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Luminescent chiral triangular prisms capable of forming double helices for detecting traces of acids and anion recognition[†]

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Orchestrating stimuli-responsiveness, molecular recognition, and self-assembly in macrocycles plays a vital role in the preparation of multifunctional materials. Herein, we report a pair of triangular prismatic enantiomers (*R*)- and (*S*)- Δ based on tetraphenylethylene (TPE), which assemble into left- and right-handed double helices, respectively, in their single-crystal X-ray superstructures. Due to the geometrical constraint of triangles to the phenylene rotation within TPE, their emission is enhanced significantly. In addition, their fluorescence and circularly polarized luminescence change from blue to yellow in response to traces of acids in solvents, which, combined with enhanced intensity, makes them sensitive indicators for the quantitative detection of the acid present in common organic solvents such as CDCl₃, CH₂Cl₂, and CHCl₃, with a detection limit as low as 5×10^{-7} M. Furthermore, the cavity of triangles can bind l₃⁻ to form host–guest adducts by single-crystal-to-single-crystal transformation.

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Introduction

Multi-purpose application requirements have brought challenges to the design of functional molecules. By embracing macrocyclic chemistry, especially in relation to molecular recognition and selfassembly, chemists can design sophisticated host molecules with multifunctionality. Thanks to their rigid geometries and enforced cavities, shape-persistent macrocycles, such as calixarenes,^{1–3} cucurbiturils,^{4,5} pillararenes,⁶ and cycloparaphenylenes,^{7,8} as well as various cages,^{9–19} have been investigated extensively. Among them, molecular triangles^{20–23} have exhibited considerable potential in storage, sensing, separation, catalysis, *etc.* on account of their triangular cavities being able to encapsulate selectively guest molecules of matching size and stereoelectronic properties.^{24–29} In addition, the small internal angle inside the triangles enforces interactions between their side units enhancing³⁰ some of their properties.

Aggregation-induced emission (AIE) of tetraphenylethylene (TPE) has been applied widely in fluorescent probes,^{31,32} bioimaging agents,^{33,34} chemical sensors,^{35–37} and circularly polarized luminescnece (CPL).^{38,39} The D_{2h} symmetry and tetratopic reaction positions of TPE make it an ideal building block for constructing macrocycles and cages.^{40–44} Employing TPE as an organic linker enhances significantly the luminescence of these molecules as a result of the restriction of the phenylene rotation by either geometrical constraints or guest complexation.^{45–48} Taking advantage of the enhanced emission intensity of TPE, chiral macrocycles and cages possess hierachical structural complexity and functionality involving chiral recognition and sensing, CPL signal, and chiral superstructures.^{16,49}

Herein, we report (Fig. 1) a pair of TPE-incorporated enantiomeric molecular triangles (*RRRRRRRRRR*)- Δ [(*R*)- Δ] and (*SSSSSSSSS*)- Δ [(*S*)- Δ], which, as confirmed by single-crystal X-ray diffraction (SCXRD), can (i) assemble into left-(*M*) and righthanded (*P*)-double helices, respectively, as a result of chirality transfer. (ii) The enhanced emission intensity of the geometrically constrained TPE, combining with (iii) the trace acid-responsive emission, makes them (iv) sensitive indicators for the detection of the amount of acid in organic solvents to a limit of 5×10^{-7} M. Moreover, their triangular prismatic cavities can also (v) bind triiodide anions (I₃⁻) to form host–guest adducts through a singlecrystal-to-single-crystal (SCSC) transformation.

Results and discussion

Enantiomeric (*R*)- and (*S*)- Δ were synthesized (Fig. 1) by a [3+6] cycloimination between tetrakis(4-formylphenyl)ethylene (1) with (1*R*,2*R*)- or (1*S*,2*S*)-diaminocyclohexane (2) in 88% and 90% yields, respectively. Their formation was confirmed by ¹H

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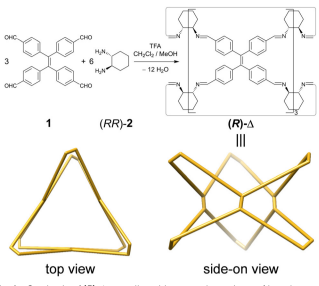


Fig. 1 Synthesis of (R)- Δ as well as side-on and top views of its schematic representation.

and ¹³C NMR spectroscopies (Fig. S1–S4, ESI†), high-resolution mass spectrometry (Fig. S5, ESI†), and FT-IR spectroscopy (Fig. S6, ESI†).

Single crystals of (*R*)- and (*S*)- Λ suitable for X-ray crystallography were obtained by slow liquid–liquid diffusion of MeOH into their solutions in CHCl₃. SCXRD Analysis revealed (Fig. 2a and Table S1, ESI[†]) their triangular prismatic geometry wherein three TPE units are linked, in a compact manner, by six diaminocyclohexanes to form a triangular channel (diameter = ~ 4 Å), thus preventing the phenylene rotation of TPE. In the X-ray superstructure (Fig. 2b) of (*R*)- Δ , adjacent triangles bite together through complementary van der Waals interactions (Fig. S7, ESI[†]) between the sides of the triangular prisms to form a (*M*)-single helix. Furthermore, two single helices with complementary configurations are intertwined (Fig. 2c) to form stereocontrolled (*M*)- and (*P*)-double helices from (*R*)- and (*S*)- Δ respectively, wherein each pitch is composed of eight triangles with a length of 5.00 nm and a diameter of 2.5 nm.

The photophysical properties of (*R*)- and (*S*)- Δ in their crystal states were first investigated under ambient conditions. Upon irradiation with 365 nm UV light, block crystals of both (*R*)- and (*S*)- Δ exhibit green-yellow emission (Fig. S8, ESI†). As shown in Fig. S9 (ESI†), their steady-state photoluminescence spectra showed a green-yellow emission band at around 530 nm with lifetime of 4.59 and 4.49 ns (Fig. S12 and Table S2, ESI†) as well as quantum efficiencies of 27.7% and 26.2% (Table S3, ESI†) for (*R*)- and (*S*)- Δ , respectively. When cleaning a flask containing (*R*)- Δ using commercial grade CH₂Cl₂, we discovered serendipitously that the fluorescence emission of the solution upon irradiation with 365 nm-UV light changed from the original blue to yellow. This observation inspired us to measure the fluorescence spectra of (*R*)- Δ in commercial grade CH₂Cl₂ and CHCl₃ at concentrations as low as 1 × 10⁻⁴, 1 × 10⁻⁵, and

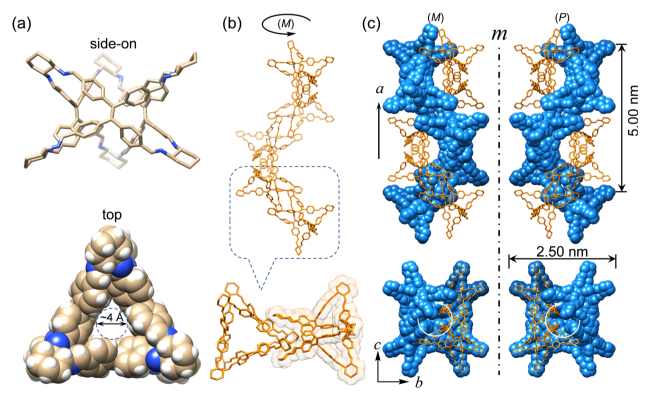


Fig. 2 Single-crystal X-ray (super)structures of triangles. (a) Side-on (tubular) and top views (space-filling) of (\mathbf{R}) - Δ . C, tan; N, blue, H, white. (b) (M)-single helix (top) assembled through complementary bite (down) between two adjacent (\mathbf{R}) - Δ . (c) Mirror-symmetrical (M)- and (P)-double helices formed by entangling two single helices assembled from enantiomeric (\mathbf{R})- and (\mathbf{S})- Δ , respectively. One strand in both double helices is depicted in orange tubular representation, while the other in blue space-filling. Hydrogen atoms are omitted for the sake of clarity.

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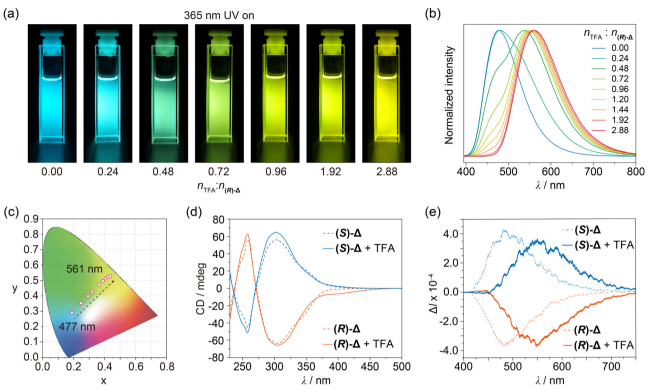


Fig. 3 Photophysical properties of (**R**)- and (**S**)- Δ . (a) Luminescence photographs taken upon shinning 365 nm UV light. (b) Fluorescence spectra upon excitation at 380 nm. (c) Trajectory of tuning colors shown in the CIE coordinate diagram of solutions (1×10^{-5} M) of (**R**)- Δ in CHCl₃ containing different amounts of TFA relative to (**R**)- Δ ranging from 0 to 2.88 equiv. (d) CD and (e) CPL spectra of solutions (2×10^{-4} M) of (**R**)- and (**S**)- Δ in deacidified CHCl₃ (dash line) and in acidified CHCl₃ with TFA (2.88×10^{-5} M, solid line).

 1×10^{-6} M, which exhibit (Fig. S10, ESI⁺) a clear trend in emission color from blue (\sim 480 nm) to yellow (\sim 560 nm) upon decreasing concentrations. We believe that the change in emission color might result from traces of acid in these solvents. Next, we measured the fluorescence spectra of (R)- Δ in deacidified CH₂Cl₂ and CHCl₃ with K₂CO₃ at the same concentrations and discovered that they show no change in emission, confirming that traces of acid in solvents can change the emission of (R)- Δ . We performed a series of photophysical characterizations to compare properties of (R)- and (S)- Δ in deacidified and quantitatively acidified CHCl₃, respectively. UV-Vis Spectra of solutions $(1 \times 10^{-5} \text{ M})$ of (R)- and (S)- Δ in deacidified CHCl₃ show (Fig. S11, ESI[†]) characteristic peaks at 272, 310, and 345 nm, while the same solutions exhibit (Fig. 3a) bright blue emission upon irradiation with 365 nm-UV light. Fluorescence spectra recorded on the same solutions demonstrate (Fig. 3b) an emission band at \sim 477 nm upon excitation at 380 nm, which has (Fig. S13 and Table S4, ESI⁺) a typical fluorescence feature with lifetimes of 2.29 and 2.52 ns and high quantum efficiencies of 25.5% and 23.4% (Table S5, ESI⁺) for (*R*)- and (*S*)- Δ , respectively. The significantly enhanced emission efficiencies can be ascribed to the restriction of the phenylene rotation within triangles.

Enantiomeric (*R*)- and (*S*)- Δ in deacidified CHCl₃ lead (Fig. 3d) to mirror-symmetrical negative and positive Cotton effects in circular dichroism (CD) spectra, respectively, while CPL signals of (*R*)- and (*S*)- Δ are (Fig. 3e) also mirror-symmetrical at 484 nm

with dissymmetry factors (g_{lum}) of -5.8×10^{-4} and 6.6×10^{-4} , respectively.

Trifluoroacetic acid (TFA) was chosen to prepare quantitatively acidified CHCl₃. Upon increasing the molar ratio of $n_{\text{TFA}}: n_{(R)-\Delta}$ from 0 to 2.88 in solutions $(1 \times 10^{-5} \text{ M})$ of (*R*)- Δ in quantitatively acidified CHCl₃, their UV-Vis spectra exhibit (Fig. S11, ESI[†]) a decrease at 345 nm in concert with an enhancement at 365-500 nm, while their emissions show (Fig. 3a and b) a significant redshift from 477 to 561 nm, which covers a multicolor region from blue to green to yellow as shown (Fig. 3c) in the CIE chromaticity diagram. We envision that the protonation of (R)- Δ might narrow the energy gap and thus leading to the redshifted emission. Meanwhile, the more rigid cage stablized by hydrogen bonding of NH upon protonation can reduce the non-radiative transition of intramolecular excitons and thus exhibiting (Fig. S13 and Table S4, ESI[†]) a longer fluorescence lifetime of 4.87 ns. In order to gain insight into the change in photophysical properties, absorption spectra of (R)- Δ and protonated (R)- Δ were calculated using time-dependent density functional theory (TD-DFT). TD-DFT Data indicate that, upon protonation, the HOMO-LUMO energy gap decreases (Fig. S14, ESI⁺) from 3.53 eV in the case of (*R*)- Δ to 3.02 eV for the protonated counterpart, an observation which is in line with the redshifts in absorption and emission spectra upon protonation. CD Signals of (R)- and (S)- Δ solutions in a molar ratio of n_{TFA} : $n_{(R)-\Delta} = 2.88$ were (Fig. 3d) slightly enhanced, while their CPL signals were red-shifted (Fig. 3e) from 484 to 550 nm with increased g_{lum} of -7.5×10^{-4} and 7.6×10^{-4} ,

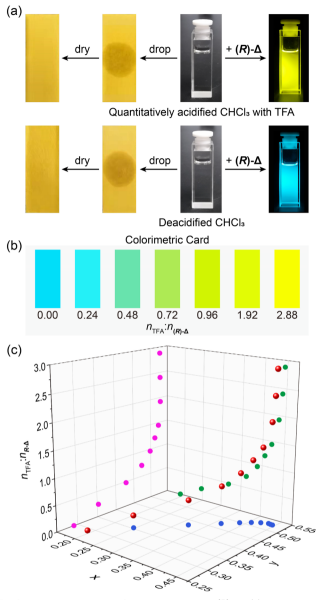


Fig. 4 Detecting traces of acids employing (*R*)-**Δ**. (a) Fluorescence changes of solutions (right) of (*R*)-**Δ** in CHCl₃ with and without TFA upon shinning 365 nm-UV light in comparison with detecting trace acid in the same solvents using pH papers (left). (b) A colorimetric card showing the ability of (*R*)-**Δ** to detect visually amounts of acid in solutions with different $n_{\text{TFA}}: n_{(R)-\Delta}$ ratios. (c) The fitting function based on the PL emission of (*R*)-**Δ** upon changing the $n_{\text{TFA}}: n_{(R)-\Delta}$ ratio from 0.00 to 2.88. The concentration of (*R*)-**Δ** is 1×10^{-5} M.

respectively, thus realizing trace acid-responsive and color tunable CPL signals.

While trace amount of acids in organic solvents can cause undesired consequences, they are difficult to quantify.⁵⁰ Based on the trace acid-responsive fluorescence change of (*R*)- Δ , we have explored its potential as an indicator to detect trace acids in organic solvents. Upon dropping (Fig. 4a) quantitatively acidified CHCl₃ with TFA (2.88 × 10⁻⁵ M) and deacidified CHCl₃ on to two pH papers, respectively, colors of both pH papers did not undergo change. In contrast, when shining Journal of Materials Chemistry C

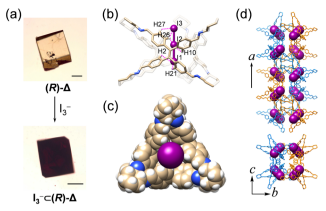


Fig. 5 Anion recognition in (R)- Δ . (a) Photographs of single crystals of (R)- Δ and [$I_3^- \subset (R)$ - Δ]. Scale bars represent 200 µm. (b and c) Superstructure of [$I_3^- \subset (R)$ - Δ] (b) in tubular and (c) space-filling representations exhibiting the encapsulation of I_3^- inside the cavity of (R)- Δ by 18 [C-H···] interactions. C, tan; N, blue; H, white; I, purple. (d) Side-on (top) and top (down) views of the double helical superstructure of [$I_3^- \subset (R)$ - Δ]. Hydrogen atoms are omitted for clarity. One strand in the double helix is depicted in orange tubular representation and the other in blue, while I_3^- is represented in a space-filling form.

365 nm-UV light on two solutions $(1 \times 10^{-5} \text{ M})$ of (*R*)- Δ prepared from the above two solvents, the former $(n_{\text{TFA}}: n_{(R)-\Delta} = 2.88)$ exhibits yellow emission, while the latter $(n_{\text{TFA}}: n_{(R)-\Delta} = 0)$ is blue, suggesting that the emission of (R)- Δ is highly sensitive to traces of acid. Upon increasing the molar ratio of $n_{\text{TFA}}: n_{(R)-\Delta}$ from 0 to 2.88 in solutions (1 \times 10⁻⁵ M) of (*R*)- Δ in CHCl₃, their fluorescence colors could be discerned by the naked eye, demonstrating the potential of (R)- Δ as a visual colorimetric card (Fig. 4b) for rapidly detecting traces of acid in solvents. Moreover, trace amounts of various acids in solvents can be detected quantitatively (Fig. 4c) as equivalent TFA by correlating the relationship between the molar ratio of n_{TFA} : $n_{(R)-\Delta}$ and the CIE chromaticity coordinates. We detected the acid amount in two commercial CDCl₃ samples [one with silver foil (C) and the other without (E)] using (R)- Δ . A solution of (R)- Δ (1 \times 10⁻⁵ M) in CDCl₃ C exhibited (Fig. S15, ESI⁺) a yellow-green emission in CIE coordinates upon excitation at 380 nm, which is equivalent to a n_{TFA} : $n_{(R)-\Delta}$ ratio of 0.49, namely, 0.49 \times 10⁻⁵ M of TFA. In striking contrast, the amount of acid in a solution of (R)- Δ $(1 \times 10^{-5} \text{ M})$ in CDCl₃ E is too much and exceeds the detection threshold. When we diluted CDCl₃ E 100 times with deacidified CHCl₃ to make a solution of (**R**)- Δ (1 × 10⁻⁵ M) in solvent D, the emission of which was measured (Fig. S16, ESI⁺) to correspond to a n_{TFA} : $n_{(R)-\Delta}$ ratio of 0.63—that is, 0.63 \times 10⁻⁵ M of TFA. Considering the 100-fold dilution, the amount of acid in CDCl₃ E without silver foil is equivalent to 0.63 \times 10^{-3} M of TFA and 129 times higher than that in CDCl₃ C stabilized with silver foil, suggesting that the addition of silver foil stabilizer can decrease significantly the amount of acid in CDCl₃. Similarly, traces of acid in commercial grade CH₂Cl₂ and CHCl₃ were also quantified (Fig. S16, ESI[†]) to be equivalent to 0.30×10^{-5} and $0.46 \times$ 10^{-5} M of TFA, respectively. Finally, the detection limit was measured (Fig. S17, ESI^{\dagger}) to be 5 × 10⁻⁷ M.

The channel of ~ 4 Å inside (*R*)- Δ can act as an ideal host for binding linear guest molecules. We have investigated the

complexation between linear I_3^- anions and (R)- Δ through SCSC transformation. Upon slow diffusion of a solution of [Bu₄N][I₃] in MeOH into the mother liquor containing single crystals of (R)- Δ , yellow crystals changed (Fig. 5a) to black after one day. SCXRD Analysis reveals (Fig. 5) that one I_3^- anion fills completely the tubular cavities of (R)- Δ to form a host-guest adduct $[I_3^- \subset (R) \cdot \Delta]$ which assembles into a (*M*)-double helix in a similar cubic unit cell to the one with (*R*)- Δ . I₃⁻ Anions are stabilized (Fig. 5b, Fig. S18 and Table S8, ESI⁺) inside the channel through up to 18 weak [C-H···I] hydrogen bonding interactions (mean [H···I] separation $d_{[H···I]} = \sim 3.4$ Å), while the Bu_4N^+ cations are disordered. Although the disordered cations in the single-crystal superstructure cannot be observed, we believe that the Coulombic interaction between cations and I_3^- is also responsible for the stable binding of I_3^- anions to the cavity of the triangular prism.

Conclusions

We have prepared a pair of tetraphenylethylene-based enantiomeric molecular triangles (R)- and (S)- Δ . X-Ray crystallographic analysis reveals the stereocontrolled formations of left- and right-handed double helices from triangular prismatic (R)- and (S)- Δ , respectively, as a result of chirality transfer, wherein the cavities in the triangles can bind I₃⁻ anions to form host-guest adducts through SCSC transformation. Both enantiomers exhibit mirror symmetrical CD and CPL signals, as well as significantly enhanced fluorescence as a result of chiral induction together with the restriction of the phenylene rotation by geometrical constraints. Moreover, trace acid-responsive changes in fluorescence and CPL from blue to yellow have been applied to detect quantitatively the acid content in common organic solvents involving CH₂Cl₂, CHCl₃, and CDCl₃, with a detection limit as low as 5×10^{-7} M. This research demonstrates the integration of self-assembly, molecular recognition, and stimuli-responsiveness in a judiciously designed triangle while, at the same time, providing insight into fabricating multifunctional materials.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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