pi-Dimers of end-capped oligopyrrole cation radicals

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**Communications**

**π-Dimers of End-Capped Oligopyrrole Cation Radicals**


The conductivity of doped polymers with conjugated π-electron systems is directly related to their redox states and part of it has been rationalized in terms of polarons (cation radicals) and bipolarons (dications) migrating along the conjugated chain.11 The general understanding of the electronic structure and redox states of semiconducting polymers has benefited from studies on well-defined conjugated oligomers that allow functional properties to be investigated as a function of (effective) conjugation length.12-15 In recent years, numerous studies have addressed the redox states of oligothiophenes,12-15 and it was found that in addition to cation radicals and dications, spinless π-dimers of cation radicals can be formed.13-15 These interchain cation radical π-dimers have been proposed as an alternative to diamagnetic bipolarons as an explanation for the weak ESR signal for highly oxidized polythiophenes.13-15 Although several organic ion radicals form π-complexes in solution, dimerization has never been reported for conjugated oligomers other than oligothiophenes. The availability of well-defined oligopyrrolole (Table 1). The chemical stability of the first two oxidation states of PhPy,Ph is a result of the phenyl end caps, which prevent polymerization over the α-positions. The successive electrochemical oxidation of the oligomers to cation radicals (PhPy,Ph3+) and dications (PhPy,Ph22+) at controlled potentials was monitored by in-situ UV/VIS/NIR absorption spectroscopy (Fig. 1). Several optical transitions, listed in Table 1, are observed. The number and relative ordering of the absorption bands closely resembles the spectral characteristics of oxidized oligothiophenes.12-15 For the oligopyrrole cation radicals two bands (M1 and M2) appear at low and high energy, respectively. Each of the M1 and M2 bands has an associated second transition (D1 and D2), shifted to higher energy by about 0.3 eV. These second transitions are attributed to π-dimers of oligopyrrole cation radicals. In contrast, for the oligopyrrole dications, only a single electronic transition (DC) is observed, with one or two associated vibronics at higher energy. Each of the optical transitions (N, M1, M2, D1, D2, DC; Table 1) shifts to lower energy for longer oligomers, in accord with an approximately linear dependence with the inverse length of the oligomer (taken as 1/m; where m = n + 2 is the total number of aromatic rings).16 The various bands can be assigned to specific transitions using the molecular symmetry of PhPy,Ph. Assuming an all-transoid conformation of the bonds between pyrrole rings, the molecular symmetry is either C2v (n = 2, 4) or Cn (n = 3). The symmetry of the frontier π orbitals is given in Figure 2a; the dipole-allowed transitions among these orbitals are indicated. Because these excitations are all polarized along the oligomer chain axis they are expected to be intense.17 The π-interaction between two cation radicals to give a π-dimer will cause a splitting of frontier orbitals into bonding (+) and antibonding (−) levels (Fig. 2b). In this dimer the D1 (−−) and D2 (+−) transitions (originating from the states M1 and M2, respectively, in the cation radical) are dipole-allowed and polarized along the

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**Table 1.** Electrochemical data [a] and optical transition energies [b] of redox states of PhPy,Ph oligopyrroles.

<table>
<thead>
<tr>
<th></th>
<th>n = 2</th>
<th>n = 3</th>
<th>n = 4</th>
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</thead>
<tbody>
<tr>
<td>E0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E1</td>
<td>0.37</td>
<td>0.13</td>
<td>0.01</td>
</tr>
<tr>
<td>E2</td>
<td>1.05</td>
<td>0.60</td>
<td>0.37</td>
</tr>
<tr>
<td>N</td>
<td>3.05</td>
<td>2.89</td>
<td>2.78</td>
</tr>
<tr>
<td>M1 [c]</td>
<td>1.60</td>
<td>1.79</td>
<td>1.13</td>
</tr>
<tr>
<td>M2</td>
<td>2.55</td>
<td>2.26</td>
<td>2.10</td>
</tr>
<tr>
<td>D1</td>
<td>1.91</td>
<td>1.65</td>
<td>1.39</td>
</tr>
<tr>
<td>D2</td>
<td>2.86</td>
<td>2.63</td>
<td>2.42</td>
</tr>
<tr>
<td>DC [c]</td>
<td>2.21</td>
<td>2.34</td>
<td>1.88</td>
</tr>
</tbody>
</table>

[a] Potentials in CH2Cl2/NBu4PF6 (0.1 M) solution at 100 mV s−1 and oligomer concentration c = 0.1 – 1.0 mM vs SCE, internally calibrated vs Fe/Fe3+. [b] Transitions of redox states labeled as follows: N: neutral oligomer; M1, M2: monomer cation radical; D1, D2: dimer of cation radicals; DC: dication. [c] Second and third entries are attributed to different vibronics of the same electronic transition.

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* Dr. R. A. J. Janssen, J. A. E. H. van Haare, L. Groenendaal, Dr. E. E. Havinga, † Prof. E. W. Meijer
Laboratory of Organic Chemistry, Eindhoven University of Technology
P. O. Box 513, NL-5600 MB Eindhoven (The Netherlands)
Fax: Int. code + (40)245036
E-mail: tgoerl@chem.tue.nl

† Also at Philips Research Laboratories, Eindhoven (The Netherlands)

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(Notes and references have been omitted for brevity.)
It was observed that the D1 and D2 transitions are shifted to higher energy than M1 and M2 by about 0.3 eV, comparable with shifts found for oligothiophene cation radical dimers and other n-radical ion dimers. Such a blue shift is a well-known phenomenon for ionic n-dimers and has been attributed to the interaction between transition dipole moments on the adjacent cation radicals (Davydov shift), although other phenomena may contribute such as different splittings of frontier orbitals upon dimerization. In addition to D1 and D2, two charge transfer transitions CT1 and CT2 are possible in the n-dimer, polarized along the short axis connecting the two cation radicals. As a consequence, the CT1 and CT2 bands are expected to be less intense than D1 and D2.

The temperature dependence of the electronic spectrum of PhPy,Ph⁺ (Fig. 3), obtained by chemical oxidation with FeCl₃ in CH₂Cl₂ solution between 295 and 245 K (c = 0.2 mM, path length 10 mm), reveals that the intensities of the M1 and M2 bands decrease at lower temperatures in favor of the intensities of the D1 and D2 bands. These changes with temperature are reversible and consistent with the proposed equilibrium between two oligopyrrole cation radicals and a diamagnetic π-dimer.

In conclusion we have shown that phenyl end-capped oligopyrroles can be oxidized reversibly to the corresponding oligomers in a 1:1 mixture of CH₂Cl₂ and trifluoroacetic acid (TFA). Figure 4 shows the ESR spectrum of PhPy,Ph⁺ in CH₂Cl₂/TFA. The ESR spectrum is fully symmetric, indicating the formation of a single doublet species and arguing against the presence of different conformations of the PhPy,Ph⁺ cation radical (e.g. s-transoid and s-cisoid), as has been found for oligothiophene cation radicals. The intensity of the ESR signal decreases by one order of magnitude upon lowering the temperature from 295 K to 200 K (Fig. 4). This temperature dependence is reversible and in agreement with the expected behavior for an equilibrium between two oligopyrrole cation radicals and a diamagnetic π-dimer.

The in situ electrochemical or chemical (FeCl₃) oxidation of PhPy,Ph to the cation radical produces a strong ESR signal at g = 2.0025. A subsequent decrease of ESR signal intensity is observed upon further oxidation, leading to the loss of the signal after complete conversion of the cation radical to the (diamagnetic) dication. Highly resolved ESR spectra of the oligopyrrole cation radicals can be obtained by oxidizing the PhPy,Ph oligomers.

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**Fig. 2.** Schematic energy diagram and symmetry of the frontier orbitals for the neutral, cation radical, and dication states of PhPy,Ph (a), and cation radical π-dimers (b). Dipole-allowed transitions are indicated. Experimentally D1 and D2 exhibit an additional blue shift as a result of a Davydov interaction.

**Fig. 3.** Electronic spectrum of PhPy,Ph⁺ cation radicals, obtained by oxidation with FeCl₃ in CH₂Cl₂ solution between 295 and 245 K (c = 0.2 mM, path length 10 mm). The arrows mark the decay of the M1 and M2 transitions and the increase of D1 and D2 transitions with decreasing temperature.

**Fig. 4.** ESR spectrum of the PhPy,Ph⁺ cation radical generated by dissolving PhPy,Ph (0.4 mM) in CH₂Cl₂/TFA (1:1) at 295 K. The inset shows the temperature dependence of the ESR intensity. The ESR spectrum can be simulated by using the following parameters: α_H(4H) = 0.614, α_H(2H) = 0.853, α_H(4H) = 0.328, α_H(2H) = 0.703, α_D(2H) = 0.395, α_D(2H) = 0.470, and D_H = 0.03 mT. These couplings are tentatively assigned to o-PhH, p-PhH, m-PhH, CH₂, C₄H, and NH based on a semiempirical quantum chemical calculation using the PM3 Hamiltonian.
cation radicals and dications. Variable temperature UV/Vis/NIR and ESR spectra give direct evidence for the \(\pi\)-dimerization of oligopyryrole cation radicals. The results on oligopyrryles are in full analogy to \(\pi\)-dimerization of oligothiophene cation radicals.\(^1\)\(^2\)\(^3\)\(^4\) Therefore, the present results together with those obtained on oligothiophenes, give credence to the proposition that \(\pi\)-dimers have an important role as interchain conducting entities for conducting polymers in general.

**Experimental Procedure**

The preparation of PhPy, Ph oligomers and conditions for cyclic voltammetry have been described elsewhere.\(^8\) All experiments were conducted under rigorously inert conditions. UV/Vis/NIR spectrophotometry data were obtained with a Perkin Elmer Lambda 9 spectrophotometer in CH\(_2\)Cl\(_2\)/NBu\(_4\)PF\(_6\) (0.1 M) solution with a thin-layer cell consisting of two ITO glasses separated by a spacer of 0.1 mm as the optical transparent working electrode, a platinum disk as a counter electrode, and an Ag/AgCl wire as reference electrode.\(^4\) ESR spectra were recorded on a Bruker ER200D SRC X-band spectrometer with a ER4111 variable temperature unit.

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The UV/Vis/NIR spectrum of PhPy, Ph in CH\(_2\)Cl\(_2\)/TFA exhibits the same transitions as observed after electrochemical or chemical oxidation with FeCl\(_3\).


\(^14\) From the one order of magnitude decrease of the ESR intensity it is possible to estimate the enthalpy of dimerization to be on the order of \(\Delta H^0 = -25 \text{ to } -35 \text{ kJ mo}^{-1}\). Further studies on the equilibrium and thermodynamic parameters are currently in progress.

**Solid-Phase, Parallel Syntheses by Ugi Multicomponent Condensation**

Paul A. Tempest, S. David Brown, and Robert W. Armstrong*

Dedicated to Professor Ivar Ugi on the occasion of his 65th birthday

The combinatorial synthesis of chemical libraries of small organic molecules represents a new tool for investigating the effects of structure on biological activity.\(^1\)\(^2\) Recent efforts have focused on the solid-phase syntheses of a core set of atoms (or a pharmacophore), in which diversity is introduced sequentially at each step of a linear synthesis.\(^1\)\(^2\) The theoretical number of compounds that can be generated in these libraries is related to the number of steps required for their synthesis and the availability of chemical starting compounds (inputs) for each step. An alternative strategy for the generation of chemical libraries involves the use of multicomponent condensation reactions (MCCs).\(^1\)\(^2\) MCC-derived compounds require a single synthetic transformation. The potential size of these libraries is dependent on the number of components in the MCC and their availability—not on the number of synthetic steps. Described herein is the solid-phase synthesis of a library of \(\alpha\)-acylamino amides by a four-component condensation (4CC) and a strategy for increasing the structure diversity of one of its inputs by a pre-MCC condensation of two substrates in solution.

The Ugi 4CC is ideally suited for the construction of chemical libraries based on the \(\alpha\)-acylamino amide core structure (Scheme 1). When the synthesis is conducted on a solid sup-

![Scheme 1. A 4CC reaction on solid support (P) to generate compounds with an \(\alpha\)-acylamino core.](image)

They are readily available. A secondary amide results upon cleavage of the amino group since a variety of amino-functionalized matrices (aldehyde, carboxylic acid, isonitrile, and amine) can be tethered to the solid support. While any of the four inputs (aldehyde, carboxylic acid, isonitrile, and amine) can be used, the four-component condensation of an amine, and 1 isocyanide were the starting compounds in a pre-MCC condensation of two substrates in solution. The combinatorial synthesis of chemical libraries of small organic molecules represents a new tool for investigating the effects of structure on biological activity.\(^1\)\(^2\) Recent efforts have focused on the solid-phase syntheses of a core set of atoms (or a pharmacophore), in which diversity is introduced sequentially at each step of a linear synthesis.\(^1\)\(^2\) The theoretical number of compounds that can be generated in these libraries is related to the number of steps required for their synthesis and the availability of chemical starting compounds (inputs) for each step. An alternative strategy for the generation of chemical libraries involves the use of multicomponent condensation reactions (MCCs).\(^1\)\(^2\) MCC-derived compounds require a single synthetic transformation. The potential size of these libraries is dependent on the number of components in the MCC and their availability—not on the number of synthetic steps. Described herein is the solid-phase synthesis of a library of \(\alpha\)-acylamino amides by a four-component condensation (4CC) and a strategy for increasing the structure diversity of one of its inputs by a pre-MCC condensation of two substrates in solution.

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\[^*\] Prof. R. W. Armstrong, P. A. Tempest, S. D. Brown

Department of Chemistry and Biochemistry

University of California, Los Angeles

Los Angeles, CA 90095-1569 (USA)

Fax: Int. code +(310)825-0767

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