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Cyclobutene based macrocycles†‡

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Nanoscale macrocycles could have unusual magnetic, optical, and electronic properties when compared to their linear counterparts. Conjugated π -systems in unsaturated macrocycles are particularly interesting as they have no end groups that limit electronic delocalization in equivalent linear oligomers. The rigid four-membered ring structure in 3,4-bis(methylene)cyclobutene with vicinal connections provides a vertex with an angle slightly less than 90° , which promotes macrocycle formation. We report herein a facile high-yielding synthesis of a series of 3,4-bis(methylene)cyclobutene-based π -conjugation macrocycles. The structure–property studies reveal that the smaller macrocycles are rigid crystalline frameworks and display symmetrical conformations in solution. The electrochemical, photophysical and magnetic properties of these macrocycles were also studied with a framework of characterization methods, revealing their size- and linkage-dependent properties. Density functional theory (DFT) calculations and molecular dynamics (MD) simulations at the molecular level suggest that several possible configurations are possible for macrocycles with larger ring sizes.

Introduction

The concepts of supramolecular chemistry provide a natural translation of molecule constructions to create functional materials.^{1–3} Supramolecular assemblies containing extended π -conjugated macrocycles are particularly attractive for applications in organic electronics/devices,^{4,5} examples of note include cyclic porphyrins,^{6,7} annulenes,^{8–10} phenylenes^{11–15} and oligothiophenes.^{16–18} The most common building blocks of π -extended macrocycles are phenylenes, ethynylenes, thiophenes and pyrroles. Methods to achieve high yields of macrocycles include the use of templates,¹⁹ and post cyclization reactions are needed to release the macrocyclic products from the templates. Most macrocyclic precursors are rigid structures that create vertices with angular orientations larger than 90° at their reacting sites. An advantage of the cyclobutene structure is that it can create a vertex angle of about 90° which will be

more favorable for cyclic structures, but has rarely been explored in macrocycle synthesis.

As a highly strained and air-sensitive isomer of benzene, 3,4-bis(methylene)cyclobutene has drawn much attention since it was first synthesized in the 1960s.²⁰ The air-stable analogues were developed by Toda *et al.*, by the thermal conversion of diallenes in the crystalline state.^{21–23} The introduction of aromatic substituents on the exocyclic methylenes and electron-withdrawing groups into the cyclobutene skeleton provides improved stability. Four-membered ring compounds that contain a bis(methylene)cyclobutene base structure have been extensively studied,^{23–26} and we have been interested to extend this unique structure as a vertex in macrocyclic compounds with different conjugation linkages to produce a series of fully conjugated macrocycles with persistent defined shapes.

Herein, we report the efficient template-free synthesis of a series of cyclobutene-based π -conjugation macrocycles with different ring sizes in high yields benefiting from the 3,4-bis(methylene)cyclobutene core structure bearing angular orientations less than 90° . Two-dimensional nuclear magnetic resonance (2D) NMR and variable temperature (VT)-NMR analysis demonstrated that these macrocycles are highly symmetrical in solution and single crystal X-ray crystallography revealed the packing mode and the planarity of the main-chain carbon atoms. The synthesized macrocycles can be chemically oxidized and the formation of radical cations (that can provide polarons in the solid state) was monitored by electron paramagnetic resonance (EPR) spectroscopy. SEM images revealed the relatively rigid crystalline macrocyclic framework. The optical and

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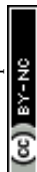
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† Dedication: To the inspiring example of Fred Wudl and his many pioneering
contributions in the design, synthesis and study of organic electronic materials.

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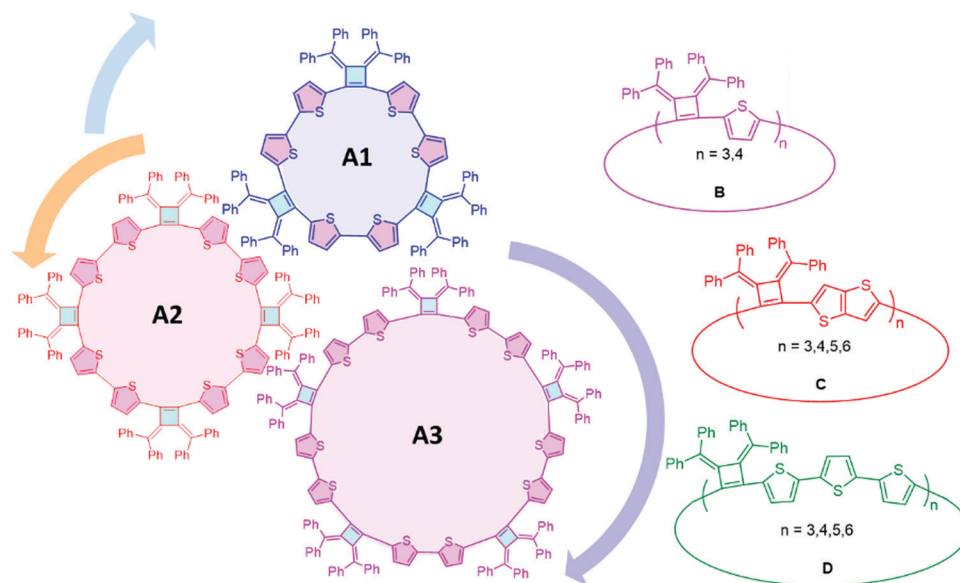


Fig. 1 Schematic structures of synthesized cyclobutene based macrocycles.

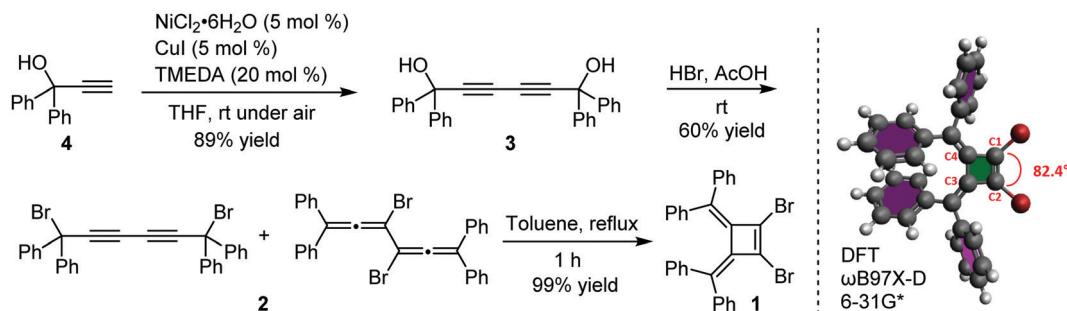
electrochemical properties were investigated, revealing the size- and linkage-dependent properties of these macrocycles (Fig. 1).

Materials design and synthesis

Our interest in the target macrocycles were guided by the computed geometries of 1,2-dibromo-3,4-bis(diphenylmethylene)cyclobutene **1**, using DFT (ω B97X-D/6-31G*). The optimized geometry displays notable intramolecular π - π interactions between the two inner phenyl rings on the two double bonds. The distance of C3–C4 is 1.52 Å, and the distance of C1–C2 is 1.35 Å (Scheme 1). The angles between C1–Br bond and C2–Br bond is computed to be 82° , a value similar to that obtained in a single crystal structure of an analogue.²² The rigidly fixed molecular geometry of **1** is poised to bias away from the formation of linear oligomers in cross-coupling sequences. This geometric relationship provides a rational and practical synthesis of macrocycles *via* cross-coupling reactions of 1,2-dibromo-3,4-bis(diphenylmethylene)cyclobutene with mono-thiophene, fused thiophene, bi-thiophene or tri-thiophene comonomers.

We synthesized 1,2-dibromo-3,4-bis(diphenylmethylene)cyclobutene **1** *via* a three-step synthetic route as shown in Scheme 1. Glaser-Hay oxidative coupling reaction of the

corresponding terminal alkyne **4** gave compound **3** in 89% yield. Following Toda's synthetic method,²² upon treatment of **3** with hydrobromic acid in acetic acid provided the brominated products **2** as a mixture. Thermal rearrangement in refluxing toluene afforded pure cyclobutene **1** in quantitative yield within 1 h. We explored cyclization reactions using Stille-coupling of cyclobutene **1** with 5,5'-bis(trimethylstannyl)-2,2'-bithiophene (**2THSn**). As shown in Table 1, using $[\text{Pd}(\pi\text{-cinnamyl})\text{Cl}]_2$ as the precatalyst and 'BuXPhos as the ligand, the Stille-coupling reaction in toluene after 24 h gave a 34% yield of soluble red solids and side products that are not soluble in common organic solvents (Table 1, entry 1). MALDI-TOF mass spectrometry analysis reveals that the soluble portion are primarily macrocycles with 3 repeating units. Changing the ligand from 'BuXPhos to XPhos increased the yield of the soluble portion to 43% (Table 1, entry 2). Microwave reaction conditions with a shortened reaction time to 4 h, increased the yield of soluble products to 83% (Table 1, entry 3). The MALDI-TOF mass spectrum analysis of macrocycle mixtures indicate the presence of four different ring-size macrocycles, namely the 3-mer **A1**, 4-mer **A2**, 5-mer **A3** and a trace of the 6-mer **A4** (From MALDI-TOF mass spectrometry: **A1** ($m/z = 1632.223$, $n = 3$), **A2** ($m/z = 2176.527$, $n = 4$),



Scheme 1 Synthesis and optimized structure of 1,2-dibromo-3,4-bis(diphenylmethylene)cyclobutene.



Table 1 Stille-coupling reactions to make macrocycles

Entry ^a	Ligand	Yield (%) ^b	MALDI-TOF ^c
1	^t BuXPhos	37	3-mer and other pieces
2	XPhos	43	3-, 4-, 5-, 6-mers
3 ^d	XPhos	83	3-, 4-, 5-, 6-mers

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^a Reaction conditions: 0.15 mmol scale, 0.05 M, under N₂ atmosphere. ^b Yield of the soluble mixture of macrocycle portion. Macrocycles were purified by passing through a pad of Celite and precipitated from MeOH. ^c 2-[(2*E*)-3-(4-*tert*-Butylphenyl)-2-methylprop-2-enylidene]malononitrile (DCTB) was used as the matrix in MALDI-TOF mass spectrometry with a PEG3000 calibration standard. ^d The reaction was carried out in a microwave reactor at 110 °C for 4 h.

A3 ($m/z = 2720.659$, $n = 5$), **A4** ($m/z = 3265.815$, $n = 6$). The observed isotope distributions of all the different ring-size macrocycles from MALDI-TOF mass spectrometry are in agreement with the calculated values.

The relative constitutions of macrocycles **A1**, **A2** and **A3** were quantified and isolated using high-performance liquid chromatography (HPLC) analysis (Fig. 2c). **A4** was present only in trace quantities and wasn't studied in detail. UV-vis spectroscopic analysis (Fig. 2d) shows that **A1** has the maximum absorption band at 426 nm, whereas **A2** is red shifted to 440 nm as a result of its longer effective conjugation length. **A3** shows an absorption band at 427 nm similar with **A1**, suggesting a twisted configuration of the larger ring. Our observations suggest that there is no full delocalization of the whole π -system as the ring size increases, thus limiting the conjugation length in the ring. The molar absorption coefficients of the three macrocycles are around $7.0 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ to $8.2 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ at 427 nm, and the relative concentrations in the soluble reaction products were roughly estimated based on the integrated peak areas from HPLC analysis. As can be seen in Fig. 2c, the HPLC trace area with monitoring at 430 nm indicates the relative percentages of **A1**, **A2**, **A3** to be 55.2%, 26.4% and 18.4% respectively. All of the macrocycles are emissive when dissolved in tetrahydrofuran (THF) and show a similar emission profile around 630 nm (Fig. 2e and f).

¹H NMR spectra of the three isolated macrocycles **A1–A3** indicate these macrocycles are highly symmetric in solution that only one set of proton signals of the repeating unit was observed. The combined information from 2D NMR heteronuclear single quantum coherence (HSQC) and heteronuclear multiple bond correlation (HMBC) analysis, allows for the assignment of all the proton signals (Fig. S1–S3, ESI[†]). The protons on thiophene ring display different chemical shifts for

each ring-size macrocycle (Fig. 3). For example, as shown for **A1**, the proton **H^a**, which is the closest to the cyclobutene ring, is shielded with a chemical shift at 5.67 ppm, whereas the proton **H^b** is de-shielded and resonates at 6.58 ppm. The different chemical shifts of the different ring-size macrocycles are indicative of the differences in their electronic and conformational structures.

X-ray crystal structure and self-assembly

The structure of macrocycle **A1** was elucidated by X-ray crystallography. The single crystal was prepared by vapor diffusion of methanol into a chlorobenzene solution of **A1**. The single crystal structure (Fig. 4a) reveals a macrocycle in an isosceles triangular shape with cyclobutene rings are located at each vertex and thiophene rings on sides. The internuclear distances of C13–C2 and C14–C7 are 9.152 Å and 9.162 Å respectively, while the distance of C1–C8 is 8.894 Å. The structure produces a large inner cavity with potential applications in host–guest chemistry.^{2,27} The bithiophene connector favors a *trans* configuration on the two longer sides of the isosceles triangle, with a *cis* configuration providing the reduced length on the other side. It is noteworthy that we do not observe ¹H NMR resonances for the different conformations in solution even under low temperatures (0 °C to –90 °C in VT-NMR, Fig. S4, ESI[†]), indicating that either there is one single conformer in solution, or there is a very rapid interconversion of the two isomers. The conformations also prevent the main-chain atoms of the macrocyclic structure from being in plane of the isosceles triangle. The root-mean-square deviation (RMSD) of the main-chain carbon atoms from the calculated plane is 0.172 Å. The phenyl rings on the cyclobutenes are twisted out of plane to reduce steric repulsions to each other. The two inner phenyl rings twist to promote face-to-face π -stacking with a distance of



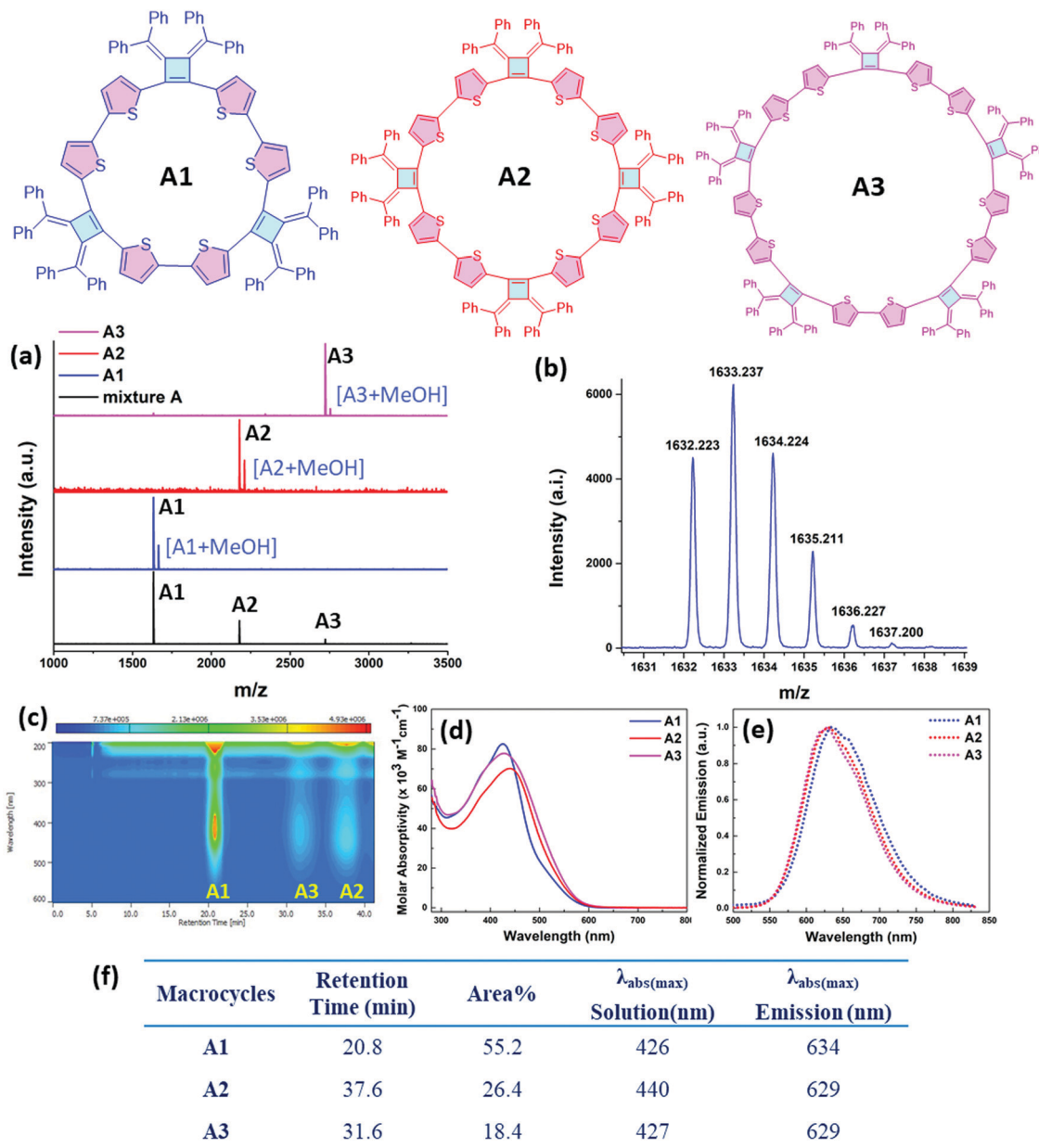


Fig. 2 (a) MALDI-TOF mass spectrometry data of macrocycle mixtures **A** and macrocycles **A1**, **A2** and **A3** after HPLC separation. *trans*-2-[3-(4-*tert*-Butylphenyl)-2-methyl-2-propenylidene]malononitrile (DCTB) was used as the matrix with PEG3000 as a calibration standard. (b) Isotope distribution of **A1** in the MALDI-TOF mass spectrometry. (c) 2D contour map UV-vis spectrum for the HPLC separation in heptane/THF using a ZORBAX-CN prep-column. (d) UV-vis spectra and (e) normalized emission spectra of the isolated macrocycle **A1**, **A2** and **A3** in THF solutions. The emission spectra were measured after excitation at the maximum of each absorption wavelength. (f) A summary table of the HPLC separation results, and photophysical properties of the compositions in macrocycle **A** mixtures.

3.419 Å between the planes of the phenyls (the distance between C615 and the mean plane of C511, C513 and C515) (Fig. 4a and Fig. S12a, ESI†). **A1** forms stacked dimers with a distance of 4.223 Å between the planes of the isosceles triangles (Fig. 4c). These dimers further organize in a 1-D 'slipped-stacked' fashion with a distance of 9.454 Å along *a* axis (Fig. S12b, ESI†).

Macrocycle **A1** crystallizes more readily than the other macrocycles and displays interesting self-assembled morphologies when grown from different solvents.²⁸ As visualized by scanning electron microscopy (SEM), crystals of **A1** tend to form cylinders in

various diameters when grown in chlorobenzene (Fig. 4d) and form fibrous structures when grown in THF (Fig. 4e). These fibrils have a smaller diameter that assemble into balls.

Macrocycles using linkers with different conjugation lengths

Linkers adjoining the cyclobutenes play an important role in controlling structure, effective conjugation length and photophysical properties. Our synthetic method affords modular access to macrocycles with monothiophene, bithiophene, fused thiophenes and terthiophene linkages in good to excellent



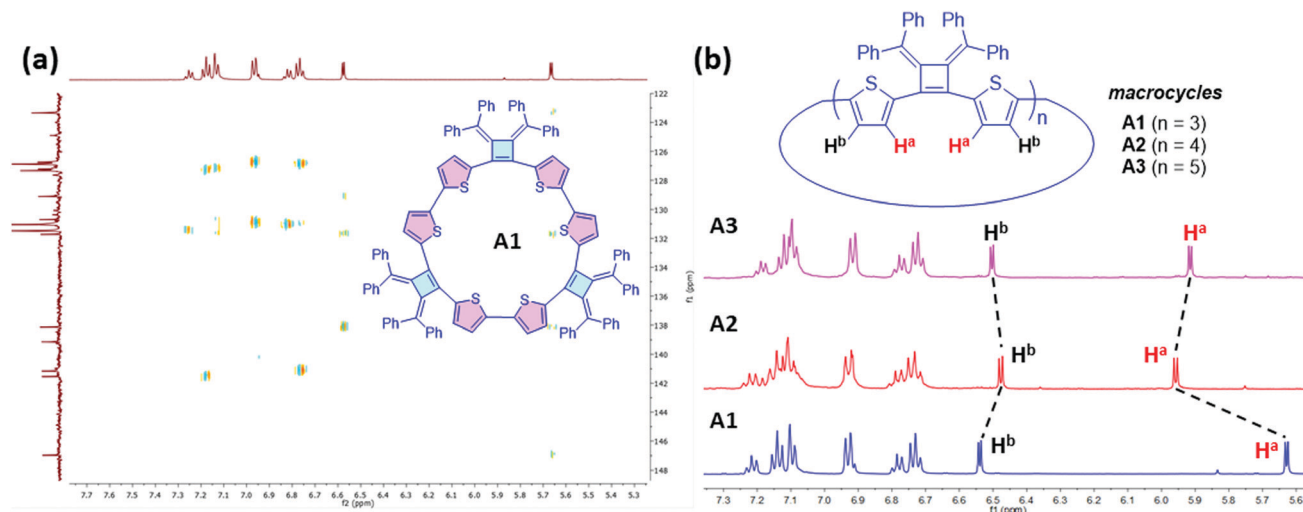


Fig. 3 Structural elucidation of Macrocycles **A1**, **A2** and **A3** by NMR spectroscopy. (a) 2D NMR HMBC (500 MHz, 25 °C, THF- d_6) spectrum of **A1**. (b) ^1H NMR (500 MHz, 25 °C, THF- d_6) spectra of **A1**, **A2** and **A3**.

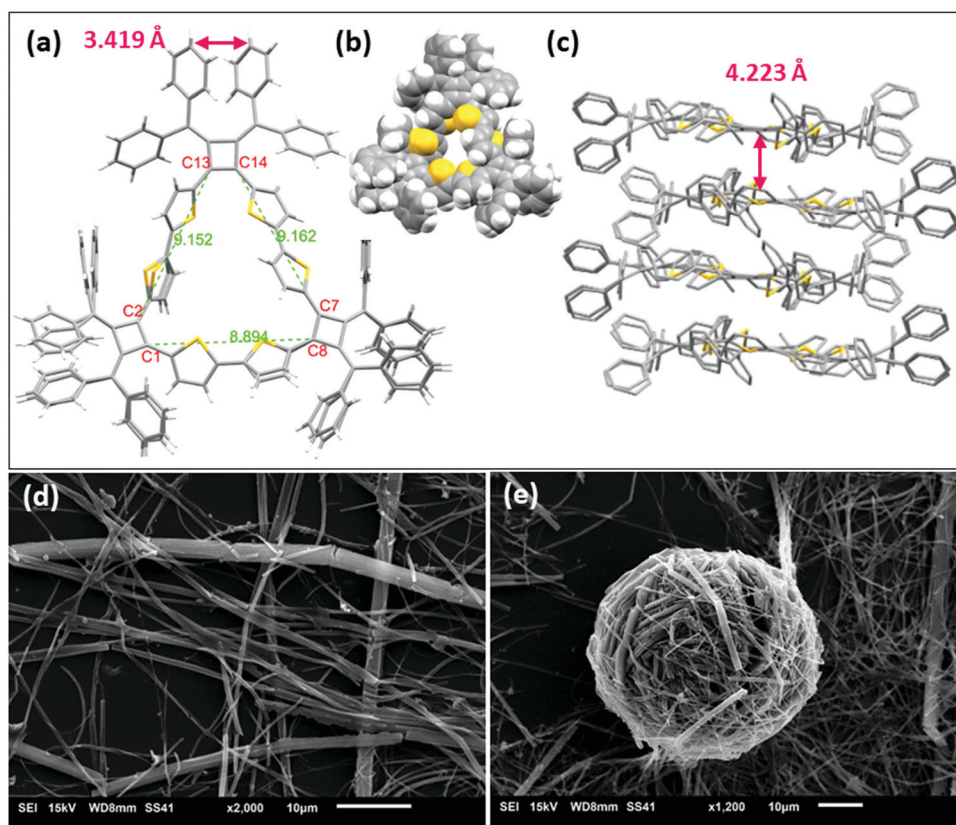
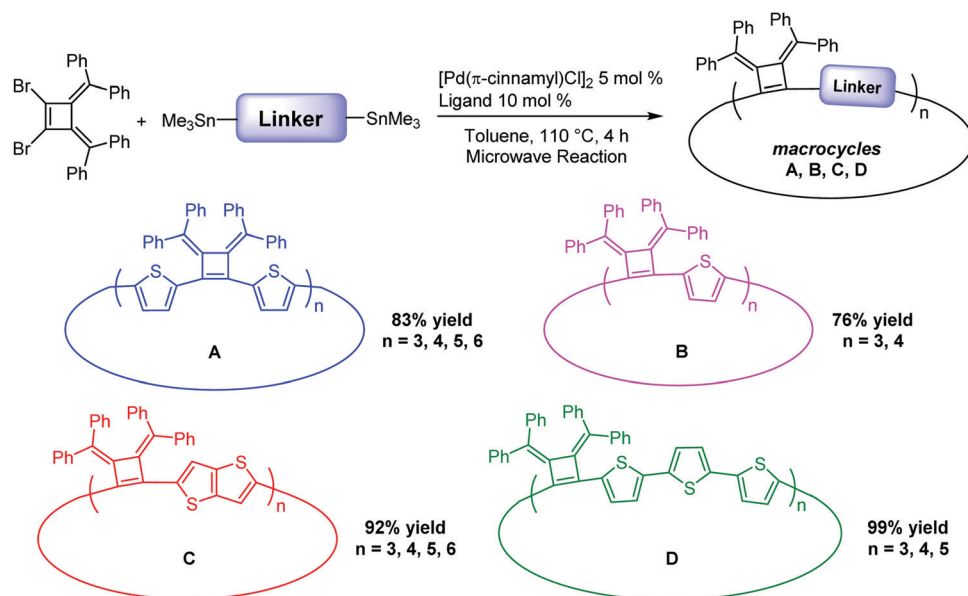


Fig. 4 X-ray crystal structure of macrocycle **A1**, (a) top view, stick model; (b) space filling model; (c) side view of the stacked model; (d) SEM images of crystals of **A1** grown in chlorobenzene and (e) crystal fibrils of **A1** grown in THF.

yields (Table 2). Single thiophene linkages produce macrocycles **B** with 3 to 4 repeating units; fused thiophene rings give macrocycle **C** with 3 to 6 repeating units; and terthiophenes produce the largest macrocycle **D** with 3 to 5 repeating units. Their compositions contain 3- to 6-mers as confirmed by MALDI-TOF mass spectrometry (Fig. S5–S7, ESI†).

As summarized in Table 1, thermal gravimetric analysis (TGA) reveals that the macrocycles display high thermal stability with initial decomposition temperatures ranging from 328–374 °C (Fig. S9, ESI†). Differential scanning calorimetry (DSC) analysis of the macrocycle mixtures shows no obvious thermal transitions between 100 °C to 350 °C as expected as a result of the

Table 2 Synthesis of other macrocycles with different linkages



Macrocycles ^a	Yield ^b (%)	T _d ^c (°C)	λ _{abs(max)} solution ^d (nm)	λ _{em(max)} emission ^e (nm)	λ _{abs(max)} film ^f (nm)	Quantum yield ^g (%)
A	83	374	422	628	445	5.9
B	76	339	352	688	380	6.5
C	92	328	387	660	383	7.6
D	99	339	440	603	440	7.3

^a Standard reaction conditions: 0.15 mmol scale, 0.05 M, under N₂ atmosphere. ^b Yield of the soluble products. The macrocycles were purified by passing through a pad of Celite and precipitated from MeOH. ^c Initial decomposition temperature, onset temperature in TGA analysis. TGA analysis was conducted in N₂ atmosphere with a ramping rate 20 °C min⁻¹ to 900 °C. ^d In THF solution. ^e The excitation is at the maximum absorption wavelength. ^f Spin-coated films on glass slides with macrocycles dissolved in THF solution. ^g Quantum yields were measured in THF solutions with 1,4-bis(5-phenyloxazole-2-yl)benzene (POPOP) as the standard.

mixture of macrocycles and conformations, thereof appears to produce an amorphous state.

The macrocycle mixtures show different UV-vis absorption behaviors, which are related to their effective conjugation

Effective conjugation length in macrocycles:

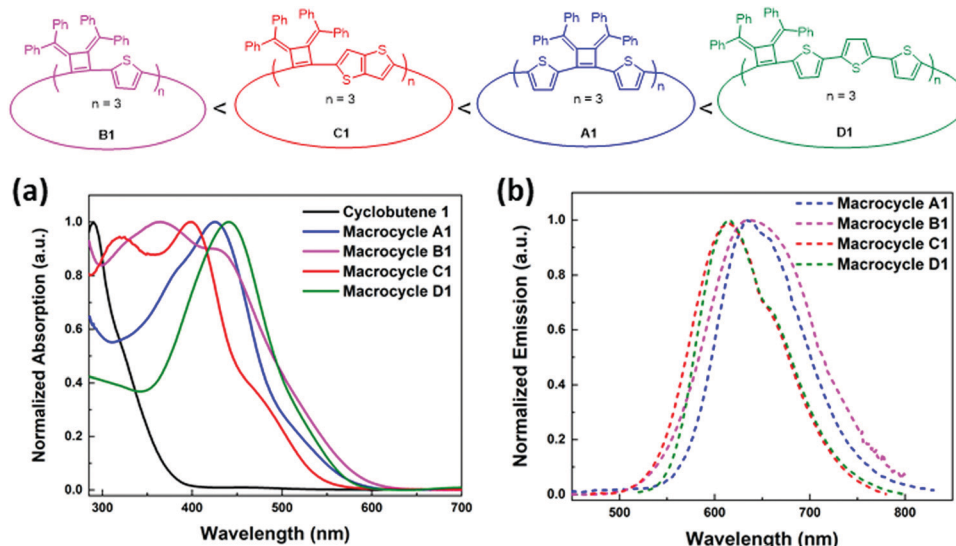


Fig. 5 Normalized (a) UV-vis spectra and (b) the emission spectra of the isolated macrocycles **A1**, **B1**, **C1** and **D1** in THF solutions. The emission spectra were measured after excitation at the maximum of each absorption wavelength.



lengths (Fig. S8, ESI†) and modest emissive quantum yields ranging from 5.9% to 7.6%. Macrocycles with three repeating units have been separated successfully by HPLC prep-column, giving macrocycle **A1** (3 repeating units), **B1** (3 repeating units), **C1** (3 repeating units) and **D1** (3 repeating units) as single pure products. Their UV-vis and emission spectra were also studied in THF solutions. As shown in Fig. 5, **D1** with a terthiophene linker has the longest conjugation length with absorption and emission maximum at $\lambda_{\text{abs(max)}}$ of 440 nm and $\lambda_{\text{em(max)}}$ of 615 nm respectively, while **A1** displays a $\lambda_{\text{abs(max)}}$ of 426 nm and a $\lambda_{\text{em(max)}}$ of 634 nm. **C1** displays two absorption bands at 319 nm and 397 nm, and its emission spectrum shows a $\lambda_{\text{em(max)}}$ of 634 nm and a shoulder peak at 658 nm that are related to the absorption and emission of two different transition states. Similarly, **B1** presents two absorption bands at 365 nm and 428 nm, and a broad emission band around 637 nm. The difference in conjugation lengths observed in these macrocycles with the same repeating units (three repeating units) could be attributed to the effective conjugation between the cyclobutene monomer and the different linkers. The conformations between the linkers and the cyclobutenes may vary and therefore affect the conformations in macrocycles. Based on the trends in the λ_{max} values, it appears that the length of the linking heterocycles dominates the conformation structures: the terthiophene linker in **D1** having the longest conjugation length and the single thiophene of **B1** having the lowest degree of conjugation.

The synthesized macrocycles readily absorb I_2 to create “doped” samples. Thin films were prepared by spin coating with macrocycles in THF solution (5 mg mL^{-1} , 1500 rpm) and were then exposed to iodine vapor in a sealed chamber for 60 h at room temperature. After doping, the $\lambda_{\text{abs(max)}}$ of the thin film

of macrocycles mixtures **A** shifts from 445 nm to 395 nm, and a new strong absorption band arises at 793 nm that is related to the absorption band of polaronic or bipolaronic states of the macrocycles²⁹ (Fig. 6a). The iodine treated samples also display an induced electron paramagnetic resonance (EPR) signal in solution phase (Fig. 6b). Upon adding excess amount of iodine to the dry toluene solution of macrocycle **A1**, a broad EPR spectrum was observed with a g -value of 2.0089, indicating the formation of polarons that have localized spin distribution in the macrocycles.^{30,31} Its spin intensity decreased over time, as the solution undergoes to higher doping levels, creating bipolaronic states (with charges of $+4e$ and $+6e$) which are spinless.

To further study the redox behavior of these macrocycles, cyclic voltammetry (CV) was conducted on the macrocycle mixtures. A model compound **RP1** with two single thiophenes attached to the cyclobutene was synthesized by Suzuki coupling reaction. The CV of **RP1** dissolved in dichloromethane (DCM) solution produced that one quasi-reversible redox wave at $E_{1/2} = 0.48 \text{ V}$ (vs. Fc/Fc^+) (Fig. S10a, ESI†). The CV of macrocycles **A1** spin-coated on the indium tin oxide (ITO) glass displays a single redox wave at $E_{1/2} = 0.40 \text{ V}$ (vs. Fc/Fc^+) similar to **RP1** (Fig. S10b, ESI†). The CVs of the macrocycle mixtures of **A**, **B**, **C** and **D** were also studied as thin films on ITO glass and all had quasi-reversible redox waves at slightly lower potentials and thereby establishing the semi-conductive potential of these materials (Fig. S10c, ESI†).

Computational studies of macrocycles

In order to further understand the structural and electronic properties of these macrocycles, a series of DFT calculations were performed on **A1** ($n = 3$), **B1** ($n = 3$), **C1** ($n = 3$), and **D1** ($n = 3$). For each macrocycle, starting from an initial estimate of

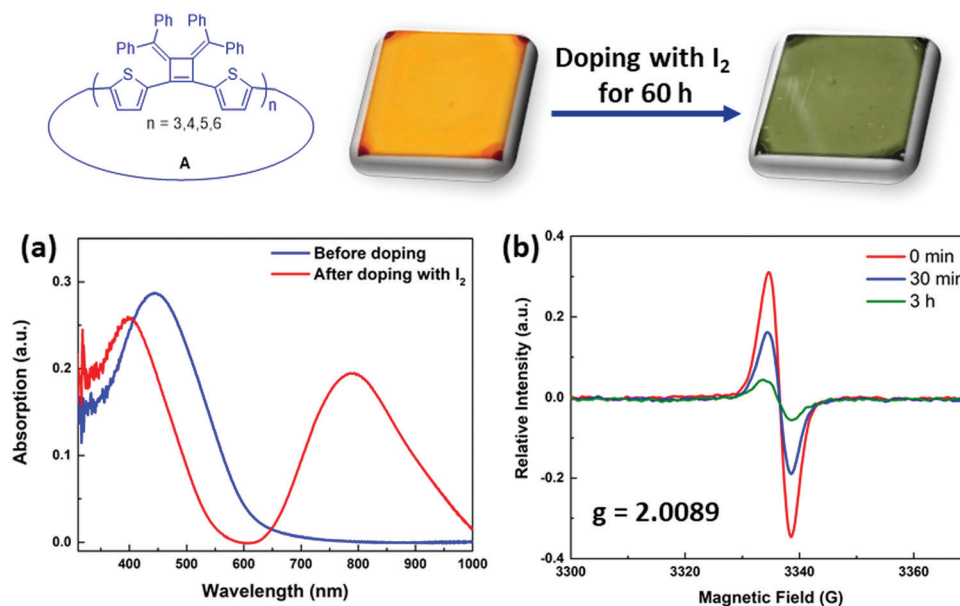


Fig. 6 Iodine vapor doping with thin films prepared by spin-coating (5 mg mL^{-1} , 1500 rpm). Macrocycle **A** was dissolved in THF and spin-coated on glass slides. (a) UV-vis absorption with thin films before and after doping with iodine in a sealed chamber for 60 h. (b) EPR spectra of macrocycle **A1** doping with iodine in dry toluene solution.

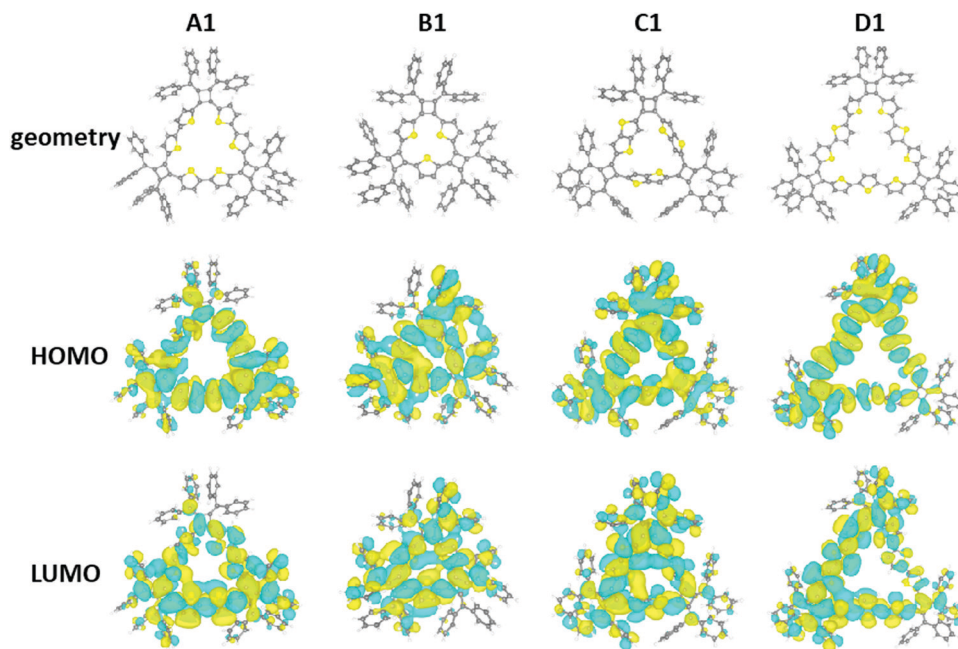


Fig. 7 DFT results: structure and frontier orbitals of the **A1**, **B1**, **C1**, and **D1** compounds. In the structural models, yellow, grey, and white spheres respectively correspond to sulfur, carbon, and hydrogen atoms.

the molecular structure, the geometry was optimized in order to minimize the total energy and atomic forces using the Vienna *Ab Initio* Simulation Package (VASP),^{32,33} and the Perdew–Burke–Ernzerhof (PBE) functional.³⁴ In a subsequent step, the electronic structure was determined at a higher level of theory, using the ORCA program³⁵ with the PBE0 hybrid functional.³⁶ All details of the DFT calculations are summarized in the ESI.†

The DFT results are presented in Fig. 7. The structure of the **A1** macrocycle differs slightly from the crystal structure, and the computed structure has all thiophene rings oriented towards the center of the molecule. This is not the case for all macrocycles and in **D1** the thiophene units show alternating orientations. The macrocycles all share two main geometrical features: (1) the inner phenyl rings stack face-to-face with π – π interactions, and (2) the triangular ring formed by the thiophene groups is corrugated. Computed structures reveal similar orientations of the outer phenyl groups of the X-ray crystal structure. Both the HOMO and LUMO orbitals of the macrocycles are significantly delocalized throughout the π -conjugated network for all macrocycles. Finally, the calculated HOMO–LUMO gaps of the **A1**, **B1**, **C1**, and **D1** molecules are 2.40, 2.57, 2.54, and 2.42 eV, respectively. It is important to note that the calculated HOMO–LUMO gap is the difference between the vertical ionization potential and the vertical electron affinity, which is typically larger than the actual band gap in the condensed phase as a result of interactions between π -conjugated molecules.³⁷ The DFT calculations are in agreement with the experimental observation of these macrocycles. MD simulations for larger macrocycles were also performed at the molecular level in the gas phase, which suggest a few possible configurations of the isolated macrocycles exist at room temperature (300 K). Here the calculated configurations for macrocycles with higher

repeating units ($n > 3$) demonstrated a tendency to buckle and fold (Fig. S14, ESI†).

Conclusion

In summary, a series of cyclobutene-based π -conjugation macrocycles were synthesized by a facile template-free modular method using a cyclobutene building block that provides a vertex angle of less than 90° . The synthesized macrocycles were found to be highly symmetrical in solution by ^1H NMR and VT-NMR. X-ray crystallography of one macrocycle ring confirmed the structure and provided insights into the conformations. DFT calculations and MD simulations suggest that several possible configurations are possible for macrocycles with larger ring sizes. The redox activity and nature of the charged species were investigated by cyclic voltammetry and EPR measurements. Our results suggest that the cyclobutene geometry provides facile access to a number of cyclic topologies by incorporating different linkages for the creation of new organic electronic materials.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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