Chiral molecular face-rotating sandwich structures constructed through restricting the phenyl flipping of tetraphenylethylene†

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Chiral tetraphenylethylene (TPE) derivatives have great potential in chiral recognition and circularly polarized luminescence. However, they were mainly constructed through introducing chiral substituents at the periphery of the TPE moiety, which required additional chemical modifications and limited the variety of chiralities of products. Herein, we constructed a series of chiral face-rotating sandwich structures (FRSs) through restricting the phenyl flipping of TPE without introducing any chiral substituents. In FRSs, the complex arrangements of TPE motifs resulted in a variety of chiralities. We also found that non-covalent repulsive interactions in vertices caused the facial hetero-directionality of FRSs, and the hydrogen bonds between imine bonds and hydroxy groups induced excited-state intramolecular proton transfer (ESIPT) emission of FRSs. In addition, the fluorescence intensity of FRSs decreases with the addition of trifluoroacetic acid. This study provides new insights into the rational design of chiral assemblies from aggregation-induced emission (AIE) active building blocks through restriction of intramolecular rotation (RIR).

Introduction

Tetraphenylethylene (TPE), as an aggregation-induced-emission chromophore (AIEgen),¹ has been extensively applied in fluorescent probes,² optoelectronic materials,³ and bioimaging agents.⁴ As a pseudo-C₄ symmetric building block, TPE has been assembled into covalent-organic frameworks (COFs),⁵ molecular cages⁶ and supramolecular fibres.⁷ Among those TPE-based materials, the chiral ones are particularly interesting because they exhibit extensive functions such as chiral recognition⁸ and circularly polarized luminescence.⁹ Generally, chiral TPE derivatives are constructed through introducing chiral substituents at the periphery of the TPE moiety.¹⁰ Nevertheless, such strategies required additional chemical modifications on TPE, and the chiral substituents limited the variety of chiralities of products. Interestingly, the propeller-like TPE core can exhibit inherent two-dimensional (2D) chirality in P or M helical configurations¹¹ through the restriction of intramolecular rotation (RIR).¹² In addition, the spatial orientation of vinyl bonds (Fig. 1a), either vertical or horizontal, also influences the chirality of assembled structures. Therefore, the complex arrangements of the four TPE motifs (Fig. 1b) could generate numerous chiral diastereoisomers without introducing any chiral substituents. But such a strategy to construct chiral TPE-based assemblies was rarely investigated and challenging due to difficulties in chiral separation.

Recently, we have constructed a series of molecular face-rotating polyhedra (FRP),¹³ which display a special form of

Fig. 1 (a) TPE motif exhibits two rotational modes (P or M) of the phenyl groups and two orientational modes (horizontal or vertical) of the vinyl bonds in two dimensions. (b) Four 2D motifs of TPE could generate numerous diastereoisomers through special arrangements. (c) Structural formula of THFPE, TREN and the corresponding sandwich structures formed by dynamic imine chemistry.
supramolecular chirality originating from the arrangements of 2D chiral building blocks. In TPE-based FRP,\textsuperscript{14} we found that the phenyl flipping of TPE faces was significantly restricted and hence FRP exhibited remarkable stability of chirality and fluorescence in diluted solution. To further control the chirality of FRP, we modulated the rotational patterns of faces with several features. On one hand, the subtle difference in building blocks could bring a huge change in the resultant assemblies and their chirality. For instance, Fujita et al. demonstrated that only a $3^\circ$ difference in the bending angles of two ligands can critically switch achiral Platonic or Archimedean structures into chiral Goldberg polyhedra.$^{15}$ More recently, our group reported that different positions of aldehyde in triazatrunexene (TAT) lead to distinct assembly behaviour and geometry.$^{16}$ On the other hand, vertices also determined the chirality of FRP. We reported that chiral vertices sufficed to control the facial directionality and the conformations of other achiral vertices in FRP assembled from truxene.$^{17}$ Nevertheless, how achiral vertices influence the rotational patterns of TPE units remains unclear.

Herein, we assemble the meta-aldehyde TPE (THFPE, Fig. 1c) with tris(2-aminoethyl)amine (TREN) to form a series of chiral face-rotating sandwich structures (FRSs) without introducing any chiral substituents on TPE. The subtle difference in TPE (the positions of aldehyde groups) changes the resultant structures from previously reported molecular cubes$^{18}$ into FRSs, wherein only four FRSs emerge as products out of twelve possible diastereoisomers. In FRSs, three TPE units exhibit hetero-directional configurations, which are controlled by the repulsive interactions in TREN vertices as indicated by theoretical calculations. In addition, the hydroxyl groups form hydrogen bonds with imine bonds and induce the excited-state intramolecular proton transfer (ESIPT) phenomenon. In conclusion, the fluorescence intensity of FRSs decreases with the addition of trifluoroacetic acid.

Results and discussion

The facial building block THFPE was prepared from 4,4’-hydroxybenzophenone through multi-step synthesis (Fig. S1†). In the Schiff-base condensation, THFPE (3 equiv.) was mixed with TREN (4 equiv.) and catalytic amounts of trifluoroacetic acid in chloroform at ambient temperature (Fig. 1c). High-resolution mass spectrometry identified the molecular weight to be 1894.21 (Fig. S2†), corresponding to THFPE$_3$TREN$_4$. However, the initial nuclear magnetic resonance (NMR) spectrum showed notable overlap of proton signals in the aromatic region (Fig. S3†), suggesting that more than one stereoisomer existed in the products.

High-performance liquid chromatography (HPLC) was employed to separate the mixed products by using a chiral column (Daicel Chiralpak IE) and a mixed solvent as the mobile phase (toluene/chloroform/methanol). After 24 hours of reaction, the HPLC spectrum confirmed that only four fractions eluted in a 1 : 2 : 2 : 1 ratio (Fig. 2a, top). Further characterization revealed that the four fractions correspond to two pairs of hetero-directional enantiomers, i.e., $MPM$-1, $MPM$-1, $PMP$-2 and $PMP$-2, which respectively eluted at 17.5, 32.9, 15.5 and 44.7 min. However, as the reaction was heated at 50 °C for five days, the fraction of FRS 2 gradually decreased (Fig. S4†) and eventually disappeared (Fig. 2a, bottom), suggesting that FRS 2 was a kinetic product which finally converted into thermodynamic FRS 1. The reaction yield of THFPE$_3$TREN$_4$ was calculated to be 97.6% according to the full HPLC spectrum (Fig. S5†).

Slow evaporation of FRSs in chloroform was allowed to afford prismatic crystals after a month. Although we used enantiopure fractions to grow crystals, single-crystal X-ray diffraction analysis revealed a pair of enantiomers co-crystallized in a unit cell, indicating racemization of FRSs 1 or 2 during the crystallization process. Further time-dependent chiral-HPLC analyses revealed that only 2% of $MPM$-1 or $MPM$-2 can change into their enantiomers after five days in solution (Fig. S6†), indicating that the racemization of each enantiopure fraction was quite slow in solution. All crystal structures of FRSs bear a tetra-capped hetero-directional sandwich structure, wherein one TPE unit was stuck between the other two TPE units with a different rotational configuration and TREN occupied four vertices to link three TPE units.

The crystal structure of $PMP$-1 is $C_2$ symmetric according to the central axis of each TPE face (Fig. 3a). The $M$ face in the middle is 5.29 Å from two $P$ faces (Fig. 3b). The vinyl orientations of $P$ faces are parallel whereas they form a 51° angle with the vinyl bond of the middle $M$ face (Fig. 3a, inset). In addition, three phenyl rings linked by the same TREN have highly twisted conformations, wherein two phenyl rings from two $P$ faces are nearly parallel and they are perpendicular to that from the $M$ face as shown in Fig. S7†. Similar to $PMP$-1, $PMP$-2 has only a subtle difference in the vinyl orientations of two $P$ faces (Fig. 3c): they are noncoplanar perpendicular to each other and form a 45° angle with the vinyl bond of the middle $M$ face (Fig. 3d and more structural details are in Fig. S7†).

The imine bonds in FRSs form hydrogen bonds (O–H···N) with hydroxyl groups and are conjugated with phenyl rings, resulting in complete coplanarity with the adjacent phenyl rings. In our previous work, the imine bonds and the adjacent phenyl rings, without the assistance of hydrogen bonds, have a 7° dihedral angle.\textsuperscript{14} To investigate the role of hydrogen-bonding interactions, we substituted the hydroxyl groups in
THFPE with ethoxy groups (TOMPE, Fig. S8a†) to replace hydrogen bonds with imine bonds. TOMPE was reacted with TREN under the same conditions and mass spectrometry revealed that the condensation reaction only produced some uncharacterized oligomers (Fig. S8c†), indicating that the hydrogen bonds assisted in the imine formation and subsequently stabilized the resultant structures.

All TREN vertices in FRSs influence the stacking of TPE because they exhibited clockwise or anti-clockwise configurations (Fig. S9a†) and strong non-covalent repulsive interactions inside TREN. In a previous report, Bein et al. demonstrated that the TPE entities in highly crystalline COFs are energetically favourable for homo-directional stacking rather than alternating stacking of $P$ or $M$ configurations to form successive layers. However, in FRSs, TPE units take the alternating stacking of $P$ or $M$ configurations. We attributed this unusual stacking in FRSs to two types of non-covalent repulsive interactions in TREN vertices. The first type is repulsive interactions among the three alkyl chains of TREN (Fig. S9c†), and the second one is repulsive interactions of atoms in the same alkyl chain (Fig. S9a,† top). In further density functional theory (DFT) calculations, we constructed a structure of $PPP-1$ through substituting the middle $M$ face into the $P$ face in $PMP-1$ (Fig. S9b†). The energy of $PPP-1$ was much higher than that of $PMP-1$. In contrast to the anti-clockwise rotational configurations of TREN in $PMP-1$, TREN in the optimized structure of $PPP-1$ has an extremely distorted configuration. This suggested that the first type of repulsive interactions led to an anti-clockwise rotational configuration of TREN. We also varied the torsional angle ($\theta$, Fig. S9†) in the N-C-C-N gauche conformation of $PMP-1$ from 0° to 180° in one alkyl chain to construct the potential energy surfaces (PESs). In PESs, due to the second type of repulsive forces in one alkyl chain, the energy of $PMP-1$ was the lowest when $\theta$ was 65° (Fig. S9d†), which is consistent with the crystal structure of $PMP-1$. Therefore, the second type of repulsive interactions restricted the movement of alkyl chains and thus transferred the chirality from rotational conformations of TREN to TPE entities, resulting in facial hetero-directionality of FRSs.

Similarly, the crystal structures of $MPM-1$ or $MPM-2$ were the enantiomers of $PMP-1$ or $PMP-2$ as shown in Fig. S10,† respectively. In addition, the theoretical calculations also confirmed that the energy of thermodynamic product 1 was 67.01 kJ mol$^{-1}$ lower than that of kinetic product 2.

The circular dichroism (CD) spectra of $PMP-1$ and $MPM-1$ showed mirror-image Cotton effect signals at 250 nm (molar ellipticity of $5.1 \times 10^3$ deg cm$^2$ dmol$^{-1}$, Fig. 2b), which belong to the absorption region of TPE, and thus confirmed that phenyl flipping was restricted in diluted solution. $MPM-2$ and $PMP-2$ also displayed mirror-like Cotton effect signals at the similar intensity of FRSs 1 as shown in Fig. 2b. More importantly, their CD spectral signals at 250 nm were dominated by the major rotational faces of TPE units, that is, the number of $M$ or $P$ faces in FRSs. For instance, FRSs with more $P$ faces exhibited negative signals at 250 nm, which is consistent with previous work by our group. All four CD spectra of FRSs well matched the (ZINDO/S)-predicted CD spectra (Fig. S11†), confirming the absolute configuration of each diastereoisomer.

The $^1$H-NMR spectrum of THFPE only displayed one peak for protons in the aldehyde groups (Fig. S12†), indicating high freedom of phenyl rings in solution. In contrast, the $^1$H-NMR spectrum of $PMP-1$ exhibited complex proton signals of imine bonds because the restriction of phenyl flipping in $PMP-1$ lowers the symmetry of THFPE (Fig. S14†). The 2D NMR spectra allowed us to assign all the protons in $PMP-1$ (details in ES†), which exhibited six peaks for protons in 12 imine bonds (Fig. S16†). Therefore, those protons of $PMP-1$ are individually interchangeable through $C_2$-symmetric operations, which is consistent with its single-crystal structure. Particularly, two protons of methylene in TREN, namely $H_a$ and $H_b$ (Fig. S17†), are in a distinct chemical environment and split into two individual peaks, confirming that the non-covalent repulsive interactions restricted the movement of flexible TREN in solution. In addition, the nuclear Overhauser effect (NOE) cross-peaks between $H_a$ and $H_b$ reflected the $E$ configuration of imine bonds and the structural rigidity of $PMP-1$ (Fig. S17c†).

Compared with the fluorescence of THFPE and some similar compounds, FRSs 1 and 2 exhibited an emission peak at 550 nm with a large Stokes shift (~200 nm) in diluted solution (Fig. 4b). In addition, the fluorescence quantum yields of FRSs 1 and 2 were measured to be 2.08% and 1.56% in toluene, respectively. The emission at 550 nm can be rationalized by restriction of phenyl rotation of TPE units and the ESIPT phenomenon induced by the intramolecular hydrogen bonds (Fig. 4a). ESIPT emission involves a four-level photophysical cycle, in which the enol–imine tautomerism from $E*$ to cis-K* occurs upon photoexcitation and then cis-K returns to the initial enol–imine form through ground state intramolecular proton transfer (GSIPT). However, there are two competitive processes for cis-K* relaxation: radiative decay to the ground state (cis-K) and rotation-induced non-radiative decay to trans-K. Generally, E/Z isomerization from cis-K* to trans-K is forbidden on
aggregation and it will increase the intensity of ESIPIT emission.20 In FRSs, the rigidity of TREN vertices can also prevent the E/Z isomerization of cis-K* and strong ESIPIT emission of FRSs can be observed in diluted solution. In addition, the ESIPIT emission of FRSs can be tuned by pH and solvent, which can remarkably block the enol–keto tautomerism in the excited state.21 By adding TFA into FRSs 1 in chloroform incrementally, the fluorescence intensity decreased dramatically at 550 nm (Fig. 4c), whereas the fluorescence can recover to the initial intensity by using trimethylamine to neutralize TFA (Fig. S19c†). As expected, increasing the solvent polarity can also switch off emission at 550 nm with a red shift (Fig. S19b†).

Conclusions

To conclude, we have constructed a series of FRSs with emergent chirality and fluorescence through restricting the phenyl flipping of TPE units in diluted solution. Due to the non-covalent repulsive interactions, TREN vertices exhibited rotational configurations and transmitted their chirality to the TPE entities, resulting in facial hetero-directionality of FRSs. In addition, the hydrogen bonds between imine bonds and hydroxy groups induced ESIPIT properties, which can be tuned by pH and solvents. This study provides a strategy to construct chiral supramolecular assemblies from AIEgens by restricting intramolecular rotation, which can be applied in luminescent materials, molecular recognition or asymmetric catalysis.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank the support from the NSFC (No. 21722304, 91427304, 21573181, and 91277111), the 973 Program (No. 2015CB885605), the Top-Notch Young Talents Program of China and the Fundamental Research Funds for the Central Universities of China (No. 20720160050). We thank Wei Zhang, Hao Li, Haitao Peng, and Jonathan Nitschke for helpful discussions. We also thank Linlin Yang, Wenting Deng, Kai Wu and Jijun Jiang for assistance in experiments.

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