Correlation of morphology and charge transport in poly(3,4-ethylenedioxythiophene)–polystyrenesulfonic acid (PEDOT–PSS) films

C S Suchand Sangeeth, Manu Jaiswal and Reghu Menon

Department of Physics, Indian Institute of Science, Bangalore 560012, India
E-mail: manujaishwal81@gmail.com and manu@physics.iisc.ernet.in

Received 10 October 2008, in final form 11 December 2008
Published 5 January 2009
Online at stacks.iop.org/JPhysCM/21/072101

Abstract
A wide variation in the charge transport properties of poly(3,4-ethylenedioxythiophene)–poly(styrenesulfonate) (PEDOT–PSS) films is attributed to the degree of phase segregation of the excess insulating polyanion. The results indicate that the charge transport in PEDOT–PSS can vary from hopping to the critical regime of the metal–insulator transition, depending on the subtle details of morphology. The extent of electrical connectivity in the films, directly obtained from a temperature-dependent high-frequency transport study, indicates various limiting factors to the transport, which are correlated with the phase separation process. The low temperature magnetotransport further supports this morphology-dependent transport scenario.

(Some figures in this article are in colour only in the electronic version)

1. Introduction
Current progress in organic/polymer optoelectronics and device technology relies significantly on the ability to tune the various device transport parameters by means of morphology, interface modification, controlling the extent of disorder and varying the charge carrier density and mobility. The performance of these devices is strongly interlinked with the morphology and structure of the constituting organic material [1]. Significant enhancement in performance has been achieved by tuning the morphology, notably in the case of polythiophene transistors [2] or polymer/methanofullerene blended organic solar cells [3]. Among the important conducting polymers for device applications is poly(3,4-ethylenedioxythiophene)–poly(styrenesulfonate) (PEDOT–PSS), consisting of a conducting polythiophene derivative that is electrostatically bound to a PSS polyanion [4, 5]. Since this polymeric system simultaneously has high electrical conductivity, processability, high stability, flexibility and good transparency, it has found a large variety of applications including polymeric anodes for organic photovoltaics [6, 7], light-emitting diodes [8–11], flexible electrodes [12], supercapacitors [13], electrochromic devices [14], field-effect transistors [4] and antistatic coatings [15]. Usually, the appropriate combination of surface morphology and conductivity is required for various applications of PEDOT–PSS. The processing of PEDOT–PSS depends on a variety of factors including the fraction of PSS, the fraction of solid content, the particle size and the viscosity—which can lead to a wide range of morphological and electronic characteristics in the dried films. Antistatic coating applications require high conductivity for the PEDOT–PSS layer. In organic light-emitting diode (OLED) applications, the PEDOT–PSS layer serves as an efficient hole-injecting layer into the active semiconductor. This requires lower conduction for PEDOT–PSS, but the interface between the active layer and the polymer must be smooth in order to prevent microshorts leading to dark spots [4]. Furthermore, the surface roughness of highly conducting PEDOT–PSS (PH500 from HC
was 5% for PH510 solution and 25% for P solution. The glass substrates used were ~2 cm square and these were pre-cleaned by ultrasonication in acetone, triple distilled water and isopropyl alcohol for 30 min each. After stirring, the films were solution cast on glass substrate held at ~70 °C and annealed for 24 h, and then peeled off after drying to obtain free-standing films. Tapping-mode atomic force microscopy (TM-AFM) measurements were performed on freshly prepared films in a Veeco Instruments/CPR2.

The transport measurements were performed using a standard four-probe dc technique, and conductive carbon paint was used to make contacts (~1 mm apart) on the sample. For the conductance measurement, the current was sourced through a Keithley 220 current source and the voltage measured through a Keithley 2000 multimeter. The current values were typically in the range of 0.2–1 μA, chosen to avoid any sample heating at low temperatures. The sample is immersed in helium vapor/liquid in a Janis variable-temperature cryogenic system equipped with a superconducting magnet. The measurements are performed in magnetic fields up to 11 T and temperatures down to 1.3 K. The ac impedance is measured in the standard four-terminal configuration from 100 kHz to 10 MHz using an Agilent-4285A LCR meter in a Janis continuous-flow cryogenic system down to 4.2 K. These measurements were further extended to a lower frequency (40 Hz) using a lock-in amplifier (Stanford Research 830) relying on a resistor-divider setup to obtain the sample conductance at low frequencies.

3. Results and discussion

The tapping-mode atomic force microscope (TM-AFM) images of the films are shown in figure 1 in the topography mode. The morphology of the PEDOT–PSS films reveals granular structures which can be assigned to the polymer nanoparticles of the dispersion solution, consistent with previous observations [7, 30]. The extent of aggregation of the PEDOT-rich islands varies for the two films—with PH510 showing significantly larger size aggregates in comparison to P films. The average grain sizes in P and PH510 films are 40 and 110 nm, respectively (table 1). This is further verified by considering the surface roughness of the films. The PH510 film has almost twice the surface roughness on a sub-micron scale which originates from the deeper intervening gaps between the large-sized grains. While these films may have similar disorder on the molecular scale, the increased size of the grains may determine an improved transport process in PH510. The observed extent of granularity is an indicator of the extent of the phase-segregation process for the excess PSS, and this is more evident from the detailed transport study. Note that both these films have nearly the same ratio for PEDOT–PSS.

The temperature dependence of normalized conductivity of freestanding films (thickness ~15–40 μm) of P and PH510, processed with DMSO, is plotted in figure 2(a). In both cases the room temperature conductivity values, typically between 50 and 100 S cm−1, are sensitive to the processing conditions and thickness. Nevertheless, the temperature dependence of conductivity consistently shows the significant difference between P (stronger) and PH510 (weaker). In the case of...
P films, the conductivity decreases by nearly three orders of magnitude, whereas in PH510 it decreases only by a factor of five, as shown in figure 2(a). Recent investigations have suggested a hopping form, of stretched exponential type $R = R_0 \exp(T_0/T)^\beta$, for the low temperature transport in PEDOT–PSS films. However, these investigations were limited to $T > 60$ K, which only marks the onset of sharp change in conductance; suggesting that the transport at $T < 60$ K is more relevant for determining the role of disorder. To investigate this in detail, the reduced activation energy $[W = \Delta \ln \sigma / \Delta \ln T]$ is plotted as a function of temperature, as shown in figure 2(b) [31]. The $W$-plot slope suggests a distinctly different transport regime for PH510. For the P film, the stretched exponential form is justified (rather than the simple exponential form of fluctuation-induced tunneling models) and the value of the exponent ‘$p$’ is directly obtained from the negative slope of the graph at low temperature (typically $T < 40$ K), $p \sim 0.46$. This suggests that quasi-one-dimensional (1D) variable range hopping (VRH) is the dominant transport mechanism in P films. The scenario for quasi-1D VRH is mainly due to the important role played by the highly disordered quasi-1D regions in the system, especially at low temperatures; also indicative of the weak interchain transport [32]. Interestingly, the evidence for this type of quasi-1D transport has been observed in recent high-electric-field experiments in PEDOT–PSS [29]. The PH510, however, has a temperature-independent $W(T)$ at low temperatures indicating that this system is in the critical regime of the metal–insulator ($M$–$I$) transition, as has been observed in several conducting polymers [33]. This is further ascertained by considering the resistivity ratio $R_{4.2 \, K}/R_{300 \, K}$, which is lower by two orders in magnitude in PH510 film. At low temperatures ($T < 30$ K), the resistivity of PH510 can be described by a power-law form, $\rho(T) \sim T^{-\beta}$, with $\beta \sim 0.31$, distinct from the stretched exponential form for VRH. For the critical regime, the resistivity of disordered systems has been shown to follow a power-law dependence rather than the usual activated behavior, and it is expressed in terms of $\rho(T) = (e^2 p_F/\hbar^2)(k_B T/E_F)^{-\beta}$, where $p_F$ is the Fermi momentum, $e$ is the electronic charge and $0.3 < \beta < 1$ [34]. The value of the exponent $\beta$ being close to its lower limit (0.3) is indicative that electron–electron interaction effects may be important for the PH510 system. Furthermore, PH510 being in the critical regime of $M$–$I$ transition suggests that the extent of delocalization is much larger, resulting in enhanced interchain transport.

It is well known that in low-dimensional disordered systems, the temperature dependence of conductivity by itself is not adequate to fully understand the mechanism of charge transport; and the complementary magnetoresistance (MR) data is often required to determine the nanoscopic scale transport parameters. The MR of both systems at low temperatures is compared in figure 3. There is a significant positive MR for P–films (figure 3(b)), but this is considerably

---

**Table 1.** Comparison of transport parameters for P and PH510 PEDOT–PSS films.

| PEDOT–PSS   | Average grain size (nm) | Low temperature transport $(T < 40 \, K)$ | $R_{4.2 \, K}/R_{300 \, K}$ | $\Delta R(B)_{sat}/R$ (%) at $4.2 \, K$ | Onset frequency (MHz) |
|-------------|-------------------------|----------------------------------------|----------------------------|----------------------------------------|-----------------------|
| Baytron P   | 40                      | 1D VRH                                | 285                       | 56                                      | 3                     |
| Baytron PH510 | 110                    | critical regime                       | 2.8                       | 3                                      | 6.3                   |

$\Delta R(B)_{sat}$ denotes the saturation value of MR, at 11 T.
less in the case of PH510 film (figure 3(a), and table 1). The positive MR has a quadratic dependence at low fields and a tendency towards saturation at high fields, suggesting an origin in the wavefunction shrinkage mechanism under magnetic field. Further, this effect is more pronounced at lower temperatures. The localization length $\xi$ can be estimated from the low-field data for P film, using the following expression valid for 1D Mott type VRH, $\ln\left[\rho(B)/\rho(0)\right] = K(T_0/T)^{y(eB\xi^2/\hbar)^2}$, where $K = 0.0015$ is a numerical coefficient and $y = 3/2$ [35]. A quadratic fit to the low-field MR data at 1.3 K gives a value for the localization length of $\xi \sim 9–10$ nm, wherein the value of $T_0$ ($\sim 275$ K) from the temperature dependence of resistance has been used to estimate the above value. This value of localization length favors transport across conducting regions limited by insulating PSS lamellae in a matrix comprising both the regions, as also indicated by recent works [29, 36, 37]. Due to the carriers being sufficiently delocalized even at low temperatures, any shrinkage of the overlap of the electronic wavefunctions under magnetic field for the PH510 film does not have a discernible impact on the resistance; whereas in the case of P even minor shrinkage of the overlap of wavefunctions can dramatically decrease the conductivity of the system. The absence of a microscopic theory for magnetotransport in the critical regime, however, prevents a precise calculation of the length scale of the delocalized state. This sharp contrast between the two systems has interesting implications for understanding the transport, especially in large-grained film (PH510), and this is described in detail in the section below.

In figure 4, the normalized conductance of two PEDOT–PSS films is plotted as a function of frequency at various temperatures. The frequency dependence of conductance as a function of temperature simultaneously considers the intra-grain and inter-grain processes in the system. At low frequencies, the conductance is nearly constant and it increases as $\sim \omega^s (s < 1)$ above a certain threshold frequency, denoted as the onset frequency, $\omega_0$. This frequency response is characteristic of many other disordered systems. Whereas the $\omega^s$ form holds approximately for the PEDOT–PSS system, the onset frequency $\omega_0$ is quite a useful parameter for probing the extent of nanoscale ‘connectivity’ in the films. A correlation length $\lambda$ can be defined for any disordered system and this length scale corresponds to the distance between connections (e.g. junctions, nodes, etc) in the system. The charge carrier travels a distance $L$ at a frequency $\omega L$ and this distance traveled is lower at higher frequencies. At the onset frequency $\omega_0$, the carrier travels the distance $\sim \lambda$. Below this frequency the carriers have to travel across the conducting clusters, which are separated by the insulating PSS lamellae. The onset frequency therefore separates the two regimes of transport [38]. A higher value of onset frequency implies a smaller correlation length and shorter connections in the system [38]. For our system, we define the onset frequency as the value where $\sigma(\omega_0) = 1.1\sigma(0)$, where $\sigma(0)$ is the dc conductivity. The PH510 films have superior transport characteristics in comparison to P films at room temperature, as seen from the significantly higher value of the onset frequency (see table 1). The onset frequency of PH510 films changes by only a small factor, in sharp contrast
to the P films (see figure 4(c)). Moreover, the increase in conductance as a function of frequency, at all temperatures, is rather weak (well below a factor of 2) in the case of PH510; whereas it increases considerably for P samples. The onset frequency for PH510 is nearly temperature independent, whereas in P type films it is strongly temperature dependent, as in figure 4(c). The presence of ‘shorter network connections’ together with a very weak temperature dependence down to \(\sim 5\) K, suggest that the limitation on transport in PH510 arises from the connectivity within the PEDOT-rich grain rather than transport via the PSS barriers. This implies that the PH510 system can be electrically modeled as comprising closely packed junctions and nodes, determined by the intrinsic disorder of the conducting regions. Such a model also explains the rather weak temperature variation of frequency-dependent transport. On the contrary, if the transport was determined by the insulating barrier a significant frequency-dependent transport could have been observed. In P films, however, the strong frequency dependence of ac conductance indicates a significant decrease in the onset frequency \(\omega_0\) as the temperature is lowered (see figure 4(b)), indicating the weakening of the temperature-activated transport across the insulating PSS lamellae intervening between the conducting PEDOT-rich grains [29]. This difference in behavior, considered together with the magnetotransport data, points to the following surprising result for transport in PEDOT–PSS systems. In both PEDOT–PSS systems the carriers are confined at least to within the span of a single PEDOT-rich domain. In the case of small-grain size systems (P), the conduction is limited by the insulating PSS barriers, with the ‘connections’ weakening with decrease in temperature. At lower temperatures, the phonon-mediated process between the conducting regions becomes hampered. However, in the case of PH510, no such weakening of network connectivity is observed since the intervening barriers are sufficiently thin and no longer inhibit carrier motion, even at very low temperatures. Therefore, it is the intrinsic disorder in the conducting domains that ultimately limits the conduction in the PH510 system. A comparison of the morphology and transport parameters for these two types of conducting PEDOT systems is shown in table 1.

4. Conclusion

In summary, experimental investigation of the charge transport in thin films of poly(3,4-ethylenedioxythiophene)–polystyrenesulfonic acid (PEDOT–PSS) reveals significant correlation with the morphology of the system. This work compares the magnetotransport and temperature-dependent ac-transport parameters across different conducting grades of PEDOT–PSS processed with dimethyl sulfoxide (DMSO). Depending on the subtle details of morphology, the transport at low temperatures is shown to vary from the hopping regime (P) to the critical regime of the metal–insulator transition (PH510). It is shown that the conduction mechanism in PEDOT–PSS thin films depends strongly on the extent of aggregation of the constituent polymer nanoparticles. From the magnetotransport analysis, it is shown that the characteristic length scale for transport correlates with the PSS-barrier-limited picture. The onset frequency for the increase in ac conduction as well as its temperature dependence is a tool that directly probes the extent of connectivity in the network; and these data are a confirmation that distinctly different conduction mechanisms are present, depending on the morphology. The evidence from ac transport indicates that the limitation on conductivity arises from the insulating PSS barriers in one system (P), and from the intrinsic disorder present within the conducting regions in the other system (PH510). Whereas both the conduction processes are consistent within the phase-segregation model, the extent of such segregation determines different conduction processes. The results suggest that the present development of the enhancement in room temperature conductivity in PEDOT–PSS, relying on segregating the excess insulating PSS, is already close to an optimum scenario, and future enhancements in PEDOT conductivity must instead address the reduction of intrinsic disorder within PEDOT-rich highly conducting domains.
Acknowledgments

The magnetic field measurements were performed at the DST National facility for Low Temperature and High Magnetic Field, Bangalore (we thank Dr V Prasad). SSCS and MJ thank CSIR, New Delhi for financial assistance.

References

[1] Jaiswal M and Menon R 2006 Polym. Int. 55 1371
[2] Sirringhaus H, Brown P J, Friend R H, Nielsen M M, Bechgaard K, Langeveld-Voss B M W, Spiering A J H, Janssen R A J, Meijer E W, Herwig P and de Leeuw D M 1999 Nature 401 685–8
[3] Shaheen S E, Brabec C J, Sariciftci N S, Padinger F, Fromherz T and Hummelen J C 2001 Appl. Phys. Lett. 78 841
[4] Kirchmeyer S and Reuter K 2005 J. Mater. Chem. 15 2077
[5] Groenendaal L, Jonas F, Freitag D, Pielartzik H and Reynolds J R 2000 Adv. Mater. 12 481
[6] Zhang F, Johansson M, Andersson M R, Hummelen J C and Inganas O 2002 Adv. Mater. 14 662
[7] Snaith H J, Kenrick H, Chiesa M and Friend R H 2005 Polymer 46 2573
[8] Crispin X, Jakobsson F E, Crispin A, Grin P C M, Andersson P, Volodin A, van Haesendonck C, Van der Auweraer M, Salaneck W R and Berggren M 2006 Chem. Mater. 18 4354
[9] Cao Y, Yu G, Zhang C, Menon R and Heeger A J 1997 Synth. Met. 87 171–4
[10] Brown T M, Kim J S, Friend R H, Cacialli F, Daik R and Feast W K 1999 Appl. Phys. Lett. 75 1679
[11] Ho P K H, Kim J-S, Burroughes J H, Becker H, Li S F Y, Brown T M, Cacialli F and Friend R H 2000 Nature 404 481–4
[12] Aernouts T, Vanlaeke P, Geens W, Poortmans J, Heremans P, Borghs S, Martens R, Andriessen R and Leenders L 2004 Thin Solid Films 451–452 22
[13] Ghosh S and Inganas O 1999 Adv. Mater. 11 1214
[14] Argun A A, Cirpan A and Reynolds J R 2003 Adv. Mater. 15 1338
[15] Jonas F and Morrison J T 1997 Synth. Met. 85 1397
[16] Zhou Y, Zhang F, Tvingstedt K, Barrau S, Li F, Tian W and Inganas O 2008 Appl. Phys. Lett. 92 233308–3
[17] http://www.hitech-projects.com/cuprojects/olla/index.html
[18] Samiutsu S, Shimomura T, Ito K, Fujimori M, Heike S and Hashizume T 2005 Appl. Phys. Lett. 86 233103
[19] Bagchi D and Menon R 2006 Chem. Phys. Lett. 425 114
[20] Kim J Y, Jung J H, Lee D E and Joo J 2002 Synth. Met. 126 311
[21] Ouyang J, Xu Q, Wei C-W, Yang Y, Li G and Shinar J 2004 Polymer 45 5443
[22] Nardes A M, Kemerink M and Janssen R A J 2007 Phys. Rev. B 76 085208
[23] Ashizawa S, Horikawa R and Okuzaki H 2005 Synth. Met. 143 281
[24] Jonsson S K M, Birgerson J, Crispin X, Greczynski G, Osikowicz W, Denier van der Gon A W, Salanek W R and Fahlman M 2003 Synth. Met. 139 1
[25] Louwet F, Groenendaal L, Dhaen J, Manca J, Luppen J V, Verdonck E and Leenders L 2003 Synth. Met. 135 115
[26] Zuppiroli L, Bussac M N, Paschen S, Chauvet O and Forro L 1994 Phys. Rev. B 50 5196
[27] Crispin X, Marciniak S, Osikowicz W, Zotti G, Denier van der Gon A W, Louwet F, Fahlman M, Groenendaal L, De Schryver F and Salanek W R 2003 J. Polym. Sci. B 41 2561
[28] Higgins A M, Martin S J, Jukes P C, Geoghegan M, Jones R A L, Langridge S, Cubitt K, Kirchmeyer S, Wehrum A and Gruzzi J 2003 J. Mater. Chem. 13 2814
[29] Nardes A M, Kemerink M, Janssen R A J, Bastiaansen J A M, Kiggen N M M, Langeveld B M W, van Breemen J J M and de Kok M M 2007 Adv. Mater. 19 1196
[30] Ionescu-Zanetti C, Mechter A, Carter S A and Lal R 2004 Adv. Mater. 16 385–9
[31] Jaiswal M, Wang W, Fernando K A S, Sun Y-P and Menon R 2007 J. Phys.: Condens. Matter 19 446006
[32] Aleshin A N, Williams S R and Heeger A J 1998 Synth. Met. 94 173
[33] Menon R, Väkiparta K, Yoon C O, Moses D and Heeger A J 1994 Synth. Met. 65 167
[34] Larkin A I and Khmel’nietskii D E 1982 Sov. Phys.—JETP 56 647
[35] Long Y, Chen Z, Shen J, Zhang Z, Zhang L, Huang K, Wan M, Jin A, Gu C and Duvail J L 2006 Nanotechnology 17 5903–11
[36] Greczynski G, Kugler T, Keil M, Osikwicz W, Fahlman M and Salanek W R 2001 J. Electron Spectrosc. Relat. Phenom. 121 1
[37] Hwang J, Amy F and Kahn A 2006 Org. Electron. 7 387
[38] Kilbride B E, Coleman J N, Frayssy J, Fournet P, Cadek M, Drury A, Hutzler S, Roth S and Blau W 2002 J. Appl. Phys. 92 4024