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# Tripyrrin-armed isosmaragdyrins: synthesis, heterodinuclear coordination, and protonation-triggered helical inversion†

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Oxidative ring closure of linear oligopyrroles is one of the synthetic approaches to novel porphyrinoids with dinuclear coordination sites and helical chirality. The spatial arrangement of the pyrrolic groups of octapyrrole (**P8**) affected the position of the intramolecular oxidative coupling of the pyrrolic units; tripyrrin-armed isosmaragdyrin analogue (**1**) containing a  $\beta,\beta$ -linked bipyrrole moiety was synthesized regioselectively in a high yield by using  $\text{FeCl}_3$ .  $\text{Ni}^{\text{II}}$ -coordination at the armed tripyrrin site of **1** allowed the formation of diastereomeric helical twisted complexes (**2A** and **2B**) and succeeding  $\text{Cu}^{\text{II}}$ -coordination at the macrocyclic core afforded heterodinuclear  $\text{Ni}^{\text{II}}/\text{Cu}^{\text{II}}$ -complexes (**3A** and **3B**). Each of them comprised a pair of separable enantiomers, exhibiting *P*- and *M*-helices, respectively. Notably, diastereomeric interconversion from **2A** to **2B** was quantitatively achieved as a consequence of helical transformation under acidic conditions.

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## Introduction

Recently, increasing attention has been focused on expanded porphyrin analogues bearing large  $\pi$ -conjugated oligo-pyrrolic scaffolds because of their diverse structural properties.<sup>1</sup> The expanded porphyrins can serve as unique ligands for multi-metal coordination and adopt tunable three-dimensional conformations (*e.g.*, figure-eight, triangle, spiral twisted, *etc.*) with aromaticity-dependent near-infrared optical properties.<sup>2</sup> In some cases, topologically twisted metallo-porphyrinoids serve as chiral sensing components.<sup>3</sup> The metalation-induced switchable function of expanded porphyrinoids (such as changes in stereoisomeric structures) should be fascinating for applications in environmentally-responsive molecular machines.<sup>4</sup>

To build such large  $\pi$ -conjugated porphyrinoids, various approaches have been developed, mostly by acid-catalyzed condensation and organometallic cross-coupling reactions.<sup>5</sup> Among them, oxidative ring closure of linear oligopyrroles could produce bipyrrole-containing porphyrinoids directly linked at the two terminal pyrroles in the  $\alpha$ - $\alpha'$  or  $\alpha$ -N modes (Fig. 1a).<sup>6</sup> However, less investigation has been carried out on oxidative coupling between  $\beta,\beta'$ -pyrrolic C-Hs using a rational approach, which may be due to the intrinsic low reactivity of pyrrolic  $\beta$ -CHs compared to  $\alpha$ -CHs and the lower acidity of CHs than typical N-Hs.<sup>7</sup> Moreover, the oligopyrroles commonly used in the oxidative ring closure reactions are usually too flexible to regulate the targeted reactive sites close to each other.

Inspired by the oxidative coupling reactions of tetraaza[8]circulenes (depicted in Fig. 1b)<sup>7c</sup> and triply fused porphyrin tapes<sup>7d</sup> where the reaction sites (*i.e.*,  $\beta$ -pyrrole CHs) reside spatially in close proximity, we here report the synthesis of a novel  $\beta,\beta$ -linked isosmaragdyrin-based macrocycle (**1**) from a fully conjugated linear octapyrrole (**P8**)<sup>8</sup> with terminal  $\alpha$ -acyl substituents by oxidative aromatic coupling using  $\text{FeCl}_3$ . The octapyrrole **P8** adopts a spiral conformation through its intrinsic hydrogen-bonding network, which resulted in positioning the specific CHs of pyrroles in close proximity. Indeed, two pyrrolic  $\beta$ -carbon atoms in **P8** on rings A and E are close to each other with a short distance of 3.7 Å in the crystal (Fig. 1 and S24†), which enables a regioselective internal ring closure through the oxidative C–H coupling to afford a tripyrrin-armed pentapyrrolic macrocycle containing a  $\beta,\beta$ -linked bipyrrole moiety in high yield (Fig. 1c). Notably, the appended tripyrrin moiety and

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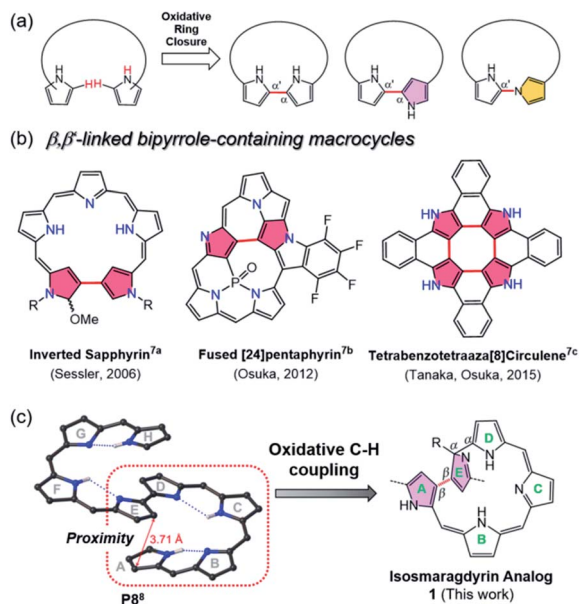


Fig. 1 (a) Typical synthetic approaches for constructing bipyrrole-containing porphyrinoids linked in different modes through oxidative coupling of oligopyrroles. (b) Representative macrocyclic compounds containing  $\beta,\beta'$ -linked (red) bipyrrole moieties. (c) The oxidative coupling reaction of a helical oligopyrrole, **P8**, where the appropriate pyrrole rings are present spatially in close proximity, giving a new isosmaragdyrin analogue **1**. Meso-aryl rings are omitted for clarity.

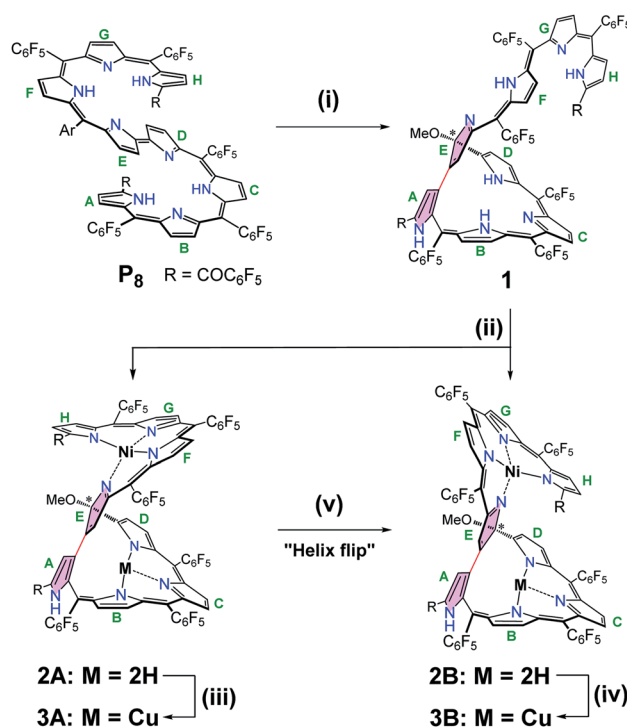
one of the outward-pointing pyrrolic nitrogen atoms of the macrocycle constitute a peripheral N4 coordination site and the isosmaragdyrin core serves as an internal N3 donor site, hence **1** provides hetero-bimetal chelation spheres. Taking advantage of this unique structure, metal-dependent complexation (with  $\text{Ni}^{\text{II}}$  and  $\text{Cu}^{\text{II}}$  ions) at both the peripheral and the isosmaragdyrin cores successfully afforded helical diastereomeric mononuclear  $\text{Ni}^{\text{II}}$  complexes (**2A** and **2B**) and heterodinuclear  $\text{Ni}^{\text{II}}\text{Cu}^{\text{II}}$  complexes (**3A** and **3B**). Each of these complexes contains a pair of enantiomers exhibiting *P*- and *M*-helices, where all four optically pure isomers (*i.e.*, **M,R-x**, **P,R-x**, **P,S-x**, and **M,S-x**;  $x = 2$  or  $3$ ) have been successfully separated. Unprecedentedly, diastereomeric interconversion from **2A** to **2B** was quantitatively achieved by the addition of trifluoroacetic acid (TFA), thus enabling transformation between the *P*- and *M*-helices. These results provide a practical approach for developing novel porphyrinoids through oxidative ring closure at the spatially adjacent  $\beta$ -positions of elaborately designed conjugated oligopyrroles.

## Results and discussion

The oxidative cyclization of linear octapyrrole **P8** was performed using 10 equivalents of anhydrous  $\text{FeCl}_3$  in  $\text{CH}_2\text{Cl}_2/\text{MeOH}$  for 24 h at room temperature. The  $\beta$ -positions of the spatially adjacent pyrrole units were regioselectively linked to afford **1** in a high yield of 88% (Scheme 1). The resulting macrocycle can be regarded as a novel isosmaragdyrin analogue (Fig. 1c).<sup>9</sup>

In the macrocycle **1**,  $\pi$ -conjugation was disrupted, as evident from the blue-shifted absorption spectrum ( $\lambda_{\text{max}} = 1073$  and  $638$  nm for **P8** and **1**, respectively) (Fig. S11†). The high resolution mass spectrum (HRMS) of **1** showed a molecular ion peak at  $m/z = 2010.1192$  ( $\text{M}^+$ ), consistent with a molecular formula of **1**:  $\text{C}_{89}\text{H}_{22}\text{F}_{40}\text{N}_8\text{O}_3$  (indicating the formal addition of a methoxy group to **P8**, *i.e.*, loss of two hydrogens by oxidation with  $\text{FeCl}_3$ , followed by the addition of  $\text{MeOH}$ ). The  $^1\text{H}$  NMR spectrum of **1** in  $\text{CDCl}_3$  indicates the unsymmetrical structure and shows two sets of signal patterns in the entire region, suggesting the presence of two conformational isomers in solution (Fig. S1 and S4†).<sup>10</sup> For example, the signal of the  $\text{OCH}_3$  group appeared at  $\delta = 3.19$  and  $3.09$  ppm with a ratio of  $0.85 : 1$  (calculated from the integral) for the two isomers. On the basis of the preliminary crystal structure of **1**, a tripyrrin appended pentaphyrin structure consisting of a direct  $\beta,\beta'$ -linkage between the terminal ring A and the middle ring E was verified (Fig. S25†).<sup>11</sup> The unusual  $\beta,\beta'$ -bonding mode tends to exhibit a significant distortion of the pentapyrrolic core similar to that observed in the tetrapyrrolic corrin.<sup>6d</sup> Notably, a methoxy group was found at the  $\alpha$ -position of one of the directly linked pyrroles (ring E), thus generating an  $\text{sp}^3$  hybridized carbon linkage. The isosmaragdyrin macrocycle of **1** can be thus regarded as a nonaromatic derivative of modified isosmaragdyrin.

Owing to the intrinsic pentaphyrin cavity and flexible oligopyrrole arm, isosmaragdyrin derivative **1** may bind two metal



Scheme 1 Stepwise syntheses of pentaphyrin **1** and metal complexes **2** and **3**. The labels, A–H, are defined to identify each pyrrole ring. Conditions: (i)  $\text{FeCl}_3$ ,  $\text{CH}_2\text{Cl}_2/\text{MeOH}$  (1 : 50), 88%; (ii)  $\text{Ni}(\text{OAc})_2 \cdot 4\text{H}_2\text{O}$ ,  $\text{CH}_2\text{Cl}_2/\text{MeOH}$ , 63% for **2A**, 19% for **2B**; (iii)  $\text{Cu}(\text{OAc})_2 \cdot \text{H}_2\text{O}$ ,  $\text{MeOH}$ , 81%; (iv)  $\text{Cu}(\text{OAc})_2 \cdot \text{H}_2\text{O}$ ,  $\text{MeOH}$ , 91%; (v) TFA,  $\text{CH}_2\text{Cl}_2$ .

ions and exhibit unexpected properties, such as external-stimuli induced helical transfer.<sup>12</sup> Actually, **1** allows the coordination of metal cations at the tripyrrin side chain (Scheme 1). When a methanol solution of **1** and 50 equiv. of  $\text{Ni}(\text{OAc})_2 \cdot 4\text{H}_2\text{O}$  was heated at reflux in the presence of  $\text{NaOAc}$  under anaerobic conditions, two greenish products, that is, isomeric mononuclear  $\text{Ni}^{\text{II}}$  complexes **2A** and **2B** were obtained in 63% and 19% yields, respectively (Scheme 1). The HRMS of **2A** and **2B** showed essentially identical values of  $m/z = 2066.0459$  and  $2066.0432$  (Fig. S18–S19†), respectively, indicating that the molecular formula is  $\text{C}_{89}\text{H}_{20}\text{F}_{40}\text{N}_8\text{O}_3\text{Ni}$ , which is consistent with the mononuclear  $\text{Ni}^{\text{II}}$  chelate. Later the X-ray crystallographic analysis disclosed the structures of **2A** and **2B** with helical twists (Fig. 2a–d).<sup>13</sup> The  $\text{Ni}^{\text{II}}$  cation is wrapped with three N atoms from the tripyrrin unit (rings F–H) and the fourth N atom from the confused pyrrole (ring E), while the inner cavity of the macrocycle remains uncoordinated. The directly linked pyrrole rings A and E are located almost coplanar, and the rings C and H are at opposite sides of the mean plane defined by rings A and E in **2A**. However, the rings C and H in **2B** are arranged on the same side of the plane. Thus, **2A** and **2B** can be regarded as pseudo-*trans* and pseudo-*cis* conformations, respectively. The methoxy-attached pyrrolic  $\alpha$ -carbon (*i.e.*, C20) for **2A** is  $\text{sp}^3$  hybridized, which can be evidenced by the C20–C21, C20–C19, C20–O, and C20–N distances of 1.495(8), 1.536(7), 1.410(6), and 1.498(7) Å, respectively, typical values for C–C, C–O, and C–N single bonds.<sup>13</sup> Similar bond lengths around C20 were also observed in **2B** and the rough crystal structure of **1** (Fig. S25†). The pseudo-tetrahedral geometry of the  $\text{sp}^3$  carbon center can be further elucidated by the bond angles at C20 lying in the range from  $103.7(4)$  to  $111.6(4)^\circ$ , which are close to  $109.5^\circ$ , a typical value for  $T_d$  carbon atoms. Thus, C20 is a chiral center. Meanwhile, the helical structure of the tetrapyrrole (biliverdin-like)  $\text{Ni}^{\text{II}}$ -complex leads to another source of molecular chirality. As a result, there are four possible chiral isomers, namely, **M,R-2**, **P,R-2**, **M,S-2**, and **P,S-2**, where **R** and **S** represent the chirality of C20, and **M** (left-handed) and **P** (right-handed)<sup>14</sup> denote the handedness of the biliverdin-like unit. Indeed, a pair of enantiomers, **M,R-2** and **P,S-2**, was present in the crystals of the racemic compound **2A**. Likewise, the racemic crystal **2B** comprises enantiomers, **P,R-2** and **M,S-2**. Hence, compounds **2A** and **2B** can be regarded as diastereomeric isomers.

Further metalation of **2A** and **2B** with  $\text{Ni}^{\text{II}}$  ions did not proceed under the above conditions. Instead, the reaction of **2A** and **2B** with 10 equiv. of  $\text{Cu}(\text{OAc})_2$  in a mixture of  $\text{CH}_2\text{Cl}_2/\text{MeOH}$  (1 : 1) at reflux for 2 h afforded hetero-dinuclear  $\text{Ni}^{\text{II}}/\text{Cu}^{\text{II}}$  complexes, **3A** and **3B**, in the yields of 81% and 91%, respectively (Scheme 1). The molecular ion peaks at  $m/z = 2126.9590$  and  $2126.9608$ , respectively, agree with the dinuclear  $\text{Ni}^{\text{II}}\text{Cu}^{\text{II}}$  coordination structures (Fig. S20 and S21†). The broad active EPR spectra of complexes, **3A** and **3B**, are consistent with the coordination of the paramagnetic  $\text{Cu}^{\text{II}}$  ion (Fig. S22 and S23†).<sup>15</sup> Incorporation of a  $\text{Cu}^{\text{II}}$  ion into the inner N atoms of the macrocyclic core was also identified by X-ray crystallographic analysis (Fig. 2e–h).<sup>11</sup> Notably, the  $\text{Cu}^{\text{II}}$  cations in **3A** and **3B** coordinate to the nitrogen atoms of pyrrole rings B, C, and D in a T-shaped manner and the vacant site of the square-planar  $\text{Cu}^{\text{II}}$

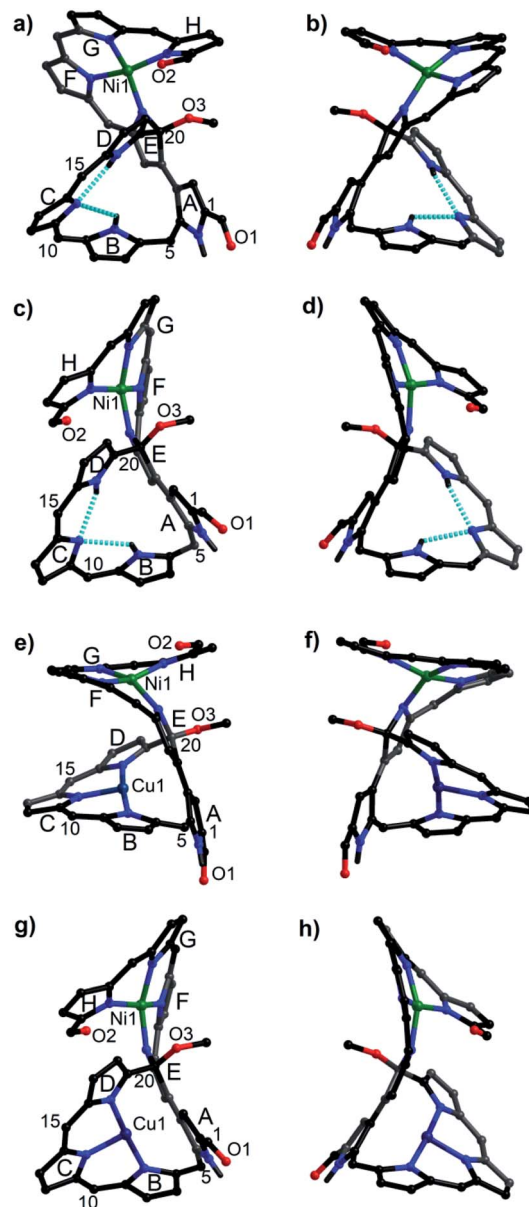


Fig. 2 Molecular structures of **2A** (a and b), **2B** (c and d), **3A** (e and f), and **3B** (g and h). (a) and (b) A pair of enantiomers in the racemic crystals of **2A**, specifically **M,R-2** for (a) and **P,S-2** for (b).<sup>16</sup> Similarly, **P,R-2** for (c), **M,S-2** for (d), **M,R-3** for (e), **P,S-3** for (f), **P,R-3** for (g), and **M,S-3** for (h).  $\text{C}_6\text{F}_5$  groups and the hydrogen atoms attached to carbon atoms are omitted for clarity.

ion seems to be occupied by neighboring  $\text{sp}^2$  carbons ( $\text{Cu}-\text{C}$ : 2.380–2.748 Å) presumably through  $\text{Cu}^{\text{II}}$ -arene interactions (Fig. S27 and S28†).<sup>16</sup> From the viewpoint of molecular chirality, complex **3A** has the same chirality as **2A**, consisting of enantiomers **M,R-3** and **P,S-3** (Fig. 2e and f). Likewise, the crystals of **3B** comprise enantiomers **P,R-3** and **M,S-3** (Fig. 2g and h).

The aforementioned addition of a methoxy group introduces a chiral center into the macrocycle of **1**. To understand the chiroptical properties of the isosmaragdyrin derivatives, enantiomeric separation was conducted using a chiral HPLC column with a mixture of *n*-hexane/2-propanol as the eluent. A





pair of enantiomers of **1** (**S-1** and **R-1**, respectively) was thus separated roughly in a 1 : 1 ratio (Fig. S2a†). The first fraction **1-I** and the second **1-II** displayed opposite Cotton effects in the circular dichroism (CD) spectra; they can be assigned as **R-1** and **S-1**, respectively, judging from the data obtained experimentally and theoretically using time-dependent density functional theory (TD-DFT) calculations (Fig. S2b†). The absolute configuration of C20 remains intact during metalation; that is, coordination of **R-1** with Ni<sup>II</sup> ions afforded a diastereomeric mixture of **M,R-2** and **P,R-2**, while the corresponding reaction of **S-1** likewise afforded **P,S-2** and **M,S-2**. These diastereomers can be separated on conventional silica gel columns. Along this way, subsequent complexation of a Cu<sup>II</sup> ion produced a pair of pure chiral isomers (**M,R-3**, **P,R-3**) and (**P,S-3**, **M,S-3**), respectively. According to the CD spectra of the resulting enantiomers **2** and **3** (Fig. S34–S37†), the intense Cotton effects were observed in the series of **M,R-x** and **P,S-x** ( $\Delta\epsilon \approx 400 \text{ M}^{-1} \text{ cm}^{-1}$ ) compared to those of **M,S-x** and **P,R-x** ( $\Delta\epsilon \approx 100 \text{ M}^{-1} \text{ cm}^{-1}$ ). These stereostructure-dependent spectral features could be rationalized by theoretical simulations (Fig. S39–S42†). The different strengths of  $\Delta\epsilon$  may reflect the extent of magnetic dipole transition moments for each electronic transition of the complexes.<sup>17</sup>

Unprecedentedly, it was found that the isomer **2A** could be quantitatively converted to **2B** upon treating with 1 equiv. of TFA in CH<sub>2</sub>Cl<sub>2</sub> for 12 h (Scheme 1). TFA-induced stereo-transformation was further analyzed by CD, absorption and <sup>1</sup>H NMR spectroscopy. Upon treatment with TFA, the CD spectrum of **M,R-2** in CH<sub>2</sub>Cl<sub>2</sub> was spontaneously changed to the same spectra obtained from the acidic solution of **P,R-2** while retaining the stereochemistry (Fig. 3). Absorption spectroscopy also showed the clear conversion from **2A** to **2B** in the presence of TFA in CH<sub>2</sub>Cl<sub>2</sub> or MeOH (Fig. S12, S13, and S15†). Whereas, the profile of **2B** remained intact under the same conditions. The <sup>1</sup>H NMR spectra also assisted in tracking the helix inter-conversion processes between **2A** and **2B**. When the solution of

**2A** in CD<sub>2</sub>Cl<sub>2</sub> was treated with TFA, the signals of **2A** gradually disappeared and subsequently, a new set of signals appeared identical to what is seen for **2B** under acidic conditions (Fig. S8†). In CDCl<sub>3</sub> or CD<sub>2</sub>Cl<sub>2</sub>, broadening of a specific NH peak for **2B** was observed upon addition of TFA (Fig. S7 and S9†).

Although the mechanism for the helical inversion is yet unknown, TFA-assisted protonation at the most probable imino nitrogen site of the isosmaragdyrin core of **2A** and **2B** could occur, which affected the hydrogen bonding network in the core and caused a conformational change, as inferred from the broadening of the NH signals of the resulting species in the <sup>1</sup>H NMR spectrum (Scheme S1†). Intervening of the associated counter anion in this helical inversion sterically or through metal coordination could not be excluded.<sup>18</sup> The solution of **2A** or **2B** in CD<sub>3</sub>CN reached an equilibrium state (the molar ratio **2A** : **2B**  $\approx$  3 : 2 from the integrals of the signals) by treatment with TFA (Fig. S10†). The solvent basicity may affect the protonation/deprotonation dynamics of the above system. In the case of Cu<sup>II</sup>/Ni<sup>II</sup> complexes, **3A** and **3B** reached equilibria in various media in the presence of acid. Demetallation of Cu<sup>II</sup> ions of **3A** and **3B** seems to have occurred, judging from the resulting absorption spectra, especially in CH<sub>3</sub>CN (Fig. S14–S16†).

The calculated energies of all conformers enabled the interpretation of the relative stability of the isomers.<sup>19</sup> In particular, the pseudo-*trans* conformation of **2A** has a favorable van der Waals interaction, whereas the pseudo-*cis* conformation of **2B** has better coplanarity between the coordinating and non-coordinating moieties. As a result, the pseudo-*cis* conformation for **2B** is more stable than the pseudo-*trans* one for **2A** (Fig. 4), which is in contrast to the observation that the yield of **2A** is much higher than that of **2B** during the nickel complexation of **1**. The nickel complex **2A** could thus be a kinetically favorable product, while **2B** is a thermodynamically favorable one. The conformation of the helical structure of **2** is unaffected by further metal coordination. For complexes **3**, **3B** is also more

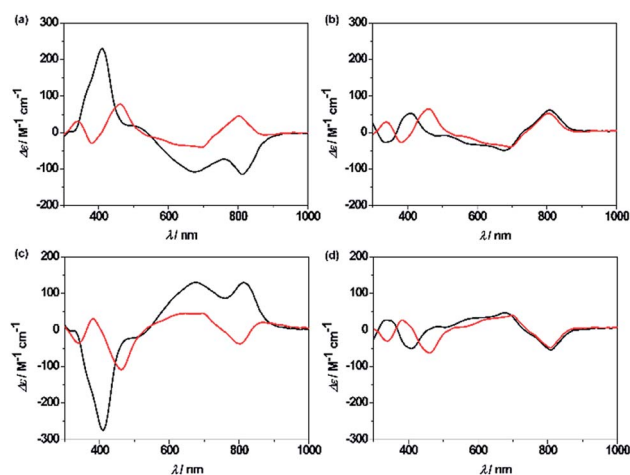


Fig. 3 CD spectra of the complexes in CH<sub>2</sub>Cl<sub>2</sub> in the absence (black line) and presence (red line) of TFA, (a) **M,R-2**, (b) **P,R-2**, (c) **P,S-2**, and (d) **M,S-2**.

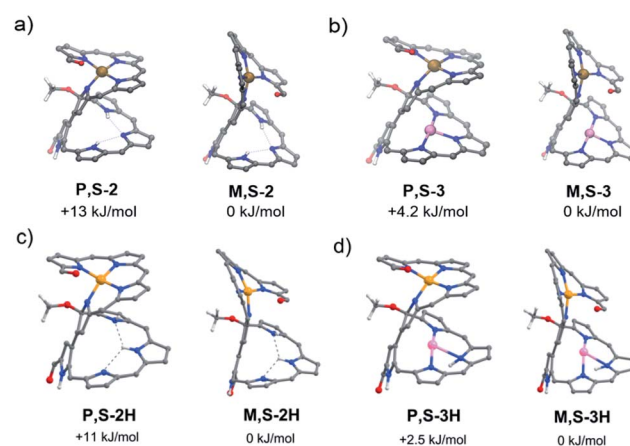


Fig. 4 Relative energies between two conformers of (a) **2A** and **2B**, (b) **3A** and **3B**, (c) **2AH** and **2BH**, and (d) **3AH** and **3BH**. The energies of **P,S-2**, **P,S-2H**, **P,S-3**, and **P,S-3H** (corresponding to **2A**, **2AH**, **3A**, and **3AH**) were calculated relative to those of **M,S-2**, **M,S-2H**, **M,S-3**, and **M,S-3H** (corresponding to **2B**, **2BH**, **3B**, and **3BH**), respectively.



stable than **3A**, but the net stabilization effect for **2** is more remarkable than that for **3** ( $\Delta E = 13 \text{ kJ mol}^{-1}$  vs.  $4.2 \text{ kJ mol}^{-1}$ ), which agrees with the ease of transformation for **2**. The activation energy for this helical inversion process should be rather high, judging from the unidirectional helical changes, which might be relevant to the unusually high rotation barriers of the 3,3'-bipyrrole system.<sup>20</sup> Consistently, **2A/2B** and **3A/3B** were not observed to interconvert while using temperature-dependent CD measurements. On the other hand, we can see from Fig. 4c and d that the protonation of the free-base compound **2** and corresponding Cu complex **3** does not cause visible changes in the conformational structures of the initial non-protonated species, but the energy differences between **P,S-2/3H** and **M,S-2/3H** become smaller relative to the non-protonated cases (Fig. 4a and b). Hence, we can assume that protonation might also decrease the activation barrier for the helical inversion processes leading to easier equilibria between the protonated **2A/2B** and **3A/3B** species.

## Conclusions

In summary, the  $\beta$ -positions of two pyrrole rings close to each other in the pre-organized framework of conjugated octapyrrin **P8** were linked selectively and efficiently by the  $\text{FeCl}_3$ -promoted oxidative cyclization reaction. As a result, a tripyrrin appended doubly N-confused isosmaragdyrin analogue **1** has been obtained with concurrent addition of a methoxy group to one of the  $\alpha$ -positions of the bipyrrole moiety. Stepwise coordination of **1** with  $\text{Ni}^{\text{II}}$  and  $\text{Cu}^{\text{II}}$  led to the formation of diastereomeric mononuclear  $\text{Ni}^{\text{II}}$  complexes **2A/2B** and hetero-dinuclear  $\text{Ni}^{\text{II}}$ - $\text{Cu}^{\text{II}}$  complexes **3A/3B**. X-ray crystal structures clearly revealed the unique helical oligopyrrin-Ni complex appended pentaphyrin structures of **2** and **3**. The optical resolution of **1** followed by stepwise coordination afforded all four separable isomers, **M,R-x**, **P,R-x**, **P,S-x**, and **M,S-x** ( $x = 2$  or  $3$ ). Intriguingly, the diastereomeric transformation between *P*- and *M*-twists from **2A** to **2B** was achieved upon simple treatment with acid (e.g., TFA). The combination of an open-chain oligopyrrin and a chiral porphyrinoid backbone has been demonstrated to be effective for achieving tunable chirality with external stimuli, and may be applied as a component of molecular machines or motors.<sup>21</sup> In brief, oxidation of conjugated oligopyrroles paves a new way to the synthesis of a diversity of porphyrinoids, particularly through unique  $\beta,\beta$ -linkages.

## Conflicts of interest

There are no conflicts to declare.

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