Supramolecular Assemblies Showing Thermally Activated Delayed Fluorescence

Thomas A. Comerford and Eli Zysman-Colman*

Supramolecular assemblies based on luminescent components offer significant advantages over their discrete counterparts, including improved quantum yields, stability, and tunability. Following interest as advanced optoelectronic materials, thermally activated delayed fluorescence (TADF) emitters are incorporated into a range of supramolecular structures. Herein, a summary of the known examples of emissive supramolecular systems that operate via a TADF mechanism with comparisons, where possible, with their discrete counterparts is presented. While the types of supramolecular structures are diverse, there are limited examples shown for each class. With the increase in photophysical performance and/or emergence of new photochemical properties upon going from molecular to supramolecular, the potential that these systems hold becomes apparent.

1. Introduction

Thermally activated delayed fluorescence (TADF) materials have attracted considerable interest due to their strong potential in replacing phosphorescent complexes as emitters in state-of-the-art organic light-emitting diodes (OLEDs). The ability of organic TADF materials to harness singlet and triplet excitons, the latter via reverse intersystem crossing (rISC), to achieve up to 100% internal quantum efficiency in the device is key to the attention these advanced optoelectronic emitters have received. High-efficiency blue and deep-blue OLEDs have been recently reported with external quantum efficiencies of 25.7% and 28.2% respectively, overcoming the 25% spin-statistical bottleneck for these emitters by harvesting triplet excitons via rISC. TADF compounds are also increasingly being used as photocatalysts (Figure 1). The presence of triplet states in the TADF mechanism makes these systems useful as sensors for oxygen, and the temperature-dependent nature of TADF emission allows for their use as molecular thermometers.

TADF emitters coupled with their lower-toxicity profiles than organometallic complexes have been factors driving their exploration as bioimaging reagents. In particular, their long lifetimes permit a clear distinction between the TADF marker and short-lived emission from biological autofluorescence. As the use of these materials increases and the scope of their use diversifies, so does the need to achieve greater control of their properties. Incorporation of TADF materials into supramolecular assemblies, unlike other classes of luminescent compounds, is underexplored and may fundamentally change their properties via intra/intermolecular interactions, reducing unwanted nonradiative decay and giving new reactivity when used in photocatalysis. In this Review, we focus on examples where TADF occurs from organic units within supramolecular systems. There are examples of supramolecular organometallic complexes that display TADF, for example, the Cu(I)-containing structures reported by Lescop and coworkers; however, these are outside the scope of this Review.

2. Supramolecular TADF: A Diverse Lineup

2.1. Supramolecular TADF Photocatalysis

A subclass of photocatalysts comprising a photosensitizer linked to a catalyst, the so-called chromophore-catalyst assembly, have been used as hydrogen-evolving/CO2 reduction cooperative catalyst systems. The photosensitizing units undergo excited-state electron transfer to the appended catalysts, with common photosensitizers being Ir(III), Ru(II) with the use of organic species such as BODIPY dyes, and fluorescein also known. Chao and Zhao reported a TADF chromophore comprising a 1,2-dicyanobenzene core with four methoxy pyridincarbazole groups, 1. This species can coordinate to four cobalt(III) chloropyridine)cobaloxime species to afford the supramolecular assembly, 1-Co, which they showed engaged in catalytic acceptor-less dehydrogenation (CAD) (Figure 2). CAD involves the two-electron oxidation of organic precursors to give oxidized products and two equivalents of hydrogen, allowing for the generation of hydrogen gas from organic species without the need for stoichiometric inorganic oxidants. CAD is an attractive route for green hydrogen production.

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and featureless, characteristic of emission from a charge-transfer (CT) state, and is centered at $\lambda_{PL} = 591$ nm; the compound likewise shows positive solvatochromism that is associated with CT excited states. The photoluminescence quantum yield, $\Phi_{PL}$, is only 7.2%. Density functional theory (DFT) calculations confirmed the donor–acceptor nature of 1, showing a highest occupied molecular orbital (HOMO) that is largely distributed over the carbazole moieties with the lowest unoccupied molecular orbital (LUMO) localized on the electron-poor phthalonitrile core. The $\Delta E_{ST}$ value for 1 is small, quoted at 0.13 eV, suggesting that TADF is plausible.[35] With the measured photoluminescence lifetime in degassed CH$_2$Cl$_2$ at 298 K found to be $\tau_{PL} = 17.4$ $\mu$s, the excited state of 1 is sufficiently long to allow for its engagement in photocatalytic processes. Compared with 1, 1-Co had a lower photoluminescence quantum yield and decreased lifetime ($\Phi_{PL} = 2.9\%$, $\tau_{PL} = 13.8$ $\mu$s), both attributed to photoinduced electron transfer (PET) from 1 to the cobalt centers. Investigation of the redox potentials of 1-Co showed electronic separation in the ground state between the cobalt centers and the organic core, with distinct irreversible redox waves seen for Co$^{III}$/Co$^{II}$ and 1$^+/1$ redox couples at $-0.63$ and $+1.38$ V versus saturated calomel electrode in acetonitrile, respectively. Following confirmation of electron transfer from 1 to the cobaloxime groups, the assembly was used in the CAD of secondary amines. Irradiation with blue LEDs (450 ± 10 nm, 3 W) of a degassed, dry THF solution of dibenzylamine, S1, with 0.04% loading of 1-Co at room temperature gave a turnover number (TON) of 305 for hydrogen production after 12 h. In contrast, using a mixture of cobaloxime and 1 in solution afforded a significantly lower TON of 53. This increase in efficiency for the supramolecular system over the mixture of catalysts is attributed to more efficient electron transfer between the two components as a result of the covalent bond between them. A control experiment without irradiation was
conducted and showed no H₂ production, demonstrating the photochemical nature of this reaction. The imine product, P1, of the CAD of S1 was observed by ¹H NMR, which provided evidence that S1 was the source of H₂. The highest TON achieved was 895 when using 1,2,3,4-tetrahydroisoquinoline as the substrate and with a 0.04% catalyst loading of 1-Co. PET was confirmed by in situ monitoring of the UV−vis absorption spectra, which showed the emergence of a CoII species arising from electron transfer from 1* to CoIII; there was also an increase in absorption between 550 and 700 nm, attributed to the formation of a CoI species. The proposed mechanism is shown in Figure 3. Upon PET from 1*-CoIII, 1+-CoII is formed. Oxidized 1+ can then oxidize S1 to give a radical cation, I1, which then undergoes hydrogen abstraction to give a strongly reducing radical species I2 followed by oxidation to the iminium ion I3 yielding CoI. Stephenson and coworkers demonstrated the nucleophilic addition to and oxidative alkylation of photogenerated iminium species to give a wide range of products, making this intermediate valuable in organic synthesis. Deprotonation of I3 gives the product P1. CoI is protonated by I3 to give 1-CoIII/H and P1. Finally, 1-CoIII/H is protonated a final time to regenerate 1-CoIII and release H₂. The authors note that CoIII cannot be reduced by 1* due to the less negative potential of E(1+/1*) versus E(CoII/CoI/H) (−0.71 vs −1.12 V), supporting their proposed mechanism; however, other plausible pathways exist such as homolysis of CoIII/H.[40]

2.2. Supramolecular Gels

Zysman-Colman and coworkers reported the first example of a TADF gel formed from a carbazole−benzophenone TADF acceptor, 3.[41] These were formed from the mixture of a TADF emitter (Figure 4a) with diacids (Figure 4b); the structurally related compound 2 did not form any gel when mixed with either succinic or tartaric acid. DCM solutions of 2 and 3 showed broad absorption bands at 356 and 326 nm, respectively, attributed to intramolecular charge transfer (ICT) from carbazole to benzophenone that was corroborated by DFT calculations (Figure 4c). The emission maxima of 3 in degassed DCM and in PMMA-doped films (10 wt%) at 298 K are 477 and 449 nm with associated quantum yields under N₂ measured at 52% and 21%, respectively. These quantum yields decreased significantly to ΦPL = 10% and 17%, respectively, in the presence of O₂, confirming that triplet states are involved in the emission of this compound. A yellow/green gel formed with enhanced emission at λPL = 500 nm, when one equivalent of succinic acid was added to a solution of 3. This gel was weak and flowed when the vial was inverted, with a critical gel concentration (CGC) of 5 mg mL⁻¹. When using L-tartaric acid, the viscosity increased markedly, showing that a stronger gel formed with a CGC of 3 mg mL⁻¹. This was attributed to the increased hydrogen bonding possible with (L)-tartaric acid. The emission maximum for the (L)-tartaric acid gel was red-shifted compared with the succinic acid-containing gel at λPL = 510 nm. The emission intensity is

![Figure 3. Photocatalytic cycle for the CAD of dibenzylamine to give hydrogen gas. Reproduced with permission.[33] Copyright 2019, Royal Society of Chemistry.](image-url)
dependent on the amount of L-tartaric acid, increasing by 11-fold with 0.5 equivalent and 60-fold with 1 equivalent of diacid compared with free 3. However, using more than 1 equivalent of diacid disrupted the gel and decreased the emission intensity (Figure 4d). The authors propose that H bonding between pyridine of 3 and acid units is responsible for formation of the gel, confirmed by fourier transform infrared spectroscopy. Transient photoluminescence measurements revealed that the 1:1 3:(L)-tartaric acid gel possessed biexponential decay kinetics consistent with the gel being TADF, with lifetimes $\tau_{PL} = 20$ ns and $2.3 \mu$s, representing the prompt and delayed emission, respectively. The gel fibers were shown to be amorphous, with no change in the structure shown under high vacuum. The photoluminescence quantum yield of the xerogel is six times greater than of the neat film ($\Phi_{PL} = 36\%$ vs $6\%$). As the fibers were not damaged by high vacuum and retained their structure, these gels have a potential application in OLEDs.

### 2.3. TADF Supramolecular Metalloca ges

Supramolecular metalloca ges comprise metallic vertices linked by organic bridging ligands, encompassing a wide range of shapes and sizes depending on the coordination geometry of the linker and the choice of metal. Luminescent metalloca ges may be emissive due to their metallic ions, with the most common examples being cages based on Ir(III) and Ru(II) or through fluorescent organic linkers as such as those seen from Stang and coworkers with aggregation-induced emission (AIE) from tetraphenylethylene linkers. Zysman-Colman et al. explored the use of 3 in the construction of a supramolecular metalloca ge using Pd(II) ions as vertices. The cage could encapsulate organic dye molecules. Depending on the nature of the dye, either energy (PEnT) or electron (PeT) transfer to encapsulated guests could be observed. As 3 has a vector angle between pyridine rings of 93.5°, it is an appropriate ligand to make $M_6L_{12}$ cages, as evidenced by the work of Fujita and coworkers with ligands having similar vector angles of around 90°. DFT modeling of the metalloca ge 3-Pd (Figure 5c) predicted cubic structure with an internal volume of 6400 Å. The emission spectrum of 3-Pd in degassed DCM was red-shifted and broader compared with that of 3 and with a significantly lower photoluminescence quantum yield ($\lambda_{max} = 577$ nm, $\Phi_{PL} = 52\%$, 3-Pd; $\lambda_{max} = 555$ nm, $\Phi_{PL} = 4\%$ in DCM). While 3 shows triexponential photoluminescence decay with a prompt fluorescence, $\tau_{PL} = 15.1$ ns, and biexponential delayed fluorescence, $\tau_{PL} = 0.69$ and $9.80 \mu$s, components in degassed DCM, 3-Pd shows no delayed component, with a biexponential lifetime of $\tau_{PL} = 3$ and $30$ ns. DFT modeling of 3-Pd revealed that while the HOMO remains largely centered on the carbazole donor moieties, the LUMO is no longer distributed over the benzophenone...
acceptor and is instead centered on the Pd(II) and the coordinated pyridine ligands. The implication of an ligand-to-metal charge-transfer state to formally antibonding palladium d-orbitals is the origin of the significant quenching of the emission in the metallocage. The 3-Pd cage was used to encapsulate two xanthene dyes: fluorescein (F) and rose bengal (RB) (Figure 5d). Analysis of the host–guest complexes via electrospray ionization mass spectrometry (ESI-MS) showed that up to three molecules of neutral F and up to two molecules of RB in its quinoid dianionic form could be encapsulated. Upon addition of a solution of 3-Pd to a solution of F or RB, there were marked changes in the UV–vis absorption spectra of the dyes. Addition of 3-Pd (from 0.1 to 3.0 equiv.) to a solution of F resulted in the reduction of the absorption band at 522 nm of the open quinoid form, eventually producing a colorless solution, which suggests that the cavity of the 3-Pd favors the neutral form of F. No change in the absorption spectrum of RB was observed upon addition of 3-Pd (from 0.1 to 3.0 equiv.) to a solution of RB, thus confirming the presence of the open dianion quinoid form of RB, as corroborated by ESI-MS. For the 3-PdCF host–guest complex,
PET was observed, resulting in the formation of $[\text{F}]^{+} + [\text{3-Pd}]^{-}$, with complete quenching of the emission of both F and 3-Pd. In contrast, when 3-Pd was added to a DMSO solution of RB, photoinduced energy transfer was observed, with gradual quenching of the green emission from 3-Pd and a concomitant enhancement of the orange emission from RB at 582 nm. Stern–Volmer quenching analysis of the host–guest complex showed efficient photoinduced energy transfer, with a Stern–Volmer constant of $K_{SV} = 1.22 \times 10^{4} \text{M}^{-1}$ and a quenching rate constant of $k_q = 4.07 \times 10^{11} \text{M}^{-1} \text{s}^{-1}$. Förster energy transfer was proposed as the energy transfer mechanism.

2.4. Mechanically Interlocked Rotaxane–Emitter Species

Rotaxanes are a class of mechanically interlocked molecules, typically consisting of a linear guest species and a cyclic host species held together by noncovalent forces. Recently, Zysman-Colman, Goldup, Penfold, et al. reported the fine modulation of photophysical properties of a carbazole–benzophenone-based TADF emitter 4, a compound similar to 3 but with triazole groups in place of pyridine, upon interaction with macrocycle 5 to give rotaxanes 4⊂5 and 4⊂5₂ (Figure 6a).$^{[48]}$ $^1$H NMR and single-crystal X-ray diffraction analyses showed the triazole protons to be hydrogen bonded to the macrocycle bipyridine moiety (Figure 6b). Photophysical measurements in toluene probed the effect of mechanical bond formation on the optoelectronic properties. Complexation to the macrocycles led to an increase in the photoluminescence quantum yield under N₂ and versus the uncomplexed emitter (4: $\Phi_{PL} = 11\%$; 4⊂5: $\Phi_{PL} = 31\%$, 4⊂5₂: $\Phi_{PL} = 30\%$) along with a decrease in the $\Delta E_{ST}$; in addition, there was an observed increase in the photostability of these systems under UV irradiation. DFT calculations of these systems show that while the benzophenone-centered LUMO remains largely unchanged

![Figure 6. a) Structures of emitter 4 and macrocycle 5, along with host-guest systems 4⊂5 and 4⊂5₂. b) Close-contact interactions between the bipyridine nitrogen atoms of 5 and the C–H groups on the triazole group in 4⊂5 combined with other intramolecular interactions and a space filling model of 4⊂5. Reproduced under the terms of the CC-BY 4.0 license. Copyright 2021, The Authors, published by Wiley-VCH.](image)
upon complexation in 4⊂5 and 4⊂5₂, the HOMO is destabilized due to hydrogen bonding between the C–H bond on the triazole arms and the bipyridine nitrogen atoms of 5. In summary, the mechanical bond formed between 4 and 5 allowed for fine tuning of the emission, a decrease in the ΔEST, an increase in photoluminescence quantum yields, and greater photostability, all traits that are desirable when designing emitters, particularly for electroluminescent devices.

2.5. TADF Metal–Organic Frameworks (MOFs)

Metal–organic frameworks (MOFs) are extended supramolecular systems comprising metal ions or clusters held together with organic linkers via coordination bonds.¹⁴⁹ Their porous nature combined with the near-limitless diversity of the structure associated with different combinations of metals and organic linkers has led to significant interest in their exploitation in numerous applications, including gas storage and separation⁵⁰–⁵² and catalysis.⁵³,⁵⁴ A large subset of MOFs are luminescent and these have been used in sensing,⁵⁵–⁵⁸ bioimaging,⁴⁸,⁴⁹,⁵⁹,⁶⁰ light-emitting devices,⁶¹–⁶⁴ and as photocatalysts.⁶⁵–⁶⁹ Adachi et al. reported the first example of a TADF MOF. The MOF links a TADF emitter, 6, containing two carboxylate groups with zirconium-based vertices (Figure 7).⁷⁰ The linker 6 was chosen due to its small calculated ΔEST of 0.2 eV. For 6, there is a broad CT emission at 481 nm. The ΔEST of 6 was estimated to be 0.24 eV, in line with the calculations. Emission of 6 in 2 wt% PMMA-doped films showed a biexponential decay, with prompt and delayed lifetimes, respectively, of 18 ns and 199 ms, and a ΦPL of 39%, which decreased to 32% in the presence of air. The photophysical properties of MOF-1 were measured following activation of the MOF-1 at 400 K under vacuum. Compared with 6, the emission of MOF-1 is red-shifted and broader, centered at around 501 nm (Commission internationale de l’éclairage (CIE); 0.16, 0.18), with a slightly reduced ΦPL = 30%; the ΦPL decreased to 18% in the presence of air. The transient PL under N₂ showed biexponential decay with lifetimes of 17 ns and 180 μs, representing the prompt and delayed emission, respectively. The reduction in photoluminescence quantum yield in air coincides with the loss of the delayed component of emission due to the quenching of the triplet state via oxygen. The decrease in the lifetime and red-shifting upon MOF formation were suggested to arise from coordination to the electron-poor zirconium centers.

Halder and Wöll et al. reported the formation of MOFs from a diphenylamine–tetraphenylethylene TADF emitter, 7 (Figure 8a), which, upon incorporation into oriented thin films, displayed green electroluminescence.⁷¹ By orienting emitters within a film to give channels for charge carriers, the conductivity of a system can be improved to be closer to that of a single crystal.⁷² The formation of the MOF from 7 allows for a high concentration of emitters. The presence of the tetraphenylethene moiety, a well-known AIE building block, ensures that no concentration quenching is observed in the MOF. Dilute ethanol solutions of 7 are poorly emissive, attributed to nonradiative decay, resulting from the free rotation of the phenyl groups, with a lifetime of 1.5 ns. Powder samples of 7, however, showed delayed fluorescence (τPL = ≈200 μs). Upon forming a crystalline, oriented, thin-film MOF with zinc ions (MOF-2), the same delayed component is seen but with a significantly higher emission intensity. The TADF nature of this emission was demonstrated by variable temperature PL studies that showed an increase in intensity of emission for MOF-2 with increasing temperature, which is linked to the thermal upconversion of triplet excitons to form radiative singlets. The ΦPL of MOF-2 is 14%. The thin-film MOF was integrated into an OLED, the composition of which is shown in Figure 8b. The device showed a high turn-on voltage of 5.8 eV, demonstrating a rather low luminance of 270 cd m⁻² at 14 V. Time-dependent density functional theory

![Figure 7. a) Structure of 6 with calculated HOMO and LUMO plots. b) Structure of MOF-1 with 6 binding via carboxylic acid groups to zirconium centres. Adapted with permission.⁷⁰ Copyright 2018, Royal Society of Chemistry.](image-url)
(TD-DFT) calculations provided support for the nature of TADF emission from MOF-2. The authors propose that a hot-exciton mechanism is the source of TADF, whereby an exciton forms at the T2 state and undergoes RISC to the S1 state; this is possible due to the small energy gap between T2 and S1 compared with the much larger energy gap between T2 and T1. Incorporation of 7 as a component of the crystalline MOF-2 results in decreased molecular vibrations, thus giving rise to high ISC and rISC rates of $6.94 \times 10^6 \text{s}^{-1}$ and $1.91 \times 10^7 \text{s}^{-1}$, respectively, at 300 K.

2.6. TADF Metallocycles

Photodynamic therapy (PDT) uses photosensitizers in conjunction with light to generate reactive oxygen species which oxidize and damage tumor cells. PDT is viewed as being less invasive and more targeted than some other anticancer therapies. Incorporation of platinum and ruthenium in PDT photosensitizers has the added benefit of the cytotoxicity these metals display, allowing them to also act as chemotherapy drugs in conjunction with PDT. Song, Liu, and coworkers designed a novel organic photosensitizing ligand, 8, and incorporated it into a triangular metallocycle with Pt(II) vertices (Figure 9). The resulting metallocycle, 9, was then used to form nanoparticles and their use as anticancer agents was evaluated in vitro. The photo-physical properties of the ligand and metallocycle were assessed in degassed ethanol solution; upon excitation at 508 nm, 8 was found to be highly fluorescent, with $\lambda_{PL}$ at 569 nm, $\Phi_{PL}$ of 78%, and $\tau_{PL}$ of 8.65 ns. Even in aerated solution, the emission remains strong with $\Phi_{PL} = 60\%$ and $\tau_{PL} = 8.30$ ns. The high $\Phi_{PL}$ was largely preserved upon metallocycle formation at 69%. There is an accompanying blue-shift of the emission to 550 nm and a double-exponential lifetime with an average $\tau_{PL} = 8.65$ ns.

Transient absorption spectroscopy studies ($\lambda_{exc}$ at 532 nm) revealed lifetimes of 1.87 and 1.76 $\mu$s for 8 and 9, respectively, suggesting that these species could generate reactive oxygen species. The $\Delta E_{ST}$ value of 8, measured in 2-methyltetrahydrofuran, was determined experimentally to be 0.085 eV, prompting a temperature-dependant emission study of the organic ligand, which revealed that the emission intensity for 8 increases with temperature, suggesting a TADF mechanism. This same effect was seen for 9, which suggests that this TADF property is preserved upon complexation, which the authors propose is the source of the relatively long luminescence lifetime of the emitters. The ability of 8 and 9 to generate singlet oxygen was assessed in ethanol solution versus a TPPS standard (TPPS = meso-tetrakis(p-sulfonato-phenyl) and both displayed high efficiency for oxygen generation with quantum yields of $\Phi = 95\%$ and 86%, respectively. It is suggested that the small $\Delta E_{ST}$ value increases the rate of ISC to the triplet state; this in turn increases the rate of $^{1}O_2$ production. For in vitro studies, 30 nm nanoparticles of 9 were prepared via nanoprecipitation. These nanoparticles were stable for 4 days in a culture medium, and in blood no significant hemolysis was seen; both were promising results for the biocompatibility of these species. The uptake of 9 nanoparticles by HeLa cells was confirmed by confocal laser scanning microscopy with fluorescent dying of the cell nuclei (cells dyed with 4',6-diamidino-2-phenyl-indole = DAPI). Strong fluorescence was seen from the HeLa cells after 12 h, with the green emission of the nanoparticles overlapping with the blue emission from DAPI, confirming uptake of the nanoparticles into the nuclear region of HeLa cells. The accumulation of the nanoparticles within the HeLa cells is attributed to the positive nanoparticles being attracted to the negatively charged cell membrane. Singlet-oxygen generation by the nanoparticle containing HeLa cells upon irradiation was confirmed using...
singlet-oxygen sensor green. The half-maximal inhibitory concentration (IC50) of 8 when irradiated in cells was 1.0 μg/mL, and for 9 nanoparticles it was 0.5 μg/mL, which is lower than other PDT agents containing BODIPY dyes and porphyrins.[77,78] The greater toxicity of the metallocycle nanoparticles is due to the strong binding interactions of the Pt(II) centers with DNA. The combination index of the PDT/chemotherapy combination offered by the nanoparticles, a measure of how synergistic the two approaches are, was calculated to be 0.13. This value indicates that there is good synergy between the PDT offered by the 8 ligands and chemotherapy effects of the Pt(II) centers.

2.7. Confining Carbon Dots within Zeolites to Switch on TADF

Since their discovery in 2004, carbon quantum dots have gained significant interest due to their remarkable photoluminescence and electrochemical properties combined with their low cost and toxicity/environmental impact.[79–83] Emissive carbon dots comprise a carbon nanoparticle core, whose surface is passivated with organic ligands, with the choice of surface groups being key to the emissive nature of these materials.[84] Yu and Li demonstrated that the confinement of carbon dots within zeolites allowed these materials to display efficient and long-life TADF emission (Figure 10).[85] Three systems (CD1-3) were synthesized via solvothermal conditions, forming carbon dots in situ as the zeolite crystallized, which then resulted in embedded carbon dots within the zeolite structure framework. CD1 used triethylamine (TEA) as the structure-directing agent (SDA) in combination with aluminum triisopropoxide, phosphoric acid, triethylene glycol, and hydrofluoric acid to give hexagonal crystals, with carbon dots of diameter 3.7 nm monodispersed within the structure as confirmed by transmission electron microscopy imaging. CD2 was prepared similarly but with 4,7,10-trioxa-1-13-tridecanediamine as the SDA and no hydrofluoric acid. Here the dispersed carbon dots possessed an average diameter of 3.5 nm. CD3 was prepared similarly to CD2 but with the addition of dimagnesium phosphate, giving dispersed carbon dots of an average diameter of 3.4 nm. Following excitation at 370 nm, the three structures show deep blue emission (C1: 430 nm, PL = 15%, CIE (0.17,0.13); C2: 440 nm, PL = 52%, CIE (0.17,0.14); C3: 425 nm, PL = 23%, (0.17,0.13)). The larger photoluminescence quantum yields observed for CD2 and CD3 compared with CD1 were attributed to the use of 4,7,10-trioxa-1-13-tridecanediamine as the SDA, which was shown previously.
to passivate the surface of carbon dots, improving photostability and enhancing photoluminescence quantum yields.\textsuperscript{[86,87]} All three systems displayed delayed emission with lifetimes on the order of milliseconds (CD1: $\tau_{\text{PL}} = 350$ ms, CD2: $\tau_{\text{PL}} = 197$ ms, CD3: $\tau_{\text{PL}} = 216$ ms). The TADF character of these systems was confirmed by temperature-dependent transient photoluminescence decay measurements that showed enhanced emission of the delayed component with increasing temperature. The $\Delta E_{\text{ST}}$ values for the three complexes were measured and found to be $\Delta E_{\text{ST}} = 0.22$ eV for CD1, $\Delta E_{\text{ST}} = 0.23$ eV for CD2, and $\Delta E_{\text{ST}} = 0.22$ eV for CD3. The emergence of TADF following the confinement of the carbon dots within the zeolite matrices was attributed from n to $\pi^*$ transitions in carbonyl and imine bonds within the carbon dots. These transitions associated with triplet states are of different orbital types than those associated with the CT singlet state. This leads to greater spin–orbit coupling that facilitates rISC. The emission from these systems remains long in air as the zeolite matrix impedes oxygen from quenching the triplet states of the embedded carbon dots.

Yu and Li went on to report the synthesis of zeolite-confined carbon dots, which, depending on the template used in the synthesis of the CDs, showed either RTP or TADF.\textsuperscript{[88]} The TADF zeolite-confined carbon dot, CD4, was prepared using a 4,7,10-trioxa-1-13-tridecanediamine template. Upon excitation at 370 nm, CD4 showed emission at $\lambda_{\text{PL}} = 440$ nm, with an associated quantum yield of $\Phi_{\text{PL}} = 29\%$. The photoluminescence from CD4 showed some excitation dependence, with the strongest emission seen upon excitation at 350 nm (Figure 11). The authors assigned TADF to this material based on the long-lived delayed lifetime ($\tau_{\text{PL}} = 153$ ms) at room temperature under air atmosphere, which was confirmed by temperature-dependent transient photoluminescence decay measurements, which showed an increase in the delayed emission component with increasing temperature (no prompt lifetime quoted). The $\Delta E_{\text{ST}}$ was measured to be 0.18 eV, which is compatible with rISC at room temperature. The $\Delta E_{\text{ST}}$ of the unconfined carbon dot was slightly larger at 0.21 eV in the mother liquid; however, no long-life emission is observed at room temperature.

The authors attribute the switching on of TADF upon encapsulation of the carbon dot to be due to both the suppression of nonradiative decay pathways and stabilization of the triplet state by the zeolite host by suppressing nonradiative decay processes such as molecular vibrations and rotations.

More recently, Yu and Li reported the encapsulation of two different carbon dots within the same zeolite matrix, whereby varying the ratio of the two within the framework allowed for the tuning of the emission wavelength.\textsuperscript{[84]} The ratio of the two carbon dot precursors, m-phenylenediamine and 4,7,10-trioxa-1-13-tridecanediamine, was varied from 0, 0.007, 0.0014, and 0.042 to give four different compositions of carbon dots encapsulated within zeolites, CD5-8, respectively. Scanning electron microscopy revealed similar polyhedral morphology for all compositions, whereas transmission electron microscopy showed good dispersion of the carbon dots within the frameworks. Increasing the ratio of m-phenylenediamine to 4,7,10-trioxa-1-13-tridecanediamine led to a red-shifting of the emission (CD5-8: $\lambda_{\text{PL}} = 425$, 484, 498, 515 nm, respectively). All four composites displayed very long-life delayed fluorescence, with photoluminescence quantum yields and delayed lifetimes increasing but prompt lifetime decreasing as the m-phenylenediamine content increased (CD5: $\tau_{\text{PL}} = 24.43$ ns, 271 ms, CD6: $\tau_{\text{PL}} = 37.54$ ns, and 578 ms, CD7: $\tau_{\text{PL}} = 11.76$ ns and 801 ms, CD8: $\tau_{\text{PL}} = 7.70$ ns and 860 ms. CD5-8: $\Phi_{\text{PL}} = 20.9\%$, 25.1\%, 37.1\%, 42.0%, respectively). Temperature-dependant time-resolved emission studies of the composites revealed the expected thermal activation of the delayed emission, confirming a TADF mechanism. The $\Delta E_{\text{ST}}$ values for CD5 and CD8 are 0.20 and 0.14 eV, respectively. A mechanism was proposed by the authors, as shown in Figure 12, whereby within systems CD6, CD7, and CD8 Förster resonance energy transfer (FRET) from the 4,7,10-trioxa-1-13-tridecanediamine carbon dots to the m-phenylenediamine carbon dots within the zeolite framework occurs, which then emit via a TADF mechanism. The FRET efficiency increases as the ratio of m-phenylenediamine to 4,7,10-trioxa-1-13-tridecanediamine increases.

2.8. Organic Macrocycles and Cages

TADF materials commonly comprise twisted donor–acceptor species, with the separation of frontier molecular orbitals leading...
to a reduced exchange integral and consequently a small $\Delta E_{ST}$ that results in efficient rISC. Huang and coworkers reported the synthesis of two supramolecular donor–acceptor-based structures, which enforce highly twisted conformations, decreasing $\Delta E_{ST}$, while also providing an increasing rigidity to the emitter moieties that result in narrow emission.

Huang and coworkers reported the one-pot synthesis of a family of cyclic organic species showing deep-blue TADF in doped thin films and room-temperature phosphorescence (RTP) as a crystalline powder. The cyclic species is made up of two triphenylamines linked by two sulfonyl groups. The donor groups may be substituted with a tert-butyl or methoxy group to allow tuning of the emission ($10 = H$, $11 = 'Bu$, $12 = OMe$).

**Figure 13.** DFT calculations on these systems showed the LUMO to be largely distributed over the diphenyl sulfone acceptors and the HOMO localized over the triphenylamine donors. The three emitters showed photoluminescence in DCM at 427, 443, and 469 for 10, 11, and 12, respectively, with measured $\Delta E_{ST}$ values of 0.34, 0.31, and 0.24 eV, respectively. The observed red-shift of the emission corresponds with the progressive strengthening of the donor, which also leads to a destabilized HOMO. Transient photoluminescence decays of doped thin films [5 wt% mCP (mCP = 1,3-bis(carbazol-9-yl)] of 10, 11, and 12 all showed prompt (3.78 ns, 5.55 ns, and 9.97 ns) and delayed (10.9, 16.1, 101.3 μs) fluorescence under inert atmosphere. The delayed contribution to fluorescence increases with increasing electron-donating ability of the donor ($10 = 10.5\%$, $11 = 22\%$, $12 = 80\%$), attributed to greater localization of the HOMO on the donor, leading to decrease in the $\Delta E_{ST}$. The photoluminescence quantum yields of the emitters are between 8% and 14% under N₂. Interestingly, emitters 11 and 12 also display RTP in the crystalline powder, with lifetimes of 4.55 and 5.09 ms, respectively. The origin of the RTP was rationalized by the authors using TD-DFT calculations to intermolecular interactions between adjacent donor and acceptor units.

The same group later reported the one-pot synthesis of an organic cage that displayed TADF, 13. The structure of 13 was unambiguously determined by X-ray crystallography, having $C_{3v}$ symmetry with two triphenylamine molecules linked by three sulfonyl groups, in an analogous manner to the cyclic compound discussed earlier. This structure places the donor triphenylamines next to the acceptor diphenylsulfone units in a twisted fashion. This system showed narrow (full width at half maximum: 34 nm/0.24 eV) deep-blue emission centered at 414 nm with a small Stokes shift of 43 nm, both linked to the rigidity of the cage. The measured $\Delta E_{ST}$ of 13 is 0.18 eV, which is significantly smaller than the macrocycles 10-12. The photoluminescence quantum yield of 13 is only 15%. The time-resolved transient photoluminescence decay shows both prompt ($\tau_{PL} = 8.65$ ns) and delayed ($\tau_{PL} = 8.6$ μs) fluorescence at room temperature, with the delayed component increasing in proportion at elevated temperatures, confirming TADF. OLED devices fabricated of 13 showed a maximum external quantum efficiency (EQE_{max}) of 2.63%. The electroluminescence spectrum, peaking at 413 nm, retains a narrow FWHM (35 nm/0.25 eV).

**2.9. Organic Dots**

Organic dots (also referred to as organic nanoparticles) are systems comprising hydrophobic organic cores, decorated with hydrophilic chains, which allow for high water solubility.
Over the past few years, multiple groups have reported examples of such micelle-like species that are emissive via a TADF mechanism. There are multiple examples of 4CzIPN and structurally similar emitters,[9,92–95] as well as phenoxyamine derivatives[14,96–97] being used as the organic cores, along with other TADF emitters, which may be decorated with hydrophilic chains, as shown in Figure 15. The hydrophilic chains can be covalently linked to the organic emitter or the two components may be suspended in solution and mixed to self-assemble the separate components into a micelle. The hydrophilic chains used vary, with a popular choice being derivatives of polyethylene glycol, boasting a number of benefits including low cost, low toxicity, and excellent water solubility.[9,92–94,98] An example from Tang and Zhao shows the use of peptide chains to facilitate passage through cell and nuclear membranes,[96] which illustrates the main application of these TADF organic dots as emissive biological probes. The use of hydrophilic chains is not essential, with examples where nanoparticles/nanorods formed due to aggregation of organic emitters in aqueous media[95,99] or the use of Zn(II) ions to initiate a complexation of an organic species to give a water-soluble complex.[97] Biological autofluorescence makes it hard to distinguish emission from the probe from background cell emission when the excited state of the probe is too short-lived. The use of phosphorescent complexes that possess long-life triplet excited states is somewhat problematic due to the toxicity of the heavy metal complexes. TADF provides an answer to this problem, being both long-lived and not as toxic as heavy metal complexes; however, the presence of oxygen in biological systems quenches triplet states and can lead to a reduction in photoluminescence quantum yield. Shielding of the TADF emitters within organic dots allows for high photoluminescence quantum yields under ambient atmosphere in solution that are insensitive to the presence of oxygen. The wide range of emitters and, where used, water-solubilizing groups, demonstrates the tunability and versatility of this class of supramolecular materials.

2.10. Self-Assembled Fibers
Qi et al. reported a novel TADF emitter, 14, that displays AIE and can self-assemble into microfibers (Figure 16).[100] Increasing the water fraction beyond 40% of a DMF solution results in an increase in the emission intensity attributed to aggregation of 14 as it becomes less soluble in solution. By varying the water content, the size of the aggregates can be adjusted, from 20 nm in 40% water content up to 10.0 μm in 90% water content solution. The self-assembly of 14 into fibrils was achieved by slowly evaporating a THF:CH₂Cl₂ (1:1 v/v) solution of 14 on a quartz substrate. Differences in the photophysical properties
can be seen between the powder and thin film, with a small red-shift in the emission from 640 to 645 nm and an increase in the photoluminescence quantum yield from 13% to 26%. Scanning electron microscopy (SEM) and polarizing optical microscopy (POM) showed that the powder formed needle-like crystals, whereas the thin film contained microfibers that were up to several hundred micrometers long and several hundred nanometers wide (Figure 16b). The TADF properties of the thin film were confirmed by transient photoluminescence decay measurements that showed the presence of prompt (2.3 ns) and delayed (10.0 μs) fluorescence and temperature dependence of the delayed component of the emission. The ΔE<sub>ST</sub> of free 14 was calculated via DFT to be 0.61 eV, which, despite the large energy difference, is of similar magnitude to that of a previously reported TADF compound.[101]

3. Conclusion

We have presented a diverse set of examples of supramolecular systems that exhibit TADF. This area, however, remains rather underexplored, and the potential of these systems is not yet fully realized. What is clear is that the photophysical properties can be finely tuned within the self-assembly. This sometimes leads to new reactivity, to a loss of TADF, or in other examples, to a turn-on of the TADF mechanism. We hope that as the chemical space associated with compounds that exhibit TADF continues to expand, so too will related supramolecular systems. The examples presented in this mini-Review demonstrate the potential of supramolecular TADF systems and we hope that this will spur further development in field.

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Conflict of Interest

The authors declare no conflict of interest.

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Thomas Comerford graduated from Newcastle University with M.Chem. in 2018, spending his final year working under the supervision of Dr. Michael Hall on the synthesis of helically chiral BODIPY dyes. He then took up a research associate position at Biome Bioplastics, working on the scale-up of biodegradable polymer synthesis at the Green Chemistry Centre of Excellence at the University of York. Tom joined the Zysman-Colman group in 2019 as part of the EaSI-CAT CDT and is interested in the design and synthesis of supramolecular metalallocages as molecular flasks for photocatalysis.

Eli Zysman-Colman obtained his Ph.D. in Chemistry from McGill University in 2003 with Professor David N. Harpp. He then completed two postdoctoral fellowships, one in supramolecular chemistry with Professor Jay Siegel at the University of Zurich and the other in inorganic materials with Professor Stefan Bernhard at Princeton University. He joined the Université de Sherbrooke as an assistant professor in 2007. In 2013, he moved to the University of St Andrews, where he is currently professor of Optoelectronic Materials, fellow of the RSC, and the holder of Royal Society Leverhulme Trust Senior Research Fellowship.