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# Bithiophene-based polybenzofulvene derivatives with high stacking and hole mobility $\ensuremath{\dagger}$

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Four new benzofulvene derivatives bearing bithiophene chromophores at two different key positions of the phenylindene scaffold were prepared in order to evaluate the role of different chromophores in the optoelectronic features of polybenzofulvene derivatives. The results of the photophysical studies showed that the optical properties of the newly-synthesized bithiophene-functionalized polymers were affected by both the polymer enchainment and the substitution topology of the monomeric units. On the other hand, the hole-mobility appeared to be affected to a lesser extent, but the best performances were obtained in poly-6-HBT-**BF3k** showing the strongest bithiophene side chain packing. This work demonstrates that the optoelectronic properties of polybenzofulvene derivatives can be optimized by a targeted chemical design such as side chain engineering.

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## Introduction

The development of high performance  $\pi$ -conjugated polymers has allowed the fabrication of optoelectronic devices [e.g. organic light-emitting diodes (OLEDs), field effect transistors and photovoltaic appealing (FETs), cells] showing properties.<sup>1-3</sup> In this field,  $\pi$ -stacked polymers, such as polydibenzofulvene derivatives<sup>4-6</sup> and cyclophanes,<sup>7,8</sup> may offer an alternative macromolecular platform with novel photophysical and photoelectronic characteristics, due to through-space noncovalent interactions among the pendant chromophore residues aligned along the polymer backbone. These peculiar spatial arrangements account for a new promising way to transport both charge and energy<sup>9</sup> and allow one to tune the electronic properties of the polymer system by modifying the geometry of  $\pi$ -packing<sup>10,11</sup> (e.g. the distance and the spatial orientation of the interacting chromophore planes). However, the potential optoelectronic applications of these materials

polysystems stacked along a single polymer chain.<sup>14</sup> The most studied  $\pi$ -stacked polymer is poly(*N*-vinylcarbazole) (**PVK**, Fig. 1), which has been shown to transfer charge *via* intrachain nonstacked orbitals resulting from face-to-face conformation of the carbazole moieties, and they show photoconductive, photorefractive, hole-transporting, energy-donating, and carrierswitching properties.<sup>15</sup> Similarly, polybenzofulvene derivatives were found to show g the significant hole-transporting capabilities linked to the presence of methoxy substituents in the structure of poly-6-MO-**BF3k** (Fig. 1) and combined with other peculiar features such as the prompt and almost quantitative formation by spontaneous thermoreversible polymerization, tunable solubility,

aggregation behavior, *etc.*<sup>16–21</sup> Furthermore, the replacement of the methoxy group at position 6 of the 3-phenylindene structure of poly-6-MO-**BF3k** with a 9,9-dimethylfluorene residue (as in poly-6-DMFL-**BF3k**) was the determinant in the obtainment of highly emissive polymeric supramolecular assemblies for optoelectronic applications.<sup>12,13</sup>

have been poorly investigated in the past few years. We have

recently demonstrated that  $\pi$ -stacked polybenzofulvene derivatives (*i.e.* poly-6-DMFL-**BF3k** in Fig. 1) can be considered as

intriguing materials for the production of optoelectronic devices such as OLEDs.<sup>12,13</sup> Polybenzofulvenes are through-

space conjugated polymers formed by arrays of  $\pi$ -electron

Based on the intriguing results obtained, we conceived a systematic investigation of different chromophores in the substitution of the previously employed fluorene moiety in order to explore the potentialities of polybenzofulvene derivatives in

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Fig. 1 Structures of PVK, poly-6-MO-BF3k, poly-6-DMFL-BF3k, and of the newly designed polybenzofulvene derivatives bearing bithiophene chromophores.

providing materials for optoelectronic applications. Thus, we designed four benzofulvene derivatives bearing bithiophene chromophores at two different key positions (6 and 4') of the phenylindene scaffold that were found to retain the spontaneous polymerization mechanism. By means of spectroscopic and electrical characterization, we show that the bithiophene chromophore substitution induces in the polybenzofulvene derivatives a quenching of the emission combined with an improvement of the hole-transporting features.

### **Experimental section**

#### Synthesis

The details of the preparation of benzofulvene derivatives and their spontaneous polymerization are described in the ESI.† NMR spectra were recorded with a Bruker DRX-400 AVANCE, Bruker DRX-500 AVANCE, or a Bruker DRX-600 AVANCE spectrometer in the indicated solvents (TMS as the internal standard): the values of the chemical shifts are expressed in ppm and the coupling constants (J) in Hz. An Agilent 1100 LC/MSD operating with an electrospray source was used in mass spectrometry experiments.

#### SEC-MALS

The molecular weight distribution (MWD) characterization of the newly prepared polybenzofulvene derivatives was performed by using a MALS light scattering photometer as a detector of a SEC chromatographic system. The SEC-MALS system and the corresponding experimental conditions were identical to those used in our previous studies<sup>17,18</sup> and are not reported in detail here. The dn/dc values of the polymers were measured off-line in THF at 35 °C using a Chromatix KMX-16 differential refractometer. The introduction of different functional groups in the repeating unit of the polybenzofulvene derivatives produced slight changes (increase) in the dn/dc value. In particular the measured values are the following: 0.195 mL g<sup>-1</sup> for poly-**BF3k**; 0.192 mL g<sup>-1</sup> for poly-6-MO-**BF3k**; 0.198 mL g<sup>-1</sup> for poly-6-BT-**BF3k**; 0.198 mL g<sup>-1</sup> for poly-4'-BT-6-MO-**BF3k**; 0.198 mL g<sup>-1</sup> for poly-6-HBT-**BF3k**; 0.225 mL g<sup>-1</sup> for poly-4'-HBT-6-MO-**BF3k**.

#### MALDI-TOF mass spectrometry

MALDI-TOF mass spectra were recorded in reflectron or linear delayed extraction mode, using a Voyager-DE STR instrument (PerSeptive Biosystem) mass spectrometer, equipped with a nitrogen laser ( $\lambda$  = 337 nm, pulse width = 3 ns), working in a positive ion mode. The accelerating voltage was 20 kV, and grid voltage and delay time (delayed extraction, time lag) were optimized for each sample to achieve a higher mass resolution. Laser irradiance was maintained slightly above the threshold. The best spectra were recorded using *trans*-2-[3-(4-*tert*-butylphenyl)-2-methyl-2-propenylidene]malononitrile

(DCTB) 0.1 M in THF as the matrix. Samples for the MALDI analyses were prepared as follows: 10  $\mu$ L of the polymer solution (5 mg mL<sup>-1</sup> in THF) were mixed with 10  $\mu$ L or 30  $\mu$ L of a matrix solution. Then 1  $\mu$ L of each analyte/matrix mixture was spotted on the MALDI sample holder and slowly dried to allow analyte/matrix co-crystallization. For the analysis of polymer SEC fractions 1  $\mu$ L of each selected concentrated SEC solution was mixed with 1  $\mu$ L or 3  $\mu$ L of the matrix solution (DCTB).

#### **SEC fractionation**

The polymeric material obtained by the dehydration of indenol 3a under the usual conditions was accurately fractionated by means of a SEC system consisting of an integrated Waters 515 apparatus (a pump and an injector) and a Waters R401 DRI detector, using THF as the mobile phase (1 mL  $min^{-1}$  of the flow rate). The column set was composed of four Ultrastyragel HR columns (ID = 7.8 mm, L = 300 mm, 5  $\mu$ m of the particle size) in the order HR-4, HR-3, HR-2 and HR-1 connected in series. The SEC traces were recorded and processed using the Clarity-GPC software provided by DataApex. In a typical fractionation, 100 µL of a polymer solution in THF (ca. 5 mg mL<sup>-1</sup> of the concentration) was injected and fractions were taken at every prefixed interval time. In the range of elution volume  $(V_e)$  within 16 and 28 mL (peak 1 in Fig. 3) the solutions were taken every 1 mL collecting 12 SEC-fractions, whereas in the  $V_e$  range within 28 and 38 mL (peak 2 in Fig. 3) were taken every 0.25 mL collecting 40 SEC-fractions.

#### **Optical measurements**

UV-vis absorption spectra were obtained with a Perkin Elmer Lambda 900 spectrometer. PL spectra were obtained with a SPEX 270 M monochromator equipped with a  $N_2$  cooled charge-coupled device excited with a monochromated 450 W Xe lamp. The spectra were corrected for the instrument response. PL QY of the solutions were obtained by using quinine sulfate as the reference. PL QY of solid powders were measured with a home-made integrating sphere according to the procedure reported elsewhere.<sup>22</sup>

Time resolved measurements were performed by using a mode locked titanium:sapphire laser system operating at 80 MHz and delivering 100 fs long pulses (FWHM) with a cw mean power of about 500 mW at 780 nm. Optical excitation around 390 nm was obtained using a lithium iodate (LiIO<sub>3</sub>) crystal as a frequency doubler. Photoluminescence was spectrally dispersed in a single spectrometer and temporally resolved with a visible 2D-streak camera (Hamamatsu C5680). The temporal resolution in the configuration we employed was of the order of 10 ps. Decay traces were extracted from streak camera spectrograms by integrating over a ~50 nm-wide spectral window.

#### Device preparation and characterization

Glasses covered with indium tin oxide (ITO, 15  $\Omega$   $\Box^{-1}$ ) were cleaned ultrasonically in distilled water, acetone, and isopropanol. On the substrates, a water solution of poly-(3,4-ethy-lenedioxythiophene)-poly-(styrenesulfonic acid) (PEDOT:PSS, Clevios P VP AI 4083, H. C. Starck) was spincoated through a nylon filter (pore size 0.45 µm), creating a layer with a thickness of 50 nm. Subsequently, the substrates were annealed for 10 min at 100 °C under a nitrogen atmosphere. On such prepared substrates, 10 nm of MoO<sub>3</sub> were evaporated at a pressure of  $10^{-7}$  mbar, creating an ohmic contact for hole injection. Then, inside a nitrogen filled glovebox, organic films were deposited from a chloroform solution with a concentration of

15 mg mL<sup>-1</sup>. The obtained layers showed thickness values between 200 nm and 300 nm. Finally, the top electrodes consisting of 7 nm of MoO<sub>3</sub> and 80 nm of Al were evaporated. The current density-voltage curves were measured with a Keithley 2602 source meter. The device characterization was performed under a nitrogen atmosphere. The thickness of the organic films was evaluated by using a Dektak XT (Bruker) profilometer.

#### **Results and discussion**

#### Synthesis and spontaneous polymerization of the newly designed benzofulvene derivatives bearing bithiophene chromophores

The synthesis of the newly designed polybenzofulvene derivatives was based on the chemistry developed for the preparation of the previously published fluorene members of this subseries of polybenzofulvenes bearing pendant chromophore residues.<sup>12</sup> In particular, triflate **1** (ref. 12) was used as the starting material in the multistep procedure described in Scheme 1 to obtain polybenzofulvene derivatives poly-6-BT-**BF3k** and poly-6-HBT-**BF3k**.

Suzuki–Miyaura cross-coupling of **1** with the suitable bithiopheneboronic acid pinacol esters (commercially available from Aldrich) gave indenones **2a,b**, which were transformed into the corresponding indenols **3a,b** by methylation with trimethyl-



Scheme 1 Synthesis and spontaneous polymerization of benzofulvene derivatives 6-BT-BF3k and 6-HBT-BF3k. Reagents: (i) 2,2'-bithiophene-5-boronic acid pinacol ester for 2a (or 5'-hexyl-2,2'-bithiophene-5-boronic acid pinacol ester for 2b), Pd(PPh\_3)<sub>2</sub>Cl<sub>2</sub>, PPh<sub>3</sub>, Cs<sub>2</sub>CO<sub>3</sub>, CH<sub>3</sub>OH, THF; (ii) Al(CH<sub>3</sub>)<sub>3</sub>, CH<sub>2</sub>Cl<sub>2</sub>: (iii) PTSA, CHCl<sub>3</sub> (Method B); (iv) solvent evaporation. Substituents: R = H in 2a, 3a, 6-BT-BF3k, and poly-6-BT-BF3k; R = *n*-hexyl in 2b, 3b, 6-HBT-BF3k, and poly-6-HBT-BF3k.

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aluminium.<sup>17–21</sup> The dehydration of indenol **3a** under the usual conditions (**Method A:** 0.05 M solutions of **3a** in CDCl<sub>3</sub> containing catalytic amounts of *p*-toluenesulfonic acid, PTSA, see the ESI†) led to the formation of a new polybenzofulvene derivative showing the unusual enchainment displayed in poly-5 (Scheme 2) as suggested by SEC-MALS, SEC-MALDI, and NMR spectroscopy (see below).

We assumed that the product of the first step of the dehydration reaction (*i.e.* carbonium ion 4) was capable of performing an electrophilic attack on the distal thiophene ring of the bithiophene moiety that could be depressed under diluted conditions. In fact, monomer 6-BT-**BF3k** was obtained in good purity by carrying out the dehydration reaction of **3a** under diluted conditions (**Method B**, concentration *ca.* 0.001 M), which depressed the intermolecular events (*i.e.* the oligomerization process driving to poly-5) in favour of the intramolecular hydrogen ion elimination leading to 6-BT-**BF3k**. Thus, the diluted conditions of **Method B** became the standard methodology for the dehydration of indenol derivatives bearing the bithiophene chromophores. As expected, benzofulvene monomers 6-BT-**BF3k** and 6-HBT-**BF3k** polymerized in the apparent absence of catalysts, upon solvent removal, to give the corresponding polymers poly-6-BT-**BF3k** and poly-6-HBT-**BF3k**. A similar procedure (Scheme 3) starting from bromophenylindenone **6** (ref. 12) was used for the synthesis of the polybenzofulvene derivatives bearing the bithiophene chromophores at position 4' of the phenylindene scaffold.

# Macromolecular characterization of polybenzofulvene derivatives bearing bithiophene chromophores

An absolute multi-angle laser light scattering (MALS) detector on-line to a size exclusion chromatography (SEC or GPC) system (THF as the mobile phase) was employed in the characterization of molecular weight distribution (MWD), size of macromolecules (radius of gyration  $R_g$ ), and conformation of the polybenzofulvene derivatives bearing bithiophene chromophores.

Very interestingly, the polybenzofulvene derivatives obtained by spontaneous polymerization of the newly-designed benzofulvene derivatives (Table 1) featured particularly high average molecular weight values and broad dispersity ( $D = M_w/M_n$ ) similar to the ones shown by the parent polybenzofulvenes. These results confirmed that the presence of bithiophene chromophores at two different key positions of



Scheme 2 Dehydration of indenol 3a by Method A.



poly-4'-HBT-6-MO-**BF3k** 

Scheme 3 Synthesis of polybenzofulvene derivatives bearing the bithiophene chromophores at positions 4' of the phenylindene scaffold. Reagents: (i) 2,2'-bithiophene-5-boronic acid pinacol ester for 7a (or 5'-hexyl-2,2'-bithiophene-5-boronic acid pinacol ester for 7b), Pd (PPh<sub>3</sub>)<sub>2</sub>Cl<sub>2</sub>, PPh<sub>3</sub>, Cs<sub>2</sub>CO<sub>3</sub>, CH<sub>3</sub>OH, THF; (ii) Al(CH<sub>3</sub>)<sub>3</sub>, CH<sub>2</sub>Cl<sub>2</sub>; (iii) PTSA, CHCl<sub>3</sub>; (iv) solvent evaporation. Substituents: R = H in 7a, 8a, 4'-BT-6-MO-BF3k, and poly-4'-BT-6-MO-BF3k.

the phenylindene scaffold is compatible with the spontaneous polymerization mechanism leading to ultrahigh molecular weight polybenzofulvene derivatives so that the mass values observed in the new polymers were significantly higher than those observed in the parent macromolecules poly-6-MO-**BF3k** and poly-**BF3k**. This observation could be rationalized in terms of additional stacking interactions due to the bithiophene moiety stabilizing the polybenzofulvene backbone. Finally, the slope of the conformation plot was in the range (0.5–0.6) usually found for linear polymers with a random coil conformation in a relatively good solvent (THF) as already observed in the parent polybenzofulvene derivatives poly-6-MO-**BF3k** and poly-**BF3k**.

On the other hand, SEC-MALS analysis of the samples obtained by dehydration of indenol 3a by Method A (performed in two different runs to evaluate the reproducibility of the results) showed the presence of two components differing in molecular weights. In particular, the chromatogram obtained with the first run sample (MA-1) showed the presence of a dominant (about 80%) oligomeric component and a minor (about 20%) polymeric component showing a higher  $M_{\rm W}$  value (Fig. 2).



Fig. 2 SEC-MALS signals (DRI = concentration detector, MALS  $90^{\circ}$  = MALS photodiode at  $90^{\circ}$ ) recorded for the first run sample (MA-1) obtained by the dehydration of indenol **3a** by Method A.

Table 1	Macromolecular	features	of the	polybenzofulvene	derivatives	obtained	by spontaneous	polymerization	of the	newly-designed	l benzo-
fulvene d	lerivatives bearing	bithioph	ene chr	romophores compa	ared with th	ose shown	by the parent pe	olymers poly-6-1	MO-BF	3k and poly-BF3	k

Polymer	$M_{\rm p}  ({\rm kg \ mol^{-1}})$	$M_{\rm w}^{\ a}$ (kg mol <sup>-1</sup> )	$M_{ m w}/M_{ m n}^{\ b}$	$R_{\rm g}^{\ c} ({\rm nm})$	$K^{d}$ (nm)	$\alpha^d$
Poly-6-BT-BF3k	2968	2012	3.5	43.7	$1.38 \times 10^{-2}$	0.53
Poly-6-HBT-BF3k	4735	4596	2.5	72.7	$2.16 \times 10^{-2}$	0.51
Poly-4'-BT-6-MO-BF3k	3884	2840	5.2	55.3	$2.32 \times 10^{-2}$	0.50
Poly-4'-HBT-6-MO-BF3k	3817	3438	2.9	66.2	$2.53 \times 10^{-2}$	0.51
Polv-6-MO-BF3k	312	347	4.3	19.4	$5.75 \times 10^{-3}$	0.61
Poly-BF3k	1900	1506	3.4	49.9	$6.63 \times 10^{-3}$	0.60

 ${}^{a}M_{w}$ : weight-average molecular weight.  ${}^{b}M_{w}/M_{n}$ : dispersity where  $M_{n}$  denotes the numeric-average molecular weight.  ${}^{c}R_{g}$ : radius of gyration *i.e.* dimension of the macromolecules.  ${}^{d}K_{r}\alpha$ : intercept and slope of the conformation plot.

Moreover, the SEC-MALDI analysis of the second run sample (MA-2) showed an analogous composition (Fig. 3).

The MALDI-TOF mass spectrum of sample MA-2 obtained from the dehydration of indenol 3a by Method A, and recorded in linear mode using trans-2-[3-(4-t-butyl-phenyl)-2-methyl-2propenylidenelmalononitrile (DCTB) as the matrix, is displayed in Fig. 1 (ESI<sup>†</sup>). The spectrum showed a series of repeating families of peaks spanning from m/z 400 up to m/z6000 having a 440.6  $\pm$  0.3 Da transition, corresponding to the mass of the repeating unit (440.6 g  $mol^{-1}$ ). The most intense peaks correspond to the oligomers terminated with hydrogen at both ends. Since the SEC analysis (Fig. 3) showed that the polymer sample was composed of about 60% of the oligomer component and about 40% of the polymeric component with a higher  $M_{\rm W}$  value, we assumed that the mass spectrum was representative only of the oligomeric component. It is well known that in MALDI-TOF MS analysis of polydisperse polymers, macromolecular species with molar masses lower than 10 000-15 000 g mol<sup>-1</sup> are detected, owing to several discrimination effects.<sup>23–28</sup> To obtain more reliable information on the chemical composition of the polydisperse polymers, the offline SEC/MALDI-TOF MS coupling method was applied.<sup>23,24,28,29</sup> Thus, the second run sample (MA-2) was fractionated into narrow molar mass fractions by analytical SEC analysis, by collecting 12 fractions between 16 and 28 mL (peak 1 in Fig. 3) and 40 fractions within 28 and 38 mL (peak 2 in Fig. 3). Fig. 3 shows the SEC curve and the MALDI-TOF mass spectra of some representative fractions. The assignments of the most intense mass peaks are also reported in Fig. 3. As expected, the mass spectra of the fractions corresponding to the peak 2 show that they are constituted of oligomers with a narrow molar mass dispersity: the most intense peaks belong to the protonated ions of the species formed according to Scheme 2. The average molar masses of these fractions increased with the decrease of the corresponding elution volume, in agreement with the SEC behaviour and the SEC-MALDI recorded for other polymers.<sup>23,24,28,29</sup> However, a surprising result was obtained by analyzing the SEC fractions corresponding to peak 1, at a lower elution time. Contrary to the expectation, they gave the same mass spectra independently of their elution volume (see the mass spectra of fractions F3, F5 and F7 in Fig. 3). In fact, they show peaks only in the mass range within m/z 400 and m/z 4000, corresponding to the species formed according to Scheme 1. This behavior could be explained by taking into account the following hypotheses: (i) aggregation phenomena could occur under the SEC conditions and the molar mass of the component eluted at each elution volume value was apparent; (ii) fragmentation



Fig. 3 SEC-MALDI chromatogram obtained with the second run sample (MA-2) obtained by the dehydration of indenol 3a by Method A. The SEC curve (dotted line) of the re-injected fraction F5 is also reported.

processes could occur during the MALDI-TOF MS analysis of high molar mass poly-6-BT-**BF3k**. The first hypothesis was excluded because both the elution volume and the mass distribution of some fractions re-injected in the SEC apparatus did not change with respect to that observed during the analysis of the neat sample, as well as the fraction F5 reported in Fig. 3 (dotted line curve). Therefore, the assumption that polybenzofulvenes afforded fragmentation under the MALDI-TOF conditions is reasonable. This behaviour was also confirmed by the MALDI mass spectrum of the poly-6-BT-**BF3k** (Fig. 2) (ESI†) that showed peaks due to the molecular ions of di-hydrogen ended linear oligomers from dimers (m/z 882) up to octamers (m/z 3526), together with the other fragmentation products.

This finding suggests that a fragmentation process could occur during the MALDI-TOF MS analysis of polybenzo-

fulvenes already published, because generally mass peaks were observed only in the mass range of m/z 400–9000. At present, a series of studies are carried out to investigate the mechanisms of fragmentation processes occurring during the MALDI-TOF MS analysis of polybenzofulvenes.

#### NMR characterization of the polymerization of the newly designed benzofulvene derivatives bearing bithiophene chromophores

The spontaneous polymerization of benzofulvene derivatives 6-BT-**BF3k**, 6-HBT-**BF3k**, 4'-BT-6-MO-**BF3k**, and 4'-HBT-6-MO-**BF3k** was studied by NMR spectroscopy by comparing the spectra of the polymers with those of the corresponding monomers after a careful assignment work. In particular, the <sup>1</sup>H and <sup>13</sup>C NMR spectra of the monomers were assigned by means of



Fig. 4 Comparison of the <sup>13</sup>C NMR spectra of poly-4'-BT-6-MO-BF3k and poly-4'-HBT-6-MO-BF3k with that of the parent macromolecule poly-6-MO-BF3k.

correlation experiments (see the ESI<sup>†</sup> for details). This work allowed the retention of the original vinyl(1,2) polymerization mechanism to be evaluated on the basis of the presence of the diagnostic peak at around 57 ppm attributed to the aliphatic quaternary carbon (C-1) of the 1,2-repeating indene unit (see for example Fig. 4 and 5).<sup>17,18,30</sup> Based on the comparison of the <sup>13</sup>C NMR spectra of poly-4'-HBT-6-MO-**BF3k** and poly-4'-BT-6-MO-**BF3k** with that of the parent macromolecule poly-6-MO-**BF3k**, (Fig. 4) the sharpest signals were assigned to the carbon atoms belonging to the bithiophene chromophores, while the phenylindene signals were as broad as in the parent macromolecule poly-6-MO-**BF3k**.

Moreover, the comparison of the <sup>13</sup>C NMR spectra of the four polybenzofulvene derivatives bearing the bithiophene chromophores confirmed that the presence of the bithiophene moiety at position 6 of the indene nucleus produced a signifi-

cant broadening of the signal in the spectra of both poly-6-BT-BF3k and poly-6-HBT-BF3k (Fig. 5).

Overall, these observations were rationalized in terms of different dynamics of the monomeric units in the polybenzofulvene backbone, as proposed for the previously reported fluorene-based polybenzofulvenes.<sup>12</sup> The introduction of the bithiophene residues at position 6 of the indene nucleus enhances the degree of  $\pi$ -packing among the aromatic moieties of both poly-6-BT-**BF3k** and poly-6-HBT-**BF3k** resulting in a reduction of the dynamic features of the monomeric units, although the hexyl side chains of poly-6-HBT-**BF3k** hold significant symptoms of local mobility. On the other hand, the bithiophene chromophores of poly-4'-BT-6-MO-**BF3k** and poly-4'-BT-6-MO-**BF3k** were assumed to freely rotate around their attachment bond at the pendant phenyl group in the apparent lack of stacking constraints.



Fig. 5 Comparison of the <sup>13</sup>C NMR spectra of poly-6-BT-BF3k and poly-6-HBT-BF3k with that of the parent macromolecule poly-BF3k.

Both the samples **MA-1** and **MA-2** obtained from the dehydration of indenol **3a** by **Method A** were studied by NMR spectroscopy in order to establish the structure and the formation mechanism of this unusual polybenzofulvene derivative. Thus, <sup>1</sup>H and <sup>13</sup>C NMR spectra of **3a** were assigned by means of homonuclear <sup>1</sup>H–<sup>1</sup>H-COSY, <sup>1</sup>H–<sup>1</sup>H-NOESY and heteronuclear <sup>1</sup>H–<sup>13</sup>C-HSQC, <sup>1</sup>H–<sup>13</sup>C-HMBC 2D correlation experiments and the assigned spectra were used in the interpretation of the corresponding ones obtained with the unusual polybenzofulvene derivative.

Interestingly, <sup>13</sup>C NMR spectra of both the samples were characterized by the presence of sharp and well-resolved resonances, but the most significant structural information could be obtained from <sup>1</sup>H-<sup>13</sup>C correlation experiments. The analysis of the HMBC (Heteronuclear Multiple Bond Correlation) spectrum of sample **MA-2** (Fig. 6) showed the presence of three species in different proportions: a Major one (*i.e.* the Major monomeric unit, **Mmu**, Fig. 6) with two minor ones (*i.e.* the minor monomeric unit, **mmu**, and the Terminal monomeric unit, **Tmu**, Fig. 6).

The comparison of the <sup>13</sup>C NMR spectrum (Fig. 5 ESI<sup>†</sup>) of sample **MA-2** with that of its precursor **3a** suggested that the structure of this intermediate was conserved in the Major monomeric unit (**Mmu**) with the most significant change being the loss of the indenol hydroxyl group as suggested by a dramatic change in the chemical shift value of the C-1 carbon atom.

In the HMBC of sample MA-2 (Fig. 6), the signal at  $\delta_{\rm H} = 2.10$  ppm (directly related to the signal at 24.2 ppm in the <sup>13</sup>C dimension and assigned to methyl D of Mmu species) showed four cross peaks at  $\delta_{\rm C} = 55.5$ , 139.4, 145.9, and 154.8 ppm, respectively. This suggested that  $CH_3$  D of Mmu was correlated through two bonds with C-1 ( $\delta_{\rm C} = 55.5$  ppm) and through three bonds with C-7a ( $\delta_{\rm C} = 154.8$  ppm) and C-2 ( $\delta_{\rm C} = 139.4$  ppm), and with a resonance at 145.9 ppm that is a quaternary carbon atom (roughly corresponding to thiophene C–E of 3a, as the chemical shift value), which correlates in the HMBC spectrum with the thiophene methine group K. Therefore, the resonance at 145.9 was assigned to the carbon atom C–L of the thiophene ring, meaning that the indene group at the C-1 position is directly bonded to a thiophene ring as in Mmu in Fig. 6 and in



Fig. 6 HMBC spectrum (CDCl<sub>3</sub>) of sample MA-2 obtained by the dehydration of the indenol derivative **3a** by Method A. In the inset: the possible structures of the monomeric units assumed to compose poly-5 obtained from the dehydration of indenol **3a** by Method A as determined by NMR spectroscopy. HMBC experiments were carried out with a 2-fold low-pass J-filter, a delay of 100 ms for the evolution of long-range coupling, and delays corresponding to  ${}^{1}J_{CH}$  = 130 and 160 Hz. 90° pulse – 8.05 ms, –2.0 dB.

poly-5. The structure elucidation of the minor species was complicated by the partial overlap and the low intensities of their signals, but the careful analysis of NMR data suggested the structures reported in Fig. 6 for the Terminal monomeric unit (**Tmu**) and the minor monomeric unit (**mmu**).

#### Photophysical properties of polybenzofulvene derivatives bearing bithiophene chromophores

The photophysical features of the new polybenzofulvene derivatives (Fig. 7), summarized in Table 2 for both the solid state and the solution, gave information useful in evaluating their possible application as active materials in electronic devices.

The strong attitude of thiophene oligomers towards emission aggregation quenching processes (see ref. 31) induced a sharp reduction of the PL efficiency in the solid state in the case of sample **MA-1** and for polymers with a bithiophene moiety at position 6 (*i.e.* poly-6-BT**-BF3k** and poly-6-BT**-BF3k**). In particular, for sample **MA-1** the nearly vanishing emission in the solid state is accompanied by a red-shift of the emission typical of low emissive excimer states. The PL QY values observed in solution were comparable to those of the parent polymer poly-6-MO-**BF3k**.<sup>19</sup> Interestingly, the emission of the corresponding polymers bearing the chromophore at position 4' (poly-4'-BT-6-MO-**BF3k** and poly-4'-HBT-6-MO**BF3k**) was higher and quite independent of the aggregation state, as previously observed for other polybenzofulvene derivatives and in agreement with the assumption that emission originates from the recombination from similar excimer/aggregate states both in solution and in the solid state.

This result is suggestive of different aggregation behaviors in the solid state for polybenzofulvene derivatives showing different polymer enchainments and substitution topologies of the monomeric units. Thus, the most efficient side chain packing in the solid state was obtained in poly-6-BT-**BF3k** and poly-6-HBT-**BF3k** bearing the bithiophene chromophore at position 6, whereas in the corresponding polymers bearing the chromophore at position 4' (poly-4'-BT-6-MO-**BF3k** and poly-4'-HBT-6-MO-**BF3k**), the packing was less strong. The PL time decay of the polymers is biexponential, with a dominant fast component of about 150 ps (100 ps) and a slow one of about 1.5 ns (1 ns), in solution (in the solid state) suggestive of a complex excimer recombination pathway (see Table 3 and Fig. 8).

The hole mobilities were measured for the polymers with alkyl substituted bithiophene derivatives (see Table 2) and



Fig. 7 Optical absorption and emission spectra of poly-6-HBT-BF3k (left panel), and poly-4'-HBT-6-MO-BF3k (right panel) in dichloromethane solutions (blue lines) and in the solid state (black lines).

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		Solution			Solid		
Polymer	Mobility (cm <sup>2</sup> V <sup>-1</sup> s <sup>-1</sup> )	$\lambda_{ab} (nm)$	$\lambda_{\rm em}  ({\rm nm})$	QY (%)	$\lambda_{ab}^{a}(nm)$	$\lambda_{\rm em}^{\ b} ({\rm nm})$	$\mathrm{QY}^{b}\left(\% ight)$
Sample MA-1		380	575	5	387	$\frac{640}{580^a}$	<0.1
Poly-6-BT-BF3k		380	490	7	386	555	< 0.1
Poly-6-HBT-BF3k	$5.81 \times 10^{-5}$	386	493	3	394	558	< 0.1
Poly-4'-BT-6-MO-BF3k		346	460	12	348	490	9
Poly-4'-HBT-6-MO- <b>BF3k</b>	$3.95 \times 10^{-5}$	354	460	14	360	490	13

<sup>*a*</sup> Spin coated films. <sup>*b*</sup> Powders.

	Solution <sup><i>a</i></sup>				Solid <sup>b</sup>			
Polymer	$\overline{\tau_1 (ps)}$	$ au_2 (\mathrm{ps})$	$A_1$	$A_2$	$\tau_1 (\mathrm{ps})$	$ au_2 (\mathrm{ps})$	$A_1$	$A_2$
Poly-6-HBT-BF3k	140	1550	0.72	0.28	90	1000	0.87	0.13
Poly-4'-HBT-6-MO- <b>BF3k</b>	160	1550	0.62	0.38	110	950	0.83	0.17
Poly-4'-HBT-6-MO-BF3k	140 160	1550	0.62	0.38	110	950	0.83	

Table 3 Lifetimes of two polybenzofulvene derivatives bearing bithiophene chromophores in dichloromethane solutions and cast films

<sup>1</sup> Dichloromethane solutions. <sup>*b*</sup> Cast films.



Fig. 8 PL decays of poly-6-HBT-BF3k and poly-4'-HBT-6-MO-BF3k films (black and green lines) and dichloromethane solutions (blue and red lines).

compared with that obtained for the standard hole-transporting polymer PVK measured under the same conditions. PVK is widely used in optoelectronic applications since its hole-transporting properties make this polymer an optimal matrix for emissive molecules to be used in OLEDs. Since different methods are used for the measurement of the mobility of conducting polymers and the results are often quite different according to the method used, we measured the mobility of two polybenzofulvenes and PVK by using a simple hole only device structure ITO/PEDOT/MoO<sub>3</sub>/polymer/MoO<sub>3</sub>/Al that gives good ohmic contacts. Both the polybenzofulvene derivatives showed a higher mobility with respect to **PVK**  $(4.84 \times 10^{-6} \text{ cm}^2)$ V<sup>-1</sup> s<sup>-1</sup>) and the parent polymers.<sup>19</sup> The slightly higher value measured for poly-6-HBT-BF3k was in agreement with its higher chromophore  $\pi$ -stacking and with the higher PL emission quenching observed.

#### Conclusion

In order to evaluate the effects of different chromophores on the optoelectronic features of polybenzofulvene derivatives, four new monomers were designed to carry bithiophene chromophores at two different key positions of the phenylindene scaffold. These benzofulvene derivatives were demon-

strated to be retaining the spontaneous polymerization propensity to produce polybenzofulvene derivatives showing hole-transporting performances higher than the reference PVK. Moreover, the optoelectronic features of the polymers were affected by both the polymer enchainment and the substitution topology of the monomeric units, with the hole-mobility being affected to a lesser extent, but with the best performance being obtained in poly-6-HBT-BF3k showing the strongest bithiophene side chain packing. The results of the present work suggest that the optoelectronic properties of polybenzofulvene derivatives can be optimized by an appropriate chemical design (i.e. side chain engineering). Furthermore, owing to the interesting results obtained in these exploratory approaches, the investigation of the chemical space around polybenzofulvene derivatives could afford new optoelectronic materials endowed with unprecedented features.

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