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Structural Feature and Optical Property of a Carbazole-Containing Ethene as Highly Emissive Organic Solid

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Carbazole derivatives are a kind of versatile materials especially for optoelectronic applications in light of their activity in both electronics and optics. To suppress the luminescence quenching effect in the condensed phase, we constructed a carbazole derivative with aggregation-induced emission characteristic. The highly emissive organic solid of carbazole-substituted ethane was facilely prepared and thoroughly tested. Through inspection of the geometric structure and packing motifs of the crystalline materials, the severe twisted conformation and the absence of strong intermolecular $\pi-\pi$ interactions are found to account for the extreme solid state quantum yield. The highly blue emissive crystal fibres exhibit optical waveguide property. Electroluminescence (EL) study reveals the hole-transporting nature of the material.

Introduction

Carbazole is a nitrogen-bridged biphenyl isoelectronic to fluorene. Oligomers and polymers based on carbazole have been widely investigated as materials for optoelectronic applications such as xerography, organic transistors, organic light-emitting diodes (OLEDs) and photovoltaic devices. They possess higher HOMO energy levels than that of fluorones, which is favourable of hole transporting abilities. A number of carbazole derivatives have been designed and synthesized for hole-transporting layers and host materials for red, green and even blue triplet emitters. However, the carbazole derivatives themselves used as light-emitting materials have been suppressed by the aggregation-caused quenching (ACQ) effect. This problem is also identified as a major concern for most of the common organic chromophores. From a chemical standpoint, the problem can be solved or alleviated by designing sterically hindered compounds. However, because the luminophores have an inherent tendency to aggregate in the condensed phase, the approaches are basically working against a very natural process and have thus ended up with only limited success and partial control. In a sharp contrast, using the idea of aggregation-induced emission (AIE), the intrinsic aggregation is no longer a suffering process. Oppositely it takes advantage of the aggregation process, and thus does not have any issues of temporal or spatial instabilities. We have explored high-tech applications of the AIE luminogens as, for example, fluorescence sensors (for explosive, ion, pH, temperature, viscosity, pressure, etc.), biological probes, and active layers in the OLEDs.

Although luminogens with AIE characteristic are no longer novel staffs after years of massive research, the development of more fantastic and practical ones suitable for different applications is significant. This indeed depends on the fully understanding of the mechanism and the structure particularity of the discovered ones. Analysing the structure and packing characteristics of them will help us to deepen the understanding and further facilitate the design of new highly emissive organic solids. In this report, we have successfully designed and synthesized a carbazole-containing tetrasubstituted ethene through a facile strategy. It showed typical AIE property and extremely high solid-state fluorescence efficiency. By digging into its structure feature, the propeller molecular geometry-caused loose packing was considered to respond for the highly luminescent ability in the solid state. Such a strategy is universal for most of the chromophores those originally being ACQ to become AIE.
Results and Discussions

Synthesis

Scheme 1 Synthetic route to 1,2-bis(N-ethyl-3-carbazole)-1,2-diphenylethylene (DCDPE).

Scheme 1 shows the chemical structure and synthetic route of the carbazole-containing AIE lumigen 1,2-bis(N-ethyl-3-carbazole)-1,2-diphenylethylene (DCDPE). Firstly, carbazole was incorporated into the ketone intermediate 9-ethyl-3-benzoylcarbazole by an AlCl$_3$ catalysed Friedel-Crafts acylation. Then the Zn/TiCl$_4$ catalysed Mcmurry coupling reaction produced the target in a high yield. Details of the synthetic procedures and characterization data are presented in the Supporting Information. The dye molecule was characterized by NMR spectroscopy and gave high signal:noise $^1$H and $^{13}$C NMR spectra. The reaction product gave M+ peaks at their high-resolution mass spectra, confirming the occurrence of the coupling reaction and the formation of the expected product. The purity of the product was also confirmed by elemental analysis with satisfactory result. McMurry coupling of BOC results in two different isomers with a ratio of about 1:1 calculated from the $^1$H NMR spectra. We obtained pure trans-isomer by recrystallization of the purified product via column chromatography in DCM/ethanol mixture. In the later experiments of thermal and electrochemical properties and EL performance tests, we used trans-form of DCDPE.

Optical properties

DCDPE is soluble in common organic solvents such as chloroform and THF but insoluble in water. It is practically nonemissive when molecularly dissolved in the solutions. As shown in Figure 1, the PL spectrum of DCDPE in THF is basically a flat line parallel to the abscissa. The fluorescence quantum yield in THF is as low as 0.22%, revealing that DCDPE is genuinely weak emitters as the free isolated molecular species. When a large amount of water is added into its THF solution, intense PL signals are recorded at 502 nm under the identical measurement conditions. The emission intensity starts to rise when the water fraction is larger than 75%, where the solvating power of the mixture is worsened to such an extent that the luminogen molecules begin to aggregate. From the molecular solution in THF to the aggregate suspension in 95% aqueous mixture, the fluorescence intensity increases ca. 412-fold and the fluorescence quantum yield value increases ca. 182-fold. The photograph given in the inset of Figure 1B clearly shows the changing from the nonemissive to emissive nature of the molecular species and the aggregative particles of DCDPE. Apparently, its emission is induced by aggregate formation. The aqueous mixtures are, however, macroscopically homogenous with no precipitates, suggesting that the aggregates are of nanodimension, as proved by the example of transmission electron microscope (TEM) images shown in Figure S6 and the level-off tails at the long wavelength region of their absorption spectra in the aqueous mixtures with high water contents due to the scattering effect of the dye nanoparticles (Figure S5). The measurement of the fluorescence quantum yields of DCDPE in the real solid state further validates its AIE activity, which is as high as 99% in the solid thin film determined by integrating sphere.

Structural analysis

In an effort to understand the mechanism operating in the AIE system, we checked the geometric structure and packing motifs of DCDPE in the crystalline state. The molecules crystallize easily from the mixture solvent of dichloromethane and ethanol.

Figure 2 shows the X-ray determined molecular structure. The molecule adopts a quite twisted conformation, with the four substituted aromatic rings arranged in a propeller-like shape along the ethene. The dihedral angles between each two aromatic planes of DCDPE are listed in Table 1, which are in the range from 54.5° to 83.3°. For example, the two phenyl
rings (represented by Cg(5) and Cg(8)) make a dihedral angle of 83.3°; while the two carbazole rings (represented by Cg(13) and Cg(14), not listed in Figure 2) are with a dihedral angle of 62.5°. The quite severe intramolecular torsion indicates such non-coplanar propeller-shaped structure will prevent the luminogens from packing in a close π–π stacking mode. This is proven by the packing motif of DCDPE in crystals. As can be seen from Figure 3, the intermolecular π–π interactions and other strong intermolecular interactions are totally absence. At the same time, the intramolecular bond rotational and vibrational freedom is still restricted by the intermolecular C–H•••π hydrogen bonds formed between the aromatic hydrogens and the aromatic rings. From figure 3 we can see one DCDPE molecule forms six C–H•••π hydrogen bonds with four of its neighbour molecules. Table 1 lists all the parameters of the intermolecular C–H•••π hydrogen bonds. The distances range from 2.75 Å to 2.86 Å and the angles (γ) of C(x)–H(xA) to Cg (j) (vector and normal to plane Cg (j)) range from 8.06° to 15.77°, within the standards of C–H•••π interactions of 3.0 Å and 30°.13 These kinds of weak interactions rigidify the molecular conformation and lock the vibrational freedom of the chromophores, but do not afford any energy decay channels such as the π–π interactions or excimer formation. The weak intermolecular interactions plus the severe twisted intramolecular geometry are the factors account for the extremely high quantum yield of DCDPE solids. Figure 4 shows the fluorescent microscopic images of the crystalline clusters of DCDPE under UV excitation. The crystalline samples are highly luminescent, emitting intense blue light upon photoexcitation. Glaring light emissions are seen at the ends of the microfibers indicating that an optical waveguide effect is operating in the light transmission process of the microfibers. The waveguide effect is usually observed in an optically anisotropic system, such as crystalline assembly. The crystallinity has significant effect on the emission behavior of DCDPE, which results in its fluorescence wavelength shifting from 483 nm in crystalline to 511 nm in amorphous, as shown in Figure S7. The fluorescence quantum yields in both states are as high as 99%, while in crystalline fibers it seems brighter due to the light scattering of the crystal facets. Such kind of efficient luminescent crystals are desired for light-emitting and laser applications.

**Fig. 2** Molecular structure of DCDPE obtained by X-ray diffraction. Cg(i) stands for aromatic planes (π rings).

**Fig. 3** Packing arrangement in the crystal structure of DCDPE, with C–H•••π hydrogen bonds marked by dashed lines. dx stands for the distance between the H atom of C(x)–H(xA) to plane Cg(j).

**Fig. 4** Fluorescence microscopic images of crystalline clusters of DCDPE with different scales.

### Thermal and Electrochemical properties

The thermal properties of DCDPE are investigated by the Thermogravimetric Analysis (TGA) and Differential Scanning Calorimetry (DSC). The dye is thermally stable up to 380 °C ($T_d$, the decomposition temperature) as determined by the TGA measurement shown in the inset of Figure 5. DSC scan result represented in Figure 5 shows that the initial crystalline sample melted at 252 °C to give an isotropic liquid in the first run of the DSC measurement. Upon cooling, the isotropic liquid changed into a glassy (amorphous) state. As the glassy sample was reheated for the second run, a glass transition was observed at 119 °C, which is defined as the glass transition temperature ($T_g$) of DCDPE. Thus although the molecular weight of DCDPE is not so high, its $T_g$ is much higher than that of the commonly used hole-transporting material (e.g., 98 °C for NPB),14 which may be attributed to the rigid structure. (The $T_m$ of NPB is ~280 °C and $T_d$ is 370 °C) The high morphological stability coupled with the efficient solid-state emission render the dye a promising EL material. It is well known that carbazole derivatives usually possess higher HOMO energy levels and good charge transport abilities. Thus to determine its...
energy level, the electrochemical property of DCDPE was studied by the cyclic voltammetry. The electrochemical potential was calibrated with the ferrocene/ferroenium (Fc/Fc+) standard (4.8 eV below the vacuum level). As shown in Figure 6, the compound shows a reversible oxidation process in CH3CN solution. The oxidation peak occurs at 0.50 V. From the onset of the oxidation, its HOMO energy level is estimated at 5.2 eV, slightly higher than that of NPB. The LUMO energy level of DCDPE deduced from the difference between the HOMO level and the optical band gap is 2.1 eV. As shown in Figure 7, comparing with NPB, it possesses higher-lying LUMO level, which is beneficial to blocking of electron leakage to the anode. On the other hand, its higher HOMO level is more close to the work function of the ITO anode, which can lower the charge injection barrier and the device operating voltages. These electronic properties indicate that besides light-emitting, DCDPE can also play a role as hole-transporting material. Thus according to the energy level diagrams, two different device architectures are adopted to validate its optoelectronic property. One is the typical device structure with both hole-transporting and electron-transporting layers (HTL & ETL) and the other one is only with of ETL (Figure 7).

Table 1 Analysis of dihedral angles between intramolecular aromatic planes (π rings) and C-H…π intermolecular interactions in the crystal packing of DCDPE.

<table>
<thead>
<tr>
<th>Planes (Cg(i), Cg(j))&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Angles (°)</th>
<th>Atoms to planes (C(x) –H(xA) → Cg(j))&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Distances (Å)</th>
<th>γ&lt;sup&gt;b&lt;/sup&gt; (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cg(5), Cg(8)</td>
<td>83.3</td>
<td>d&lt;sub&gt;2&lt;/sub&gt; C(2)–H(2A) → Cg(8)</td>
<td>2.81</td>
<td>8.06</td>
</tr>
<tr>
<td>Cg(5), Cg(13)</td>
<td>83.2</td>
<td>d&lt;sub&gt;10&lt;/sub&gt; C(10)–H(10A) → Cg(7)</td>
<td>2.75</td>
<td>11.78</td>
</tr>
<tr>
<td>Cg(5), Cg(14)</td>
<td>54.5</td>
<td>d&lt;sub&gt;11&lt;/sub&gt; C(11)–H(11A) → Cg(2)</td>
<td>2.86</td>
<td>13.63</td>
</tr>
<tr>
<td>Cg(8), Cg(13)</td>
<td>57.7</td>
<td>d&lt;sub&gt;18&lt;/sub&gt; C(18)–H(18A) → Cg(3)</td>
<td>2.86</td>
<td>14.12</td>
</tr>
<tr>
<td>Cg(8), Cg(14)</td>
<td>81.8</td>
<td>d&lt;sub&gt;25&lt;/sub&gt; C(25)–H(25A) → Cg(3)</td>
<td>2.77</td>
<td>6.00</td>
</tr>
<tr>
<td>Cg(13), Cg(14)</td>
<td>62.5</td>
<td>d&lt;sub&gt;31&lt;/sub&gt; C(31)–H(31A) → Cg(5)</td>
<td>2.85</td>
<td>15.77</td>
</tr>
</tbody>
</table>

<sup>a</sup> Planes number of Cg(i), Cg(j) and atom number are from Figure 2; Cg(13) and Cg(14) stands for the two carbazole planes. <sup>b</sup> γ is the angle of C(x)–H(xA) to Cg(j) vector and normal to plane Cg(j).

Fig. 6 Cyclic voltammograms of DCDPE in 0.1 M Bu₄NPF₆/CH₃CN.

Fig. 7 Energy level diagrams and device configurations of multilayer EL devices of DCDPE.

Fig. 8 (A) PL and EL spectra of DCDPE. (B) Current density and luminance versus voltage characteristics of devices based on DCDPE.
better electronic property, the simple configuration OLED balanced than that in the device with HTL. Subsequently, the turn-on voltage of the simple device (4.5 V) is lower than that with dedicated HTL (5.0 V). Profiting from the improved hole transporting material, the former device emits bluish green light with an EL maximum (λ_{EL}) of 512 nm, which is close to that of the PL in thin film. For the simple device, the charge transfer in the simple device is more efficient and device shows much better electronic character. To reach a same current density/brightness, the applied bias needed for the latter device is much higher than that for the simple one. That means the charge transfer in the simple device is more efficient and balanced than that in the device with HTL. Subsequently, the turn-on voltage of the simple device (4.5 V) is lower than that of the one with dedicated HTL (5.0 V). Profiting from the better electronic property, the simple configuration OLED radiates more brightly with a maximum luminance (L_{max}) of 5060 cd/m² at 15 V, which is much higher than the one with HTL of 1410 cd/m²; and also, the simple device exhibits much higher efficiencies. Figure 9 shows the current efficiency and external quantum efficiency versus voltage characteristics of the two kinds of devices. We can see that with the simple configuration (no dedicated HTL), the device shows much better electronic character. To reach a same current density/brightness, the applied bias needed for the latter device is much higher than that for the simple one. That means the charge transfer in the simple device is more efficient and balanced than that in the device with HTL. Subsequently, the turn-on voltage of the simple device (4.5 V) is lower than that of the one with dedicated HTL (5.0 V). Profiting from the better electronic property, the simple configuration OLED radiates more brightly with a maximum luminance (L_{max}) of 5060 cd/m² at 15 V, which is much higher than the one with HTL of 1410 cd/m²; and also, the simple device exhibits much higher efficiencies. Figure 9 shows the current efficiency and external quantum efficiency versus voltage characteristics of the two kinds of devices. The efficiency of the simple device is more than double times than that of the multilayer one in the whole bias range, e.g., 5.7 cd/A Vs 1.7 cd/A of the maximum current efficiency (CE), and 2.3% Vs 1.5% of the external quantum efficiency (EQE). (For details, see Table 2.) These results clearly demonstrated the intrinsic hole-transporting ability of the carbazole-containing luminogen.

**Table 2** Electroluminescence performances of DCDPE.

<table>
<thead>
<tr>
<th>HTL</th>
<th>λ_{EL} (nm)</th>
<th>V_on (V)</th>
<th>I_max (cd/m²)</th>
<th>CE_{max} (cd/A)</th>
<th>PE_{max} (lm/W)</th>
<th>EQE_{max} (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>✓</td>
<td>512</td>
<td>5.0</td>
<td>1410</td>
<td>1.7</td>
<td>1.7</td>
<td>1.5</td>
</tr>
<tr>
<td>x</td>
<td>512</td>
<td>4.5</td>
<td>5060</td>
<td>5.7</td>
<td>3.4</td>
<td>2.3</td>
</tr>
</tbody>
</table>

* Device structures: with HTL, ITO/NPB(40nm)/DCDPE(20nm)/TPBi(10nm)/Alq3(30nm)/LiF/Al(200nm); without HTL, ITO/DCDPE(60nm)/TPBi(10nm)/Alq3(30nm)/LiF/Al(200nm). Abbreviations: i_{EL} = EL maximum, V_on = turn-on voltage at 1 cd m⁻², L_{max} = maximum luminance, CE_{max} = maximum current efficiency, PE_{max} = maximum power efficiency, EQE_{max} = maximum external quantum efficiency.

**Conclusions**

In this paper, a carbazole-containing tetraphenylethene (DCDPE) is designed and synthesized. Its optical, thermal and optoelectronic properties are investigated. The dye exhibits typical aggregation-induced emission property: being nonluminescent when molecularly dissolved in solutions while intensely emissive in aggregate form. Close inspection of the geometric structure and packing motifs of DCDPE in the crystalline state reveals a loose packing characteristic due to the severe twisted conformation and the absence of strong intermolecular π–π interaction. On the other hand, the multiple weak intermolecular C–H•••π hydrogen bonds rigidify the molecular conformation and lock the vibrational and rotational freedom of the four aromatic substituents, accounting for the extremely high solid state quantum yield. The dye crystallizes into strong blue emissive fibres showing optical waveguide effect. The OLED devices using it as both light-emitting and hole-transporting material show superior performances. In a word, both enhanced optical and electronic properties were achieved in DCDPE. The present results demonstrate the promise of the carbazole-containing AIE luminogen as optoelectronic candidates especially for the future laser applications.

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