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Paraphenylene dimers with diphenylamine donor groups: synthesis and photophysics

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ABSTRACT

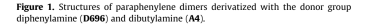
A novel paraphenylene dimer (**D696**) with electron donating diphenylamine side chain groups has been prepared. Optical absorption measurements were theoretically and experimentally determined showing a red shift in the absortivity relative to dialkylamine functionalized benzophenone dimers. The fluorescence of **D696** was quenched in a concentration dependent manner in the presence of reduced graphene oxide (RGO).

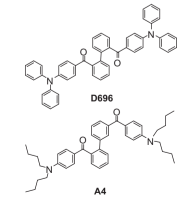
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The large absorption coefficients, high charge mobility, excellent thermal and mechanical stability, and affordable large scale production costs of semiconducting polymers position these materials to outperform their silicon-based counterparts for application in light emitting diodes, photovoltaic devices, and nonlinear optics.^{1–5} However, applications have not been fully realized due to charge transport inefficiencies in these materials.

Polyparaphenylenes and oligomeric paraphenylenes (OPPs) have lagged behind other semiconductor materials because of intrinsic limitations in flexibility, symmetry breaking, or distortion properties, and hole injection difficulties as a result of high HOMO levels. These limitations can be addressed by the selective introduction of sterically demanding and electronically rich or favorable substitution groups.

We and others continue to show an interest in model paraphenylene oligomers derivatized with donor and acceptor groups for the study of photoinduced charge transfer.^{6–11} Our previous experimental and theoretical substitution strategies involved the formation of butyl amine side chain groups.⁶ In those studies, we demonstrated the potential to red shift the theoretical band gap of paraphenylene based dimers by the substitution of electron donating groups including various alkyl amine and diphenyl amine units. While earlier studies report that the spectral absorption and emission of oligomeric semiconductors are influenced by structural modification arising from the introduction of different electron donating and withdrawing groups,^{12–14} confirmation of our promising theoretical results with OPP functionalized with diphenylamine donor groups was previously unobtainable as the dimers were not synthetically available. We describe here the synthesis of the 4-(*N*,*N*-diphenylamino)benzoyl substituted paraphenylene dimer (**D696**). The structure of **D696** is illustrated and compared with the dibutylamino substituted dimer **A4** reported earlier (Fig. 1). Important issues that we address in this work include the role of the substitution chemistry on the absorption and emission properties of the dimer. In addition, we address the role of coupling the dimers in a head-to-head (HH) conformation where the dimer contains a mirror plane of symmetry upon the union of the monomers (i.e., the two carbonyl groups face each









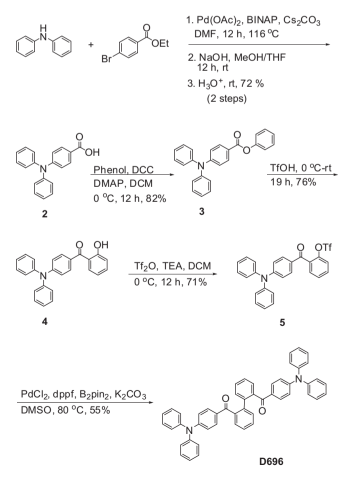
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other in a planar view of the dimer). Furthermore, we observe efficient quenching of the fluorescence of **D696** by reduced graphene oxide, a photo-induced electron transfer process reported in the literature.¹⁵

As shown in Scheme 1, the synthesis of D696 was accomplished in six-steps (see Supplementary data). Briefly, diphenylamine was coupled with ethyl 4-bromobenzoate by using the Buchwald amination chemistry.¹⁶ The resulting yellow viscous oil underwent alkaline hydrolysis¹⁷ to provide the corresponding acid as a white solid in 72% yield for two-steps. Subsequent reaction with phenol in the presence of DCC and DMAP produced the ester 3 in good vield. Initial attempts to convert the ester **3** into the ketone **4** using a Fries rearrangement reaction with Lewis acids such as AlCl₃, scandium triflate, and BF3. OEt2 in different solvents resulted in poor vields.¹⁸ The rearrangement was however successfully carried out, in a vield of 76%, when the reaction was performed in neat triflic acid at 0 °C followed by warming to room temperature. Finally, the hydroxyl group of **4** was activated by triflic anhydride to produce triflate 5 which was then dimerized in the presence of a palladium catalyst to afford D696 as a yellow solid in 55% yield.

The redox behavior of **D696** was investigated by cyclic voltammetry (CV). The studies were performed using a solution of **D696** (1.64×10^{-4} M) prepared in DMF with *n*-Bu₄NPF₆ (0.1 M) as a supporting electrolyte. As shown in Figure 2, the dimer exhibits a reversible anodic wave at ~0.75 V versus AgNO₃/Ag which is attributed to the electron donating tendency of the diphenylamine group. The energy associated with this oxidation is taken to be the HOMO energy level. **D696** exhibits no observable cathodic wave. Thus, the LUMO energy level of **D696** was calculated by subtracting the HOMO energy level and the zeroth-zeroth energy (E_{0-0})



Scheme 1. Synthesis of D696.

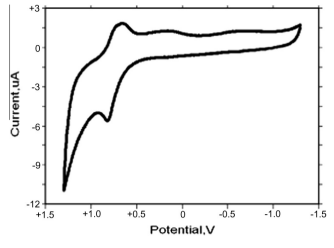


Figure 2. CV curve of D696 in acetonitrile.

extrapolated from the intersection between the absorption and emission spectra (see Fig. S1 in Supplementary data).

The absorption spectra of **D696** taken in DMF solution at a concentration of 5.49×10^{-5} M is shown in Figure 3. This is very close to our theoretically determined UV–vis spectra previously reported for diphenylamine functioned OPP.⁶ The spectrum features a relatively broad absorption in the ultraviolet and slight trailing in the visible region with maxima of 290 nm and 368 nm. These two intense absorption maxima suggest π – π * transition and intramolecular charge transfer between the diphenylamine donor group and the carbonyl group which acts as a modest acceptor.

The absorption, emission, and electrochemical properties of **D696** are listed in Table 1. From our experimental data, we calculated the band gap of **D696** to be 2.90 V. This band gap is lower than unsubstituted paraphenylene and polypyrrole (3.1 eV) but higher than other polymers such as polythiophene and poly(2-vinylpyridine).

The geometry of the ground state molecular orbitals of **D696** was fully optimized by the density functional theory (DFT) with Becke's three-parameter (B3) exchange functional along with the Lee–Yang–Parr (LYP) nonlocal correlation functional implementing the modest 6-31G(d,p) basis set. The calculations were carried out in WebMO environment interfaced with GAUSSIAN 94/98/03 software package. The plots of the HOMO and LUMO of the dimer are given in the Supplementary data (Fig. S2) and the respective electron densities images are presented in Figure 4. The absorption spectra

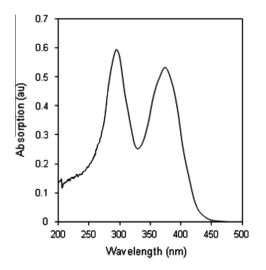


Figure 3. Absorption spectra of D696 in DMF.

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