Conjugated Polymers with Large Effective Stokes Shift: Benzobisdioxole-Based Poly(phenylene ethynylene)s

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Abstract: Phenyleneethynylene-based conjugated copolymers using benzo[1,2-d:4,5-d′]bis[1,3]dioxole (BDO) in the repeating unit are reported. The electronic structure of the BDO unit imparts a localized HOMO topology while the LUMO is delocalized over the polymer backbone, so that the lowest optical absorption band of the polymer has considerable intramolecular charge transfer character. This contrasts with published donor−acceptor polymers with localized LUMO and delocalized HOMO. The very large Stokes shifts of the monomers, which are due to the small oscillator strength of the lowest optical transition, are largely retained in the polymers as a result of covalently constrained dihedral angles in the substituents (not the backbone), as predicted/explained by calculations.

1. Introduction

Poly(phenylene ethynylene)s, PPEs, are conjugated polymers (CP)1,2 that are studied as active components in organic (opto)electronic devices such as LEDs,3 FETs,4 and solar cells5 as well as a common tool to control morphological order and to tune optoelectronic properties.8−10 Along the backbones of various CPs is a common tool to control morphological order and to tune optoelectronic properties.8−10

Recently we reported PPE’s built from alternating perfluorinated (acceptor) and peralkoxylated (donor) benzenes, as well as a transition-metal-free route for their synthesis exploiting fluoride ion catalyzed nucleophilic aromatic substitution.14,15

Surprisingly, the optical properties of these polymers built from highly oxygenated 1,2,4,5-tetraalkoxy benzenes (TAB) seemed fairly typical of PPEs. However, there were some slight differences between the different TAB-based polymers which were tentatively assigned to differing dihedral angles between the donor side chains and the polymer π-system, that is, mesomeric and inductive effects. As reported here, the electronic structure of the requisite 1,2,4,5-tetraalkoxy-3,6-diethynylbenzene monomers results in disjoint frontier molecular orbitals.16,17

This constrained geometry is provided by benzo[1,2-d:4,5-d′]bis[1,3]dioxole (BDO) units. Based on the widely varying (opto)electronic properties of 3,4-diakloxy- and 3,4-alkylene-dioxothiophene polymers, one would expect significant differences in BDO and TAB-based CPs.18 The electronic structure of BDO was exploited to study equilibria between electron- and charge-transfer states in donor−acceptor pairs and to measure the photoinduced charge separation distances in donor-spacer-acceptor systems.19 BDO has also been used as the aryl unit in persistent trityl radicals for EPR imaging of biological systems.19

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systems. In this work we have introduced BDO into the conjugated backbones of two poly(phenylene ethynylene)s (PPE) and a poly(phenylene diethynylene) (PDD) resulting in polymers with relatively little difference in the optoelectronic properties.21 Unlike here, LSS in conjugated polymers is typically attributed to large conformational differences between the ground and excited states. The very large effective Stokes shifts (LSS) demonstrated by the BDO polymers (0.7 to 0.9 eV), may be useful to minimize self-absorption and light scattering in optical materials.22 Possible applications include laser dyes, molecular imaging, scintillators, solar collectors and white light emitting materials.20–22 In a quantum-chemical approach using density-functional theory we fully elucidate the reasons for the electronic and optical properties of the new BDO-based polymers and compare them to the analogous TAB PPEs.

2. Experimental Section

Characterization. Solution photoluminescence (10–8 M THF) and UV−vis absorption spectra (10–4 M THF) were measured on a Fluorolog-3 fluorometer and Varian Cary 1 spectrophotometer. Solid state spectra were recorded from thin films spin-coated from solution (∼0.5 mg/mL toluene) onto quartz plates.

Electrochemical measurements were performed under nitrogen atmosphere using a BAS-CV-50W voltammetric analyzer and three electrode cell (platinum working, silver wire counter and pseudo-reference electrodes). The supporting electrolyte solution (0.1 M (n-Bu)4NPF6, anhydrous CH3CN) was thoroughly purged with atmosphere using a BAS-CV-50W voltammetric analyzer and three Fc+/Fc before each measurement, and ferrocene/ferrocenium (Fc+/Fc) was used as reference. Polymers were evaluated as thin films solvent cast directly onto the working electrode while small molecules were evaluated in dilute CH3CN solution. Detailed procedures for monomer and polymer syntheses and their characterization are included in the Supporting Information.

M3 and polymer TADBFPPE (Table 1, R = 3,7-dimethyloctyl) were prepared as published.

Theoretical Methodology. The geometry and electronic structure of the molecules were calculated at the density functional theory (DFT) level using the B3LYP functional and the 6-311G* basis set. Comparative calculations showed that the dioseole ring is slightly nonplanar in agreement with experiment,30 thus no symmetry restrictions were imposed. Vertical optical transition energies and oscillator strengths were obtained by time-dependent (TD)DFT. All calculations were carried out within the Gaussian03 program package.31 Orbital pictures were produced with Molekel 4.3.32

3. Results

Synthesis. An alkylated benzo[1,2-d:4,5-d’]bis[1,3]dioxole (BDO, 3, Scheme 1) was synthesized by modified published procedures in good yield.20 BDO 3 was intentionally prepared as a complex mixture of stereoisomers to enhance solubility of the resulting polymers. R and R’ may be syn or anti about the benzene ring plane and the 2-ethylhexyl chains are derived from a racemic starting material. Our first PPEs based on highly symmetric or slightly lower symmetry BDOs carrying linear side chains (e.g., BDOFPPE, scheme 2, R = R’ = n-hexyl; or R = n-hexyl, R’ = ethyl, not shown) were soluble in common organic solvents only at elevated temperature. Lithiation of 3 with BuLi/TMEDA and quenching with I2 produced the diodo BDO 4, from which monomer M1 was synthesized by Pd-catalyzed Negishi coupling.

Polymer BDOFPPE was prepared via nucleophilic aromatic substitution,24 with the fluoride-activated silyl-acetylene groups of monomer M1 acting as nucleophiles and hexafluorobenzene acting as electrophile (Scheme 2). Multinuclear NMR spectra (19F, 1H, 13C, Supporting Information) indicate high structural purity for BDOFPPE, similar to previously reported TAB-based PPEs. Polymers BDOPEE and BDOPE were synthesized using Pd-catalyzed cross- and homocoupling polymerizations, respectively.

Optical Properties. The solution UV/vis absorption spectra (Figure 1a,b) of the polymers TABFPPE, BDOPEE, BDOPE and BDOFPPE exhibit similar features, dominated by a strong absorption maximum (Amax) around 400 nm, similar to many other PPEs.1,2 Also similar to most PPEs, the separation between Amax and the photoluminescence maximum (PLmax), or Stokes shift, is relatively small for TABFPPE (0.15 eV). The absorbance spectra of the three BDO-based polymers, however, contain a more or less pronounced shoulder (A1, Figure 1a, b) at the low energy side of Amax. This is intrinsic to the diluted polymers (i.e., contamination and aggregation as a possible source for A1 were excluded), and is thus assigned to the ground to first excited state transition (S0→S1). The PL spectra of the BDO-based polymers are strongly red-shifted against the main absorption band (Figure 1a). This gives large effective Stokes shifts, varying between 0.75 and 0.95 eV (Table 1) and larger than those reported for most conjugated polymers so far.26,27,33 The Stokes shifts for the monomers M1, M2, and M3 are even larger at 1.45, 1.89, and 1.12 eV, respectively. While the LSS for M1 and M2 are mainly due the large separation between Amax and A1 (1.14 and 1.55 eV), the shift for M3 is mainly due to the large distance between A1 and PLmax (0.72 eV), see Table 1. The latter is attributed to the larger steric effects of...
the alkoxy chains in M3 compared to M1, and stronger Franck-Condon activity of torsional modes. A clear correlation of the PL max energy with the $A_1$ position is observed for the polymers, yielding an energy spacing of 0.4–0.6 eV (Table 1), as might be expected since PL should originate from $S_1$. The polymers show a distinct solvatochromism of the PL spectra, as exemplified for BDOFPPE in Figure 2. With increasing solvent polarity, the emission is shifted to longer wavelength and PL intensity decreases, suggesting intramolecular charge-transfer (ICT) contributions to the emitting state. In the solid state, the optical properties are essentially preserved (Supporting Information). The absorption and emission spectra are red-shifted by ca. 0.05 eV against solution and the effective Stokes shift is of the same magnitude as in solution. The colors of the solid polymers range from yellow for BDOPEE and BDOPPE to red-orange for BDOFPPE. The last is quite different from PPEs based on dialkoxybenzenes 1,2 and the more structurally similar TABFPPE (Figure 1a), which rather emit in the blue-green.

4. Discussion

The electronic differences between BDOs and TABs become clear during the synthesis of the monomers. Bislithiation at the 3,6-positions of BDOs (step “v”, Scheme 1) requires more forcing conditions (TMEDA, $n$-BuLi) compared to TABs ($n$-BuLi, hexanes). This may be a combination of the differing abilities of the TAB or BDO ether oxygens to coordinate to lithium cations, the differing inductive and mesomeric effects of the pendant oxygen atoms, and deviation in hybridization, from sp$^2$ toward sp$^3$, of the carbons to be deprotonated. The C–C–C bond angles around the protonated benzene carbons of BDO are 115.5° (cif file, Supporting Information) compared to the approximately 120° angles of TABs. This is also reflected in the differences in the $^1$H NMR chemical shifts of the phenyl protons, that is, 6.2 ppm for BDO (3) versus 6.5–6.6 ppm for TABs.

Monomer M1 is isolated as a complex mixture of stereo-isomers. Thin layer chromatography of other asymmetric BDOs, namely bearing ethyl and hexyl chains (not shown) indicated that syn- and anti-isomers can be separated. This additional structural variability provides another means to engineer solubility and thermal properties of the polymers. A single stereoisomer crystallizes from the oily mixture of M1 stereoisomers on storage at low temperature, and its crystallographic information can be found in the Supporting Information. The BDO ring system is essentially flat, with acetal carbons deviating just 0.17 Å from the plane of the benzene ring, in contrast to TABs, where the $O$-$\alpha$-carbon bonds of the side chains are nearly

<table>
<thead>
<tr>
<th>Backbone structure</th>
<th>$E(A_1),^a$</th>
<th>$E(A_{\text{max}}),^b$</th>
<th>$\Delta E(A_{\text{max}}-A_1),^c$</th>
<th>$f_A/f_{\text{max}},^d$</th>
<th>$E(\text{PL}_{\text{max}}),^e$</th>
<th>$\Delta E_{\text{Stokes}},^f$</th>
<th>$\Delta E_{\text{AE}},^g$</th>
</tr>
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<tbody>
<tr>
<td>BDOPEE</td>
<td>2.69 [461]</td>
<td>3.04 [407]</td>
<td>0.35 [544]</td>
<td>0.17 [544]</td>
<td>2.28 [544]</td>
<td>0.76 [544]</td>
<td>0.39 [544]</td>
</tr>
<tr>
<td>BDOPPE</td>
<td>2.96 [419]</td>
<td>3.21 [386]</td>
<td>0.25 [517]</td>
<td>0.18 [517]</td>
<td>2.40 [517]</td>
<td>0.81 [517]</td>
<td>0.56 [517]</td>
</tr>
<tr>
<td>BDOFPPE</td>
<td>2.70 [459]</td>
<td>3.07 [404]</td>
<td>0.35 [585]</td>
<td>0.11 [585]</td>
<td>2.12 [585]</td>
<td>0.95 [585]</td>
<td>0.58 [585]</td>
</tr>
<tr>
<td>TABFPPE</td>
<td>--</td>
<td>2.96 [419]</td>
<td>--</td>
<td>--</td>
<td>2.81 [442]</td>
<td>0.15 [442]</td>
<td>--</td>
</tr>
</tbody>
</table>

$^a$ Energies of absorption preband. $^b$ Energies of absorption maximum. $^c$ Energy separation. $^d$ Ratio of oscillator strengths. $^e$ PL maximum. $^f$ Stokes shift $\Delta E_{\text{Stokes}} = E(A_{\text{max}}) - E(\text{PL}_{\text{max}})$. $^g$ Energy separation $\Delta E_{\text{AE}} = E(A_1) - E(\text{PL}_{\text{max}})$. For polymers, the position of the $A_1$ state was extracted from a Gaussian fit to the filled part of the spectrum (Figure 1a), where the same half-width was applied for all polymers.

perpendicular to the benzene plane. In addition to ring-strain effects, this difference must significantly change the mesomeric/inductive influence of the substituents, imparting the (opto-)electronic properties of the BDO monomers and polymers.

To elucidate the nature of the absorbing and emitting states, quantum-chemical calculations were carried out on the corresponding oligomers and polymer properties were obtained by an extrapolation procedure. For consistency in comparing unsubstituted parent homopolymer PPE to the newly reported alternating copolymers having the same size $\pi$-system, a variable length $n$ was defined as the number of alkyne linkages within a series of oligomers. The number of benzene units then equals $n + 1$. For example, the PPE and BDOFPPE structures in Figure 3 both correspond to $n = 3$, despite their differing numbers of constitutional repeating units.

In unsubstituted oligomers and polymers of the PPE type and others including poly(phenylene vinylene) or poly(thiophene), the first excited state ($S_1$) is usually optically strongly allowed, as indicated by the calculated oscillator strengths ($f$) for PPE (Table 2). In these classes of materials, the $S_1$ state is mainly composed of the HOMO to LUMO (H→L)

Figure 1. PL (10 $^{-8}$ M THF) and UV/vis absorption spectra (10 $^{-8}$ M THF). Filled areas are the deconvoluted $A_1$ states (see text). BDO side chains omitted for clarity. All spectra normalized.

Scheme 1. Monomer Synthesis

Scheme 2. Polymer Synthesis

Figure 2. PL solvochromeism of BDOFPPE (10 $^{-8}$ M). All spectra normalized. (Inset) Optical photograph of solutions under 365 nm irradiation.

transition, where both H and L are delocalized along the polymer backbone. This is illustrated in the orbital pictures for an oligomer of the parent PPE at the top of Figure 3. This also accounts for the optical characteristics of most of the substituted species studied so far.

For BDOFPPE oligomers, the situation is very different. The \( S_1 \) state is still mainly composed from the \( H \rightarrow L \) transition, but is only weakly allowed, as seen from the TD-DFT calculated oscillator strengths (\( f \), Table 2). This is due to the very different topology of the HOMO orbital, which is here entirely localized in the BDO unit, whereas the LUMO is delocalized along the backbone (Figure 3, bottom), thus imparting pronounced ICT character to \( S_1 \). The first strongly allowed transition, which gives rise to \( \Delta E \) (Figure 1a), is a higher \( S_1 \) state (\( \pi \rightarrow \pi^* \) mainly composed of \( H-x-1 \rightarrow L \), with \( n \) being the number of alkyne linkages as defined above (Table 2). The H-x-1 orbital for each oligomer is delocalized along the backbone and therefore is the highest occupied delocalized molecular orbital (HOMO). The \( S_1 \) state corresponds to the \( S_1 \) state in unsubstituted PPE and is very similar in energy.

The localized nature of the HOMO orbital can be rationalized by a simple exercise in combining the frontier orbitals of subunits, e.g. BDO and a fluorinated phenylethylenylene (FPE) (Figure 4). The LUMOs of BDO and FPE are relatively close in energy, combining to give delocalized MOs (L, L+2). On the other hand, the energy separation of the HOMOs of the subunits is large, \( \Delta E = 2.03 \text{ eV} \). Thus, the original HOMOs of both BDO and FPE remain localized, whereas the H-1 of BDO interacts with the energetically close H-1 of FPE forming a delocalized MO (H-1). Note that a cross-combination of H of one unit with H-1 of the other unit is not possible due to symmetry mismatch. It is thus the unusually high-lying HOMO of BDO which leads to the localized HOMO topology in the oligomers and polymers.

Calculations show that with increasing chain length \( n \), the energy spacing \( \Delta E(S_1 \rightarrow S_1) \) and thus \( \Delta E(A_{\text{max}} \rightarrow A_1) \) becomes smaller (Table 2 and Figure 5) in agreement with experiment. For example, the experimentally measured \( \Delta E(A_{\text{max}} \rightarrow A_1) \) values are approximately 1.55 eV for monomer M2, but only 0.35 eV for the corresponding polymer BDOFPPE (Figure 1b).

The decrease of \( \Delta E(S_1 \rightarrow S_1) \) with \( n \) is readily seen from the energetic positions of the frontier orbitals plotted in Figure 5. The energy of the delocalized LUMO decreases, while the HOMO energy increases with \( n \) due to the extension of the conjugated \( \pi \)-system, leading to a steep decrease of \( E(S_1) \) with \( n \). On the other hand, the localized HOMO remains almost unchanged in energy, thus yielding a smaller decrease of \( E(S_1) \) compared to \( E(S_1) \) with increasing chain length.

Extrapolating to the polymer limit, the calculation yields an energetic separation between \( A_1 \) and \( A_{\text{max}} \), of ca. 0.2 eV, not far away from experiment \( \Delta E(S_1 \rightarrow S_1) = 0.35 \text{ eV} \) (Table 1). The deviation is ascribed to the shortcomings of the used B3LYP functional, which is known to overestimate the chain length dependence of transitions between delocalized orbitals, though not in our case between HOMO and LUMO, whereas the HOMO–LUMO transition is less affected due to the localized HOMO.

How important is fluorine for the ordering of the states? To answer this question, calculations were also carried out on a series of nonfluorinated BDOPPE oligomers. Within this series the HOMO is again localized, while the LUMO is delocalized. The energy difference between the HOMO and the HODMO again decreases with \( n \), and to a much smaller value, for example, 0.03 eV for \( n = 7 \) compared to 0.36 eV for the fluorinated BDOFPPE (Figure 5). The difference between the two systems results from the high electronegativity of fluorine, which generally stabilizes the orbitals. However, this is only valid for orbitals with a significant contribution from the fluorine-substituted rings, thus in particular for the delocalized orbitals (LUMO, HOMO), see Figure 3. The energy of the HOMO, which is localized on the BDO unit, decreases only slightly with fluorination of neighboring rings. Thus, the energy difference between HOMO and HODMO and thus \( \Delta E(S_1 \rightarrow S_1) \) will be larger in the fluorinated compounds, in agreement with the experimental findings, see Table 1. The virtual independence of HOMO energy on whether or not fluorine is present agrees with the electrochemical measurements of BDOPPE and BDOFPPE (Table 3). The oxidation potentials for both polymers agree to within 0.01 V (0.01 eV difference in \( E(S_1) \)), within experimental error.

How important is the BDO unit for the ordering for the states? Upon replacing BDO by TAB in an oligomer with \( n = 1 \), the localized HOMO is stabilized by \( \approx 0.2 \text{ eV} \), whereas the HODMO is destabilized by \( 0.2 \text{ eV} \), see Figure 5. Thus, an ICT emitting state is predicted for short oligomers, in agreement with a large experimental effective Stokes shift and absorbance preband for TAB-based monomer M3 (Figure 1c). On the other hand, for longer chain length \( n \) an early crossing of the HOMO with the HODMO is expected (Figure 5). At the polymer limit the lowest excited state will be determined by the HODMO–LUMO transition typical for PPEs, and no low intensity ICT preband or large Stokes shift is expected, in agreement with the experimental results for the polymer TABFPPE in solution (Figure 1a). It is important to note that the MO energies critically depend on the geometry of the TAB units. The calculations described above employed the lowest energy side-chain conformation, with the sp\(^2\) α-carbons of the other chains

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(36) For the electronic states in BDOPPEs, this leads to a crossing of the ICT state with the delocalized state, thus (TD-)DFT predicts ICT not to be the lowest state in BDOPPE in contradiction with experiment, which is again due to the shortcomings of (TD-)DFT for polymers, vide infra.

extending out of the plane of the benzene ring. For an imposed all-planar (or nearly planar) structure, the calculated MO energies and thus optical transitions are more similar to BDO-based polymers.

The localized topology of the HOMO orbital of the novel copolymers is thus a very peculiar electronic feature imposed by the BDO unit, which is not only found for short oligomers as in the TAB case, but also at the polymer limit. HOMO localization is typically not known in common all-conjugated donor—acceptor (D−A) type copolymers. For conventional materials built from units with pronounced ICT character,38 the ICT character becomes typically smaller with increasing chain

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**Table 2.** Calculated Electronic Levels ($E_{\text{HOMO}}$, $E_{\text{LUMO}}$), Band Gap $\Delta E_{\text{H-L}}$, Vertical Transition Energies $E_{\text{vert}}$ and their Composition (Configuration Interaction, or “CI” Contributions) and Oscillator Strengths $f$ of the First Two Optical Transitions $S_1$, $S_2$

<table>
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<tr>
<th>polymer</th>
<th>$n^a$</th>
<th>$E_{\text{HOMO}}$ (eV)</th>
<th>$E_{\text{LUMO}}$ (eV)</th>
<th>$\Delta E_{\text{H-L}}$ (eV)</th>
<th>$E_{\text{vert}}$ main CI contrib.</th>
<th>$f_{S_1}$</th>
<th>$f_{S_2}$</th>
<th>$\Delta E (S_1\rightarrow S_2)$</th>
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<tr>
<td>PPE 1</td>
<td>1</td>
<td>-5.963</td>
<td>-1.530</td>
<td>4.43</td>
<td>S1: 4.16 H→L</td>
<td>0.94</td>
<td></td>
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</tr>
<tr>
<td>3</td>
<td></td>
<td>-5.603</td>
<td>-2.236</td>
<td>3.37</td>
<td>S1: 2.07 H→L</td>
<td>2.73</td>
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<td></td>
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<tr>
<td>7</td>
<td></td>
<td>-5.491</td>
<td>-2.533</td>
<td>2.96</td>
<td>S1: 2.00 H→L</td>
<td>5.66</td>
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<tr>
<td>7</td>
<td>-5.417</td>
<td>-5.383</td>
<td>-2.542</td>
<td>2.84</td>
<td>S1: 2.99 H→L</td>
<td>0.04</td>
<td>0.99</td>
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</table>

**Figure 4.** Correlation diagram for the frontier molecular orbitals of a single BDOFPPE linkage from the corresponding subunits at the DFT B3LYP/6-311G* level of theory.

**Figure 5.** (TD-)DFT calculated electronic transition energies (top) and orbital energies (bottom) of BDOPPE (○), BDOFPPE (●) and TABFPPE (×) as a function of the inverse chain length $1/n$, where $n$ = number of alkyne linkages.

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$^a$ $n$ = number of alkyne linkages. The number of benzene rings = $n + 1$. See for example the structures in Figure 3 where $n = 3$. $^b$ HODMO = Highest occupied delocalized molecular orbital.
length,\textsuperscript{39} thus preserving only a partial ICT character of the frontier orbitals at the polymer limit.\textsuperscript{39} Moreover, in D–A polymers reported so far, the LUMO is generally more localized than the HOMO\textsuperscript{38,39,41} opposite to the polymers studied here. This is due to typically larger differences in the LUMO energies, compared to the HOMO energies, of the donor and acceptor building blocks.\textsuperscript{38} A PPE copolymer reported by Swager,\textsuperscript{42} composed of alternating pentiptycene and diaminobenzene units, also yields a large energy spacing between absorption and PL maxima, however with an absorption preband of much higher absorbance than in the polymers reported by us. Given the very bulky nature of the repeating units, the energy shift was reasonably attributed to large conformational differences between ground and excited states. Preliminary calculations by us (not shown) support this geometrical concept, but also stress the contribution of localized occupied frontier orbitals at the pentiptycene units in a rather complex CI description of the lowest excited state. This indicates that the right choice of the D and A building blocks for the polymer repeating units—which can be conveniently assisted by appropriate quantum-chemical methods—opens the pathway to design polymers with precisely defined orbital energies and topologies, and thus to tailor the electronic and optical properties, which govern the electron and energy transport in the active organic layer of opto-electronic devices.

5. Conclusion
We have synthesized and characterized new conjugated copolymers based on a benzodioxole (BDO) and phenylene-ethynylene (PE) repeating units. Very different to donor–acceptor copolymers reported so far, the new copolymers display a peculiar electronic structure with localized HOMO and delocalized LUMO topologies. The HOMO localization is due to the high lying HOMO of the constituent BDO unit, which cannot mix with the respective MO of the PE unit. As a consequence, the lowest optical transition shows a pronounced ICT character and shows up as a low-intensity preband below the intense main absorption band. The latter is composed from delocalized orbitals as in common conjugated polymers. Very large effective Stokes shifts of these materials both in solution and the solid state are thus observed, extending those of polymers reported in literature. Future derivatives might thus be good candidates for large Stokes shift applications, e.g., lasers, molecular imaging, scintillators, solar collectors and white light emitting materials. Further, bulky substituents which will be orthogonal to the polymer backbones when attached to the BDO acetal carbons, should lead to intrinsically nanoporous polymers as hosts/sensors for electron-accepting guests/analytes.

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Supporting Information Available: Experimental procedures, spectroscopic, electrochemical and crystallographic data. Complete refs 31 and 39. This material is available free of charge via the Internet at http://pubs.acs.org. JA9068134

Table 3. Electrochemical Data (eV)

<table>
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<tr>
<th></th>
<th>$E_{1/2}^{a}$</th>
<th>$E_{1/2}^{b}$</th>
<th>$E_{HOMO}^{c}$</th>
<th>$E_{LUMO}^{d}$</th>
<th>$E_{gap}$\textsuperscript{f}</th>
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<tr>
<td>3</td>
<td>3.6</td>
<td>0.32</td>
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<td>-1.5</td>
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<tr>
<td>M1</td>
<td>3.0</td>
<td>0.57</td>
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<tr>
<td>M2</td>
<td>3.0</td>
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<td>-2.4</td>
<td>-5.49\textsuperscript{g}</td>
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<td>BDOFPPE</td>
<td>2.4</td>
<td>0.81</td>
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<td>-3.2</td>
<td>-5.52\textsuperscript{g}</td>
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<tr>
<td>BDOPE</td>
<td>2.7</td>
<td>0.68</td>
<td>-5.5</td>
<td>-2.8</td>
<td>-5.38\textsuperscript{g}</td>
</tr>
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<td>BDOPEE</td>
<td>2.6</td>
<td>0.80</td>
<td>-5.6</td>
<td>-3.0</td>
<td>-5.47\textsuperscript{g}</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Optical energy gap from intersection of absorption and emission in solution. \textsuperscript{b} Half-wave oxidation potential referenced to Fc/Fc\textsuperscript{+} couple (0.46 V). \textsuperscript{c} $E_{HOMO} = (-E_{LUMO}^{CI} + 4.8)$ eV. \textsuperscript{d} Calculated at a first approximation from $E_{LUMO} = E_{HOMO} + E_{gap}^{CI}$. \textsuperscript{e} DFT calculations. \textsuperscript{f} Extrapolation of the DFT values in Table 2 to the polymer limit.