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Solid-state highly efficient DR mono and poly-dicyano-phenylenevinylene fluorophores

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Abstract: A highly efficient deep red (DR) emitting organic solid based on a dicyano-phenylenevinylene derivative was reported. The structural and spectroscopic properties of the solid have been described in terms of crystallographic data and time dependent DFT analysis. A noteworthy fluorescence quantum yield of 53% was observed for the brightest emitter cast into solid films. This result can be explained in terms of aggregation-induced emission (AIE) effect.

Keywords: DR; OLED/PLED; PPV; fluorophores, AIE.

1. Introduction

Deep red/near-infrared (DR/NIR) solid-state emitters have gained much attention as dopants in fabrication of OLEDs and PLEDs devices, in optical probes for bioimaging and in biomedical applications [1-3]. In white PLEDs (WPLEDs) white emission has provided using polymer–polymer blend or polymer–small-molecule blend as emissive layer and the role of the red emitting component is relevant and sensitive [4]. Compared with green and blue emitters [5-11], the development of red-light emitting materials is far behind in terms of both purity and efficiency but highly required for RGB (red-green-blue) based devices. In particular, for fluorescence bioimaging and assay applications, the optimal materials are DR/NIR photo emitters with high photoluminescence quantum efficiency, large Stoke’s shifts (for eliminating self- absorption), good chemical and thermal stability, good solubility and processability [12].

So far, the most reported DR/NIR organic emitters consists of a planar polycyclic skeleton having extended π-system or a strong donor–acceptor framework featuring a large dipole moment in the excited state [13]. NIR emitters are often designed utilizing intramolecular charge-transfer (ICT) to obtain red-shifted emission. Nevertheless, these materials usually tend to aggregate in highly concentrated solutions and in solid states, and this could result in non-uniform thin films and in concentration quenching (aggregation-caused quenching, ACQ effect), due to the strong dipole–dipole interactions. Most dyes are highly emissive in dilute solution but become weakly luminescent or even non-emissive in the solid phase due to the strong dipole–dipole interactions that causes the formation of detrimental excimers and exciplexes. Recently, new types of luminogens showing the aggregation-induced emission (AIE) effect, have received much research attention because of their unique optical properties and wide applications [14-17]. The AIE molecules display weak emission when dissolved in solutions, but strong emission in aggregate state, showing the opposite effect of ACQ. Among these molecules, the red emitters are very few, although they are the most promising, for example, in the field of bioimaging, and OLEDs [15,17].
Phenylenevinylene (PV) derivatives are among the most widely investigated DR/NIR fluorogens. Many oligo or polyphenylenevinylene (PPV) have been reported in literature as emitting layers in OLEDs and PLED [18-23]. Monomeric and polymeric systems based on PV scaffolds are simple and cheap to realize, processable and easily chemically tuned to give colors in the whole range of the visible spectrum. The introduction of the electron withdrawing cyano groups on the vinylene moiety leads to an increase in the energy of the occupied π and unoccupied π* states [19], reducing the barrier for electron injection into the conjugated system and improving the electron-transporting properties. This introduction causes also a red shift of the emission maximum compared to less substituted phenylenevinylene systems. Widely reported fully conjugated polymers based on cyano-modified PPV structures feature both good electron transport and efficient emission at high wavelengths in the visible spectrum. In many cases, they represent efficient DR probes with large Stoke's shift. The nitrile group provides high electron affinity [24] and exerts a large influence on the photoluminescent (PL) and electroluminescent (EL) properties of the final material [25,26]. Moreover, it can induce steric hindrance resulting in a twisted conformation, which enables the fluorophores to be protected from the ACQ effect [27-29]. Our approach consists in linking well-defined CN-PV emitting units to non-emitting blocks determining an interruption of the conjugation path. In this way, we can improve the polymer processability as well as prevent the aggregation-caused quenching effect.

In a previous work, we presented a series of blue fluorophores and the related polyamides with the PV units co-polymerized with non-photoluminescent co-monomers [23]. In this article, we describe the facile synthesis of a new PV based donor-acceptor system (D-A-D-A-D). This probe is symmetrically functionalized with electron donating terminal amino-groups and electron withdrawing cyano groups on the vinylene moiety. The same diamino precursor was employed to obtain both the final low molecular weight fluorophore and to realize the polyamide. In the latter case, two different dichlorides were simultaneously employed with the aim to obtain an amorphous ter-copolymer, optimizing both solubility and glass transition temperature. An accurate examination of PL properties in solution and in the solid state has been carried out. Both the fluorophore and the polymer act as DR emitters in the solid state, showing also large Stoke’s shift and relevant solvatochromic effects. Remarkably, AIE effect is dominant for the CN-PV-NHMe fluorophore, with a medium quantum efficiency in solution (30%) and a remarkable PLQY (50%) in the solid state. These results were discussed by a correlation of the X-Ray results with a computational study.

2. Results and Discussion

2.1. General Route for the Synthesis of Fluorophores

Mono and polymeric fluorophores were obtained as summarized in Scheme 1. The reduction of CN-PV-NO2 led to the monomer CN-PV-NH2. The diamino derivative was employed to prepare both the polymer CN-PPV and the compound CN-PV-NHMe, with the same D-A-D-A-D skeleton.
The identification and the evaluation of the purity degree were assessed by mass spectrometry and $^1$H NMR. Phase behavior was examined by optical observation and DSC and TGA analysis. All materials are thermally stable up to 300 °C under nitrogen flow. The ter-polymer CN-PPV was synthesized by polycondensation reaction between the chromophore CN-PV-NH$_2$ and a mixture of two different acyl dichlorides (dodecanedioyl and suberoyl). It was designed to guarantee a random and hence more soluble amorphous material, in which chromophores are diluted in a processable polymeric system. Based on $T_g$ and $\eta_{inh}$ values, the polymer can be considered an oligomer, although no evidence of COOH terminal groups was found in the $^1$H NMR spectrum. On the other hand, the low degree of polymerization guarantees good solubility in many common solvents and therefore easy processability.

2.2. Photophysical properties

Photophysical measurements were performed both in solution and in the solid state, including PLQY measurements. A strong solvatochromic effect was detected for both the low molecular weight compounds CN-PV-NH$_2$ and CN-PV-NHMe (see Table 1, columns 2 and 3).
Table 1. Optical data of synthesized compounds.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\lambda_{\text{abs-sol}}$ (nm)$^a$</th>
<th>$\lambda_{\text{em-sol}}$ (nm)$^b$</th>
<th>$\lambda_{\text{abs-film}}$ (nm)$^c$</th>
<th>$\lambda_{\text{em-film}}$ (nm)$^d$</th>
<th>PLQY% film$^e$</th>
<th>PLQY% solution$^f$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CN-PV-NH$_2$</td>
<td>436$^i$; 449$^ii$</td>
<td>536$^i$; 609$^ii$</td>
<td>476$^i$; 462$^ii$</td>
<td>530$^i$; 559$^ii$</td>
<td>5±1$^i$; 27±5$^ii$</td>
<td>26±3</td>
</tr>
<tr>
<td>CN-PV-NHMe</td>
<td>464$^i$; 470$^ii$</td>
<td>554$^i$; 622$^ii$</td>
<td>495$^i$; 474$^ii$</td>
<td>635$^i$; 598$^ii$</td>
<td>53±5$^i$; 50±8$^ii$</td>
<td>29±3</td>
</tr>
<tr>
<td>CN-PPV</td>
<td>441$^i$; 445$^ii$</td>
<td>518$^i$;$^ii$</td>
<td>458$^j$</td>
<td>613$^j$</td>
<td>18±3$^j$</td>
<td>26±3</td>
</tr>
</tbody>
</table>

$^a$ Wavelength of UV-Visible absorbance maxima in (i) dioxane (ii) DMF solution. 
$^b$ Wavelength of emission maxima in (i) dioxane (ii) DMF solution. 
$^c$ Wavelength of UV-Visible absorbance maxima on thin film obtained (j) by slow evaporation of the sample or (jj) by dispersing the sample in polystyrene at 10% by weight. 
$^d$ Wavelength of emission maxima on thin film obtained (j) by slow evaporation of the sample or (jj) by dispersing the sample in polystyrene at 10% by weight. 
$^e$ PL quantum yield on thin film obtained (j) by slow evaporation of the sample or (jj) by crystalline micro-dispersion in polystyrene, 10% by weight. 
$^f$ PL quantum yield in dioxane.

In solution, the absorption was recorded in the range from 436 to 470 nm and the emission from 518 to 622 nm with a red shift effect depending on solvent polarity. In dioxane the compounds appear yellow-green emitters. In more polar solvents as DMF, we observed a red shift of the emission peak. This is consistent with the existence of a polar excited state, which is stabilized in a polar environment and has an increased probability of decaying to the ground state by non radiative paths (Figure 1). Furthermore, quantum efficiency in DMF decreases significantly. In the case of CN-PPV polymer, we do not observe the same solvatochromic effect: the polymer exhibits a strong yellow-green emission both in dioxane and in DMF, with CIE 1931 coordinates of (0.36, 0.57) and a measured PLQY of 26% in dioxane or in DMF solution (see Figure 1).
Figure 1. Precursor CN-PV-NH\textsubscript{2}, fluorophore CN-PV-NHMe and polymer CN-PPV solution (10% w/w) in dioxane (left) and DMF (right) in natural light (left column) and under 375 nm UV light (right column).

In solution, PL quantum yields were measured by relative methods using as standard quinine sulfate QS (UQS \( \frac{1}{2} 0.546 \) in H\textsubscript{2}SO\textsubscript{4} 1 N, when excited at 365 nm according to Melhuish [30]). The values appear independent from the solvent and very similar. All compounds show deep red photoluminescence with a pronounced Stoke’s shift both in solution and in the solid state. Remarkably, in the solid state, Stoke’s shifts range from 140 to 160 nm. In order to test the effect of a dilution in the solid state, we prepared two types of solid films: by spin-coating of a solution of each compound or by microdispersion of the shattered crystals in a polystyrene matrix at 10% by weight (PS doped film). In some cases, the use of a microdispersion of the crystalline fluorophores in PS represented the right way to prevent ACQ effect in the solid state [12,22]. In Figure 2, the solid samples on quartz slides are shown.

Figure 2. Fluorophores thin film (left) and in 10% PS doped thin film (right) in natural (left column) and under 375 nm UV light (right column).

All solid compounds are red both under natural and UV lamp. CN-PV-NH\textsubscript{2} micro-dispersed in polystyrene (Figure 2, top) showed enhancement of PL emission respect to the pure thin film, reaching the same value measured in dioxane diluted solution (see also Figure 1). CN-PV-NHMe displays a strong deep red emission, with a remarkable 53% PLQY in thin film. The same compound dispersed in polystyrene emits at 598 nm with a comparable 50% PLQY, whereas in solution shows only a faint emission. The stronger emission observed in the solid state can be ascribable to the presence of cyano substituents on the conjugated system, whose effect is to restrict molecular motions and to stabilize a twisted conformation in the solid. This permits the fluorophore to emit more efficiently in the aggregated state. In fact, the twisted geometry, hindering intermolecular close stacking and intense \( \pi-\pi \) interaction, may lead to intense aggregation-induced emission effect (AIE) [14]. In order to verify this hypothesis, we carried out a crystallographic study on CN-PV-NHMe, which shows the higher PLQY in the solid and was easily obtained as single crystals.

In the solid phase, also the polymer is a DR emitter (see Figure 2), but with a PLQY value similar to those of the compounds in solution (Table 1). CN-PPV can be considered infact a diluted system of the fluorophore core. PLQY value measured for CN-PPV thin film is lower than the value of CN-PV-NHMe and close to the value of CN-PV-NH dispersed in polystyrene, so in the polymer chain the fluorophore replaces its strong electron donating amino groups with amide moieties. This result
however is not easily rationalizable because dipole–dipole interactions in the polymeric amorphous sample are conceivably very different from the same interactions acting in the crystalline samples.

2.3 X-Ray Crystallography

The molecular structure of CN-PV-NHMe was determined by single crystal X-ray analysis. Compound crystallizes in the monoclinic C2/c space group with one molecule contained in the asymmetric unit. All bond lengths and angles fall within the normal range. An ORTEP view of the molecular structure is reported in Figure 3. Details of crystallographic data and refinement parameters are reported in Table S1 of Supplementary Materials.

Figure 3. Ortep view of CN-PV-NHMe with thermal ellipsoids drawn at 30% probability level. Only major part of the disordered aliphatic group is shown for clarity.

The molecule is characterized by three phenyl rings mutually twisted each to the other. The mean planes of outlying C1/C6 and C19/C24 phenyl rings are 49.01(18)° and 31.47(16)° tilted with respect to the central C10/C15 mean plane ring. An overall elongated, almost flat shape of the molecule is observed, with aminic and alifatic groups quite in-plane. As expected, the planar geometry at the aminic N atoms shows a degree of conjugation with the binded phenyl group. The crystal packing is stabilized by face-to-edge aromatic interactions (Figure 4). No π-π aromatic stacking is found, due to the twisting of phenyl rings in adjacent molecules (Figure S1, Supp. Mat.).

Figure 4. Crystal packing of CN-PV-NHMe viewed along b axis. Ball-and-stick style, H atoms not drawn for clarity.

In the crystal packing the aminic hydrogen atoms are involved in intermolecular weak interactions with the -CN groups of adjacent molecules, forming linear ribbons of molecules in the (1 -1 0) and (1
171. 1 0) directions (N1-H···N4: 0.97(6) Å, 2.34(6) Å, 3.237(7) Å, 153(5)°, \(i = x+1/2, y-1/2, z\); N2-H···N3: 0.84(5) Å, 2.51(5) Å, 3.323(7) Å, 166(5)°, \(ii = x-1/2, y+1/2, z\)) (See Figure S2, Supp. Mat.).

2.4 Computational Studies

The excitation energies were obtained at the density functional level by using the time-dependent perturbation theory approach (TDDFT) with the adiabatic local density approximation, which have high reliability in obtaining accurate predictions for excitation energies and oscillator strengths. Calculations on CN-PV-NHMe in DMF showed the localization of HOMO and LUMO orbitals (see Figure 5).

*Figure 5. HOMO-1, HOMO and LUMO representation for CN-PV-NHMe.*

HOMO is delocalized overall the conjugated backbone, with only a moderate contribution of the cyano groups. Contrarily, the LUMO is mainly focused on the central part of the molecule and on the electron withdrawing cyano groups. The main transition, at 487 nm, is the HOMO → LUMO. The second main strong absorption at 394 nm is due to the transition HOMO-1 → LUMO. The predicted absorption maximum is in good agreement with the experimental values (see Table 2 and Table 1), the two main absorptions corresponding to the HOMO→LUMO and HOMO-1→LUMO transitions.
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Table 2. Calculated electro-optical parameters for CN-PV-NHMe.

<table>
<thead>
<tr>
<th></th>
<th>Oxidation Potential (eV)</th>
<th>Reduction Potential (eV)</th>
<th>Hole Reorganization Energy (eV)</th>
<th>Electron Reorganization Energy (eV)</th>
<th>Triplet Energy (eV)</th>
<th>( \lambda_{\text{ole}} ) (nm)</th>
<th>( \lambda_{\text{em}} ) (nm)</th>
<th>Triplet Stabilization Energy (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>-0.022</td>
<td>-1.754</td>
<td>0.229</td>
<td>0.457</td>
<td>1.477</td>
<td>487</td>
<td>587</td>
<td>3.131</td>
</tr>
<tr>
<td>Triplet Reorganization Energy (eV)</td>
<td>Electron Small Polaron Stabilization Energy (eV)</td>
<td>Hole Extraction Potential (eV)</td>
<td>Electron Extraction Potential (eV)</td>
<td>Hole Small Polaron Stabilization Energy (eV)</td>
<td>Dipole (D)</td>
<td>Scaled HOMO (eV)</td>
<td>Scaled LUMO (eV)</td>
<td></td>
</tr>
<tr>
<td>0.603</td>
<td>0.328</td>
<td>5.429</td>
<td>-1.330</td>
<td>0.116</td>
<td>0.420</td>
<td>-4.508</td>
<td>-2.776</td>
<td></td>
</tr>
</tbody>
</table>

CN-PV-NHMe exhibits hole and electron reorganization energies (HRE and ERE respectively) that are typical for these compounds. DFT calculations permit to investigate the first oxidation potential to explain the electron-donating properties of the compound. Withdrawal of the electron from the compound in a redox reaction is performed from its HOMO. The observed trend in oxidation potential for the three compounds indicates destabilization of the HOMO. The intense DR emission of CN-PV-NHMe arises from the D-A-D-A-D motif with strong donor and acceptor functionalities and at the same time largely delocalized frontier MOs, giving rise to a large oscillator strength of the emitting state. As reported for similar compounds [12], in our systems, only a minor intramolecular charge transfer character is observed, and the oscillator strength is thus large for the ground to the first excited state transition (\( \lambda = 2.27 \)) [29]. The large radiative decay and the strong fluorescence, with a quantum yield of about 50% in the solid, could also be attributed to strongly twisted equilibrium structures, as confirmed by the X-ray analysis (Figures 3, 4).

4. Materials and Methods

All starting products were commercially available. The chlorides were distilled before use.

2-(2-Ethylhexyloxy)-5-methoxyterephthalaldehyde was obtained as described in [23]. Optical observations were performed by using a Zeiss Axioscop polarizing microscope equipped with a FP90 Mettler heating stage. Phase transition temperatures and enthalpies were measured using a DSC scanning calorimeter Perkin Elmer Pyris 1 at a scanning rate of 10 °C/min, under nitrogen flow. \(^1\)H NMR spectra were recorded with Varian Gemini 300 MHz apparatus. Mass spectrometry measurements were performed using a Q-TOF premier instrument (Waters, Milford, MA, USA) equipped by an electrospray ion source and a hybrid quadrupole-time of flight analyzer. Mass spectra were acquired in positive ion mode, in 50 % CH\(_3\)CN solution, over the 400-800 m/z range. Instrument mass calibration was achieved by a separate injection of 1 mM NaI in 50% CH\(_3\)CN. Data were processed by using MassLynx software (Waters). UV-Visible and fluorescence spectra were recorded with JASCO spectrometers. Jasco F-530 (scan rate 200 nm min\(^{-1}\)) and on a spectrofluorimeter Jasco FP-750 (excitation wavelengths set at the absorption maxima of the samples, scan rate 125 nm min\(^{-1}\)).

Thin films of the amino precursor, the fluorophore and the polymer were obtained by the spin-coating technique of a micro-dispersion of shattered crystals (10% by weight) and low weight commercially available polystyrene (PS) in acetone/dichloromethane (1:2) by a SCS P6700 spin-coater operating at 800 RPM. Thin films of the sample without polystyrene were obtained by casting from a chloroform solution (about 30% by weight).

4.1 Preparation of CN-PV-NO\(_2\):

2-(2-Ethylhexyloxy)-5-methoxyterephthalaldehyde (7.60 g, 0.0260mmol) [23] and piperidine (2 mL) were dissolved in pyridine (20 mL). The solution was warmed at 40°C and then 2-(4-nitrophenyl)acetonitrile (16.2 g, 0.102 mol) was added in portions during 1 h. After this time, the
solution color turned to red and the formation of a solid occurred. After one more hour, the reaction was stopped and the suspension was poured in 40 mL of ethanol at room temperature. The bright red solid was then recovered by filtration. The compound was recrystallized from ethanol in needle shaped, dark red crystals. Yield: 55%; mp: 232 °C. 1H NMR (300 MHz, DMSO-d6, 25°C, ppm): 0.84 (m, 6H), 1.31 (m, 8H), 1.90 (m, 1H), 3.91 (s, 3H), 4.01 (d, J = 4.8 Hz, 2H), 7.81 (d, J = 3.2 Hz, 4H), 8.00 (t, J = 8.4 Hz, 2H), 8.26 (d, J = 4.0 Hz, 4H), 8.32 (m, 2H).

HRMS(ESI): m/z calcd for C₃₅H₄₇NaO₂ + H+: 581.23; found 581.12 [M + H]⁺

4.2 Preparation of CN-PV-NH₂

To CN-PV-NO₂ (1.00 g, 0.70 mmol) dissolved in DMF (20.0 mL) 5.60 mL of a 1.20 M Na₂S·9H₂O aqueous solution was added, the color turned from red to black. After 15 min, water (1.00 mL) was added and the system kept at 150°C for 15 min. The mixture was slowly cooled to room temperature leading to the formation of a red crystalline solid. The compound was recovered by filtration and washed with slightly acid water (acetic acid, pH about 5) and then with water. The red crystalline product was crystallized again from chloroform/ethanol. Yield: 90%; mp: 190 °C. 1H NMR (300 MHz, DMSO-d6, 25°C, ppm): 0.96 (m, 6H), 1.48 (m, 8H), 1.90 (m, 1H), 3.96 (s, 3H), 4.06 (d, J = 4.0 Hz, 2H), 5.19 (s, 4H), 6.80 (d, J = 8.2 Hz, 4H), 7.49 (d, J = 8.2 Hz, 4H), 7.85 (m 4H). HRMS(ESI): m/z calcd for C₃₅H₄₇NaO₂ + H+: 521.28; found 521.30 [M + H]⁺

4.3 Preparation of CN-PVNMe

To CN-PV-NH₂ dissolved in DMF (20.0 mL) 5.60 mL of a 1.20 M Na₂S·9H₂O aqueous solution was added and the system kept at 150°C for 15 min. The mixture was slowly cooled to room temperature, filtered and the solution concentrated in vacuo to 50 mL and added of 100 mL of heptane, the precipitation of a red crystalline solid. The compound was recovered by filtration and washed with slightly acid water (acetic acid, pH about 5) and then with water. The red crystalline product was crystallized again from chloroform/ethanol. Yield: 80%; mp: 160 °C. 1H NMR (300 MHz, CDCl₃, 25°C, ppm): 0.90 (m, 6H), 1.46 (m, 10H), 2.90 (s, 6H), 3.94 (s, 3H), 4.01 (d, J = 5.4 Hz, 2H), 6.80 (d, J = 8.2 Hz, 4H), 7.55 (m, 3H), 7.85 (m, 5H). HRMS(ESI): m/z calcd for C₃₅H₄₇O² + H+: 549.32; found 549.30 [M + H]⁺

4.4 Preparation of CN-PPV

To 0.160 g (0.60 mmol) of dodecandioic dichloride and 0.126 g (0.60 mmol) of suberic dichloride dissolved in 5.0 mL of THF a solution of CN-PV-NH₂ (0.607 g, 1.20 mmol) dissolved in 5 mL of dry pyridine was added. The reaction was kept under vigorous stirring at reflux for 2 h. The solution was poured into 150 mL of water and a solid product was recovered by filtration. The red crude product was dissolved in hot DMF (50 mL) and precipitated again in 400 mL of water, recovered by filtration and dried at 120 °C for 4 h. Yield: 80%; Tg = 112 °C. ηsp (DMF, 25 °C) = 0.50 dL/g. 1H NMR (300 MHz, DMSO-d6, 25°C, ppm): 0.85-0.95 (m, 6H), 1.31-1.61 (m, 20H), 2.30-2.46 (m, 3H), 3.96 (m, 5H), 7.76 (m, 12H), 10.17 (s, 2H).

4.5 PLQY measurements

Photoluminescence efficiency measurements were conducted with a setup similar to the one suggested by de Mello et al. [31]. This setup takes into account not only the exciting laser and direct photoluminescence spectrum but also the photoluminescence that originates due to the scattering effect of the integrating sphere that is part of the setup. The setup consists of a 376 nm laser whose emission does not overlap with the emitted photoluminescence spectrum, an integrating sphere...
Single crystals of CN-PV-NHMe were obtained by slow evaporation of acetone/heptane solution at ambient temperature. It was not possible to obtain very good quality crystals. One selected crystal was mounted in flowing N$_2$ at 173 K on a Bruker-NoniusKappaCCD diffractometer equipped with Oxford Cryostream apparatus (graphite monochromated MoK$_\alpha$ radiation, $\lambda = 0.71073$ Å, CCD rotation images, thick slices, $\varphi$ and $\omega$ scans to fill asymmetric unit). Semiempirical absorption corrections (SADABS [32]) were applied. The structure was solved by direct methods (SIR97 program [33]) and anisotropically refined by the full matrix least-squares method on $F$ against all independent measured reflections using SHELXL-2016 [34] and WinGX software [35]. The H atoms at aminic groups were located in difference Fourier maps and freely refined with $U_{eq}(H)$ equal to 1.2$U_{eq}$ of the carrier atom. All the other hydrogen atoms were introduced in calculated positions and refined according to the riding model with C–H distances in the range 0.95–1.00 Å and with $U_{eq}(H)$ equal to 1.2$U_{eq}$ or 1.5$U_{eq}$(Cmethyl) of the carrier atom. Difference Fourier maps showed two disordered positions of the branched aliphatic group that refined with occupancy factors 0.596 and 0.404. Some restraints were introduced in the last stage of refinement to model the disorder using DFIX, SAME and SIMU commands of Shelx program. Crystal data and structure refinement details are reported in Table S1 (Supp. Mat.). The figures were generated using ORTEP-3 [36] and Mercury CSD 3.9 [37] programs.

Crystallographic data were deposited at Cambridge Crystallographic Data Centre with assigned number CCDC 1843139. These data can be obtained free of charge from www.ccdc.cam.ac.uk/structures.

4.7 Computational studies

All quantum calculations were carried out using the Jaguar program [38] as implemented in the Schrödinger Material Science Suite [39]. Geometry optimizations were performed with the B3LYP functional and the 6-31G** basis set in which the transition metals are represented using the Los Alamos LACVP** basis that includes relativistic effective core potentials. The energies of the optimized structures are re-evaluated by additional single-point calculations on each optimized geometry using Dunning’s correlation-consistent triple-ζ basis set42 cc-pVTZ(-f) that includes a double set of polarization functions. Vibrational frequency calculation results based on analytical second derivatives at the B3LYP/6-31G**(LACVP**) level of theory were used to confirm proper convergence to local minima and to derive the zero-point-energy (ZPE) and entropy corrections at room temperature where unscaled frequencies were used.

5. Conclusions

A new DR dicyano-phenylenevinylene emitter and the ter-polymer containing the same fluorophore unit have been synthesized. A detailed study of PL properties in solution and in the solid-state compounds has been carried out. A bright red emission for the fluorophore was recorded with 50% PLQY in the solid state. Interestingly, no ACQ effect was detected on the fluorophore. A detailed X-ray investigation correlated with TD-DFT analysis was presented in order to explore this behavior. In the solid state of CN-PV-NHMe (film or doped PS film), owing to the restriction of intramolecular motions induced by the cyano substituents, the radiative channel is activated, causing intense aggregation-induced emission (AIE). Regarding the polymer, although a noteworthy quantum efficiency was not observed, this was a way of exploiting the advantages to covalently bond the fluorophores into an easily processable material retaining medium DR photoluminescence.

Supplementary Materials: The following are available online at www.mdpi.com/link, Table S1: Crystallographic data and structural refinement details of CN-PV-NHMe. Figure S1: Partial packing of CN-PV-NHMe with shortest distances involving aromatic centroids. Figure S2: Molecular ribbon of CN-PV-NHMe propagating in the (1 -1 0) direction.
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Author Contributions: B.P. and U.C. conceived and designed the experiments; R.D. and S.C. performed the experiments; S.P. and L. S. performed the theoretical analysis; A.T performed the crystallographic experiments; R.S and S. N. performed the optical measurements; B.P and R. D. analyzed the data; R.D. and S. C. wrote the paper.

Conflicts of Interest: The authors declare no conflict of interest.

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455 Sample Availability: Samples of the compounds CN-PV-NH3, CN-PV-NHMe and CN-PV are available
457 from the authors.