s-Tetrazine-Bridged Photochromic Aromatic Framework Material

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ABSTRACT: Integrating fluorescent chromophores in aromatic frameworks could not only prevent aggregation-induced quenching caused by the π−π stacking interaction between the chromophore components but also confer new fluorescence properties. Herein, we report the fabrication of s-tetrazine-bridged aromatic frameworks TzAF by the incorporation of the smallest aromatic fluorophore, s-tetrazine (Tz), into the skeleton of a tetrahedrally connected lattice of aromatic frameworks. The thin films of TzAF coated on silica gel plates were found to exhibit reversible photoswitching fluorescence characteristics under alternate UV and visible-light irradiations with excellent fluorescence stability and high on/off contrast. The repeatable "on/off" fluorescence photoswitchability of the TzAF thin films was mechanistically attributed to light-induced reversible transformation between TzAF's neutral and radical states.

1. INTRODUCTION

Stimuli-responsive materials are able to reversibly change their physicochemical states and distinguishable properties in response to externally applied stimuli, such as stress, light, temperature, moisture, pH, or electric or magnetic fields. 1–6 For example, photoswitchable fluorescent materials are photochromic fluorophores whose distinct on/off fluorescence emission could be controlled by light. 7–9 As light can be delivered instantly to precise locations of light-responsive materials, such materials have thus found wide application potentials in the areas of information encryption, anti-counterfeiting, reversible fluorescence imaging, ultrahigh-resolution bioimaging, photocontrolled biological functions, rewritable printing, and so forth. Photochromic fluorescence materials were usually obtained through a suitable molecular combination of photochromism and fluorescence, that is, chemical association between photochromic and fluorescent molecular components. 7–9 Thus far, various diarylethene (DArE)-based photochromic fluorescent materials have been well reported, thanks to the inherent fluorescence that resulted from resonance energy transfer or intramolecular electron transfer processes possessed by the DArE-based molecular components and their excellent photoswitching properties, biostability, and fatigue resistance. 7–10 Besides DArE, a few other photochromic fluorophores, including the derivatives of spiropyran, spironaphthoxazine, imidazole dimer, dipyromethene, oxazine, hydrazones, and rhodamine, have also been unveiled. 7–12 Fluorophores emit bright fluorescence in solution, but their fluorescence intensity is much weaker in the solid states as electronic coupling in well-conjugated and planar aromatic structures quenches fluorescence emission, preventing the reliable translation of fluorescence properties to applications as solid materials. 13,14 To avoid such aggregation-caused quenching (ACQ), fluorophores could be dispersed into lattice porous structures 12 or incorporated as building blocks in the structures of porous solid materials such as metal organic frameworks (MOFs), covalent organic frameworks (COFs), 15 porous organic polymers, 16 or porous aromatic frameworks (PAFs). 17 Toward this end, Laursen, Flood, and co-workers allowed cationic fluorophores cocry stallized into well-defined lattices of ion-sequestering macrocycles to keep the fluorescent dyes from dulling in the solid state. 13 Hill, Ladewig, and co-workers allowed photoswitchable fluorescent PAF materials using DArE-based fluorophores as guest molecules. 18 Klajn’s group described two families of spiropyran-based fluorescent photoswitchable frameworks in which the spiropyran units were incorporated in the framework skeleton. 17 Zhu’s group has developed photoswitchable fluorescent PAF materials by decorating the frame of PAF-SCF with DArE fluorophores. 15

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However, it is challenging to integrate large fluorophores covalently into the aromatic framework skeletons. We envisioned that it could be more straightforward to incorporate covalently smaller photochromic fluorophores into the skeletons. In our search for small non-DArE-based fluorescent photoswitchable materials, we were intrigued by 1,2,4,5-tetrazine (s-tetrazine, Tz), the arguably smallest and electron-poorest aromatic fluorophore in which the presence of four sp² nitrogen atoms makes the ring an exceptionally strong electron acceptor, comparable to tetrano-benzene. With a lower-energy π* lowest unoccupied molecular orbital (LUMO), the π-aromatic system of Tz can be reversibly reduced to a stable radical anion which possesses a distinct different fluorescence property from that of Tz. For instance, chloromethoxytel-

The as-synthesized TzAF was characterized by suspended-state ¹H nuclear magnetic resonance (NMR) and solid-state ¹³C cross-polarization magic angle spinning (¹³C CP-MAS) NMR spectroscopies, Fourier transform infrared (FTIR) spectroscopy, ultraviolet-visible (UV–vis) spectroscopy, solid-state UV–vis–near-IR (UV–vis–NIR) spectroscopy, X-ray photoelectron spectroscopy (XPS), thermogravimetric analysis (TGA), powder X-ray diffraction (PXRD) analysis, field-emission scanning electron microscopy (FE-SEM), and field-emission transmission electron microscopy (FE-TEM). The aromatic protons were obvious in the suspended-state ¹H NMR spectrum of TzAF dispersed in DMSO-d₆ (Figure 1a), while no amine NH proton signals in the range of 9–10 ppm were observed, indicating the complete conversion of 6 to TzAF. The signals at around 65 ppm, 127–155 ppm, and 160–165 ppm in the ¹³C CP-MAS NMR spectrum (Figure 1b) could be assigned to the carbons of tetrathｙmethanes, the benzene, and the Tz units, respectively. The peak at 42.7 ppm (7.5 kHz) should be attributed to the spinning sideband of the C-H carbons of 6 (Figure S7, Supporting Information). Therefore, the reaction did not yield the desired product probably due to the poor solubility of organometallic precursors (3a and 3b). Thus, instead of using Tz-containing building blocks, we decided to try a de novo synthesis during which dihydro-s-tetrazine (dihydro-Tz) units were built from nitrile and hydrazine and then oxidized to aromatic Tz units. As illustrated in Scheme 1, the attempted reaction of tetrahedral symmetric aromatic nitrile (4) and hydrazine catalyzed by sulfur powder yielded the nonconjugated dihydro-Tz-bridged cross-linked frames 6, but the sharp heat and gas release of the reaction prompted us to explore a stepwise approach (Figure S7, Supporting Information). Therefore, nitrile 4 was first transformed to amidine 5, which then reacted with hydrazine to furnish the dihydro-Tz-bridged frameworks 6. The transformation of nonrigid 6 into the desired target TzAF was realized through the oxidation of the dihydro-

Scheme 1. Synthesis of TzAF
electron transitions of the Tz units. 40 In the UV–vis–NIR spectrum of the solid TzAF thin film (Figure 1c, the insert), an absorption peak at ∼257 nm (π–π* transition of the benzene rings) and an absorption band ranging from 300 to 600 nm (n–π* and π–π* transitions of tetrazine units) were observed. The similarity between the UV–vis spectra of TzAF dispersed in the solution and TzAF film implied that TzAF is in homologous chemical environments in both the solution and solid state. TGA under a nitrogen atmosphere indicated that TzAF started to decompose at 245 °C (extrapolation, the insert of Figure 1d) probably due to the decomposition of the Tz units, 41,42 and ∼35% weight remained at 1000 °C.

The X-ray photoelectron spectrum of TzAF (Figure 2) showed three peaks at 532, 400, and 285 eV, corresponding to O 1s, N 1s, and C 1s signals, respectively (Figure 2a). The O 1s peak mainly arose from molecular oxygen or water molecules absorbed on the surface or in the pores of the frameworks, which is typical for most carbon-based materials. 30,31,43 The N 1s peak with a binding energy of 400.1 eV was from the –C=−N– of Tz rings (Figure 2b). 44 Since the charge distribution on carbon atoms in the Tz rings was influenced by the electron-deficient Tz N atoms, the C 1s peak of TzAF was deconvoluted into Gaussian–Lorentzian shapes (Figure 2c), among which the peak at 284.7 eV was assigned to quaternary carbons and phenyl carbons, the one at 285.5 eV to the carbons in the Tz units, 45–46 and the weak broad peak at 291.6 eV to the characteristic shake-up satellites of the aromatic carbons (π–π* transition). 47–49

In the FTIR spectrum of TzAF (Figure 3), the disappearance of the amide –C=−NH peak at 1676 cm−1 indicated no residual amidine left in the sample. 55,56 The IR bands at 1560, 1412, and 1072 cm−1 could arise from the C–N stretching vibrations of the Tz units, 55,56,57 and the peaks at 1609, 1493, and 1448 cm−1 corresponded to the C=C breathing vibrations of the benzene rings. Due to the effect of electron deficiency of the Tz units, the characteristic peak for the benzene rings red-shifted from 1500 cm−1 to 1493 cm−1, with its intensity reduced to a level lower than the peak at 1609 cm−1. 56 Additionally, the absorption band at 830 cm−1 was attributed the characteristic C–H out-of-plane bending vibrations of the benzene units.

From the nitrogen adsorption–desorption measurement at 77 K, a type IV adsorption isotherm, typical for mesoporous materials, was obtained (Figure 4c). 57 The Brunauer–Emmett–Teller (BET) surface area of TzAF was calculated to be 16 m2 g−1, and the pore size distribution calculated by the Barrett–Joyner–Halenda method showed that TzAF is a mesoporous material with the pore diameters ranging between 1.8 and 2.5 nm. The PXRD measurement (Figure 4b) of TzAF showed no apparent sharp peaks in 1–10° (2 theta), declaring its non-long-range ordering nature. The BET surface area of TzAF was obviously smaller than what we expected for such PAFs. 58,59 One possible reason could be structural interpenetration, which is hard to avoid in the formation of PAFs. In addition, we believe that incompleteness of the formation of the ideal diamond-shaped lattice of the frameworks could be another reason. The nonaromatic dihydro-Tz is a nonplanar twisted ring, as illustrated by a model compound 3,6-dimethyl-1,4-dihydro-1,2,4,5-tetrazine (Figure S8, Supporting Information), 60 and twisted nonaromatic dihydro-Tz should cause structural twists in the intermediate frameworks 6. Therefore, frameworks 6 could be semiporous, which could hinder NaNO2 from diffusing freely into the mesopores of 6, resulting in poor conversion.
in the incompletion of the oxidation of dihydro-Tz by NaNO2. As a result, the aromatic frameworks TzAF could also be semiporous with mesopores (pore size ~2 nm), which is consistent with a low BET surface area value.

The morphological details of TzAF were studied by FE-TEM and FE-SEM. The FE-SEM images of TzAF revealed particulates in the microscale (Figure 4a and Supporting Information), while the FE-TEM images with a relatively small thickness contrast indicated that the pore sizes were about 2 nm, which is consistent with the result of nitrogen adsorption–desorption measurement (Figure 4c and Supporting Information).

2.3. Fluorescence Properties. In order to get a better understanding of the fluorescence property of TzAF, 3,6-di-p-tolyl-1,2,4,5-tetrazine (DTTz) was prepared as a model structure of the TzAF skeleton unit. The quantum chemical calculations of DTTz revealed an absence of imaginary frequencies, which meant a true energy minimum, consistent with its crystal structure (CCDC 202204). Based on the optimized geometry of DTTz, the optical transition energies and oscillator strengths were obtained using time-dependent density functional theory (TD-DFT) calculations with the same basic set. The results of the TD-DFT calculations (Figure 5 and Table S1, Supporting Information) agreed well with the experimental UV–vis spectrum (Figure 6b).

The leading configuration of the first excited state is an \( n-\pi^* \) transition (\( S_0-S_1 \)), that is, from the nonbonding orbitals of the nitrogen atoms highest occupied molecular orbital (HOMO) in DTTz to the antibonding \( \pi^* \) orbitals of the Tz ring (LUMO). The oscillator strength is low as a result of this forbidden transition, which is also in agreement with the experimental result where the corresponding absorption band is weak (centered at 543.5 nm, \( \epsilon = 354 \text{ L mol}^{-1} \text{ cm}^{-1} \)). The shoulder-type peak range of 350–420 nm in the experimental spectrum is a weak \( \pi-\pi^* \) transition (Figure 6b) which might involve \( \pi \)-orbitals delocalized over the DTTz molecule (HOMO \(-1 \rightarrow\) LUMO). Meanwhile, an intermediate transition which involves \( n-\pi^* \) transition (HOMO \( \rightarrow\) LUMO + 1) is calculated to be with a zero oscillator strength. The absorption band at around 300 nm arises from the combination of two \( \pi-\pi^* \) transitions: one from the orbitals of the Tz ring and the other from the orbitals of the benzene-ring.

Upon excitation at the first absorption (500–600 nm) of DTTz, weak fluorescence was observed in the region of 575–620 nm (Figure 6a). In addition, a higher absorption energy leads to a higher excited (\( S_n \)) state, which could be transformed to the \( S_1 \) state through vibrational relaxation and internal conversion, accompanied by emitting \( S_1 \rightarrow S_0 \) or \( S_n \rightarrow S_0 \) fluorescence. As a consequence, the excitation of DTTz by a 330–400 nm light resulted in fluorescent emission at 560–630 nm. The inconspicuously weak fluorescence peak at around 580 nm might be due to the intermolecular orbital overlap between the electron-deficient Tz ring and the two electron-rich benzene rings in DTTz. Nevertheless, the orbital overlap between Tz units and benzene units in the frameworks TzAF was prohibited due to the blockage of quaternary carbons. Consequently, solvent-dispersed TzAF emitted fluorescence at 400–650 nm at a higher intensity with two broad excitation wavelength ranges of 270–310 nm and 350–600 nm (Figure 6c). In addition, the PL spectra of the solid-state TzAF excited
At 254, 340, and 365 nm showed two broad emission peaks: one centered at 254, 340, and 365 nm (excited by 254 nm, 340 nm, and 800 nm), respectively (Figure 6d). The fitted lifetimes of TzAF’s fluorescence at 484 and 587 nm (excited by 340 nm) were 2.278 ns (τ₁ = 0.9157 ns, 81.13%; τ₂ = 8.1241 ns, 18.87%) and 249.853 ns (τ₁ = 2.3733 ns, 23.18%; τ₂ = 324.5291 ns, 76.82%), respectively, indicating that the emission at 587 nm needs more time to be vibrationally relaxed and internally converted to its S₁ state.

2.4. Fluorescence Switching. The π-aromatic Tz ring can be reduced to a radical anion which has much weaker fluorescence emission than its neutral state, as exemplified by chloromethoxytetrazine. Thus, in TzAF, the electron-deficient Tz units conjugated to the benzene units should be able to be readily reduced to Tz radical anions by light-generated electrons, resulting in a significant decrease in the fluorescence intensity. The fluorescence intensity could be able to be regained once the Tz radical releases an electron to resume the neutral state. Therefore, we envisioned that this neutral ↔ radical state interconversion should make TzAF a good photoswitchable fluorophore with distinct on-off fluorescence emission.

Based on the above assumptions, we set out to explore the fluorescence photoswitchability of the TzAF thin films coated on silica gel plates. As shown in Figure 6d, TzAF emitted fluorescence under UV light irradiation at 365, 340, or 254 nm. To our delight, it was found that the TzAF thin film emitted bright fluorescence under a 254 nm UV light irradiation (Figure 7a), but its emission could be turned off by the irradiation of a 405 nm laser light, as was exemplified by “writing” the number “254” on a 254 nm UV-excited TzAF thin film using 405 nm laser beam (Figure 7b and S13, Supporting Information). The mark “254” on the TzAF thin film started to fade when exposed to 254 nm UV light and disappeared thoroughly within 10 min (Figure 7c–h).

UV light at 365 nm functioned similarly in restoring the bright fluorescence emission of the TzAF thin film (see Figure S16 in the Supporting Information). In the repeatability test of the TzAF thin film’s fluorescence photoswitching capability, the “writing” and “erasing” with 405 and 254 nm (or 365 nm) light irradiation were conducted for consecutive cycles without resulting in any identifiable difference in the behaviors of the thin film (Figures 8 and S17–S18, Supporting Information). It is worth noting that the mark written on the TzAF thin film by 405 nm irradiation did not last for 24 h under natural room light illumination (300–1000 nm) but lasted for no less than 70 days in the dark (Figure S15, Supporting Information).

We postulated that the on/off fluorescence emission of TzAF is attributed to the interconversion between TzAF’ neutral ↔ radical states: Tz units in the neutral TzAF were reduced to radicals by 405 nm light-generated electrons, and the radicals lost electrons to get back to the neutral form under 254 nm UV light. In order to provide evidence to support this postulation, electron paramagnetic resonance (EPR) measurement was conducted. As shown in Figure 9a,b, a clear EPR signal was observed for TzAF irradiated by 405 nm light (red curve), and such a signal could not be seen in the dark or before light irradiation (black curve). The g-factor spectrum (Figure 9b) of TzAF after 405 nm light irradiation was derived from Figure 9a and indicated that the experimental g-factor was 2.00429, typical for nitrogen-centered radicals. Therefore, the shutting off of the fluorescence emission of TzAF can be attributed to the transformation of fluorescent neutral TzAF to its nonfluorescent radical state. The photogenerated electron in the TzAF radical is considered in its singly occupied molecular orbital excited by a higher energy 365 or 254 nm...
light irradiation, followed by de-excitation to its ground state accompanied with the reproduction of fluorescence.

With the mechanism of TzAF’s “on/off” fluorescence emission elucidated, we wanted to find out whether a single unit of TzAF possesses such fluorescent photoswitchability. Thus, we conducted EPR analysis of DTTz, the model compound of a single unit of TzAF, in its solid-state (Figure 9c,d) and in CH$_3$CN solution (Figure S21, Supporting Information). The intensity ratio of characteristic signals obtained was 1.34:4.41:9.88:16.07:18.63:16.38:10.34:4.04:1.59 (Figure 9c), consistent with the theoretical value calculated for Tz-type radicals (1:4:10:16:19:16:10:4:1). However, there was a significant difference between the frameworks TzAF and the monomer DTTz: neither the DTTz thin film nor DTTz solution in CH$_3$CN showed such a switchable “on/off” emission property (Figure S20, Supporting Information). In TzAF, the Tz radical anion was stabilized by delocalization of the electron over the Tz ring and the two conjugated phenyl rings. In addition, Ph-Tz-Ph type-radical anion units in TzAF were isolated by the quaternary carbons, which further improved the stability of the radical anions in a similar way to the encapsulation strategy reported by Flood’s group. A naked DTTz radical anion does not have such a protection; thus, it lacks such fluorescence switchability.

3. CONCLUSIONS

s-Tetrazine (Tz), arguably the smallest fluorophore was incorporated in the skeleton of PAFs for the first time, resulting in photochromic aromatic frameworks TzAF. A thin film of the as-synthesized TzAF was found to possess a consecutive “writing/erasing” property through the irradiation of 405 and 254 nm (or 365 nm) light, respectively, which could be mechanistically attributed to the “on/off” fluorescence photoswitchability resulting from reversible transformation between fluorescent-neutral and nonfluorescent radical states. The rigid skeleton of TzAF provided a protective environment for stabilization of the photogenerated Tz radicals, while aggregation-caused quenching (ACQ) resulted in the loss of fluorescence photoswitchability for DTTz in the solid state. The unique fluorescent photochromic TzAF should have potential in various optoelectronic applications, including optical memories, bioimaging, and photoswitches with high sensitivity.

4. EXPERIMENTAL SECTION

4.1. General. Unless otherwise noted, the reagents and solvents were purchased from commercial sources and used as received. The $^1$H and $^{13}$C NMR spectra were recorded on a Bruker AVANCE III HD 500 NMR spectrometer or a Bruker AVANCE NEO 400 spectrometer. $^{13}$C cross-polarization MAS
NMR (13C CP/MAS NMR) spectra were recorded on a Bruker AVANCE Neo 700 MHz spectrometer. Electrospray ionization mass spectrometry analysis was conducted on a Thermo Scientific Q Exactive Focus mass spectrometer with an UltiMate 3000 RSLC high-performance liquid chromatography system. TGA was carried out on a Mettler Toledo TGA/DSC 3+ thermogravimetric analyzer, and the samples were heated to 1000 °C at a heating rate of 10 °C/min under a N2 atmosphere. The PXRD measurements were carried out on a Bruker AXS D8 ADVANCE X-ray diffractometer with Cu Kα as the radiation source (λ = 1.5418 Å) and operated at 40 kV and 40 mA. The surface area, nitrogen adsorption isotherms (77 K), and pore size distributions were measured using an ASAP 2020 HD accelerated surface area and porosimetry system, and the sample was degassed at 90 °C for 6 h prior to analysis. The Fourier transform IR (FTIR) spectra were obtained on a Thermo Scientific Nicolet iS5 FTIR spectrometer using the attenuated total reflection technique, and the sample was dried in an IR oven equipped with two 275 W lamps prior to the test. The UV–vis spectra were recorded on a UV-8000S Double Beam UV/vis spectrophotometer (Shanghai Precision Instruments) and the UV–vis–NIR spectra on an Agilent Cary UV–vis–NIR spectrophotometer. EPR/electron spin resonance (ESR) spectra were obtained on a Bruker EPR A300 spectrometer. The morphological details were studied using FE-SEM (on a FEI Inspect F50) and FE-TEM (on a JEM-2100F field emission electron microscope). The XPS analysis was conducted on a Thermo Fisher ESCALAB 250 Xi X-ray photoelectron spectrometer. Structural simulation was performed using BIOVIA Materials Studio Forcite (geometry optimization: energy, 2 × 10−5 kcal/mol; force, 0.001 kcal/(mol Å); stress, 0.001 GPa; displacement, 1.0 × 10−5 Å). Control software in fluorescence photoswitching was obtained from VigoTec, and calculations were performed at the B3LYP/6-311+G(d) level using Gaussian 09 Revision E.01.

4.2. Tetra(4-iodophenyl)-methane (2). A suspension of iodine (9.23 g, 36.0 mmol), tetraphenylmethane (3.92 g, 12.0 mmol), and bis(trifluoroacetoxy)iodobenzene (15.2 g, 30.0 mmol) in CCl4 (180 mL) was refluxed under N2 for 10 h, cooled to room temperature, and filtered to collect the solid which was washed with petroleum ether and dried at 100 °C under reduced pressure for 1 h to afford 2 (6.4 g, 65%) as a light-pink powder. 1H NMR (500 MHz, DMSO-d6): δ 7.69 (s, 8H), 6.91 (8H).

4.3. Tetra(4-cyanophenyl)-methane (4). A mixture of tetra(4-cyanophenyl)-methane (2) (1.32 g, 1.6 mmol), K2Fe(CN)6·3H2O (1.40 g, 3.3 mmol), K2CO3 (0.68 g, 4.92 mmol), and Pd(OAc)2 (3.59 mg, 0.16 mmol) in dimethylformamide (25 mL) was purged with N2, heated at 150 °C for 18 h, cooled to room temperature, poured into DCM (50 mL), and filtered to remove the solids. The filtrate was concentrated to result in a residue which was subjected to chromatography (petroleum ether:DCM = 1:1) to afford 4 as a white powder. 1H NMR (500 MHz, DMSO-d6): δ 7.83 (d, 8.4 Hz, 8H) 7.42 (d, 8.3 Hz, 8H); 13C NMR (125 MHz, DMSO-d6): δ 149.6, 132.9, 131.5, 118.8, 110.3, 65.9.

4.4. 4,4‴,4‴,4‴-Tetramethanetetrayltetrasubimide Tetrahydrochloride (5). To a solution of tetra(4-cyanophenyl)-methane (4) (353 mg, 0.84 mmol) in dry THF (20 mL) under N2 was added a solution of LiHMDS in THF (1.0 M, 7.5 mL, 7.5 mmol) at −84 °C (EtOAc-liquid nitrogen bath). The resulting mixture was warmed to room temperature and stirred for 18 h to form a yellowish-orange solution which was then cooled to 0 °C. After an ethanolic HCl solution (12 mL, 32 mL, freshly prepared by adding acetyl chloride to ethanol) was added, the precipitate collected by filtration was suspended in 10 mL of ethanol, sonicated for 1 h, filtered, and dried under reduced pressure to afford 5 as a cream-colored solid (525 mg, 98%). 1H NMR (500 MHz, D2O): δ 7.77 (d, 8.5 Hz, 8H) 7.62 (d, 8.6 Hz, 8H); 13C NMR (125 MHz, D2O): δ 165.9, 150.7, 131.3, 127.8, 126.2, 65.4.

4.5. TzAF. To a suspension of 4,4‴,4‴,4‴-methanetetrattetra(phthalimide)tetrahydrochloride (5) (64 mg, 0.1 mmol) in water (2.0 mL) was added hydrazine hydrate (46 μL, 85%, 0.8 mmol, 8 equiv), resulting in a mixture which was heated at 40 °C using MW for 1 h. The faint yellow precipitate was collected, suspended in an aqueous NaNO2 solution (0.5 M, 2.0 mL), sonicated for 5 min, and cooled down to 0 °C in an ice-water bath. After glacial acetic acid (3 × 35 μL) was added portionwise under vigorous stirring, the pale-yellow precipitate disappeared, resulting in a pink solution which gradually became cloudy pink. After water (16 mL) was added, the reaction mixture was transferred into a regenerated cellulose dialysis bag for water dialysis three times and dried under reduced pressure at 120 °C to afford TzAF as a pink powder (33 mg).

**ASSOCIATED CONTENT**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.2c00278.

1H NMR spectra, 13C NMR spectra, and additional results (PDF)

Laser writing step (MP4)

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Notes

The authors declare no competing financial interest.

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