

## Communication

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# Supramolecular Gel Phase Controlled [4+2] Diels-Alder Photocycloaddition for Electroplex Mediated White Electroluminescence

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Supporting Information Placeholder

**ABSTRACT:** Diels-Alder photocycloaddition of 9-phenylethynylanthracene results in multiple [4+2] and [4+4] cycloaddition products in solution, which can be controlled to form specific products under restricted environment. We have exploited the gel phase of a 9-phenylethynylanthracence derivative as a confined medium to specifically yield the [4+2] cycloadduct in >90% yield. The photocycloadduct (*anti*-form), exhibited a blue emission with CIE chromaticity of x = 0.16/y = 0.16. Construction of organic light emitting device with the photocycloadduct, using a carbazole based hole transporting host, resulted in white light emission with CIE chromaticity of x = 0.33/y = 0.32. This observation not only highlights the use of gel chemistry to achieve the otherwise difficult to obtain photoproducts but also underlines their potential in optoelectronic device fabrication.

Confinement of molecular systems yielding regio- and stereospecific reaction products, has been reported in several cases.<sup>1</sup> An example is the Diels-Alder photocycloaddition of anthracene<sup>2</sup> and its 9-phenylethynyl derivative.<sup>3</sup> While anthracene results in *anti* and *syn* isomers of [4+4] photocycloadducts in the absence of oxygen,<sup>4,5</sup> 9-phenylethynylanthracene in principle should form four different products (Scheme 1). Though these products are allowed by selection rules, the unfavorable geometries enable only [4+2]*anti* (64%) and [4+4] *anti* (23%) addition.<sup>3,6</sup> However, energetically unfavorable products were specifically obtained under confined conditions.<sup>7,8</sup> Majority of these reports have been limited to the synthesis of selective products on confined surfaces, however, practical application of these types of products are yet to be demonstrated.<sup>9</sup>

### Scheme 1. Photocycloaddition of 9-phenylethynylanthracene in solution.<sup>6,7</sup>



Supramolecular gels have been shown to be a suitable medium for crystallization of molecules.<sup>10</sup> H-Bonding and  $\pi$ -stacking in the gel state have advantage over solution state to obtain certain stereospecific products with high yield.<sup>11</sup> Herein, we establish that molecular self-assembly and gel chemistry can be used to control Diels-Alder photocycloaddition to achieve a specific product in high yield, which can be used for fabricating white light emitting devices. Though white light emission is a topic of current interest in lighting applications,<sup>12-14</sup> single molecular white electroluminescence is rarely observed.<sup>14,15</sup> Electromer and electroplex formation by the emitting species and the hole transporting material is the reason for white electroluminescence.<sup>14a,15</sup> Such electrically excited complexes are akin to excimers and exciplexes observed in the case of photoluminescence and has long wavelength mixed emission, providing balanced color coordinates required for white light emission.



In this study, a 9-phenylethylanthracene derivative 1 was designed which facilitates H-bonded self-assembly<sup>16</sup> in methyl cyclohexane (MCH, 3.3 µM, Figure S1a). Plots of the fraction of aggregate ( $\alpha_{agg}$ ) against the temperature could be fitted to a cooperative self-assembly pathway. The enthalpy of nucleation of the assembly  $(\Delta H_{nucl})$  was found to be -9 kJ/mol and the elongation temperature was 303 K (Figure S1b and Table S1). The molecule 1 in hot MCH (1.19 w/v) upon sonication and cooling resulted in a reversible gel (Figure 1a,b) having a storage modulus (G') of one order of magnitude greater than that of the loss modulus (G'') (Figure 1d). Atomic force microscopy (AFM) and transmission electron microscopy (TEM) analyses revealed tape like morphology (Figures 1e, S2 and S3). X-ray diffraction (XRD) of the xerogel exhibited peaks corresponding to *d*-spacing of 47.9, 21.5 and 12.5 Å. First three peaks followed the reciprocal relationship with a ratio of 1:~2:~4, which correspond to multidimensional lamellar packing. The peaks corresponding to 3.82 and 4.20 Å indicate  $\pi$ -stacking and long range ordered inter-molecular H-bonding, respectively. The peaks corresponding to 8.69, 8.28, 7.49, 6.91, 6.31, 5.80, and 5.28 Å represent the interdigitated H-bonded assembly where the anthracene moiety of the molecule is overlapped with the triple bond (Figure 1 and S4). This is evident from temperature controlled absorption changes in 230-260 nm region (Figure S5a), where the band corresponding to <sup>1</sup>B<sub>b</sub> transition dipole of anthracene, parallel to the molecular axis is not being polarized via aggregation, and remained nearly unsplit.<sup>17</sup> The steady state fluorescence spectrum (Figure S5b) of the gel did not show excimer emission,<sup>18</sup> as supported by time correlated single photon counting experiment (Figure S5c). The fluorescence decay profile of the gel was fitted with a bi-exponential decay having lifetime values of 1.95 and 4.00 ns.



**Figure 1.** (a, b) Photographs showing reversible gelation of **1** in MCH and (c) the gel after photoirradiation. (d) Rheological behavior of the gel. (e, f) TEM images of the gel before and after photoirradiation. (g) AFM image of the self-assembled **1** in MCH. (h) The height profile. (i) Molecular assembly and structural change of **1** before and after photoirradiation.

The absence of any emissive species with lifetime value of >10 ns indicates that no prominent overlap between two anthracene cores are present in the molecular assembly. Thickness of a single layered sheet evaluated by AFM analysis is 5.2 nm, corresponding to a monolayer of the assembly (Figures 1g, h). The interdigitated long-range arrangement of **1** is ideally suited for the specific [4+2] Diels-Alder photoreaction (Figure 1i) as established by the irradiation of a gel of the molecule **1**, with 365 nm light. In the process, the gel turned into a solution (Figure 1c), which could not be reversed to a gel by irradiation at shorter wavelength or by cooling.

HPLC profile of the irradiated solution showed a major peak corresponding to a 91% conversion of the starting compound (Figure S6a). The <sup>1</sup>H NMR profile of photoirradiated gel suggested 90-92% formation of photoadduct (Figure S6b), whereas, the xerogel and the solution state photoreaction showed low conversion and less stereoselectivity, respectively (Figure S7). The <sup>1</sup>H NMR and NOESY spectra confirmed the anti-[4+2] cycloaddition (Figures S8, S9). This geometry disturbs the assembly of the molecule resulting in the dissolution of the gel. AFM and TEM analyses of the photoadduct showed ill-defined particles as shown in Figure 1f and S10b, justifying the destruction of the one-dimensional H-bonded assembly of the gel phase. This observation is confirmed by the XRD analysis of the gel after photoirradiation, which shows decrease in the number of diffraction peaks (Figure S10). Comparison of the FT-IR spectra (Figure S11) before and after photoirradiation revealed that H-bonded N-H stretching frequency of the gel ( $v_{N-H}$ =  $3261 \text{ cm}^{-1}$ ) was shifted to  $3279 \text{ cm}^{-1}$  which is close to the monomeric state (3284 cm<sup>-1</sup>), indicating the breakage of the H-bonding.

The absorption and emission features of the gelator 1 and its photoadduct in chloroform  $(3.3 \ \mu\text{M})$  are shown in Figure 2a-c. The absorption spectrum of 1 showed peaks at 266, 323, 366, 386, 406 and 429 nm. However, in the case of photoadduct, the absorption bands at 429 and 406 nm were disappeared with the appearance of

two relatively less intense bands at 393 and 373 nm and an intense band at 299 nm. The gelator **1** exhibited two intense emission peaks at 450 and 475 nm, whereas the photoadduct showed an emission with maximum at 440 nm. The fluorescence quantum yields of **1** and the photoadduct were 0.79 and 0.57, respectively. The fluorescence lifetime analysis of **1** and its photoadduct in chloroform exhibited a single exponential decay with lifetime values of 2.91 and 4.28 ns, respectively (Figure 2c).



**Figure 2.** Comparative properties of **1** (red) and photoproduct (blue). (a) Absorption, (b) emission, (c) fluorescence lifetime decay profiles in chloroform and (d) cyclic voltammograms in dichloromethane.

Cyclic voltamograms of 1 and the photoadduct (Figure 2d) suggest that EHOMO of the photoadduct is (-5.54 eV) more stabilized than that of the gelator 1 (-4.79 eV) due to loss of extended  $\pi$ -conjugation (Figures S12, S13). Similar observation has been made in ELUMO of the photoadduct (- 2.59 eV) and 1 (-1.98 eV) as tabulated in Table S2. The device characteristics of the gelator 1 was investigated by fabricating solution processed single layer organic light emitting devices (Figure S14).<sup>19a</sup> The emitting layer (EML) comprised of the gelator 1, the carbazole derivatives PVCz or SB219b as the hole transporting materials (HTM) and the oxadiazole derivative PBD as the electron transporting material (ETM). PVCz was a better HTM than SB2 as indicated by the higher Lmax value of the devices with the former (Table S3). The intensity of the EL spectra increased upon increasing the applied voltage, however, the structure of the spectra remained unchanged. The  $L_{\text{max}}$  obtained for the 1+PVCz was 1244 cd/m<sup>2</sup> at 16.5 V, and 231 cd/m<sup>2</sup> for 1+SB2 at 14.5 V. In both cases, blue EL with CIE coordinates of (0.16, 0.16) and (0.16, 0.13), respectively were obtained and remained almost constant at any applied voltages (Figure 3a,c,e,f).

Interestingly, the device fabricated using the photoirradiated gelator with PVCz (or SB2) as EML and PBD as ETM exhibited broad EL spectra between 400-750 nm, with maxima at 445 and 573 nm (Figure 3b,d), resulting in white electroluminescence. The performance of the PVCz based device is better when compared to that with SB2. The structure of the EL spectra remained unchanged while the intensity was increased with increase in the voltage (Figures S15, S16). When SB2 was used as HTM, the CIE coordinate of the white light emission was x = 0.33/y = 0.32, indicating high color purity. On the other hand, the value of the CIE coordinates was x = 0.28/y = 0.25 when PVCz was used as the HTM (Figure 3e,f).

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Figure 3. Device characteristics (luminance and current density vs voltage plot) of (a) 1 and (b) photoadduct. Electroluminescence spectra of OLED made of (c) 1 and (d) photoadduct as emitter with SB2 (red) and PVCz (blue) as hole-transporting materials. (e) Photographs of electroluminescent devices fabricated with the gelator 1 (i, ii) and photoadduct (iii, iv). (f) EL color coordinates with those device made of 1 and the photoadduct.

For a deeper understanding of the origin of the white electroluminescence, the EL and the PL spectra of the devices fabricated with the gelator **1** before and after photoirradiation using PVCz and SB2 as the HTM were compared (Figure 4a-d). Both EL and PL spectra of the gelator before photoirradiation are comparable, which exhibited blue emission with maxima at 450 and 470 nm. Similarly, the PL spectra of the devices fabricated using both PVCz and SB2 as HTM showed blue emission with a maximum at 450 nm and a shoulder band at 475 nm. However, the EL spectra in both cases exhibited a broad emission at 600 nm, in addition to the sharp emission at 450 nm. Notably the intensity of the broad 600 nm emission was less when PVCz was used as the HTM.

Since the PL emission of the photoirradiated gelator was blue, we expected a similar emission characteristic for the EL spectrum as well. Instead, we observed an electrogenerated white light emission, which may not be the property of the photoadduct alone. Furthermore, film state absorption of EML did not show any new band in the presence of the photoadduct (Figure S17), which is an indication of the absence of ground state charge transfer between the emitter and the HTM. Therefore, the green and red emission required for the white emission must have generated through the interaction between the photoadduct and the transporting materials under the applied electric field. Probably, an electron transfer from the HTM to the photoadduct could generate an excited complex called electroplex, similar to a photoexcited exciplex (Figure 4e). Such an emission by the electrically excited electroplex can be visible only by an EL spectrum and not by photoexcited PL spectrum.<sup>15a</sup>

The energy level of the electroplex is lower than that of the singlet excited state of the photoadduct (Figure 4e). This electroplex

(photoadduct<sup>-</sup> + HTM<sup>+</sup>) formation is in analogy with previous reports and should be responsible for the green-to-red emission along with the blue emission of the photoadduct, leading to the white light emission (Figure 4f).<sup>15,20</sup> To establish the electroplex formation between the emitter and the HTM, we fabricated devices by mixing them together with an electron transporting layer ([1,1':3',1"-terphenyl]-4,4"-diylbis(diphenylphosphine oxide), BPOBP)<sup>21</sup> separately coated in solution processed double layer devices (Figure S18). As expected, the device with emitter 1 showed blue emission, whereas the device made out of the photoirradiated gelator exhibited white light emission, underlining the electroplex formation between the photoadduct and the carbazole based compounds (Figure S19). The role of [4+2] photocycloadduct in the electroplex formation is confirmed by the comparatively weak bluish-white electroluminescence with a photoirradiated 9-phenylethynyleneanthracene (9PEA) having a mixture of photoadducts (Figure S20).



**Figure 4.** Comparative PL and EL spectra of devices made of (a) 1+PVCz, (b) 1+SB2, (c) photoadduct+PVCz and (d) photoadduct+SB2 (e) Schematic representation of the electronic configuration in molecular orbitals of the electroplex host/emitter. (f) Proposed energy profile for the electroluminescence response of the photoadduct, exciplex, and electroplex.

In conclusion, the gel phase of the molecule **1** creates enough confinement, selectively yielding the [4+2] photocycloadduct. Light emitting devices fabricated with the gelator **1** exhibited a blue emission, whereas the device fabricated with the photoirradiated gelator showed a white electroluminescence. The generation of the green and red emission required for white luminescence is explained on the basis of an electroplex formation between the photocycloadduct and the hole transporting material within the device structure under an applied electric potential. This report highlights the role of H-bonded supramolecular assembly and gelation of functional molecules to achieve exclusive photoproducts for their use in photonic device application.

# ASSSOCIATED CONTENT

## Supporting Information

General methods, synthetic procedures, gelation, photoirradiation, OLED fabrication and additional figures. This material is available free of charge via the Internet at <u>http://pubs.acs.org</u>.

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