

Strain-Induced Stereoselective Formation of Blue-Emitting Cyclostilbenes

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Supporting Information

INTRODUCTION

stilbene subunits.

ABSTRACT: We describe the synthesis of two conjugated macrocycles that are formed from the end-to-end linking of stilbenes. We have named these macrocycles cyclostilbenes. The two cyclostilbene isomers created in this study differ in the configuration of the double bond in their subunits. These macrocycles are formed selectively through a stepwise reductive elimination from a tetraplatinum precursor and subsequent photoisomerization. Single-crystal X-ray diffraction reveals the formation of channel architectures in the solid state that can be filled with guest molecules. The cyclostilbene macrocycles emit blue light with fluorescence quantum yields that are high (>50%) and have photoluminescence lifetimes of ~0.8–1.5 ns. The breadth and large Stokes shift in fluorescence emission, along with broad excited-state absorption, result from strong electronic–vibronic coupling in the strained structures of the cyclostilbenes.

Here we describe a new conjugated belt, formed by linking stilbene subunits into macrocycles, i.e., a cyclostilbene. The molecules in this study are members of a growing class of conjugated macrocycles,^{1,2} such as cyclo-*para*-phenylenes (CPPs),^{3–5} cyclothiophenes (CTs),^{6–8} and cycloporphyrins (CPs).^{9–11} The advent of synthetic methods to form strained

conjugated macrocycles over the past few years is enabling the

formation of new cyclic structures that heretofore were difficult

to make. The three main approaches to making strained conjugated macrocycles are (1) using statistical oligomerization of precursors with curvature,^{7,12,13} (2) synthesizing an un-

strained cyclic precursor followed by an efficient reaction that introduces strain,^{4,14,15} or (3) using a template to aid in the formation of the macrocycle.^{9,16} A wealth of new materials can

result from these new types of cyclic conjugated oligomers that have well-defined sizes and shapes. For example, conjugated macrocycles are useful as supramolecular nanostructures, such as columnar tubes or inclusion complexes.^{17–20} The design

strategy to yield interesting structural, optical, and electronic

properties for cyclic conjugated macrocycles is different than

that for linear conjugated molecules.²¹ Within this context, we

investigated the formation of cyclostilbenes by the end-to-end

linking of stilbenes into macrocycles. We show in Figure 1 three

stereoisomers, 1-CTCT, 2-CCCC, and 3-TTTT, that differ in

the number of trans (T) and cis (C) double bonds in the



Figure 1. Structures of cyclostilbenes 1-CTCT, 2-CCCC, and 3-TTTT calculated at the $B3LYP/6-31G^{**}$ level.

optical and electronic properties of stilbene are extremely sensitive to both the environment and the conformation of the stilbene.^{23,24} Large cyclophanes linked with unsaturated bridges have been extensively studied for their electrochemical properties, and strong diamagnetic ring currents were observed in their dianions.^{25,26} However, their low-yielding and nonselective syntheses^{27–29} precluded their being studied fully. In this article, we describe a new selective synthesis of two cyclostilbenes, **1-CTCT** and **2-CCCC**. **1-CTCT** has alternating *cis* and *trans* double bonds and forms stereoselectively from the four-fold reductive elimination of a tetranuclear square-planar platinum macrocycle. **2-CCCC** can be formed quantitatively by UV irradiation of **1-CTCT**. Our approach gives precise control over the final product structures. We find that these molecules are strong blue-emitting materials that can act as supramolecular hosts and pack in tubular structures in the solid state.

Stilbene is a prototypical subunit used in many conjugated oligomers and polymers, such as poly(phenylenevinylene),²² yet it has not been studied extensively in cyclic structures. The

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RESULTS AND DISCUSSION

Synthesis. Scheme 1 shows the method developed to synthesize cyclostilbene 1-CTCT. We based our strategy on

Scheme 1. Synthesis of Macrocycle 1-CTCT^a



^aReagents and conditions: (a) *i. n*-BuLi, THF, -78 °C, 3 h; *ii.* SnMe₃Cl, -78 °C to rt, 12 h, 78%. (b) Pt(COD)Cl₂, 1,2-dichloroethane, 75 °C, 20 h, 84%. (c) PPh₃, toluene, 100 °C, 24 h, 28%.

those of Yamago and co-workers for the synthesis of CPPs⁴ and Bäuerle and co-workers for the synthesis of CTs.³⁰ Here, we utilize (*E*)-4,4'-bis(trimethylstannyl)stilbene in combination with Pt(COD)Cl₂ (COD = 1,5-cyclooctadiene) to form the square tetranuclear platinum complex 5 in 84% yield. Figure S13 shows the crystal structure of 5, confirming that all four stilbene units retain the *trans* geometry of the olefin. Reductive elimination (upon addition of 10 mol equiv of PPh₃) forms the cyclostilbene in 28% yield. The direct 4-fold reductive elimination would yield cyclostilbene 3-TTTT; however, the macrocycle that we isolate is 1-CTCT, in which two of the double bonds have isomerized to the *cis*-olefin. The reaction is stereoselective, in that we do not isolate or detect any of the other stereoisomers with a different arrangement of *cis* and *trans* double bonds. Under UV irradiation, 1-CTCT quantitatively converts to isomer 2-CCCC. This isomer had been previously synthesized in a poor yield (~1%) via a four-fold Wittig reaction.³¹

We isolate both 1-CTCT and 2-CCCC as light yellow solids. Unlike their unsubstituted linear counterparts,^{32,33} cyclostilbenes are soluble in common organic solvents such as dichloromethane (DCM), tetrahydrofuran (THF), and chloroform (CHCl₃). They are insoluble in toluene and methanol (MeOH). Both isomers are air-stable and can be kept in the ambient for extended periods of time without change. Neither stereoisomer (1-CTCT or 2-CCCC) shows any mass loss up to 400 °C (for the thermogravimetric analysis data, see the Supporting Information, Figures S8 and S9). The two isomers behave similarly upon electrochemical reduction. We measured their reduction potentials using cyclic voltammetry (CV) in THF solution with Bu_4NPF_6 (0.1 M) as the electrolyte. The cyclostilbenes show reversible reduction events, with a reduction onset potential of -1.26 V for 1-CTCT and -1.28 V for 2-CCCC versus the ferrocene/ferrocenium (Fc/Fc^{+}) redox couple (see Supporting Information, Figure S18). The calculated lowest unoccupied molecular orbital (LUMO) levels are essentially the same for 1-CTCT and 2-CCCC.

Crystal Structures. We were able to grow single crystals suitable for X-ray diffraction for both isomers. After extensively screening solvent combinations, we found that **1-CTCT** forms crystals by layering of toluene on a THF solution of the macrocycle and that **2-CCCC** forms crystals by layering of MeOH on a DCM solution of the macrocycle. Figure 2a shows the top view of cyclostilbene **1-CTCT**. The macrocycle crystallizes as a 1:1 inclusion complex, **1-CTCT**-toluene. The molecule adopts an ellipsoidal shape, with the long axis (19.1 Å) nearly twice the length of the short axis (9.7 Å). The two stacked *trans*-stilbene moieties deviate away from planarity by ~19° [C—C=C—C dihedral angle = 161.0(11)°]. The angles at the sp² carbons of the *cis* and *trans* double bonds are slightly expanded to between 121.8(11)° and 128.3(10)°. The biphenyl units have a wide range of Ar—Ar twist angles between



Figure 2. Single-crystal X-ray diffraction structure of (a) 1-CTCT toluene and (b) 2-CCCC from top view. Disordered DCM solvent molecules are removed for clarity. Packing structure of (c) 1-CTCT along the (100) and (302) planes and (d) 2-CCCC along the (001) and (102) planes. Hydrogen atoms and solvent molecules are removed.



Figure 3. Structures and relative energies of cyclostilbene isomers calculated at the B3LYP/6-31G** level. Energy values are in kcal/mol, and compared with the lowest energy isomer 2-CCCC.



Figure 4. Aromatic region of the ¹H NMR spectra of two reaction intermediates (a) $6-Pt_3$ and (b) $7-Pt_2$. (c) Proposed mechanism of stepwise reductive elimination to stereoselectively form isomer 1-CTCT. Hydrogen atoms in (c) and phosphine ligands in all structures are removed for clarity. Red, platinum; black, carbon; gray, hydrogen.

 $21.3(17)^{\circ}$ and $51.5(15)^{\circ}$. The packing of **1-CTCT** in the solid state is similar to that of [8]-, [9]-, [10]-, and [12]-CPP.¹⁹ The molecules stack in tubular structures along the *a*-axis direction, and the tubes pack in a herringbone pattern (Figure 2c).

Isomer 2-CCCC has a different shape and a smaller inner cavity than 1-CTCT. As shown in Figure 2b, the macrocycle is square-shaped. The cavity measures 7.9 Å from the edge of one phenyl to the edge of the other phenyl across the macrocycle (Figure 2b). The cavity is filled with disordered DCM molecules from the crystallization. The angles at the sp² carbons of the *cis* double bonds are more expanded than 1-CTCT to between $129.5(2)^{\circ}$ and $133.1(2)^{\circ}$. The biphenyl units are nearly coplanar, and the Ar–Ar dihedral angles range from $9.2(3)^{\circ}$ to $26.3(3)^{\circ}$. Similar to 1-CTCT, channel architectures form in the packing of 2-CCCC, as the molecules

stack with their cavities over each other along the *c*-axis direction. The channels are smaller in 2-CCCC than in 1-CTCT because for 2-CCCC the molecules slip stack with respect to their neighbors. This slip stacking can be seen in the alternating layers in Figure 2d. The channels in 2-CCCC are also more continuous than in 1-CTCT, which caused several challenges in the crystal structure determination. DCM easily diffuses out of the channels, which causes the crystals to shatter rapidly when in contact with oil. The continuous solvent-filled channels also cause extensive crystallographic disorder.

The π -systems around the peripheries are oriented differently in **1-CTCT** and **2-CCCC**. The p-orbitals are aligned essentially perpendicular to the plane of the macrocycle in **2-CCCC**, whereas they are aligned radially in **1-CTCT** (excepting those of its two *cis* C=C bonds). This orientation difference is

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controlled by varying the geometry of the double bond in the macrocycle and plays an important role in both the strain energies and the excited-state properties of the molecule, as discussed in the following sections.

Stereoselective Isomerization. Although examples of conjugated macrocyclic isomers containing cis (C) or trans (T) olefins are known, the stereochemistry is built in during their synthesis, using typically Wittig or McMurry reactions.^{26,34-36} Our observation of isomerization from a TTTT precursor to give stereoselectively a CTCT isomer is unusual. We used density functional theory (DFT) calculations to gain insight into the different cyclostilbene isomers. Figure 3 shows the optimized geometries and relative energies of all six possible cyclostilbene isomers. The CCCC geometry is the lowest in energy, and we use its total energy as the zero benchmark. The next most stable configuration is CTCT, which is 5.6 kcal/mol higher in energy, followed by CCCT. The least stable isomer, TTTT, is 32.7 kcal/mol higher in energy than the all cis macrocycle. We note that for 1-CTCT, there exists two conformers: one where the two trans-stilbene units mirror each other (bottom structure in Figure 3), and one where they exhibit C_2 symmetry (top structure in Figure 3). DFT calculations indicate that they are isoenergetic (see Supporting Information).

We estimated the ring strain energies for 1-CTCT and 2-CCCC using isodesmic comparisons,³⁷ and found values of 17.0 kcal/mol for the former and 0.1 kcal/mol for the latter (see Supporting Information). The large contrast between the differences in energies of the macrocycles (5.6 kcal/mol) and their strain energies (16.9 kcal/mol) highlights the greater ability of the *cis* olefin to support the ring shape. For a stilbene monomer, the *cis* isomer is more sterically hindered and higher in energy than *trans*, whereas in the macrocycle the *trans*stilbene subunit needs to bend its π -system to accommodate the curvature of the belt structure. Therefore, the larger strain in 1-CTCT accounts for the 5.6 kcal/mol difference mentioned above, as well as the intrinsically higher energy of the *cis*stilbene subunit in the acyclic isodesmic partners.

Uncatalyzed, thermal *trans*-to-*cis* isomerization is known for overcrowded ethylene systems, where steric effects lower the activation barriers due to destabilization of strained ground-state conformations.³⁸ Similarly, it is possible that the highly strained **3-TTTT** isomer is initially formed from direct reductive elimination of Pt macrocycle **5**, and then isomerizes to a more stable isomer **1-CTCT** to reach an energy minimum. However, this cannot explain the fact that **1-CTCT** is formed exclusively in the reaction and not a thermodynamically controlled ratio of different products. Therefore, instead of a direct 4-fold reductive elimination to yield **3-TTTT**, it is more likely that the stereoselective isomerization occurs during the multi-step reductive elimination process.

In order to follow the cascade of reactions that lead from **5** to the ultimate product **1-CTCT** and further understand the stereoselective isomerization process, we monitored the reaction in $C_2D_2Cl_4$ by ¹H NMR. Figure 4 shows the downfield region ($\delta = 6.5-7.0$ ppm) of the ¹H NMR spectrum that is remote from the aryl phosphine resonances. Within this region, we are able to observe the partial spectra of two intermediates during the reaction. Upon addition of PPh₃ and heating to 50 °C, three resonances emerge that have equal integrated intensities, suggesting that it is not simply the ligand-exchange product containing four platinum centers, in which the resonances would integrate 2:2:1. We assign this new species to the fan-shaped intermediate **6**-**Pt**₃, in which one platinum center has been eliminated. In this structure, there are two types of stilbene units present—*stilbene* I and *stilbene* II—and there are two of each type. The two *stilbene* I units are those that had been bonded to the now-eliminated Pt center, and they are now bonded to one another to give the arced arm of **6**-**Pt**₃. Each of the two *stilbene* II units retains both of its Pt termini, forming a straight arm of **6**-**Pt**₃. The phenyl resonances in *stilbene* II are shifted upfield to the stilbene olefin region due to PPh₃-ligated Pt(II), whereas in *stilbene* I the phenyl protons are shifted downfield, outside the clean spectrum window. The seemingly simple spectral pattern can be deconvoluted by COSY and HSQC experiments (Figures S14 and S15), and Figure 4a displays the assignment of the resonances.

When we heat 6-Pt₃ to 90 °C, we observe the formation of a second intermediate, as well as the fully eliminated product 1-CTCT (Figure 4b). This second intermediate has only one resonance in this window of the spectrum, suggesting the highly symmetrical structure 7-Pt₂. The observed resonance corresponds to both olefin protons. Each of the phenyl protons is buried underneath the phosphine region as those of *stilbene II* in 6-Pt₃. Consistent with reductive elimination from platinum biaryls being a first-order reaction,^{39,40} both the decay of 6-Pt₃ and the formation of 1-CTCT over time fit to a single exponential (Figures S16 and S17).

Therefore, by combining the NMR experimental observations and DFT calculations (see Supporting Information), we propose a stepwise platinum reductive elimination mechanism for the stereoselective formation of 1-CTCT (Figure 4c). There is a great amount of curvature in the trans-stilbene units in 7-Pt2, and the formation of an additional aryl-aryl bond would introduce even more distortion, making reductive elimination unfavorable compared to other processes such as homolytic cleavage to give linear oligomers instead of the desired cyclic product.^{30,41} The other pathway for $7-Pt_2$ is thermal isomerization of the trans double bonds to the cis conformation. The stereochemical isomerization occurs because this isomerization pattern gives rectangular-shaped 8-Pt₂ that is the most favorable pathway to release strain from 7-Pt₂. From this structure, reductive elimination proceeds rapidly to form cyclostilbene 1-CTCT, and 8-Pt, is not observable on the NMR time scale.

Photophysics. Figure 5a displays the UV-vis absorption and fluorescence emission spectra for 1-CTCT and 2-CCCC. A striking feature is that both cyclostilbenes show broad blue emission and large Stokes shifts. Table 1 summarizes the photophysical data for 1-CTCT and 2-CCCC. The maximum absorption of cyclostilbene 1-CTCT shifts slightly to the red from that of 2-CCCC. Time-dependent DFT (TD-DFT) calculations indicate that the highest occupied molecular orbital (HOMO)→LUMO transition is symmetry forbidden in both molecules. This is similar to what is observed for CPPs and other symmetrical, conjugated macrocycles.⁴² The absorption peak in the UV region arises from a $\pi - \pi^*$ excitation that is a combination of HOMO−1→LUMO and HOMO→LUMO+1 transitions with a high oscillator strength (f). Our calculations indicate that this transition occurs from primarily the olefin (the *trans* olefin in the case of 1-CTCT) to its π^* orbital (Figure S23). The lower-energy shoulder peak corresponds to the HOMO \rightarrow LUMO transition (arrow in Figure 5a). This is due to dynamic conformational changes in solution that break symmetry and result in nonzero *f* values.



Figure 5. (a) UV-vis absorption spectra of 1-CTCT (blue, solid line) and 2-CCCC (red, solid line), 1×10^{-5} M in DCM. Fluorescence spectra of 1-CTCT (blue, dashed line) excited at 340 nm and 2-CCCC (red, dashed line) excited at 320 nm, 1×10^{-6} M in DCM. The arrow indicates the shoulder peak that corresponds to the HOMO→LUMO transition. (b) Normalized photoluminescence decay kinetics of 1-CTCT (blue) and 2-CCCC (red) in DCM solution and their fits (black).

Table 1. Photophysical Data for 1	I-CTCT and	l 2-CCCC
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	absorption	emission			
	$\lambda_{\rm max}/{\rm nm}$	$\lambda_{\rm max}/{ m nm}$	$\Phi_{\mathrm{F}}^{\ a}$	$ au_{\mathrm{F}}/\mathrm{ns}$	$k_{\rm r}^{\ b}/10^8 \ {\rm s}^{-1}$
1-CTCT	344	436	0.56	1.26	4.44
2-CCCC	322	450	0.55	0.78	7.18
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^{*a*}Coumarin 1 ($\Phi_{\rm F}$ = 0.79 in air) was used as a reference for fluorescence quantum yield calculations (see Supporting Information). ^{*b*}Radiative decay rate constants from the excited singlet state, given by the equation $\Phi_{\rm F} = k_{\rm r} \tau_{\rm F}$.

Both 1-CTCT and 2-CCCC show high photoluminescence quantum yields (PLQYs) above 50%, similar to [9]- and [10]-CPP.^{42,43} The high PLQYs for the cyclostilbenes are surprising because the $S_0 - S_1$ transition is dipole forbidden, with minimal or no oscillator strength in the S₁ state. Calculations⁴⁴ revealed that, in large CPP molecules, S_1 strongly couples with vibrational modes, resulting in a self-trapped localized excitonic state (S_1^*) with a nonzero dipole transition that facilitates strong emission. Similarly, the emission from the cyclostilbenes also comes from S_1^* state (Figure 6c) with high PLQYs. The absorption and emission processes taking place from different electronic states also explains the large Stokes shift for these cyclostilbenes. The structures of 1-CTCT and 2-CCCC in their excited states are highly strained. This strain results in a strong coupling of the excited states with the vibrational modes yielding the vibronic bands observed in the emission spectra. Figure S18 shows the emission spectra of 1-CTCT that is fitted by three Gaussian functions with the peaks centered at 2.706,



Figure 6. Transient absorption spectra at various pump-probe delays (a) and exciton dynamics at different probe wavelengths (b) for 1-CTCT upon 325 nm excitation. (c) Schematic illustration of optical transition and relaxation in cyclostilbenes. The energy levels for S_1 and S_2 obtained from the calculations (see Supporting Information), but not for S_1^* . A and Em represent absorption and emission, respectively. Red arrows depict the excited-state absorption. Q is the coordinate.

2.886, and 3.049 eV. The differences in energy between these peaks are 0.18 and 0.163 eV, corresponding to the phenylene stretching vibration mode.⁴⁵

Time-resolved fluorescence decay kinetics for 1-CTCT and 2-CCCC dissolved in DCM are shown in Figure 5b. Their excited-state decay kinetics are similar to a slightly longer lifetime for 1-CTCT. The values are of fluorescence quantum yield (Φ_F), fluorescence lifetime (τ_F), and radiative decay rates (k_r) for both cyclostilbenes are summarized in Table 1. The radiative decay rates for both macrocycles are similar, and this is not surprising given that k_r is closely related to the oscillator strength.⁴⁶ It is known that excited alkenes can undergo adiabatic isomerization.⁴⁷ To examine this, we encapsulated both molecules in polystyrene film and measured the timeresolved PL decay kinetics as a function of temperature (Figure S21). The PL decay kinetics of both molecules show very little dependence on temperature (down to 88 K), precluding the isomerization process for these macrocyles.

Transient Absorption Spectroscopy. To further study the role of electron-phonon coupling in these molecules, we performed transient absorption spectroscopy on 1-CTCT. Figure 6 shows differential transmission $(-\Delta T/T)$ spectra and exciton dynamics for 1-CTCT in CHCl₃ upon 325 nm excitation. Interestingly, we observed a very broad excited-state absorption from the visible to near-IR range. The dynamics at different wavelengths are described by the same lifetime of 1500 ± 40 ps (Figure 6b), indicating that they originate from a single species. The large structural change suggested by fluorescence measurements can also explain the broad transient absorption feature, as illustrated in Figure 6c. The optically bright S₂ state from initial optical excitation can rapidly relax to the twisted S_1^* state. The broad transient absorption from S_1^* to S₂ can be attributed to the large offset in equilibrium nuclear coordinates, i.e., large reorganization energy, as illustrated by the red arrows.

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CONCLUSION

In summary, we have revealed a method to stereoselectively synthesize two members of the cyclostilbene family of conjugated macrocycles. This unusual reaction sequence is programmed by the four-fold reductive elimination of Pt centers to produce the strained, conjugated macrocycle. 1-CTCT has not previously been reported and would be difficult to make by traditional methods. This synthetic method now allows these two macrocycles to be studied that differ only in the configuration of the stilbene double bonds. We have shown here that it is possible to create materials that have different channel architectures in the solid state and different excitedstate geometries. The cyclostilbenes are suitable hosts to form inclusion complexes, which augurs well for the creation of optoelectronic devices whose properties can be varied by changing the guest that is included. The molecules begin to fill the void in studies of the material properties of conjugated macrocycles. The macrocycles emit blue light with high quantum yields due to their rigid, cyclic configuration providing strong coupling between their excited states and vibrational modes. This study also leaves a challenge to find a method to create the all trans-cyclostilbene that has heretofore been unattainable.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.5b06258.

Synthetic procedures, ¹H and ¹³C NMR spectra, TGA analysis, NMR kinetic experiments, PLQY measurements, transient absorption measurements, and computational details (PDF)

X-ray crystallographic data for 2-CCCC (CIF)

X-ray crystallographic data for 1-CTCT (CIF)

X-ray crystallographic data for 5 (CIF)

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Notes

The authors declare no competing financial interest.

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