The influence of the triplet exciton and charge transfer state energy alignment on organic magnetoresistance

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ABSTRACT

Recently, it was discovered that the current through an organic semiconductor, sandwiched between two non-magnetic electrodes, can be changed significantly (up to 25%) by applying a small (a few millitesla) magnetic field. At present, the microscopic mechanisms underlying this so-called organic magnetoresistance (OMAR) are intensively being debated. One of the mechanisms which can successfully describe the magnetic field effects on the current in pristine organic semiconductor devices uses the reactions of triplet excitons and polarons. Here, we present a proof of concept study in which we tune these interactions in the device by deliberately doping our devices with fullerene, creating additional charge transfer states (CTS). By engineering devices with different energetic alignments of the CTS and triplet exciton, we can influence the triplet exciton density in the device. We correlate pronounced changes in the magnetic field effect magnitude and lineshape to the energy of the CTS with respect to the triplet exciton.

1. Introduction

The discovery of surprisingly large, room temperature, magnetoresistance effects in organic semiconductor devices without any ferromagnetic components has puzzled the young field of organic spintronics during the past decade [1–3]. The effect is often referred to as organic magnetoresistance (OMAR). The possibility to obtain large (up to 25%) changes in the current at room temperature, while applying only small (a few millitesla) magnetic fields, makes this effect interesting for future applications and, moreover, from a scientific point of view. Studying OMAR provides a deeper fundamental understanding of spin-physics and charge transport in organic semiconductor devices.

All contemporary models explaining OMAR rely on magnetic field dependent reactions of the spin carrying particles, where a magnetic field suppresses the spin mixing of the particle pairs prior to the reaction. There is, however, an active debate about the nature of the particles (e.g. electrons, holes, bipolarons, excitons or trions) and the exact underlying mechanism influencing the current [4–13]. Currently, the possible mechanisms can be divided into two main categories: (i) mechanisms which only need one type of charge carrier or (ii) mechanisms which need both charge carriers. In the first class, the so-called bipolaron mechanism [6] has been successfully applied to explain magnetic field effects in unipolar devices [8,14,15]. In the second class, we can distinguish between models which describe the reactions of polarons with opposite charge into excitons [4] and models which rely on the spin dependent reactions of triplet excitons with polarons [5,12].

Besides the microscopic models, it is also important to be aware of the highly non-trivial role played by the device physics in order to explain experimentally observed magnetic field effects (MFEs) on the current [16,17]. Very recently, we have shown that the dominant mechanism for OMAR depends on the exact material choice and operating conditions of the device [15] and that the MFE in pristine organic semiconductor devices are predominantly governed by the reactions of (trapped) triplet excitons and polarons [12].

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To study the underlying mechanism of triplet–polaron interactions, an exquisite control over the triplet exciton density is desired. In line with earlier work [8,15], we propose polymer–fullerene blends as an ideal model system. We study the influence of triplet excitons on OMAR by introducing a low concentration of fullerene in the polymer devices. This creates additional charge transfer states (CTS) by which the number of triplet excitons can be increased, stabilized or decreased, depending on the relative energetic alignment of the CTS with respect to the triplet exciton [18]. Different energetic alignments can be engineered by choosing the right material combinations [19].

2. Theory

2.1. Magnetic field effects

As introduced in the foregoing, mixing of spin states plays a crucial role in OMAR. In this section we will show how spin mixing can lead to magnetic field effects on the current. We start our discussion with MFEs in pristine polymer devices. Fig. 1 presents an energy diagram of the relevant spin dependent transitions from free charges to the ground state. Free charges can form precursor pairs in a singlet \( ^1(\text{e} + \text{h}) \) or triplet \( ^3(\text{e} + \text{h}) \) configuration. Within the precursor pairs the separation between the two carriers is such that exchange interactions are still negligible. This enables hyperfine fields, originating from the hydrogen nuclei in the organic material, to mix the singlet and triplet precursor pairs. An external magnetic field suppresses this mixing. From the precursor pair state, the electron–hole pair can recombine into a singlet (S) or triplet (T) exciton and thereafter to the ground state.

In the absence of a magnetic field (Fig. 1b), spin mixing increases the number of triplet excitons if the formation rate of triplets is larger than that of singlets [11]. This is indicated by process ISC\(_{eh}\) in Fig. 1b. Triplet excitons can decrease the current by interacting with free charge carriers [5,12]. Increasing the magnetic field (Fig. 1a) reduces the triplet formation and thus reduces the triplet–polaron interactions and thereby increases the total current, leading to a positive MC. This magnetic field effect, which is governed by the hyperfine field strength experienced by the precursor pair, will be referred to as a low field effect (LFE) and has a linewidth of a few millitesla.

The LFE in the pristine polymer device can be accompanied by a distinct magnetic field effect at ultra-small field scales (USFE, typically \( \lesssim \text{mT} \)) as has recently been demonstrated [11,20]. Moreover, the triplet–polaron interaction is also magnetic field dependent, leading to so-called high field effects (HFE) occurring at field scales on the order of the zero-field splitting (ZFS, typically 100 mT) of the triplet exciton. When describing the triplet–polaron interactions using a mechanism where (trapped) triplet excitons temporarily capture free polarons into meta-stable trions, as proposed by Cox et al. then this HFE results in a positive MC.

2.2. Introducing additional states

To further investigate the underlying mechanism of triplet–polaron interactions, we are going to control the triplet exciton density by introducing additional states in the energy diagram. Hereby, the number of triplet excitons can be increased, stabilized or decreased. As introduced, this can be accomplished by adding fullerene to the polymer devices. At low concentration, the fullerene effectively quenches singlet excitons into weakly bound, spatially separated charge transfer states (CTS), where the hole resides on the polymer and the electron on the fullerene. The effect of fullerene on the triplet excitons depends on the energetic alignment of the CTS and triplet exciton, where three different kinds of devices can be distinguished.

First, in device type I, where the triplet exciton lies higher in energy than the CTS, the triplet exciton will also be effectively transferred into a CTS in a triplet configuration, as depicted by process \( T \rightarrow \text{CTS in the top panel of Fig. 2} \). When both states are energetically aligned, the CTS and triplet exciton are in equilibrium and an increase in the number of triplet excitons will be stabilized by the CTS, as indicated by process \( T \leftrightarrow \text{CTS in the middle panel of Fig. 2} \). Finally, in device type II, the triplet exciton lies lower in energy than the CTS and the number of triplet excitons can increase by a back-transfer from the triplet CTS, as shown by process \( \text{CTS} \rightarrow T \) in bottom panel of Fig. 2.

The introduction of a CTS can have profound effects on the MFEs, since the number of triplet excitons, ultimately responsible for the magnetic field effect on the current, can be changed significantly in this manner. We will now briefly explain the MFEs. In device type I, CTS decrease the number of triplet excitons.

![Energy diagram of relevant spin dependent transitions (indicated by the arrows) from free charges to the ground state in a pristine device. (a) In a large external applied magnetic field, free charges can form precursor pairs in a singlet \( ^1(\text{e} + \text{h}) \) or triplet \( ^3(\text{e} + \text{h}) \) configuration. From this pair state, the precursor pair can recombine into a singlet (S) or triplet (T) exciton. (b) Without a magnetic field, singlet and triplet precursor pairs can mix due to hyperfine interactions, as indicated by transition ISC\(_{eh}\).](image-url)
number of triplet excitons in the device by process T \rightarrow CTS and this reduction is independent of the magnetic field. As a result, introducing CTS in device type I leads to a reduction of the MFE. In the intermediate case, the increase in the number of triplet excitons by process ISC_{eh} is diminished by process T \rightarrow CTS, also leading to a reduction in the MFE.

Finally, type II devices will show a distinctly different behavior when introducing CTS. The CTS is basically an electron–hole pair on two different molecules and therefore the singlet and triplet CTS can also undergo mixing due to hyperfine fields, as indicated by transition ISC_{CTS} in Fig. 2. The difference with electron–hole pairs in the polymer is that CTS directly recombines to the ground state, therefore, the triplet recombination will be slower than the singlet recombination due to spin selection rules. In the absence of a magnetic field, (Fig. 2f), spin mixing of CTS decreases the number of triplet CTS back-transferred to triplet excitons if the singlet CTS recombination rate is larger than the back-transfer rate. Increasing the magnetic field leads to an increase of the number of triplet excitons as indicated by process CTS \rightarrow T in Fig. 2e. Thus, increasing the magnetic fields leads to more triplet–polaron interactions and thereby a decrease in the total current, resulting in a negative MC. This magnetic field effect is governed by the hyperfine field strength experienced by CTS (B_{hf,CTS}) rather than the electron–hole pair (B_{hf,eh}). This field strength will be significantly lower as a result of the very small hyperfine field coupling that electrons experience on the fullerene sites due to the vanishing nuclear magnetic moment of ^{12}C [15]. Thus, type II devices can give rise to two opposing MFEs with distinctly different linewidths.

In summary, the introduction of additional states, whereby the triplet exciton density can be influenced, could be a valuable tool to study triplet–polaron interactions unraveling their role in OMAR.

3. Methods

3.1. Materials

In this work we studied the magnetic field effects on the current for devices consisting of different polymer–fullerene blends. We used a variety of blends, but this study will focus on blends consisting of poly[(2-methoxy-5-(3,7-
dimethylctyloxy)-1,4-phenylenevinylene] (MDMO-PPV) and a phenyl substituted poly (1,4-phenylenevinylene) semiconducting polymer called Super Yellow PPV (SY-PPV). Both polymers were blended with [6,6]-phenyl-C61-butyric acid methyl ester (PCBM) in different concentrations. In these devices, PPV acts as hole-conducting electron acceptor and this blend is a well-known and extensively studied organic photovoltaic system [21]. The MDMO-PPV was purchased from American Dye Source Inc., the SY-PPV from Merck and the PCBM from Solenne B.V. and all materials were used as received.

3.2. Device fabrication

The devices were prepared on glass substrates with patterned indium tin oxide (ITO) anodes. After careful cleaning, followed by a UV-ozone treatment, a thin layer of poly (3,4-ethylenedioxythiophene);poly (styrenesulfonate) (PEDOT:PSS) was applied by spin coating. The MDMO-PPV, SY-PPV and PCBM were dissolved in orthodichlorobenzene, with a concentration of 10, 8 and 20 mg/ml respectively, and stirred on a hot plate at 50 °C for at least 2 h after appropriate blending. The blends were spin coated at 1200 rpm for 60 s. Subsequently, the samples were transferred to a nitrogen filled glove box where the cathode, consisting of LiF and Al, was evaporated in a high vacuum system (∼ 10⁻⁷ mbar). From this point on, the samples always remain in a dry nitrogen environment. The total junction stack thus consisted of ITO/PEDOT:PSS (60 nm)/blend (~80 nm)/LiF (1 nm)/Al (100 nm), where the blend is either MDMO-PPV(1−x)−PCBMx or SY-PPV(1−x)−PCBMx and with x the PCBM concentration in wt.%.

3.3. Measurements

Magnetic field effect measurements were performed in a cryostat that is attached to a glovebox with a dry nitrogen environment ([O₂] < 0.3 ppm, [H₂O] < 0.3 ppm). The cryostat is placed between the poles of an electromagnet, which allows us to apply an external magnetic field up to 0.5 T, with an accuracy of 50 μT. The devices were driven at a constant voltage V using a Keithley 2400 Series SourceMeter. We measured the current I through the device while sweeping the magnetic field B. From this measurement, after correction for the drift, the magnetoconductance (MC) was calculated with \( MC(B) = \frac{|I(B)| - |I(0)|}{|I(0)|} \).

3.4. Empirical lineshapes

To analyze the magnetoconductance, we used the following fitting function:

\[
MC(B) = \text{LFE} \cdot f(B, B_{\text{LFE}}, B_m, r) + \text{HFE} \cdot B^2 / (|B| + B_{\text{HFE}})^2 \\
+ \text{CTS} \cdot f(B, B_{\text{CTS}}, B_m, r)
\]

The function \( f(B, B_{\text{LFE}}, B_m, r) \), with \( f(B = 0) = 0 \) and \( f(B = \infty) = 1 \), is explained in full detail in Ref. [22]. The function correctly describes the low field effect (LFE), including the ultra-small field effect, in pristine polymer devices, as is shown in the right panel of Fig. 3a. This empirical function allows us to separately extract the role of the intrinsic hyperfine field (\( B_{\text{LFE}} \)) and the extrinsic additional broadening (\( B_m \)) induced by the microscopic mechanism. The function converges to a Lorentzian lineshape for \( f(B_m = 0) \) and a so-called non-Lorentzian for \( f(B_m \gg 0) \). The USFE is incorporated by the parameter \( r \) which describes the limit in which hopping of carriers is no longer slow compared to spin precession in the hyperfine fields. The accompanying high field effect (HFE), depicted in the left panel of Fig. 3a, is fitted with non-Lorentzian lineshape and has a characteristic linewidth \( B_{\text{HFE}} \). Given the similarity of the magnetic field effect induced by charge transfer states compared to the LFE, we will use the same empirical function \( f \), but using a necessarily different hyperfine field parameter \( B_{\text{CTS}} \).

In the fitting procedure, to reduce the number of fitting parameters, the intrinsic hyperfine strength (\( B_{\text{HFE}} \)), extrinsic additional broadening (\( B_m \)) and the USFE parameter (\( r \)) are determined for the pristine polymer devices and these values are then used for the doped devices. The intrinsic hyperfine strength of the CTS (\( B_{\text{CTS}} \)) and the high field effect linewidth (\( B_{\text{HFE}} \)) are determined in a global fit. The amplitudes of the magnetic field effects (LFE, HFE and CTS) can vary with composition and voltage.

4. Results and discussion

4.1. Magnetic field effects

To investigate the influence of additional states on magnetic field effects in the current, we have systematically measured the magnetoconductance as a function of magnetic field \( B \), bias voltage \( V \) and PCBM concentrations \( x \) for two different polymer–fullerene blends. Typical results for three different concentrations \( x \) are shown in Figs. 4 and 5 for MDMO and SY blends respectively. Both pristine devices (Figs. 4a, b, and 5a, b), show a similar lineshape with a positive LFE accompanied by a positive HFE and a USFE. When increasing the PCBM content, we observe a drastic decrease in amplitude of the MFEs. Although the LFE and HFE still show a similar shape for both polymers (Figs. 4c, e and 5c, e), the lineshape around zero applied field is completely different, as is clearly visible in Figs. 4d, f and 5d, f. We conjecture that the drastic decrease in amplitude as well as the remarkable differences in lineshape around zero applied field are caused by the introduction of CTS and their energetic alignment with respect to the triplet exciton in the polymer host. This conjecture will be put on more solid ground in the remainder of this article.

4.2. Lineshapes analysis

We performed a quantitative analysis by fitting the MC(B) data for all compositions and voltages with a superposition of a LFE (including USFE), a HFE and a contribution from the CTS (refer to the Methods section for a complete description). Thus, for each measurement, we get an amplitude of the LFE, HFE and CTS, and also the intrinsic hyperfine field scales (\( B_{\text{LFE}} \) and \( B_{\text{CTS}} \)), as well as the...
The linewidth of the HFE ($B_{\text{HFE}}$). The extracted parameters are shown in Fig. 6a and b present the amplitude of the MFEs in MDMO and SY blends respectively, while the linewidths are shown in Fig. 6c and d. We observe pronounced trends in the amplitudes of the MFEs, whereas the linewidths are approximately constant over the whole doping range.
We previously stated that triplet–polaron interactions are the dominant underlying mechanism for the magnetic field effects in the current in our devices. The positive sign of the LFE and HFE, and the corresponding linewidths are consistent with this interpretation.

4.3. Effect of additional states

We will continue our discussion by examining the triplet–exciton–CTS alignment. In both polymer–fullerene blends, the magnetic field effects caused by triplet–polaron interactions (LFE, HFE) are quenched by more than an order of magnitude (\(B_{\text{LFE}} = 0.85 \pm 0.10 \text{ mT}, B_{\text{HFE}} = 70 \pm 25 \text{ mT} \) and \(B_{\text{CTS}} = 0.45 \pm 0.05 \text{ mT}\)). We previously stated that triplet–polaron interactions are the dominant underlying mechanism for the magnetic field effects in the current in our devices. The positive sign of the LFE and HFE, and the corresponding linewidths are consistent with this interpretation.

Fig. 5. Magnetoconductance as a function of magnetic field at 3.0 V for (a, b) a pristine, (c, d) 1 wt.% doped and (e, f) 5 wt.% doped SY-PPV device. The solid line is a fit using Eq. (1). The right panels (b, d, f) show a zoom around zero applied field.

Fig. 6. Global trends of the magnetic field effects in the current (MFE) as a function of PCBM content, where the left panels (a, c) show the results for the MDMO-PPV device and the right panels (b, d) the SY-PPV device. The top panels (a, b) show the amplitude of the magnetic field effects, whereas the bottom panels (c, d) indicate the corresponding linewidths. The dashed lines provide a guide to the eye.
order of magnitude by adding only a few wt.% PCBM. However, in the SY-PPV blends, an additional MFE contribution arises when the PCBM concentration is increased. This contribution has the opposite sign as the LFE and the linewidth is significantly smaller. We therefore attribute this MFE to the CTS and assign the SY blend as type II, where the triplet exciton lies lower in energy than the CTS. The absence of an additional MFE caused by the CTS in MDMO-PPV blends, gives rise to a classification as type I. To confirm this hypothesis, we need to know the exact energy levels. However, only few reports about the energy alignments of CTS and triplet excitons in low (<20 wt.%) concentration polymer–fullerene blends are available in the literature [23–26]. Triplet energies in conjugated polymers, however, have been studied intensively [27–29]. From the literature, we can conclude that the triplet exciton and CTS in MDMO-PPV:PCBM blends are approximately aligned (1.40 eV [19] and 1.40–1.55 eV [23,25] respectively), whereas the triplet energy in SY-PPV lies slightly lower in energy (1.30 eV [30]). This is in agreement with our assignment. To further explore the effects of energy alignments on the MFEs, one would like to study devices with more pronounced differences in triplet exciton CTS energies. Therefore, we have identified the polymers PF10TBT (\(E_T = 1.35\) eV [24] and \(E_{CTS} = 1.60\) eV [24]) and P3HT (\(E_T = 1.40\) [19] and \(E_{CTS} = 0.90\) eV [31]) mixed with PCBM as good candidates.

5. Conclusion

In conclusion, in this article we presented a proof of concept study in which we tune triplet–polaron interactions in our devices by deliberately doping with fullerene which influences the triplet exciton density. We have systematically investigated the magnetic field effects on the current for two different polymer–fullerene blends. Using a quantitative analysis of the lineshapes we are able to extract the amplitudes and linewidths of the MFEs. We show that distinct changes in the lineshapes between the two polymer–fullerene blends are correlated to the energetic alignment of the triplet exciton and CTS in the blends. Our findings enable a novel method to engineer the OMAR effect at ultra-small magnetic fields and, moreover, investigate the energetic alignment of excited states in organic semiconductors.

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