Cite this: *Chem. Sci.*, 2018, 9, 6853

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 22nd June 2018
Accepted 16th July 2018

DOI: 10.1039/c8sc02739k

rsc.li/chemical-science

Conjugated double helices *via* self-dimerization of α,α' -dianilinothripyrrins

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A new motif for artificial double helices was developed on the basis of α,α' -disubstituted tripyrrin. α,α' -Dibromotripyrrin **3** was prepared by gentle bromination at the pyrrolic α -positions of 5,10-diphenyltripyrane followed by oxidation with DDQ. Nucleophilic substitution reactions of **3** with anilines proceeded efficiently to furnish a series of α,α' -dianilinothripyrrins **4–11**, which displayed monomeric and dimeric forms depending upon the solvent used for crystallization and the structures of the substituted anilines. Dimeric forms show double helical structures with smooth π -conjugation as indicated by their absorption spectra. van't-Hoff plot analyses revealed that the dimerizations in CDCl_3 are enthalpy-driven. Larger association constants of the dimerization are attained for 3,5-di-*t*-butylanilino- and 3,5-bis(trifluoromethyl)anilino-substituted tripyrrins (**7** and **8**) *via* additional multiple intermolecular interactions. In a nonpolar and aprotic solvent, tripyrrins (**9** and **10**) bearing bulkier 1-naphthylamino and mesitylamino groups do not dimerize but undergo unique tautomerization.

Introduction

An artificial double helix motif has stimulated chemists' interest not only to mimic the biological functions as seen in DNAs, but also to develop new functional devices for organic electronics and photonics.¹ Significant efforts have been made for synthetic oligomers that can form bimolecular double helices through different types of non-covalent interactions.^{2–5} One of the most useful non-covalent interactions is metal–ligand coordination. Metal ion-directed self-assemblies in which the organic ligands form double-helical complexes are called double helicates.² As a pioneering study, J. M. Lehn *et al.* employed copper(II)-chelation by bipyridyl ligands to obtain double helicates (**A**).^{2a} A variety of metallohelicates utilizing pyridyl coordinations have been extensively studied later.^{2b–d} Not only transition metal complexes, but also spiroborates (**B**) were found to form double helicates, in which coordination of the central sodium ion plays an important role in shielding the electrostatic repulsion between the two borate groups.³ As for purely organic approaches, effective salt-bridge formation was utilized for *m*-terphenylene-based double helical oligomers (**C**).⁴ Yashima and co-workers have beautifully extended this motif to well-defined double helical polymers.⁴ On the other hand, Huc and co-workers synthesized double-stranded helices of aromatic oligoamides (**D**) in which the NH site of the amide moiety and the N site of the pyridine moiety serve as the

hydrogen bonding donor and acceptor, respectively.⁵ As such, hydrogen bonding is a useful intermolecular interaction to hold the supramolecular arrangement, and a variety of hydrogen bonding donor and acceptor sites have been properly allocated to form double helical structures.⁶ However, there are only a few examples of fully π -conjugated double helices that may be promising for visible-light harvesting or effective carrier transporting along the helix axis.⁷ Recently, a large polycyclic helical molecule (**E**), so called expanded helicene, was reported to display a homochiral double helical structure in solution and in the solid-state due to the favourable π -stacking interaction.⁸ A new motif of artificial double helices having smooth π -conjugation still remains highly desired (Chart 1).

Here, we report a new motif of a fully conjugated double helix based on tripyrrin derivatives. Tripyrrin is a conjugated tri-pyrrolic chain consisting of three pyrrolic moieties and two bridging methine carbons.^{9–11} Since the tripyrrin itself is generally not so stable, the isolation of tripyrrin derivatives has been limited to its protonated salts,^{9a} metal complexes,^{9b–f} oxygenated derivatives (tripyrrinones)¹⁰ or those with steric protections at the labile α -positions.¹¹ We developed α,α' -dianilinothripyrrins as stable entities exhibiting a large absorption band reaching the NIR region. Surprisingly, some of these molecules spontaneously form doubly helical structures *via* effective intramolecular and intermolecular hydrogen bonding interactions.

Results and discussion

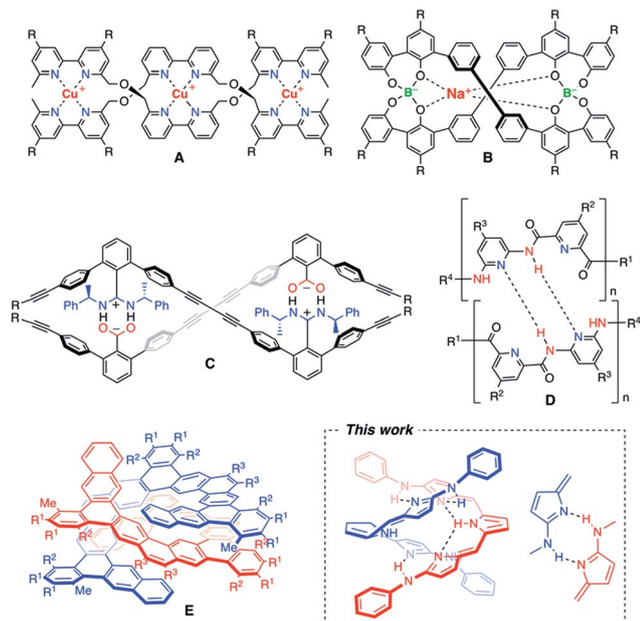
Synthesis of α,α' -dibromotripyrrin

5,10-Diphenyltripyrane **1** was easily prepared according to Gryko's report.¹² Bromination of **1** with 2.1 equivalents of

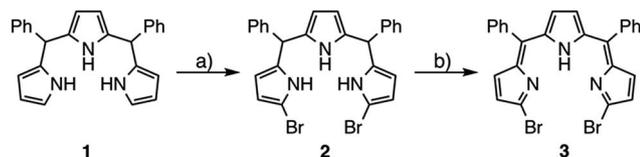
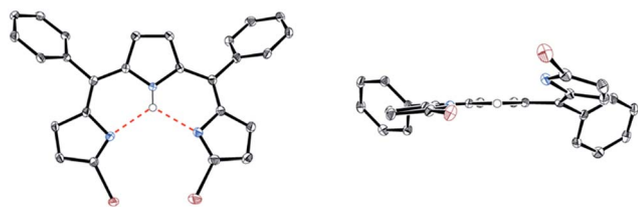
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† Electronic supplementary information (ESI) available: CCDC 1849984, 1849988, 1849990–1849996, 1850011–1850013, 1850022 and 1850026. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c8sc02739k



Chart 1 Examples of artificial double helix molecules.^{2–5}

N-bromosuccinimide at $-78\text{ }^{\circ}\text{C}$ afforded α,α' -dibrominated tripyrrane **2** in 72% yield (Scheme 1).¹² Subsequently, oxidation of **2** with 2.1 equivalents of DDQ gave a key compound, α,α' -dibromotripyrrin **3**, in 87% yield. Fortunately, **3** is fairly stable under aerobic conditions and can be purified by silica-gel column chromatography. Indeed, recrystallization from $\text{CH}_2\text{Cl}_2/\text{MeOH}$ gave crystals suitable for X-ray diffraction analysis (Fig. 1). Due to the intramolecular hydrogen bonding networks among the two imine-like pyrroles and one pyrrole-NH, **3** takes a *cis*-conformation with a slight deformation from planarity due to the steric repulsion between the two bromine atoms. The ^1H NMR spectrum of **3** is consistent with the structure, featuring two doublets at 6.75 and 6.54 ppm due to

Scheme 1 Synthesis of α,α' -dibromotripyrrin **3**: (a) NBS, THF, $-78\text{ }^{\circ}\text{C}$, 2.5 h, 72%; (b) DDQ, CH_2Cl_2 , $0\text{ }^{\circ}\text{C}$, 10 min, 87%.Fig. 1 X-ray crystal structure of **3**. Solvent molecules and hydrogen atoms except for NH are omitted for clarity. The thermal ellipsoids are scaled to 50% probability.

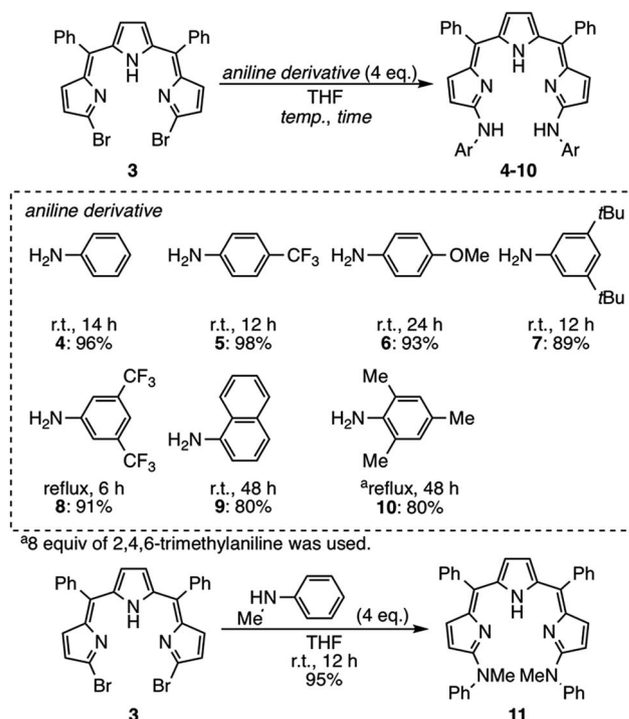
the β -protons of imine-like pyrroles, a singlet at 6.31 ppm due to the β -protons of the central pyrrole and the NH proton signal at 13.44 ppm.

Nucleophilic substitution reactions of **3** with anilines

Next, nucleophilic substitution reactions of **3** with anilines were examined (Scheme 2). With 4 equivalents of aniline in dry THF at room temperature, the substitution reaction of **3** proceeded very smoothly to give α,α' -dianilinothripyrrin **4** in 96% yield.¹³ Treatment of **3** with an equimolar amount of aniline in THF/acetonitrile at room temperature afforded a mixture, from which α -anilino- α' -bromotripyrrin **5**† was isolated in 90% yield.¹⁴ The substitution reactions of **3** were also examined with different types of aniline derivatives. *p*-Trifluoromethylaniline and *p*-methoxyaniline were reacted to give the corresponding anilinothripyrrin derivatives **6** and **7** in 98% and 93% yields, respectively. Sterically encumbering 3,5-di-*t*-butylaniline, 3,5-bis(trifluoromethyl)aniline and 1-naphthylaniline also reacted to afford **8**, **9** and **10** in 89%, 91% and 80% yields, respectively. Sterically more hindered 2,4,6-trimethylaniline did not react at room temperature, but upon refluxing in THF for 48 h, **10** was obtained in 80% yield. Secondary *N*-methylaniline also reacted smoothly to afford *N,N'*-dimethylanilinothripyrrin **11** in 95% yield.

X-ray crystallographic analysis

Single crystals of **4** were obtained from its solution in CHCl_3 and *n*-hexane (Fig. 2a). The X-ray diffraction analysis revealed that **4** formed a double helix in the solid state due to the interstrand hydrogen bonding interaction between the aniline-NH and the imine-type pyrrole (Fig. 2b, right) as well as the intramolecular

Scheme 2 Substitution reactions of **3** with aniline derivatives.

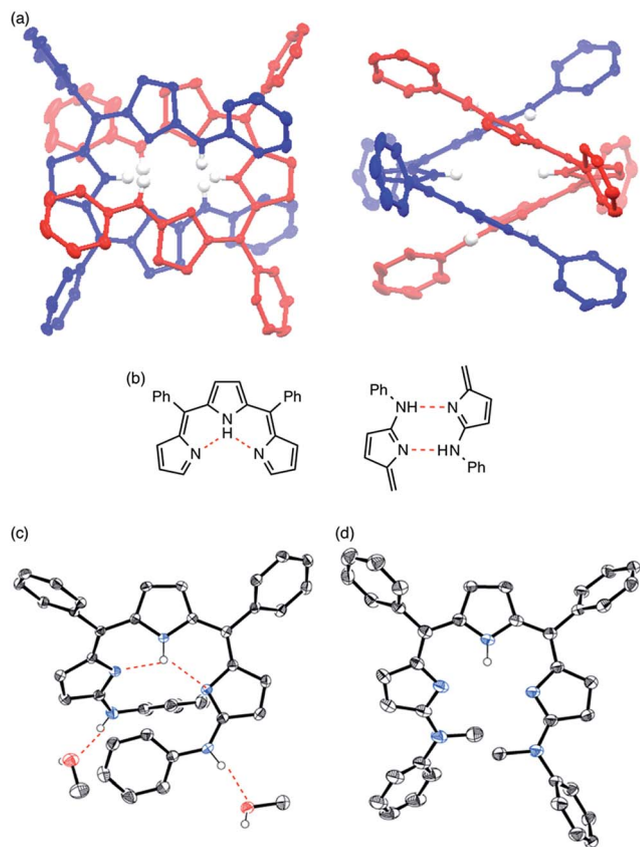


Fig. 2 (a) X-ray crystal structure of **4** obtained from CHCl_3/n -hexane. Representations for intramolecular hydrogen bonding and interstrand hydrogen bonding networks are shown in (b). (c) X-ray crystal structure of **4** obtained from $\text{MeOH}/\text{H}_2\text{O}$. (d) X-ray crystal structure of **11**. Solvent molecules and hydrogen atoms except for NHs and OHs are omitted for clarity. The thermal ellipsoids are scaled to 50% probability.

hydrogen bonding interaction among pyrrole-NH and imine-type pyrroles (Fig. 2b, left). When grown from a protic solvent system such as a mixture of methanol and H_2O , single crystals of **4** showed a monomeric helical structure featuring intermolecular hydrogen bonding interactions between the aniline-NH and methanol (Fig. 2c). While doubly helical structures were also observed in crystals of substituted anilinothripyrrins **5**, **6**, **7** and **8** (Fig. S5-4–S5-8; ESI[†]),¹⁵ N,N' -dimethylanilinothripyrrin **11** showed a monomeric structure without any intermolecular hydrogen bonding with solvent molecules (Fig. 2d). Therefore, it is obvious that the two aniline NH segments and the tripyrrin segment are both crucial to form the double stranded helix. The helical pitches are calculated to be 5.44–5.88 Å.¹⁶ The pyrrole planes are smoothly conjugated with torsion angles along the helical twist of 3.92–45.29°. It should be noted that tripyrrins **9** and **10** could not form double helical structures probably due to serious steric hindrance around the aniline NH segments. Detailed solid-state structures of **9** and **10** will be discussed later.

NMR analysis

Solvent-dependent structural changes were also evident in the ^1H NMR analysis. The ^1H NMR spectrum of **4** in CDCl_3 at room

temperature showed two sets of signals, namely monomer and dimer, whose ratio was concentration dependent (Fig. 3b). The ^1H NMR spectrum of **4** in a hydrogen-bonding-accepting polar solvent, $\text{DMSO-}d_6$, showed clear signals featuring three doublets at 6.77, 6.53 and 5.97 ppm due to the pyrrolic β -protons and two singlets at 13.06 and 9.55 ppm due to the tripyrrin-NH and aniline-NH protons, respectively (Fig. 3c), which can be assigned as a monomeric species. On the other hand, the ^1H NMR spectrum of **4** in a nonpolar and aprotic solvent, cyclohexane- d_{12} , exhibited sharp peaks but different spectral features: a quartet at 6.39 ppm and a doublet at 5.94 ppm due to the β -protons and two singlets at 12.41 and 11.95 ppm due to the tripyrrin-NH and aniline-NH protons, respectively (Fig. 3a), which can be assigned as a dimer. Therefore, collectively, the ^1H NMR spectra of **4** in CDCl_3 indicate a slow equilibrium between the monomeric and dimeric forms of **4**. Diffusion ordered spectroscopy (DOSY) in CDCl_3 supports this assignment (Fig. S3-35; ESI[†]). The association constant K_{dim} for dimerization of **4** at 25 °C in CDCl_3 was determined to be $2.7 \times 10^2 \text{ M}^{-1}$ by ^1H NMR spectral analysis. In contrast, the ^1H NMR spectra of **11** were not significantly changed by solvents. The ^1H NMR

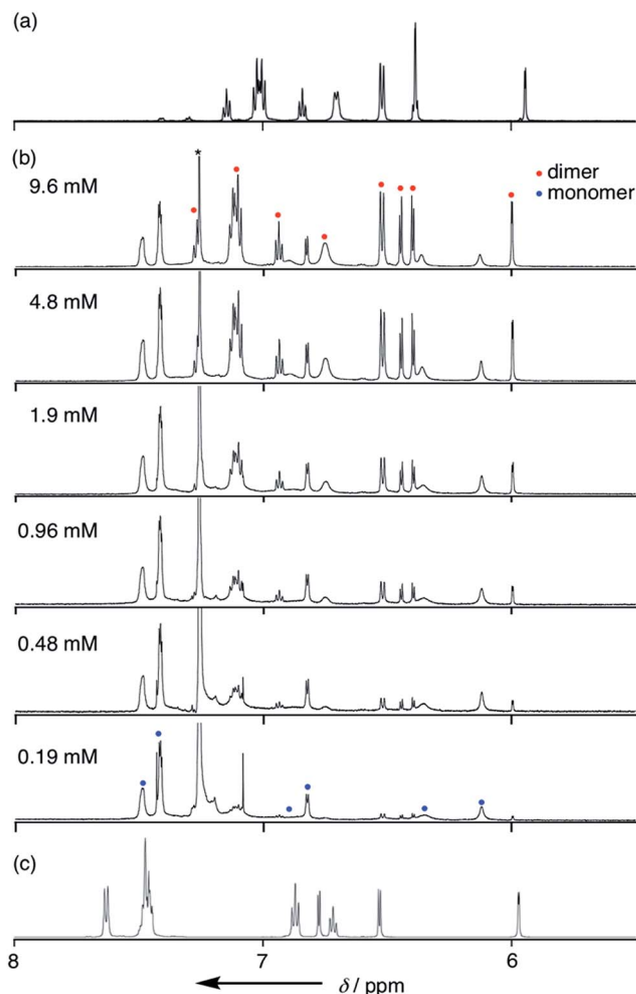


Fig. 3 ^1H NMR spectra of **4** (a) in cyclohexane- d_{12} , (b) in CDCl_3 with various concentrations and (c) in $\text{DMSO-}d_6$.



spectrum of **11** in CDCl_3 showed only a set of signals: a broad singlet peak at 6.4 ppm, doublet peaks at 6.21 and 6.68 ppm due to the β -protons and a singlet at 13.22 ppm due to the inner NH protons, indicating the predominant existence of the monomer in solution.

The dimerization equilibrium was also affected by temperature (Fig. S3-6; ESI†). At low temperature, the amount of monomeric **4** decreased and the amount of dimeric **4** increased, leading to a larger K_{dim} value. A van't Hoff plot of K_{dim} gave thermodynamic parameters, $\Delta H = -53.6 \text{ kJ mol}^{-1}$ and $\Delta S = -134 \text{ J mol}^{-1} \text{ K}^{-1}$, indicating that the dimerization is mostly enthalpy-driven.

Similar to **4**, the ^1H NMR spectra of **5**, **6**, **7** and **8** in CDCl_3 displayed concentration- and temperature-dependent equilibria between monomeric and dimeric species (Fig. S3-10, S3-11, S3-14, S3-15, S3-19, S3-20, S3-24 and S3-25; ESI†). The K_{dim} values of **5**, **6**, **7** and **8** at 25 °C in CDCl_3 were estimated to be 52, 58, 1100 and 4000 M^{-1} , respectively. The thermodynamic parameters of **5**, **6**, **7** and **8** were also determined and are summarized in Table 1. Among these, *meta*-substituted dianilinothripyrins **7** and **8** exhibited larger K_{dim} values and large negative ΔG values, while *para*-substituted dianilinothripyrins **5** and **6** were less amenable to form the double helices, as compared with non-substituted **4**. Fig. 4 displays the space-filling models of the solid-state structures of **7** and **8**. In both cases, aromatic stacking interactions are observed in the imine-like pyrrole moieties with average stacking distances of 3.53–3.60 Å for **7** and 3.48–3.52 Å for **8**. Additionally, the *meta*-substituents on the aniline moiety are in close proximity to the central pyrrolic plane to gain a certain thermodynamic

stabilization. In the case of **7**, however, the 3,5-di-*t*-butylphenyl groups are slightly tilted in order to have an appropriate arrangement to gain such stabilization, that likely resulted in some entropic losses.

Different conformations

Similar to **4**, the crystal structure of **9** obtained from acetone/ H_2O showed intramolecular and intermolecular hydrogen bonding interactions as shown in Fig. 5a. The intramolecular hydrogen bonding interactions are between the pyrrolic NH and the pyrrolic imine nitrogen atoms, and the intermolecular hydrogen bonding interactions are between the aniline-NH and solvent molecules (acetone). Curiously, the crystal structure of **9** obtained from an aprotic solvent such as a mixture of CH_2Cl_2 and *n*-hexane did not show a dimeric structure but displayed a tautomeric structure with different hydrogen-bonding interactions. Judging from the bond lengths and angles, one of the

Table 1 Summary of the thermodynamic parameters for **4**–**8** in CDCl_3

	K_{dim}^a	$\Delta H \text{ kJ mol}^{-1}$	$\Delta S \text{ J mol}^{-1} \text{ K}^{-1}$	$\Delta G_{298} \text{ kJ mol}^{-1}$
4	2.7×10^2	−53.6	−134	−13.7
5	5.2×10	−49.1	−131	−10.0
6	5.8×10	−47.1	−125	−9.85
7	1.1×10^3	−57.3	−133	−17.7
8	4.0×10^3	−56.1	−119	−20.6

^a $T = 298 \text{ K}$.

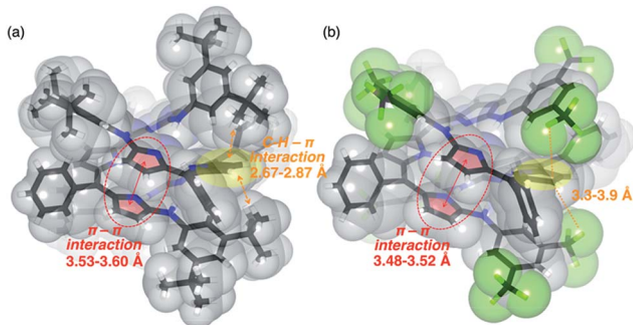


Fig. 4 Space-filling and stick representations of the solid-state structures of (a) **7** and (b) **8**. Solvent molecules included in the unit cell are omitted. Selected intermolecular short-contacts are shown.

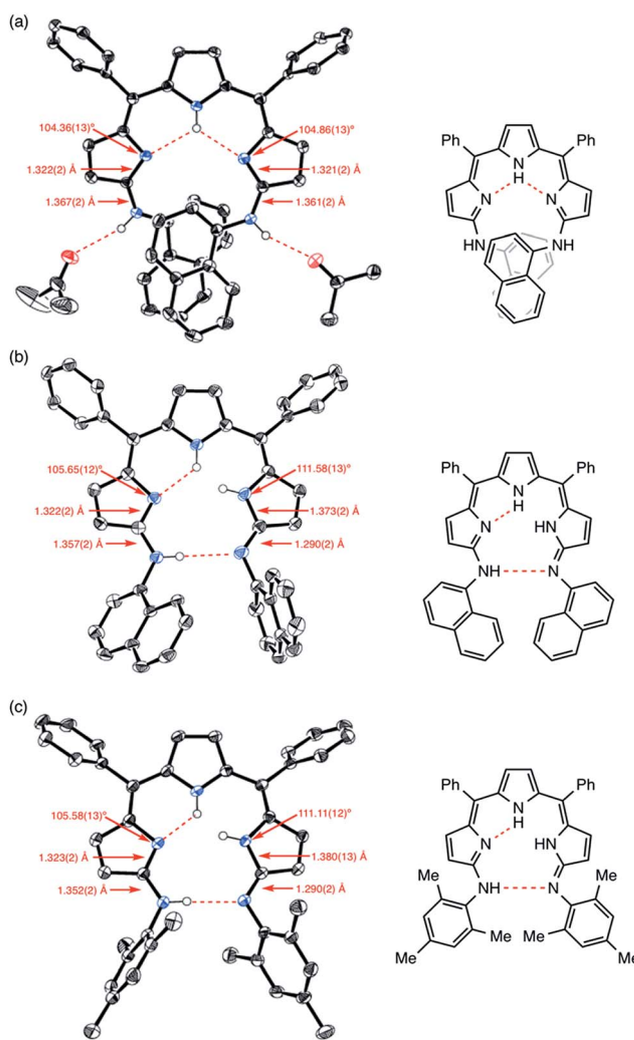


Fig. 5 X-ray crystal structures of (a) **9** obtained from acetone/ H_2O , (b) **9** obtained from $\text{CH}_2\text{Cl}_2/n$ -hexane and (c) **10** obtained from $\text{CH}_2\text{Cl}_2/n$ -hexane. Solvent molecules except for acetone and hydrogen atoms except for NHs are omitted for clarity. The thermal ellipsoids are scaled to 50% probability.



aryl-amino-substituted imine-type pyrroles was assigned as a different arylimino-substituted amino-type pyrrole tautomer (Fig. 5b). Thus, the overall structure is asymmetric, and two sets of intramolecular hydrogen bonding networks are formed. The same structure was observed in the crystal structure of **10** (Fig. 5c and S5-13–S5-16†). Presumably due to the steric hindrance of the 1-naphthyl and mesityl groups around the aniline-NHs, **9** and **10** were not able to form dimeric structures. In the case of **10**, even intermolecular hydrogen bonding with solvent molecules seemed to be prevented in the solid state (Fig. S5-15; ESI†).

The ^1H NMR spectrum of **9** in CDCl_3 showed broad peaks probably due to the slower tautomerism with regard to the ^1H NMR timescale. In contrast, the ^1H NMR spectrum of **9** in $\text{DMSO}-d_6$ showed sharp peaks featuring three doublets at 6.86, 6.82, and 5.98 ppm due to the β -protons and two singlets at 13.31 and 9.27 ppm due to the tripyrrin-NH and amine-NH, respectively. This signal pattern is similar to that of **4** measured in $\text{DMSO}-d_6$ although signals assignable to the naphthyl protons were largely shifted.¹⁷

Mixing of the two different strands in solution may give homo-sorted or hetero-sorted dimers. To test this sorting behaviour, to a CDCl_3 solution of **8** was added an equimolar amount of **7** or **10**. In the case of **7** and **8**, a new signal that can be assigned to the hetero-strand [**7** + **8**], was observed in the ^1H NMR spectrum (Fig. S3-40†). Interestingly, the peak intensity of [**7** + **8**] is larger than that of **7** or **8**, indicating a larger association constant of the hetero-strand.¹⁸ In contrast, a mixture of **10** and **8** only showed the superposition of the signal sets (Fig. S3-41†).

UV/Vis absorption spectra

The UV/Vis absorption spectra of **4–11** were measured in various solvents. These tripyrrin derivatives exhibit mainly two absorption bands around 400 nm and 570 nm and the absorption tails reach a deep visible region (~ 700 nm) due to the extended π -conjugation. These absorption features resemble those of porphyrinoids and natural chlorophyll analogs,²⁰ thus demonstrating promising applications as functional dyes. Indeed, the solution colour of **4** is vivid blue in CH_2Cl_2 owing to the large extinction coefficient (*ca.* $49\,000\ \text{M}^{-1}\ \text{cm}^{-1}$ at 398 nm and $14\,000\ \text{M}^{-1}\ \text{cm}^{-1}$ at 576 nm in CH_2Cl_2). In addition, the absorption spectra of **4** showed a slight shift in polar solvents (Fig. 6a and S6-2; ESI†). Tripyrrin derivatives **5–11** also display similar spectral features in some solvents, while the UV/Vis absorption spectra of **10** are more dramatically changed by solvent effects. In DMSO , **10** displays blue-shifted absorption bands at 539 nm. The degree of such shifts is likely solvent-polarity dependent, thus demonstrating a vivid colour change from blue to purple as shown in Fig. 6b.

Molecular orbital calculations have revealed that the effects of substituted aniline moieties are more significant in the HOMOs rather than the LUMOs (Fig. S7-1–S7-7; ESI†). Strong CT-like contributions are not evident in these systems. In **9** and **10**, the NH tautomers are proved to be the lowest-energy conformers when solvent molecules are not considered (Fig. S7-8 and S7-9; ESI†). Variable probabilities of this form in

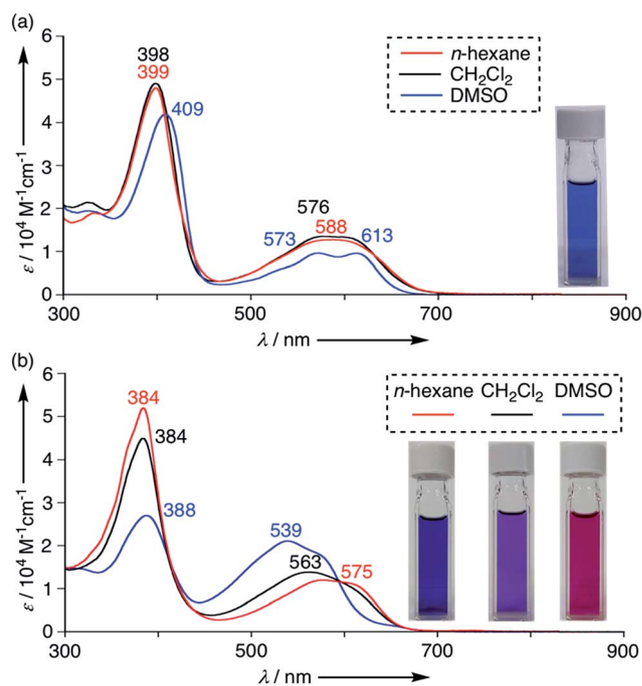


Fig. 6 UV/Vis absorption spectra of (a) **4** and (b) **10** in *n*-hexane (red), CH_2Cl_2 (black) and DMSO (blue). Concentrations of **4** are set at 1.0×10^{-5} M.¹⁹

solution may result in the observed solvent-dependent spectral shifts.

Conclusions

In summary, the nucleophilic substitution reactions of α,α' -dibromotripyrrin **3** with anilines gave dianilino-tripyrrins **4–11** in high yields. Dianilino-tripyrrins **4–8** form double helix structures in the solid states and in nonpolar and aprotic solvents. The dimerization constants are larger for **7** and **8** bearing 3,5-di-*t*-butylanilino- and 3,5-bis(trifluoromethyl)anilino-substituents, respectively, in which the dimeric structures are stabilized *via* the close contacts between the *meta*-substituents and the central pyrrolic π -plane as well as aromatic stacking interactions. In nonpolar and aprotic solvents, **9** and **10** carrying bulky substituents at the aniline moiety do not form dimers but form different tautomeric structures. These compounds retain π -conjugated networks as evinced by their absorption spectra and π -orbital calculations. Collectively, these α,α' -dianilino-tripyrrins may offer a unique platform for novel artificial double helix molecules. Further structural modifications and elongation of the helical structure are actively ongoing in our laboratory.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by JSPS KAKENHI Grant Numbers (JP26810021, JP18H03910 and JP18K14199). T. T. acknowledges



the TOBEMAKI Scholarship Foundation, Murata Science Foundation and Izumi Science and Technology Foundation. M. U. acknowledges a JSPS Fellowship for Young Scientists.

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- 14 For the characterization of α -anilino- α' -bromotripyrrin S1, see ESI.† We assume that two equivalents of aniline are necessary for one substitution because an equimolar amount of hydrogen bromide is generated.
- 15 The solid-state structures of anilinotripyrrin **5** for both the monomer and dimer were revealed by X-ray crystallographic analysis (Fig. S5-4 and S5-5; ESI†).
- 16 The helical pitch was defined as the distance between the two nitrogen atoms of the aniline segments.
- 17 Despite the different conjugation modes, the NH tautomerism does not cause a significant change in their electronic state in view of the MO aptitude (see Fig. S7 and S6; ESI†).
- 18 The ratio of **7** : **8** : [**7** + **8**] was roughly estimated to be 1 : 1.4 : 7.7 by considering the ¹H NMR peak intensities and symmetry.
- 19 In some solvents, the absorption intensity does not follow the Beer-Lambert law, implying a decent concentration effect even in 10⁻⁵ M order.
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