**Oligothiophenes**

**Inherently Chiral Macrocyclic Oligothiophenes: Easily Accessible Electrosensitive Cavities with Outstanding Enantioselection Performances**

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**Abstract:** Linear conjugated oligothiophenes of variable length and different substitution pattern are ubiquitous in technologically advanced optoelectronic devices, though limitations in application derive from insolvency, scarce processability and chain-end effects. This study describes an easy access to chiral cyclic oligothiophenes constituted by 12 and 18 fully conjugated thiophene units. Chemical oxidation of an “inherently chiral” sixthiophene monomer, synthesized in two steps from commercially available materials, induces the formation of an elliptical dimer and a triangular trimer endowed with electrosensitive cavities of different tunable sizes. Combination of chirality with electroactivity makes these molecules unique in the current oligothiophenes literature. These macrocycles, which are stable and soluble in most organic solvents, show outstanding chiroptical properties, high circularly polarized luminescence effects and an exceptional enantio-recognition ability.

We have recently reported the example of a new approach to chiral electroactive oligothiophenes, consisting in the electro-oligomerization of monomers (R)-1a and (S)-1a in which chirality results from an internal torsion in the conjugated backbone (from here the appellative of inherently chiral monomers).[1] According to this design, the same conjugated system responsible for the electro-optical properties of these materials is also responsible for molecular chirality, implying that electrochemical, chiroptical and enantio-recognition properties are strictly interdependent (Scheme 1).

The conducting films resulting from electro-oxidation of enantiopure (R)-1a and (S)-1a, constitute oligomers from the dimer (largely prevailing) to the pentamer (present in traces). They display impressive chiroptical activity, reversible potential-driven variations of the circular dichroism (CD) signal and outstanding enantioselection ability towards a chiral ferrocenylamine selected as an electrochemical probe.

Here we report the results obtained when performing the chemical oxidation of racemic 1a and its enantiopure antipodes with FeCl₃ excess, in CHCl₃ solution, under stirring, at RT, according to a procedure currently employed for preparing polythiophenes from electron-rich monomers when regioselectivity problems are not involved.[2]

The preliminary experiments have been carried out on racemic (±)-1a.[3] Extraction with THF in a Soxhlet apparatus of the crude reaction product to remove some insoluble iron-containing materials, gives an orange residue (50% yields in weight), composed of dimers (67%), trimers (27%), tetramers (2%) and pentamers (less than 1%), on the basis of peak intensities in laser desorption ionization (LDI) mass spectrum (see the Supporting Information section SI 1). High resolution LDI (HR-LDI) experiments show that the molecular weights and the isotopic

**Scheme 1.** The inherently chiral enantiomeric monomers employed in the present research.
patterns of all the components of the mixture are in agreement with formulas having two hydrogen atoms less than the expected values for linear oligomers (see the Supporting Information sections SI 2 and SI 3), as if the thiophene terminals of the oligomers had undergone a FeCl₃-promoted intramolecular coupling.

To prove this hypothesis, we submitted to the same FeCl₃ treatment the dideuterated compound (±)-1b, prepared by quenching with D₂O the dianion of (±)-1a, obtained in turn by lithiation of the latter with nBuLi excess, in the presence of N,N,N′,N′-tetramethylethlenediamine, in THF solution, at −78 °C (see the Supporting Information section SI 4). We found (HR-LDI, see the Supporting Information section SI 5) that the deuterium had been totally lost, thus proving the cyclic structure of the oligomers. These MS experiments did not give evidence of the formation of open chain compounds, which are largely present (1:1) in the electrochemically prepared material (HR-LDI, see the Supporting Information section SI 6).

These results demonstrate that the FeCl₃ oxidation of monomer 1a generates a series of cyclic oligothiophenes, constituted by 12, 18, 24 and so on, conjugated thiophene units. When produced from enantiopure starting materials, all of them must be inherently chiral macrocycles constituted by a single Dₘ symmetric stereoisomer.

Oligothiophene macrocycles are not only aesthetically charming but operationally active molecules. They idealize conducting polymers without ends, offering electroactive cavities of different sizes for selective inclusion of guest molecules. The α-cyclo[n]thiophenes have been investigated by Bäuerle and co-workers who studied both their synthetic accessibility through different strategies and their physicochemical and electro-optical properties.[1] In the present case, the oligothiophene macrocycles display non-interrupted π systems, even though not fully α-conjugated. Combination of chirality with electroactivity makes these cavities unique in the current library of oligothiophenes.

In this context, we considered as primary target the isolation of the more abundant oligomers present in the mixture resulting from the FeCl₃ oxidation of both the enantiopure anti-podes of 1a, since the oxidation of racemic (±)-1a is expected to afford diastereomeric mixtures of cyclic oligomers produced by homochiral or heterochiral couplings: a pair of Dₘ symmetric enantiomers and a Cₘₙ symmetric meso compound in the case of the dimer 2 and two diastereomeric racemates, one Dₘ and one Cₘₙ symmetric, in the case of the trimer 3.

Accurate column chromatography enabled us to isolate the enantiopure dimer (S,S)-(−)-2 and the (S,S,S)-(−)-3 trimer from (S)-(−)-1a and, specularly, the (R,R)-(−)-2 dimer and the (R,R,R)-(−)-3 trimer from (R)-(−)-1a, in 15 and 8% isolation yields, calculated on 1a, respectively, for dimers and trimers (Figure 1). The enantiomeric purity of the four compounds was checked and demonstrated by analytical HPLC on a chiral stationary phase (see the Supporting Information section SI 7).

The ¹H and ¹³C NMR spectra of enantiopure cyclodimers 2 and trimers 3 unequivocally proved the Dₘ and Dₙ symmetry of the molecules. According to stereotopism rules, only four coupled signals for the benzothiophene ring protons and four doublets for the protons in β-position of the two thiophene rings directly connected to it are observed in the ¹H NMR spectra, while only sixteen signals are present in the ¹³C NMR spectra (see the Supporting Information section SI 8).

A picture of the shapes of homochiral dimers 2 and trimers 3 (Figure 1, http://users.unimi.it/ChiralCycles) was obtained by density functional theory (DFT) calculations in the generalized gradient approximation, using the Perdew–Burke–Ernzerhof functional to handle exchange-correlation effects (see the Supporting Information section SI 9).[10]

The cavities are characterized by different internal sizes and shapes, small and elliptical in the case of 2 (the major and

**Figure 1.** a) Planar molecular formulas of cyclic dimer 2 and trimer 3 evidencing connectivity. Bonds in red highlight the 3,3’-bithiophene junctions corresponding to the stereogenic axes, while bonds in blue indicate the oligomerization sites. b) Calculated structures of (R,R)-2 and (R,R,R)-3.
minor axis lengths are about 17.7 and 5.8 Å, respectively) and quite large, equilateral triangular, in the case of 3 (side of the triangle: ~17.9 Å long; geometrical height: ~17.0 Å). It is interesting to note how bent outwards are the two facing sequences of the six α-linked thiophene units in dimers 2 under the constraint of the atropisomeric scaffolds, with consequent conjugation loss.

From the spectroscopic point of view, both 2 and 3 display absorption maxima significantly red-shifted with respect to that of the monomeric unit 1a; this is attributable to the spin allowed π-π transition ($
\lambda_{\text{max}}$ = 423 nm, $\varepsilon = 5.3 \times 10^4 \text{M}^{-1} \text{cm}^{-1}$) for 2; $\lambda_{\text{max}}$ = 447 nm, $\varepsilon = 1.5 \times 10^5 \text{M}^{-1} \text{cm}^{-1}$ for 3 and $\lambda_{\text{max}}$ = 372 nm, $\varepsilon = 4.8 \times 10^4 \text{M}^{-1} \text{cm}^{-1}$ for 1a in CH$_2$Cl$_2$ solution (Figure 2a). This feature is in agreement with the extended π-conjugated system involving the six α-linked thiophene units. However, it is interesting to note that the absorption maximum of 2 is slightly blue-shifted with respect to that of the corresponding linear α-sixthiophene ($\lambda_{\text{max}}$ = 445 nm, $\varepsilon = 5.0 \times 10^4 \text{M}^{-1} \text{cm}^{-1}$). This is mainly due to the constraint of the atropisomeric scaffolds, which bends outwards the two sequences of the six α-linked thiophene units with consequent conjugation loss. A sterically more relaxed situation characterizes trimers 3, where the rings of the α-sixthiophene sequences are coplanar, with remarkable gain in conjugation extent. However, the broad features of the absorption spectra indicate that 2 and 3 still maintain conformational flexibility in the ground state (see the Supporting Information section SI 10).

Films of enantiopure and racemic dimer 2 and trimer 3, drop-casted on screen printed electrodes are electroactive and conducting, affording reversible and repeatable oxo-reduction cycles (see the Supporting Information section SI 11). The cyclic voltammetry (CV) of enantiopure 2 displays a peculiarly sharp oxidation peak which is lost in racemic 2, pointing to a better supramolecular organization. In all cases the activation of the redox sites strongly depends on the scan rate, in accordance with slow doping/undoping processes similar to those displayed by oligothiophene films. The chiroptical manifestations (optical rotation and circular dichroism) of the antipodes of 2 and 3 are outstanding, as expected, since produced by extended inherently dissymmetric chromophores (Figure 2a).

Preliminary experiments on enantiopure 2 and 3 were focused: 1) to evaluate their circularly polarized luminescence (CPL) activity; 2) to check their enantiorecognition capacity in front of classical electrosensitive chiral probes.

Fluorescence and CPL simultaneous measurements were carried out in a toluene solution, at RT ($\lambda_{\text{mon}}$ = 450 nm for 2 and 470 nm for 3). Upon optical excitation, both 2 and 3 display intense orange-yellow vibrationally structured emission at $\lambda_{\text{em}}$ = 545, 573 nm and 542, 580 nm, respectively (Figure 2b). These structured emissions are consistent with the planar quinoid-like structure proposed for the singlet excited-state of oligo-

Figure 2. a(2), a(3): CD and UV spectra and [α]$_D^{25}$ (CHCl$_3$) of (S,S)-(−)-2 and (R,R)-(−)−2 (6.0×10$^{-4}$ M), and of (S,S)-(−)-3 and (R,R)-(−)-3 (3.0×10$^{-4}$ M), b(2), b(3): Emission spectra of (−)-2 (green) and (−)-3 (6.0×10$^{-4}$ M) and of (−)-3 (3.0×10$^{-4}$ M) in aerated toluene solution at room temperature. Emission spectra were corrected for emission spectral response (detector and grating) by standards correction curves. c(2), c(3): CPL of (−)-2 and (−)-3 and CPL of (+)-3 and (−)-3 under the same conditions. Spectra of enantiomers with S stereogenic axes figure are in green, with R configuration in red.

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The Stokes shift observed for 2 ($\Delta \lambda_{\text{max}} = 122$ nm), which is larger than that observed for 3 ($\Delta \lambda_{\text{max}} = 96$ nm), suggests that the ground state of the latter better matches the quinoid-like geometry of the excited state requiring a lesser conformational reorganization. This feature is in agreement with the more planar conformation of the \( \alpha \)-sexithiophene sequences in 3, as indicated by DFT calculations and absorption features. The extent of backbone rigidity in the excited state has a significant impact on the photoluminescence quantum yields ($\Phi_L$), which are found to be 16 and 37% for 2 and 3, respectively. The more accessible nonradiative deactivation pathways, favored by torsional vibration motions and responsible for the lower quantum yields observed for 2, match the high steric constrain of this species, both in ground and excited states.

The CPL emission is notable (Figure 2c), with an anisotropy $g$ factor close to $3 \times 10^{-3}$, comparable with other inherently chiral molecules like helicenes\(^{16}\) and higher than polythiophenes with chiral side chains.\(^{15}\) These properties suggest possible applications of these materials in chiroptical devices (see the Supporting Information section SI 12).

Tests of the enantiorecognition capacity of the enantiopure macrocycles were performed on gold screen printed electrodes in (BMIM)PF\(_6\), employing \((S)-(-)-\) and \((R)-(+)\)-N,N-dimethyl-1-ferrocenylethylamine, \((S)-4\) and \((R)-4\), as chiral probes. The same probes had been previously employed for evaluating the enantiodiscrimination properties of the electrochemically prepared materials (Figure 3).\(^{11}\)

On bare gold electrodes \((R)-4\) and \((S)-4\) result in the same CV pattern, a diffusive reversible peak at a formal potential very similar to that of ferrocene. Outstanding enantiorecognition is achieved when the electrodes are coated with the enantiomers of 2 and 3.

In particular, the electrodes coated with the enantiomeric cyclodimers \((R,R)-(++)-2\) and \((S,S)-(--)-2\) result in specular separation of enantiopure \((R)-4\) and \((S)-4\) probes, with a $E_{pa}$ separation of approximately 90 mV; a slightly larger $E_{pa}$ separation ($\sim$110 mV) is obtained with the electrodes coated with enantiomeric cyclotrimers \((R,R,R)-(--)-3\) and \((S,S,S)-(++)-3\).

It is worthwhile noting that, unlike the performances of the films obtained by 1a electro-oligomerization,\(^{11}\) the CV peak morphology resembles a reversible diffusional process (distance of 60–70 mV between forward and backward peak even at low scan rates) rather than a solid state one (specular forward and backward peaks). This could point either to a different mode in the approach of the probe molecule to the gold surface within the film and/or to the fact that the reaction occurs on the film surface, slightly charged at the potential at which the electron transfer takes place.

Finally, the enantioselective performance in the case of experiments with the \((\pm)-4\) racemic probe is spectacular; the electrodes coated with \((S,S)-(--)-2\) result in a 190 mV $E_{pa}$ separation; also in this case, the electrodes coated with enantiopure trimer \((S,S,S)-(++)-3\) perform slightly better than those coated with the corresponding dimer, resulting in an impressive 210 mV $E_{pa}$ separation.

In conclusion, a very simple strategy with high preparative value was discovered for achieving inherently chiral electroactive oligothiophene macrocycles, soluble in organic solvents and therefore easily processable, starting from inexpensive thiophenes.\(^{7}\)
monomers. This combination of properties makes the new molecules an outstanding entry in the library of dissymmetric organic cavities. The preparative enantioseparation of the monomer racemate is not a severe limitation given the capacity of modern chiral technologies to resolve racemates at the level of several kilograms. It is also probable that the crude complex of several kilograms could find application in fields where chirality is not important, like energy and optoelectronics.

The formation of cyclic systems from 1a looks general for open-blade scissor-shaped molecules (directional bond strategy), and we are convinced that the oxidative oligomerization and cyclization steps would occur on the FeCl₃ surface. Computed DFT cyclization energetics was found to be only moderately endothermic (+94.5 and 46.9 kJ mol⁻¹), for homodimer 2 and homotrimer 3, respectively, if referenced to gas-phase H₂, thereby suggesting that cyclization is indeed energetically feasible under these reaction conditions, where the removed hydrogen atoms are likely to adsorb on the oxidant phase H₂, thereby suggesting that cyclization is indeed energetically feasible under these reaction conditions, where the removed hydrogen atoms are likely to adsorb on the oxidant microparticles.

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