² triangular,^{61,64} rhombic,^{63,64} and hexagonal.⁶³

Especially, a number of reports have described the formation of double-stranded helices. These complexes can be obtained by bis(dipyrrin) ligands with spacers exhibiting modest flexibility, such as short alkyl chains,^{52–55,57,60} sulfur,⁵⁶ and 4-alkoxyphenylene spacers.^{58,59} Focusing on the chirality of double-stranded complexes, the Thompson group prepared enantiomeric bis(dipyrrin)s **14a,b** to conduct diastereoselective complexation reactions to obtain double-stranded helical bis(dipyrrinato)zinc(II) complexes **15a,b** (Fig. 7).⁵⁷ Circular Dichroism (CD) spectroscopy, NMR spectroscopy, and chiral HPLC analysis revealed that ligands **14a,b** underwent complexation with nearly a 1 : 1 mixture of diastereomeric helicates, and **15a,b** did not show stereochemical inversion in solution.

Maeda and co-workers developed *meso*-aryl dipyrrin dimers whose dipyrrin moieties are linked together at the 2 and 2'-positions to construct double-stranded helicates with zinc(II) ions (Fig. 8).⁵⁸ Although complexes without straps to interlock the aryl group **16a–d** are known to undergo racemization in solution, those with straps, *e.g.*, **17a,b**, retain their chirality. Variable-temperature CD spectroscopy and fluorescence lifetime measurements have revealed that the interlocked species exhibit a temperature-dependent spring-like motion, where the distance between the zinc centers fluctuates. The same group has developed a class of similar complexes and demonstrated that there are two conformations in the complexes, S (short, contracted) and L (long, stretched) (Fig. 9).⁵⁹ Although the latter is favored theoretically, X-ray crystallography and variable temperature absorption spectroscopy revealed that the two modes are interchangeable in **18**.

Although double-stranded complexes are not reported to be intensely emissive in general,^{56,58,59} Antina, V'yugin, and co-workers reported that several complexes exhibited bright emission.^{4,65–71} For example, zinc(II) complex **19** tethered with a methylene linker at the 3 and 3' positions fluoresces at 543 nm with a ϕ_F of 0.91 in cyclohexane (Fig. 10).⁶⁶ It is notable that even heavier d¹⁰ metal complexes show fluorescence; a Cd(II) analogue emits at 537 nm with a ϕ_F of 0.27 in cyclohexane,⁶⁹ whereas a Hg(II) counterpart luminesces at 538 nm with a ϕ_F of 0.034.⁶⁸



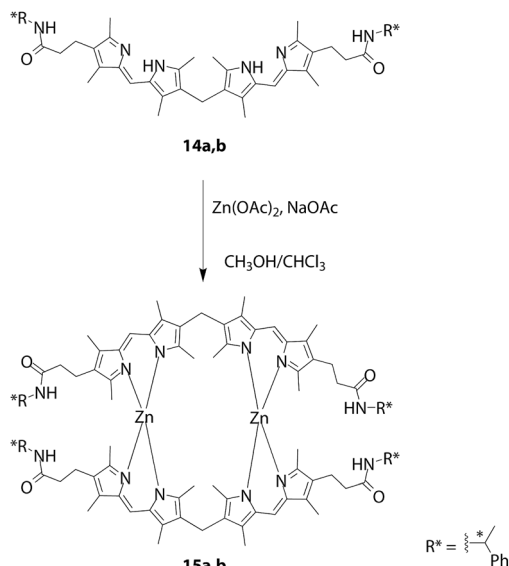


Fig. 7 Double-stranded helical zinc(II) complexes **15a,b** comprising bis(dipyrrin)s **14a,b** with an accessory chiral amide.

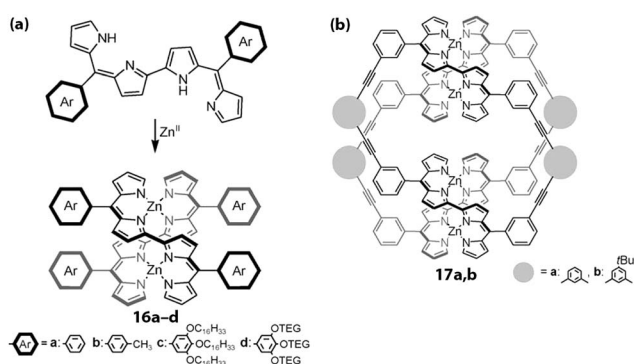


Fig. 8 Double-stranded helical complexes. (a) Double helical complexes **16a-d** without straps on the *meso*-aryl group. OTEG = a triethyleneglycol chain. (b) Double helical complexes **17a,b** with linking straps on the *meso*-aryl group. Adapted with permission from ref. 58. Copyright 2010 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

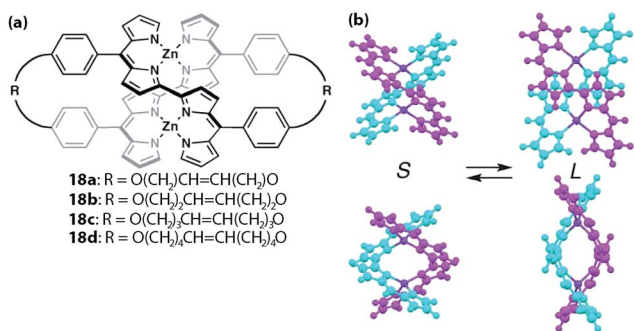
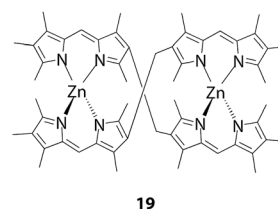


Fig. 9 (a) Double-stranded helical complex **18**. (b) Two possible conformations. Adapted with permission from ref. 59. Copyright 2013 The Royal Society of Chemistry.



As previously mentioned, supramolecular assemblies with geometries other than double-stranded helicates have also been reported. Dolphin's group synthesized ring-fused bis(dipyrrin)s **20–22** to prepare rhombic (**23–25**) and hexagonal (**26**) multi-nuclear zinc(II) complexes (Fig. 11).⁶³ The distance between the two adjacent zinc(II) centers is 8.2 Å for both complexes, but the diagonal distances are 9.19 Å and 13.80 Å for the rhombic and hexagonal complexes, respectively. However, Maeda and co-workers reported dinuclear and trinuclear bis(dipyrrinato)zinc(II) and nickel(II) complexes (e.g., **27** and **28**, Fig. 12) with dipyrrin dimers tethered at an acute angle (Fig. 12).⁶⁴ The presence of α -methyl groups alters the selectivity of the dinuclear and trinuclear configurations because of their steric repulsion around the metal center. The packing structure of **28** provided a smooth hole conduction pathway, up to a mobility of $0.11 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ (from a non-contact flash-photolysis time-resolved microwave conductivity measurement), along the aligned axis of the trinuclear assembly.

3.2 One-dimensional coordination polymers

Maeda and coworkers synthesized one-dimensional polymers **29** or **30** (Fig. 13a and b).^{73,74} By mixing the corresponding dipyrrin dimer and zinc(II) acetate in a molar ratio of 1 : 1 in THF, the desired one-dimensional coordination polymer featuring the bis(dipyrrinato)zinc(II) motif is generated spontaneously. The polymer strands are dispersible in the medium, showing uniform nanosized spherical structures with diameters measured to be approximately 0.3 μm by scanning electron microscopy (SEM) images (Fig. 13c and d), and approximately 0.1 μm by dynamic light scattering (DLS). The nanospheres contain the constitutive zinc element as shown in Fig. 13c. Additionally, the microspheres emit green fluorescence (Fig. 13d). The authors ascribed the formation of the nanospheres to three steps: (1) formation of the coordination oligomers; (2) stacking of the oligomers; and (3) conversion to spheres.

Dolphin employed a linear and rigid bridging dipyrrin dimer to create bis(dipyrrinato)cobalt(II) complex chains up to pentanuclear ones.²⁸ Single strands of bis(dipyrrinato)zinc(II) coordination polymers were synthesized and isolated by Sakamoto and Nishihara (Fig. 14).⁷⁵ This series of materials is easy to fabricate: by simply mixing two solutions, a dichloromethane solution of **31** or **32** and an ethanol solution of metal(II) acetate (metal = Zn, Ni, and Cu), coordination polymers **33–36** are obtained (Fig. 14a and b). A distinctive feature of the molecular system is that it may be crystallized: a liquid/liquid interfacial

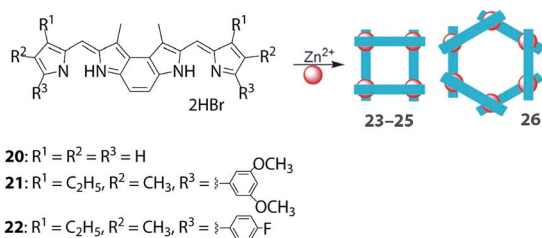


Fig. 11 Rhombic (23–25) and hexagonal (26) multinuclear bis(dipyrrinato)zinc(II) complexes using ring-fused bis(dipyrrin)s 20–22. Adapted with permission from ref. 63. Copyright 2011 The Royal Society of Chemistry.

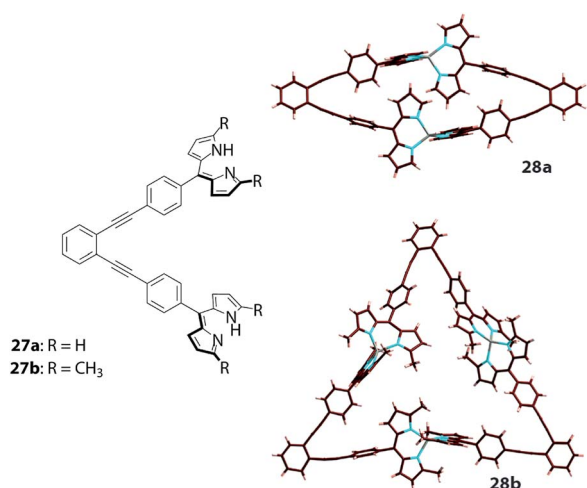


Fig. 12 Bis(dipyrrin)s 27a and 27b and corresponding multinuclear bis(dipyrrinato)zinc(II) complexes with triangular and linear structures (28a and 28b). Adapted with permission from ref. 64. Copyright 2013 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.

reaction between ligand 31 and metal(II) acetate affords single crystals of 33–35 suitable for X-ray diffraction analysis (Fig. 14c–f). In addition, the crystalline or solidified wires may be disentangled into single strands upon ultrasonication in organic solvents. A suspension of 33 showed Tyndall scattering (Fig. 14g), whereas luminescence dominated Tyndall scattering in a suspension of 36, with a quantum yield of 8% in toluene (Fig. 14h). The isolated fibers of 33 and 36 were imaged by atomic force microscopy (AFM), with the suspension sampled on flat HOPG substrates (Fig. 14i–l). Interestingly, the strands were observed as straight lines, the heights of which (0.7 and 1.9 nm) were consistent with the corresponding mononuclear complexes (0.65 and 1.8 nm).

Cohen and coworkers reported one-dimensional coordination polymers based on mono(dipyrrinato)copper(II) complexes with the acetylacetonato type of ancillary ligands.^{76–78} At the *meso*-position, additional coordination sites were implanted, such as pyridyl, 4-cyanophenyl, and 4-methylthiophenyl groups, which allowed the molecular system to adopt one-dimensional polymeric forms in the crystal phase.

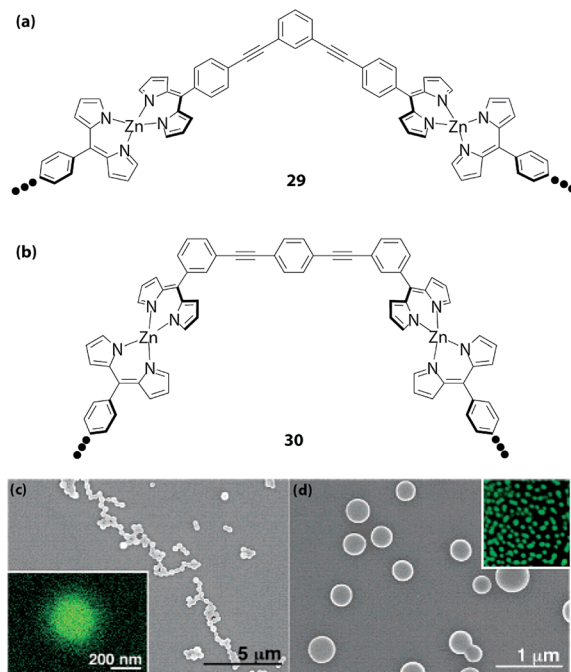


Fig. 13 One-dimensional bis(dipyrrinato)zinc(II) complex polymers (a) 29 and (b) 30. (c) SEM image of one-dimensional coordination polymer 29. The inset shows Zn mapping drawn by HRTEM/EDX. (d) SEM image of one-dimensional coordination polymer 30. The inset shows a fluorescence micrograph with a 25 μm side of each square. Adapted with permission from ref. 73. Copyright 2006 American Chemical Society.

3.3 Two-dimensional coordination polymer

Sakamoto and Nishihara created the two-dimensional polymer 37 comprising the bis(dipyrrinato)zinc(II) complex motif (Fig. 15).⁷⁹ Nanosheets 37 were composed of the three-way dipyrrin ligand 38 and zinc(II) ions and could be effectively synthesized by interfacial fabrication processes. Multi-layer 37 was synthesized by a liquid/liquid interfacial reaction, in which a dichloromethane solution of 38 was layered with an aqueous solution of zinc(II) acetate under ambient conditions (Fig. 15b and c). The spontaneous complexation between ligand 38 and zinc(II) ions then occurred at the liquid/liquid interface, such that the desired two-dimensional network structure was formed. The multi-layer nanosheets exhibited a sheet morphology both macroscopically and microscopically (Fig. 15d–g). The thickness of multi-layer 37 may be controlled by the concentration of 38, spanning from 6 to 800 nm (corresponding to 5–670 layers). The authors also indicated that a series of conventional single-phase syntheses resulted in solid materials with disordered structures. However, an air/liquid interfacial reaction gives rise to single-layer or few-layer 37 (Fig. 15h and i). In this process, a diluted dichloromethane solution of ligand 38 was gently sprayed under ambient conditions. After spontaneous evaporation of dichloromethane, spontaneous nanosheet formation occurred at the air/liquid interface. The nanosheet could be salvaged from the interface and deposited onto various flat substrates. Fig. 15j–n show a



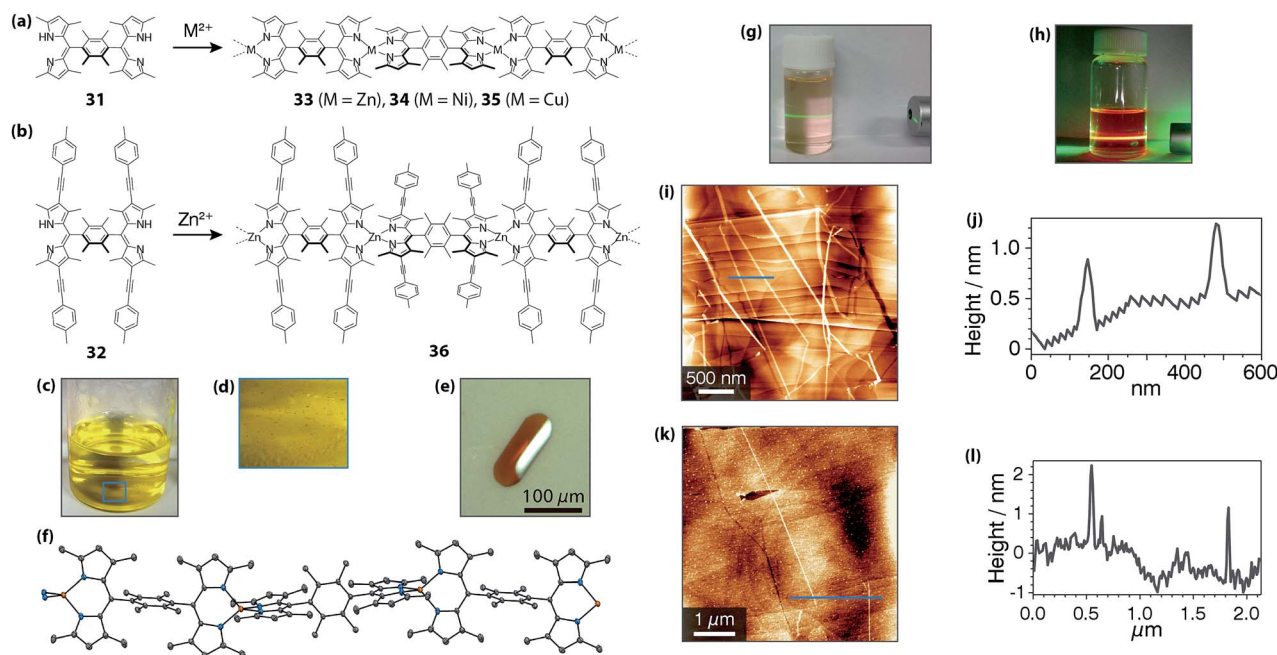


Fig. 14 (a and b) Bridging dipyrin ligands **31** and **32**, and corresponding bis(dipyrinato)metal(II) complexes **33–36**. (c) Photograph of the interfacial reaction system after the emergence of single crystals of **33**. (d) Close-up image at the liquid/liquid interface shown as a blue square in (c). (e) Representative single crystal of **33**. (f) ORTEP drawing of **33** with a thermal ellipsoid set at the 50% probability level (C, gray; N, blue; Zn, orange). (g) Tyndall scattering from a dichloromethane colloidal suspension of **33**. (h) Dichloromethane dispersion of **36** irradiated with a green laser. Tyndall scattering was concealed by orange fluorescence from **36**. (i and j) AFM height image and cross-section analysis along the blue line for **33**. (k and l) AFM height image and cross-section analysis along the blue line for **36**. Adapted with permission from ref. 75. Copyright 2015 The Royal Chemical Society.

series of AFM images of single-layer **37** transferred onto a hydrophobized silicon(111) substrate. The domain size exceeds 10 μm on one side, which is very large for molecule-based nanosheet materials. An interesting investigation was conducted using the AFM tip-scratch method (Fig. 15l–n). Upon making a scratch, only “soft” nanosheet **37** was removed, leaving the substrate surface intact. A cross-section analysis traversing one of the scratched edges revealed a step size of 1.2 nm, which is consistent with the size of the bis(dipyrinato)-zinc(II) complex motif. Therefore, the AFM tip-scratch experiment proved the single-layer nature of nanosheet **37**.

3.4 Metal-organic frameworks and porous coordination polymers

A number of coordination polymers have been developed and investigated within the context MOFs and PCPs. These polymers are expected to be promising and versatile functional materials because of their diverse structures and tunability derived from their organic ligands and metals. Tens of MOFs have been constructed using dipyrin ligands and metal ions toward the realization of photofunctional MOFs.

Baudron and Hosseini developed a series of crystalline coordination polymers.^{80–87} They utilized heterometallic bis(dipyrinato)metal complex networks using dipyrin ligands with pyridine or benzonitrile at the *meso*-position. These peripheral N-bearing moieties may be utilized as additional coordination sites for silver(I)^{80–82} and cadmium(II)^{83,84} centers to form

heterometallic coordination polymers. Moreover, silver(I) ions may experience silver(I)– π interactions, which also contribute to network formation. Because the metal centers of bis(dipyrinato)copper(II), nickel(II), and cadmium(II) complexes still have room for extra coordination, they can accept the peripheral pyridine to form polymeric structures.^{80,82,85–87} A good example is a series of luminescent MOFs comprising bis(dipyrinato)cadmium(II) complexes (**39** in Fig. 16).⁸⁶

They also revealed that heterometallic MOFs with defined structures can be obtained by a similar one-pot method. They reacted a 4-pyridine or 4-(1*H*-imidazol-1-yl)benzene-appended dipyrin ligand (**40a,b**), a cadmium(II) salt, and either of a zinc(II), copper(II), or palladium(II) salt to afford heterometallic MOFs (Fig. 17). In the MOFs, the pyridine moieties were bound to cadmium(II) centers, whereas the dipyrin moieties bound to zinc(II), copper(II), or palladium(II) centers, reflecting the difference of the affinity of the metals and the coordination sites. The MOFs synthesized by the one-pot method were identical to those prepared by a sequential method, which involves the synthesis of a bis(dipyrinato)metal complex, followed by a reaction with a cadmium(II) salt (Fig. 17).⁸⁴

Cohen also developed several series of MOFs,^{88–94} in which nitrogen-bearing heterocycles,^{88–90,94} nitriles,⁹¹ carboxylates^{92,93} or thiols⁷⁶ were introduced at the *meso*-position of dipyrin ligands. One series involved three-fold symmetric tris(dipyrinato)iron(III) and cobalt(III) complexes with heterocycles, namely, pyridyl, quinolyl, (4-pyridylethynyl)phenyl groups (*e.g.*



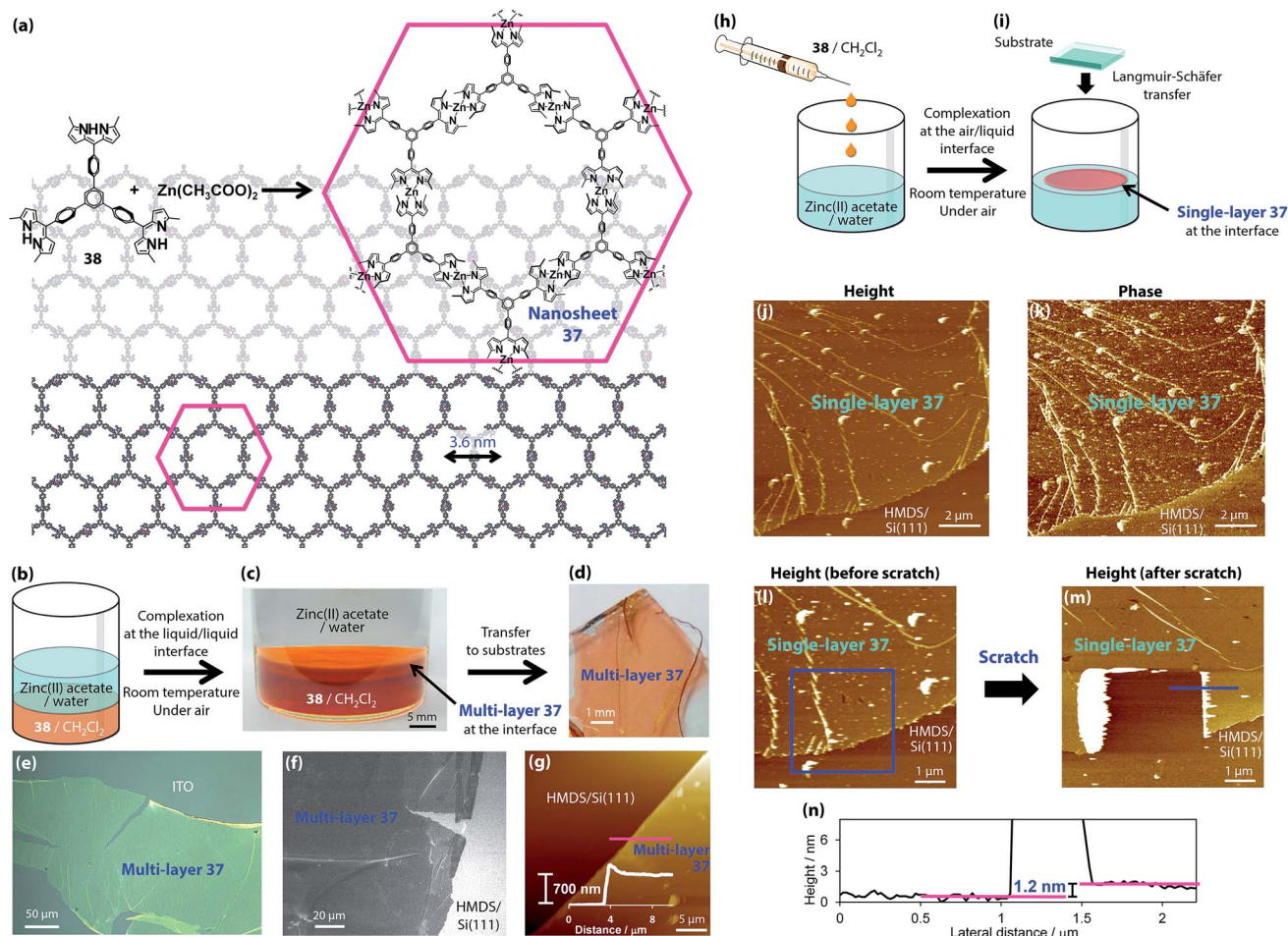


Fig. 15 Chemical structures of three-way dipyrin ligand molecule **38** and bis(dipyrinato)zinc(II) complex nanosheet **37**. (b) Schematic illustration of the liquid/liquid interfacial synthesis. (c) Photograph of the liquid/liquid interfacial synthesis holding multi-layer **37** at the interface. (d) Multi-layer **37** transferred onto an ITO substrate. (e) Optical microscopic image of an ITO substrate. (f) Field-emission scanning electron microscopic (FE-SEM) image of a 1,1,1,3,3,3-hexamethyldisilazane-modified silicon(111) [HMDs/Si(111)] substrate. (g) Atomic force microscopic image of HMDs/Si(111) and its cross-section analysis along the magenta line. (h) Schematic illustration of the air/liquid interfacial synthesis. (i) Schematic illustration of single-layer **37** at the air/liquid interface. (j and k) Height and phase images. (l and m) Height images before and after a scratch by the AFM tip. The blue square indicates the scratched region. (n) Cross-section analysis at one of the steps in the scratched region (shown as a blue line in m). Adapted with permission from ref. 79. Copyright 2015 Macmillan Publishers Limited.

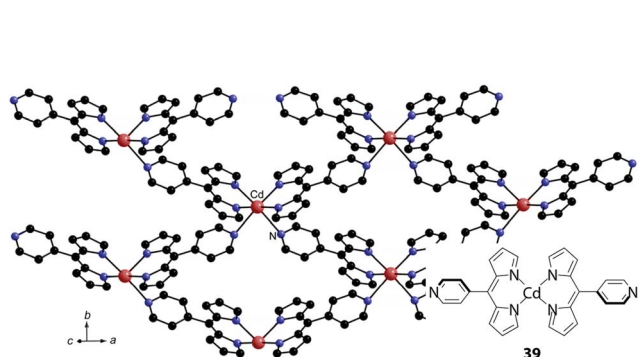


Fig. 16 Luminescent coordination polymer based on pyridine-appended bis(dipyrinato)cadmium(II) complex **39**. Adapted with permission from ref. 86. Copyright 2013 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim Chem.

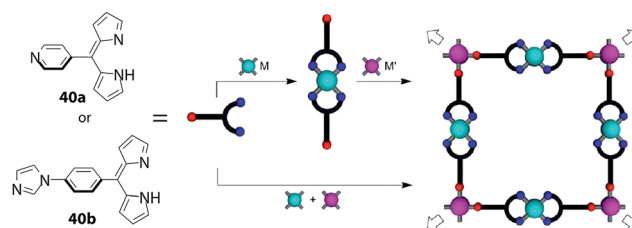


Fig. 17 2D grid heterometallic metal-organic frameworks based on heterocycle-bearing dipyrins (**40a** and **40b**). $M = \text{Cd(II)}$, $M' = \text{Zn(II)}$, Cu(II) , or Pd(II) . Adapted with permission from ref. 84. Copyright 2013 American Chemical Society.

41 in Fig. 18).⁸⁸ They referred to these ligands as “metal-ligands.” A systematic investigation of their assemblies induced by silver(I) ions was conducted, along with a structural analysis of the resultant MOFs (Fig. 18). The counter anion for



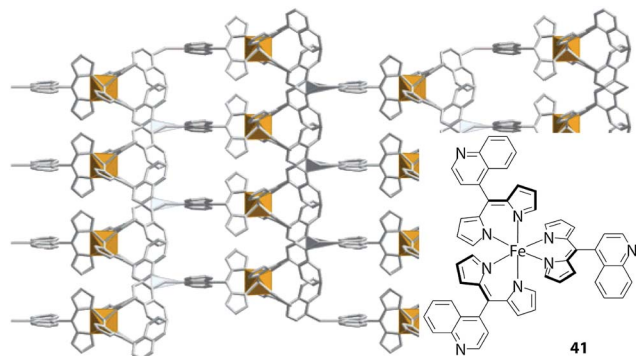


Fig. 18 Metal-organic framework comprising tris(dipyrrinato)iron(III) complex **41** with 4-quinolyl groups at the *meso*-position. Adapted with permission from ref. 88. Copyright 2006 American Chemical Society.

the silver(I) ions was observed to affect the structure and topology of the MOFs. The MOFs also underwent peculiar anion exchange reactions, preserving their structures.

4. Materials applications

In the previous section, a series of elaborate supramolecular and coordination polymeric architectures was introduced, most of which exploit spontaneous and facile coordination reactions between dipyrrin ligands and metal ions. However, the bis(dipyrrinato)metal(II) complex motif, especially that with a zinc(II) center, absorbs and emits light intensely. Therefore, the motif may also be utilized as a light-harvesting, exciton-transmitting, and emitting subunit. Thus, by integrating bis(dipyrrinato)zinc(II) complexes into supramolecular assemblies and polymer systems, the resultant architectures are expected to induce certain photofunctionalities. In the following section, a few examples of such architectures are described.

4.1 Electron and energy transfer systems

Guldi and co-workers demonstrated that a bis(dipyrrinato)zinc(II) complex serves as a light-absorbing and electron-donating unit upon combination with fullerene, an electron acceptor (Fig. 19).⁹⁵ They designed triad **42** with two fullerene units linked by a bis(dipyrrinato)zinc(II) complex. The triad may be synthesized in a self-assembled fashion from a fullerene-appended dipyrrin ligand and zinc(II) ions. A spectroscopic study of the triad demonstrated that upon photoexcitation at the bis(dipyrrinato)zinc(II) unit with 480 nm light, a rapid charge-separation process follows, in which the photoexcited bis(dipyrrinato)zinc(II) unit serves as an electron donor. The lifetime of the charge-separated state is approximately 1 ns, with slight variations between different solvents.

Another example of utilizing the bis(dipyrrinato)zinc(II) complex motif as a light absorbent was demonstrated by Lindsey and co-workers (Fig. 20).⁹⁶ They developed several triads with a general formula of porphyrin-bis(dipyrrinato)zinc(II) complex-porphyrin, such as **43** (Fig. 20). The UV/vis/NIR absorption spectrum of **43** was a simple superposition of the

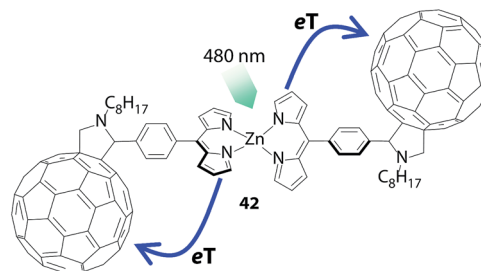


Fig. 19 Hybrid supramolecule **42** comprising fullerene and bis(dipyrrinato)zinc(II) complex units. eT denotes electron transfer.

spectra of the complex's components. In contrast, the bis(dipyrrinato)zinc(II) unit showed no contribution to the fluorescence spectrum of **43**; instead, only the zinc(II) porphyrin unit emitted luminescence. The authors interpreted this series of phenomena as quantitative energy transfer from the bis(dipyrrinato)zinc(II) complex to porphyrin units.

Sakamoto and Nishihara developed asymmetric linear assemblies of bis(dipyrrinato)zinc(II) complexes bearing anthracene, plain and π -extended dipyrrin ligands (**44**, Fig. 21a).⁹⁷ The three types of pigment moieties align in the order of photoexcitation energies from left to right, thereby an exciton ought to be transferred from the anthracene to the π -extended dipyrrin moiety through the plain dipyrrin linkers. In fact, the UV/vis/NIR absorption spectra of these dinuclear complexes are simple superpositions of the spectra of the pigment units, whereas their fluorescence spectra show features attributed to the π -extended dipyrrin at high efficiencies (up to a ϕ_F of 0.78), with the complete absence of emission from the rest (Fig. 21b). Moreover, the selective photoexcitation of each pigment converged to the identical ϕ_F and fluorescence lifetime. This series of investigations proved the presence of fast, quantitative, unidirectional energy transfer from the anthracene and plain dipyrrin to the π -extended dipyrrin unit. In conjunction with the ability to collect a broad range of UV and visible light, the series of photochemical properties allows asymmetric dinuclear complexes **44** to be fashioned into artificial light-harvesting systems and molecular photonic wires.^{98–105}

4.2 Thermoelectric conversion

As described in Section 3.2, one-dimensional coordination polymers **33** and **36** are dispersible in organic solvents,

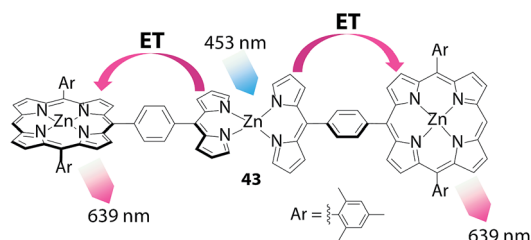


Fig. 20 Conjugate **43** between zinc(II) porphyrin and an intervening bis(dipyrrinato)zinc(II) complex. ET denotes energy (exciton) transfer.



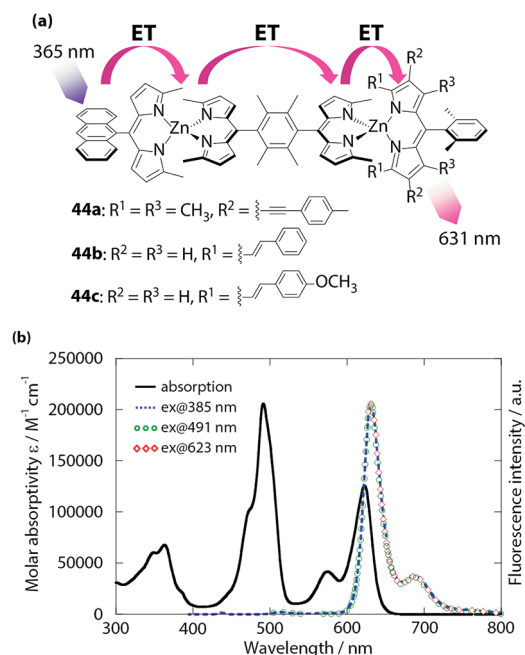


Fig. 21 (a) Asymmetric linear bis(dipyrrinato)zinc(II) complexes **44**. (b) UV/vis/NIR and luminescence spectra of **44c** in toluene. ET denotes energy (exciton) transfer. Adapted with permission from ref. 97. Copyright 2014 The Royal Chemical Society.

retaining their chain structure. Taking advantage of their processability, Sakamoto and Nishihara attempted to form a conjugate between **33** and single-wall carbon nanotubes (SWCNTs).⁷⁵ Nonoguchi and Kawai previously demonstrated that SWCNTs physisorbed with small organic molecules may control their carrier type and may enhance their thermoelectric conversion ability.¹⁰⁶ A dispersion of **33** and SWCNTs (1 : 10 w/w) in DMF was ultrasonicated and shaken and then filtered off to afford a free-standing film of the conjugate **33-SWCNT** (Fig. 22a). Transmission electron microscopy accompanied by electron energy-loss spectroscopy (TEM/EELS) demonstrated the existence of zinc, which was distributed uniformly on the **33** carbon scaffold derived from SWCNTs (Fig. 22b–e). Thus, **33** coiled around the SWCNTs uniformly. The thermoelectric conversion ability of **33-SWCNT** was evaluated based on a power factor of $\alpha^2\sigma$, where α and σ denote the Seebeck coefficient and electrical conductivity, respectively. **33-SWCNT** showed α and σ values of +66.2 $\mu\text{V K}^{-1}$ (Fig. 22f) and 76.5 S cm^{-1} , thereby demonstrating a power factor of 33 $\mu\text{W m}^{-1} \text{K}^{-2}$. These values all exceed those of pristine SWCNTs ($\alpha = +56.1 \mu\text{V K}^{-1}$, $\sigma = 29.5 \text{ S cm}^{-1}$, and $\alpha^2\sigma = 9.3 \mu\text{W m}^{-1} \text{K}^{-2}$).¹⁰⁶ The enhancement in the thermoelectric conversion ability of SWCNTs by **33** is also superior to that observed for small organic molecules ($\alpha^2\sigma < 26 \mu\text{W m}^{-1} \text{K}^{-2}$).¹⁰⁶

4.3 Photoelectric conversion

Two-dimensional polymer **37** described in Section 3.3 belongs to the “bottom-up” type of nanosheets, which are woven directly from atomic, ionic, and molecular components.¹⁰⁷ Research in this field has emerged; for example, single-layer cycloaddition-

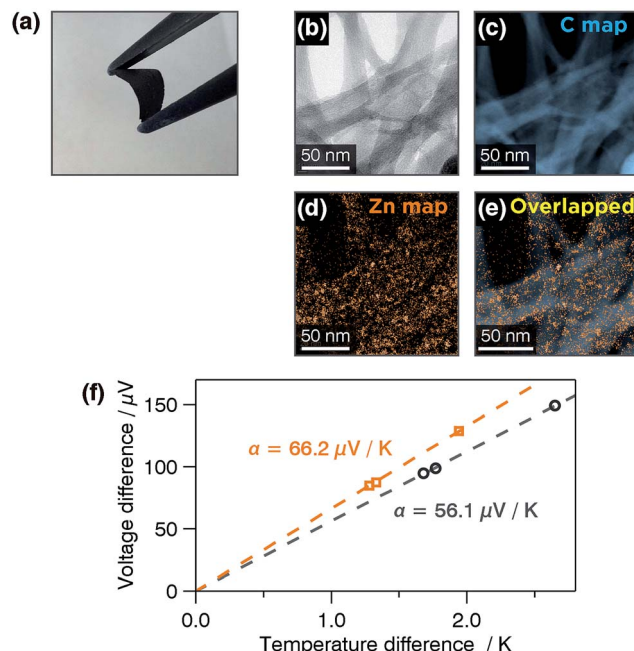


Fig. 22 (a) Photograph of a free-standing film of **33-SWCNT** with a thickness of 64 μm . (b) Bright-field-TEM image of **33-SWCNT** subjected to electron energy-loss spectroscopy (EELS) mapping. EELS mapping for: (c) carbon K edge intensity and (d) zinc M2 and M3 edge intensities. (e) Overlapped image of (b) and (c). (f) Voltage-difference/temperature difference plots for pristine SWCNTs (black) and **33-SWCNT** (orange). Adapted with permission from ref. 75. Copyright 2015 The Royal Chemical Society.

induced anthracene nanosheets featuring carbon–carbon covalent bonds,^{108–111} bis(2,2':6',2''-terpyridine)metal complex nanosheets,^{112,113} and a carbon-rich nanosheet synthesized from carbonization of an amphiphilic hexane molecule¹¹⁴ have been reported within the past five years. A distinctive characteristic of bottom-up nanosheets is that structures can be customized through the design and selection of components. This feature represents an advantage over the “top-down” type of nanosheets, such as graphene and metal dichalcogenides, the structure of which is confined by the bulk layered material from which the sheets are derived. This series of reports on the synthesis of bottom-up nanosheets, however, has concentrated on the fabrication and analysis of various two-dimensional structures, and no functionality has yet been demonstrated.^{115–119}

In this context, Sakamoto and Nishihara pursued the application of bottom-up nanosheet **37** in a photoelectric conversion system, taking advantage of the intense $1\pi-\pi^*$ absorption of the material.⁷⁹ We note that several mono(dipyrrinato)ruthenium(II)^{120–122} and copper(I)¹²³ complexes were employed as pigments in dye-sensitized solar cells (DSSCs). A transparent SnO_2 electrode was decorated with 36-layer **37**, which was used as a working electrode (photoanode). The modified SnO_2 electrode was incorporated into a three-electrode system, and a photoelectric conversion cell thereof was established using an acetonitrile electrolyte solution. Triethanolamine (TEOA) was also added as a sacrificial electron donor. An anodic current was gained only when the working electrode was irradiated with



500 nm light, corresponding to the absorption maximum of **37** (Fig. 23a). The action spectrum shown in Fig. 23b shows that the maximal photocurrent was acquired with 500 nm light, whereas no response was observed when using light outside of the range of the $^1\pi-\pi^*$ absorption. Control experiments that lacked either **37** or TEOA did not show a photocurrent response at all. These findings indicate that the observed photocurrent stemmed from the photocatalytic oxidation of TEOA, which was initiated by the photoexcitation of **37** on the SnO_2 electrode. The highest quantum yield reaches 0.86%, which is far greater than that of a mononuclear bis(dipyrrinato)zinc(II) complex sensitizer physisorbed on a SnO_2 electrode (0.030% in an aqueous electrolyte solution) and that of a sensitizer chemisorbed on the electrode through the carboxylate-metal oxide protocol (0.069%). The nanosheet structure of **37** affords suitable porosity and suppresses molecular aggregation, which presumably makes **37** a better sensitizer. In addition, single-layer **37** may be accumulated stepwise on a flat substrate to form a film with a desired thickness. For example, **37** synthesized by the air/liquid interfacial process (Fig. 15h and i) on a Langmuir-Blodgett trough was deposited iteratively on a quartz substrate at a constant surface pressure using the Langmuir-Schäfer method. The peak absorbance of the $^1\pi-\pi^*$ band at 500 nm was proportional to the number of deposition processes (Fig. 23c and d), which indicated the quantitative, layer-by-layer accumulation of single-layer **37**. Thus, **37** is the first photofunctional bottom-up nanosheet that exhibits both insolubility (to avoid redissolution into media) and tunability (to facilitate deposition and layering), making it useful for various potential applications.

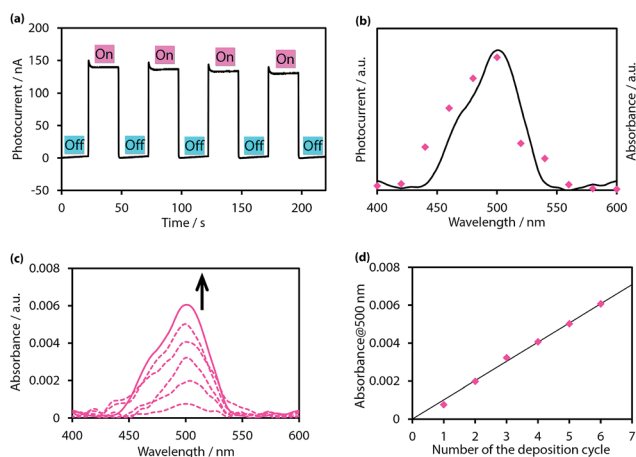


Fig. 23 (a) Typical anodic current response upon irradiation of a working electrode (SnO_2 substrate modified with 36-layer **37**) with intermittent 500 nm light in an acetonitrile electrolyte solution (0.1 M tetrabutylammonium perchlorate with 0.05 M TEOA) (b) action spectrum for the photocurrent generation (magenta dots) and absorption spectrum of **37** (black solid line). (c) Absorption spectral change upon stepwise depositions of single-layer **37** on a quartz substrate. (d) Linear relationship between the absorbance at 500 nm and the number of deposition processes. The magenta dots are extracted from Fig. 23c, and the black solid line corresponds to the least-squares linear fit of the plots. Adapted with permission from ref. 79. Copyright 2015 Macmillan Publishers Limited.

Sakamoto and Nishihara also reported that one-dimensional bis(dipyrrinato)zinc(II) coordination polymers **33** and **36** could be applied in photoelectric conversion (Fig. 24).⁷⁵ The processability of **33** and **36** or their dispersibility in organic solvents allowed the authors to fabricate thin films on SnO_2 electrodes by dropcasting their dispersions (Fig. 24a and d). A photoelectric conversion system was prepared in a manner similar to that used to produce nanosheet **37**, using a SnO_2 electrode modified as a photoanode. Again, photocurrent generation was confirmed upon excitation of **33** and **36** (Fig. 24b and e). The photoelectric conversion efficiencies were determined to be 1.0% and 0.027% for **33** (in an acetonitrile medium) and **36** (in an aqueous medium), respectively. These values are much greater than that of a mononuclear bis(dipyrrinato)zinc(II) complex sensitizer chemisorbed on the electrode (0.069% and 0% in, respectively, acetonitrile and aqueous media). It is also noteworthy that the responsive wavelength range of the photoelectric conversion may be tuned: the maximal photocurrent was observed to occur with 500 nm light in **33**, whereas that of **36** underwent a bathochromic shift by 40 nm (Fig. 23c and f). Such a redshift is esteemed in photoelectric conversion applications and can certainly be attained due to the designability and tunability of the present one-dimensional polymer system.

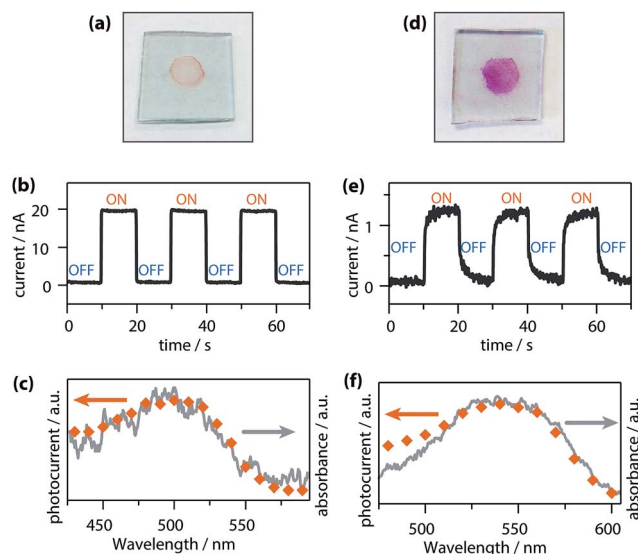


Fig. 24 (a) Photograph of a thin film of **33** on a SnO_2 electrode. (b) Typical anodic photocurrent response upon irradiation of a working electrode (SnO_2 substrate modified with **33** as shown in (a)) with intermittent 500 nm light in an acetonitrile electrolyte solution (0.1 M tetrabutylammonium perchlorate with 0.05 M TEOA). (c) Action spectrum for the photocurrent generation (orange dots) and absorption spectrum of **33** on a SnO_2 substrate (gray solid line). (d) Photograph of a thin film of **36** on a SnO_2 electrode. (e) Typical anodic photocurrent response upon irradiation of a working electrode (SnO_2 substrate modified with **36** as shown in (d)) with intermittent 550 nm light in an aqueous electrolyte solution (0.1 M sodium sulfate with 0.03 M TEOA). (f) Action spectrum for the photocurrent generation (orange dots) and absorption spectrum of **36** on a SnO_2 substrate (gray solid line). Adapted with permission from ref. 75. Copyright 2015 The Royal Chemical Society.



5. Conclusions

To date, a tremendous number of reports related to dipyrrens have been published, although most of them focus on BODIPYs, BF₂ complexes of dipyrrens, rather than other dipyrren metal complexes. Researchers have been attracted to the enticing nature of BODIPYs as fluorescent chromophores, in contrast to dipyrren metal complexes, which have long been regarded as less intriguing materials due to their negligible or low luminescence efficiencies. Herein, we described the rich chemistry of bis and tris(dipyrrenato)metal complexes developed over the past fifteen years. In Section 2, careful and elaborate molecular designs for achieving bright luminescence, including the introduction of *meso*-aryl groups with steric hindrance and the creation of heteroleptic complexes with suitable frontier orbital ordering, were discussed. Section 3 provided an overview of supramolecules, coordination polymers, and MOFs based on the bis and tris(dipyrrenato)metal complex motifs. One of the distinctive advantages of dipyrren metal complexes over BODIPYs is the spontaneous formation of the coordination bonds, which allows for the formation of a series of self-assembled supramolecular and polymeric architectures in conjunction with the appropriate design of dipyrren ligands, such as oligo(dipyrren)s and those with additional coordination sites. The favorable features and characteristic structures demonstrated to date potentially renders bis and tris(dipyrrenato)metal complexes valuable materials. Reported attempts to produce such materials have been presented in Section 4, which discussed how methods based on photoinduced charge separation, fast, quantitative, unidirectional exciton transport, and thermoelectric and photoelectric conversion have been pursued. The remarkable progress made in bis and tris(dipyrrenato)metal complex chemistry represents a milestone toward the development of profound science and real-world applications.

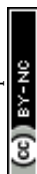
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