Ultrafast intersystem-crossing in platinum containing \( \pi \)-conjugated polymers with tunable spin-orbit coupling

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The development of efficient organic light-emitting diodes (OLED) and organic photovoltaic cells requires control over the dynamics of spin sensitive excitations. Embedding heavy metal atoms in \( \pi \)-conjugated polymer chains enhances the spin-orbit coupling (SOC), and thus facilitates intersystem crossing (ISC) from the singlet to triplet manifolds. Here we use various nonlinear optical spectroscopies such as two-photon absorption and electroabsorption in conjunction with electronic structure calculations, for studying the energies, emission bands and ultrafast dynamics of spin photoexcitations in two newly synthesized \( \pi \)-conjugated polymers that contain intrachain platinum (Pt) atoms separated by one (Pt-1) or three (Pt-3) organic spacer units. The controllable SOC in these polymers leads to a record ISC time of \( \sim 1 \) ps in Pt-1 and \( \sim 6 \) ps in Pt-3. The tunable ultrafast ISC rate modulates the intensity ratio of the phosphorescence and fluorescence emission bands, with potential applications for white OLEDs.

The dynamics of spin singlet and triplet excitations in \( \pi \)-conjugated polymers define their performance as active layer in OLEDs and organic photovoltaic (OPV) cells. For example, if both triplet and singlet excitons can be used in OLEDs to convert electrical energy to electroluminescence emission, then the fraction of excitons that potentially can emit light may reach 100\%\textsuperscript{1,2}. Similarly in OPV based on donor/acceptor (D-A) blends, the photogenerated singlet exciton in the polymer donor domains may recombine before reaching the D-A interface, because of its relatively short life time (\( \sim 100 \) ps). In contrast, because of the much longer life time (\( \sim 5 \) ms), triplet excitons could be the answer to this loss mechanism\textsuperscript{3,4}. Therefore, both OLED and OPV technologies may substantially benefit from the proper use of the spin triplet states. Alas, because the SOC in polymers is typically very weak, triplet excitons cannot be efficiently photogenerated in the donor polymers for OPV enhancement; and, similarly, cannot efficiently emit light in OLEDs. The SOC however can be enhanced by embedding heavy atoms such as platinum (Pt) in the polymer backbone chains. Such enhanced SOC, in turn may increase the ISC rate from the singlet-to-triplet manifold, which would make triplet excitons more viable for OPV applications. In addition, the enhanced SOC may also trigger substantive phosphorescence (PH) emission from the lowest triplet state\textsuperscript{5,6}, and thus the emission spectrum from such semiconductor polymers may contain both fluorescence (FL) and PH bands. In fact, these two bands span the visible spectral range, and therefore may potentially be used in designing 'white' electroluminescence emission from the same polymer in OLEDs, with internal quantum efficiency approaching 100\%\textsuperscript{7}.

These beneficial triplet characteristic properties are the main reason that a variety of Pt-containing polymers have been synthesized and studied. Most of these studies however have been focused on the photophysics of the triplet excitons\textsuperscript{8,9}. Importantly, the dynamics of the internal conversion and ISC processes have not been elucidated, and a complete description of the essential electronic excited states involved is still lacking. For example, the metal-to-ligand charge-transfer (MLCT) state, which has been studied extensively in metal-organic complexes\textsuperscript{10,11}, has not been properly addressed in Pt-containing polymers. In the present work we use a broad arsenal of linear and nonlinear optical (NLO) spectroscopies, complemented with electronic structure calculations, for studying the photoexcitations dynamics in two Pt-containing \( \pi \)-conjugated polymers with different \( \pi \)-conjugated spacer unit length between the nearest Pt atoms on the chain, which controls the spin dynamics of excited states, namely the timescales of the ISC process in this class of materials.
The backbone structures of the two Pt-polymers that we studied are shown in Figure 1(a); the synthesis routes of these polymers are described in the Supplemental Information. It is seen that one spacer unit contains a single phenyl ring, dubbed here Pt-1; whereas the other spacer unit contains three phenyl rings, dubbed Pt-3. The NLO spectroscopies used in this work include broad-band ultrafast as well as cw pump-probe photomodulation (PM), electroabsorption (EA), and two-photon absorption (TPA). We complete our investigation by comparing the NLO spectra with linear optical measurements such as absorption and photoluminescence (PL) spectra. We note that absorption and TPA spectroscopy probe electronic excited states with odd or even symmetry, respectively; and consequently are complementary to each other. Whereas EA spectroscopy is sensitive to excited states of both odd and even symmetry. These extensive experimental techniques and quantum-chemical computational results reveal fascinating ultrafast dynamics of the ISC owing to the excited state order and enhanced SOC that is controlled by the chemical composition and structure of these Pt-polymers.

Results

(i) Linear and nonlinear optical spectroscopies. The PL spectra of Pt-1 and Pt-3 films (Fig. 1(b)) show both fluorescence (FL; 2.6–3.2 eV) and phosphorescence (PH; 1.5–2.5 eV) emission bands. The FL (PH) band is assigned to transitions from the lowest singlet (triplet) excited state, $S_1$ ($T_1$) into the ground state, $S_0$ (or $1A_g$)

$^1$Bu, $m^1$Ag, and $n^1$Bu, as well as the novel state, Metal-to-Ligand Charge Transfer (MLCT) are assigned.

In contrast to the PL spectra, the absorption spectra of Pt-1 and Pt-3 films (Figs. 1(c,d)) and solution (Figs. 2(a,b)) have multiple broad overlapping peaks that indicate contributions of many optically active excited states. Consequently, in order to elucidate the nature

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**Figure 1** | Linear and NLO measurements of the Pt-polymers. (a), Chemical structures of the two Pt-polymers, where the “spacer” in Pt-1 has a single phenyl ring, whereas that of Pt-3 has three phenyl rings. (b), Normalized photoluminescence emission spectrum of Pt-1 (blue) and Pt-3 (red) films. FL (PH) stands for fluorescence (phosphorescence) emission. (c) and (d), Electroabsorption (EA) spectra of Pt-1 (c) and Pt-3 (d) films, compared to their respective absorption ($\alpha$) spectrum. The absorption of a non-Pt polymer film is also shown (broken line) in (c). The three excited “essential states” $1B_u$, $mB_u$, and $nB_u$, as well as the novel state, Metal-to-Ligand Charge Transfer (MLCT) are assigned.

**Figure 2** | Two photon absorption (TPA) spectra of Pt-1 (a) and Pt-3 (b) solutions, compared to their respective absorption spectrum ($\alpha$).
of the excited states responsible for the various absorption bands, we studied the EA and TPA spectra of the two Pt-polymers (see Methods). Quantitatively EA is a third-order NLO effect, and thus may be described by the imaginary part of the third order optical susceptibility. The EA spectroscopy, where the sample is exposed to a modulated dc electric field, \( E \), has been a sensitive tool for studying the band structure of inorganic-semiconductors\(^\text{16}\) and organic-semiconductors\(^\text{17}\) that allows for separation of absorption peaks due to tightly bound one-dimensional excitons from that of the continuum band. We note that the EA of confined exciton scales with \( F^2 \), and its spectrum is proportional to the first derivative of the absorption with respect to the photon energy; whereas the EA related to the continuum band scales with \( F^3 \), and its spectrum contains Franz-Keldysh (FK) type oscillation\(^\text{18}\). In contrast, the TPA spectroscopy has been a powerful tool to study the optically forbidden \( \Delta g \) states in \( \pi \)-conjugated polymers\(^\text{19}\).

Figures 1(c,d) show the EA spectra of Pt-1 and Pt-3 films at 80 K up to 4.5 eV. Following Mazumdar et al. model\(^\text{20}\), four essential states (\( 1A_g, 1B_u, mA_u, \) and \( nB_u \)) contribute substantially to the EA spectrum of \( \pi \)-conjugated polymers. At low energy there is a ‘first-derivative’ like feature with zero crossing at ~3.17 eV (Pt-1) and ~3.05 eV (Pt-3) that we thus assign to the \( 1B_u \) exciton, accompanied by several phonon sidebands\(^\text{20}\). In addition there is a prominent field-induced absorption band at higher energies at ~3.9 eV (Pt-1) and ~3.8 eV (Pt-3) that we assign as due to the \( 1A_g \to mA_u \) transition, which becomes partially allowed by the symmetry breaking induced by the electrical field. These \( mA_u \) states appear as inhomogeneous broadened band in the TPA spectra at ~4.03 eV (Pt-1) and 3.9 eV (Pt-3) (Figs. 2(a,b)). In Pt-1 (Fig. 1(c)), we see a ‘second derivative’-like feature with zero crossing at 4.15 eV attributed to the \( nB_u \) state, which is consistent with a weak linear absorption band at 4.2 eV. The EA spectrum of Pt-3 (Fig. 1(d)) at similar photon energies shows a modulation spectral feature with two zero crossings at 4.1 eV and 4.35 eV, respectively, which we ascribed to \( mA_u \) and \( nB_u \) that mix under the influence of the electric field\(^\text{21}\).

Importantly, compared to the absorption spectrum, the \( 1B_u \) EA spectral feature is very different in Pt-1 and Pt-3. The EA feature related to \( 1B_u \) in Pt-1 actually lies above the absorption peak at ~3.05 eV. We thus infer that the absorption onset in Pt-1 is due to a band different than the \( \pi-\pi^* \) transition, and speculate that it originates from the lowest lying \( MLCT \) singlet state (as confirmed by quantum chemistry calculations, see below); whereas the EA derivative feature points to the lowest \( \pi-\pi^* \) electronic character singlet state (\( 1B_u \)) at ~3.17 eV. A splitting of ~0.12 eV between the lowest lying \( MLCT \) state and that of the \( \pi-\pi^* \) state is in good agreement with the theory (see below). The lack of EA intensity for \( MLCT \) state could be due to its localization nature, for which the external field is too small of a perturbation to cause sizable changes in its related optical transitions. The diminished EA feature from the \( MLCT \) state that lies below \( 1B_u \) in Pt-1 indicates weak coupling between the \( MLCT \) and \( mA_u \) of the \( \pi-\pi^* \) manifold.

To further elucidate the excited state properties of Pt-1, we synthesized a \( \pi \)-conjugated polymer similar to Pt-1 but without the Pt-containing group. Compared to Pt-1, the absorption onset of a deposited film of this non-Pt polymer is at ~3.2 eV (see broken line in Fig. 1(c)), which is at the energy level of the \( 1B_u \) exciton in Pt-1 deduced from the EA spectrum. This is additional evidence that the absorption onset of Pt-1 originates from the lowest lying \( MLCT \) singlet state rather than from the \( \pi-\pi^* \) transition.

We also note that the absorption spectra in solution (Fig. 2(a)) and film (Fig. 1(c)) are very similar for the Pt-1 polymer. This shows that the lowest singlet state in Pt-1 is quite robust, in agreement with a localized \( MLCT \) state. In contrast, \( E(1B_u) \) in Pt-3 solution (Fig. 2(b)) is blue-shifted by ~0.1 eV as compared to the film (Fig. 1(d)), because in this polymer the lowest singlet exciton has \( \pi-\pi^* \) electron character. This indicates that the states related with \( MLCT \) do not show the usual solid state effect, probably due to their more localized character. Moreover compared to the robust energy of \( MLCT \) states, the obtained change in \( E(mA_u) \) between film (Fig. 1(c,d)) and solution (Fig. 2(a,b)) in both Pt-polymers gives another evidence that the \( \pi \)-electron singlet states are more spatially extended than the \( MLCT \).

Overall, no FK type oscillation related to the onset of the interband transition is seen in the EA spectra of both polymers. We thus conclude that the \( \pi \)-electron states in Pt polymers are better described in terms of excitons. Assuming the continuum band in \( \pi \)-conjugated polymers to be very close to \( E(nB_u) \)\(^\text{22}\), we may estimate the intrachain exciton binding energy, \( E_0 \) in the two polymers as \( E_0 = E(nB_u) - E(1B_u) \). This yields \( E_0 \approx 1 \) eV for Pt-1 and \( E_0 \approx 1.1 \) eV for Pt-3. This large intrachain \( E_0 \) is in agreement with other measured \( E_0 \) values in the class of \( \pi \)-conjugated polymers\(^\text{23}\). However, for Pt-1 the lowest excited state is \( MLCT \) and therefore \( E_0 \) estimate in this case is not as clear.

(ii) Theoretical calculations. Our theoretical modeling confirms the conclusions that the excited state structure in Pt-1 and Pt-3 polymers is indeed different as obtained in the experiment discussed above. Figure 3 shows schematic ordering of the singlet and triplet manifolds of Pt-1 and Pt-3 that emerge from our work. In addition, the obtained Natural Transition Orbitals (NTOs) of the relevant low-energy excited states are displayed in Fig. 4. We found that the strongly allowed optical transition \( 1A_g \to 1B_u \) (\( S_0 \)) in Pt-1 is a \( \pi-\pi^* \) transition that is characteristic for \( \pi \)-conjugated polymers\(^\text{24,25}\), where both electron and hole are delocalized over the conjugated backbone with only a small participation of Pt atoms (Fig. 4(a)). However, the lowest state in Pt-1 (\( S_1 \)) is \( MLCT \) with significant charge-transfer character as schematically shown in Fig. 3(a). Indeed, the calculated NTOs (Fig. 4(b)) show that in this excited state the electron becomes strongly localized on the Pt atom.
whereas the hole remains delocalized over the conjugated backbone. We note that the absolute values of the calculated excitation energies are systematically blue shifted compared to the experimental data, since they are strongly dependent on the hybrid DFT model used in calculations (see Methods); however the relative trends are well reproduced. For example, the splitting between $E(1B_u)$ and the lowest $E(MLCT)$ in Pt-1 is calculated to be 0.14 eV, close to the experimental value of 0.12 eV. The situation is opposite in Pt-3 system, where $E(MLCT)$ $S_n$ (Fig. 4(e)) is higher than $E(1B_u)$ $\pi-\pi^*$ ($S_1$) (Fig. 4(f)). The lowering of $\pi-\pi^*$ transition energy in Pt-3 is attributed to the larger conjugation segment length between the Pt-atoms (the latter partially break the $\pi$-conjugation, see $\pi-\pi^*$ orbitals in Fig. 4). However, $E(MLCT)$ does not change substantially when going from Pt-1 to Pt-3 polymers. While $E(MLCT)$ may red-shift as a result of better conjugation, this effect is partially cancelled due to smaller electron-hole binding energy in Pt-3 MLCT caused by the larger effective separation between charges, as compared to Pt-1.

Our calculations also suggest a more efficient ISC process in Pt-1 compared to that in Pt-3, which is confirmed by the experimental data (discussed below). Namely, the fast internal conversion within the singlet manifold in Pt-1 populates the lowest singlet MLCT state. The corresponding triplet MLCT state ($T_n$ in Fig. 3(a) and Fig. 4(c)) has the same nature and a very similar energy to the $S_1$ MLCT state. Localization of the electron on the Pt atom leads to an efficient ISC (where the electron spin is flipped) due to the enhanced SOC. Subsequently, the internal conversion process within the triplet manifold populates the lowest triplet state, $T_1$ (Fig. 3(a)), which has $\pi-\pi^*$ character (Fig. 4(d)) from where the phosphorescence emission originates. In contrast, the ISC process in Pt-3 occurs between $S_1$ (Fig. 4(f)) and $T_n$ (Fig. 4(g)) states both having $\pi-\pi^*$ character (Fig. 3(b)), since the lowest MLCT state lies higher than the lowest $\pi-\pi^*$ state in this system. Consequently, the ISC process in Pt-3 is less favorable process, since $\pi-\pi^*$ transitions have smaller effective SOC constant.

(iii) Transient and steady state photomodulation spectroscopy. For studying the ISC dynamics in the two Pt-polymers, we explored the ultrafast photoexcitation kinetics using the time-resolved...
photomodulation (PM) technique. Figure 5(a) shows the transient PM spectrum of Pt-1 solution at various delay times, t, following the pump pulse excitation. The PM spectrum contains three PA bands: PA1 at ~0.75 eV, PA2 at ~2.5 eV, and a broad PA band (TA) at ~1.9 eV, which partially overlaps with PA2. The bands PA1 and PA2 (fast components) are correlated since they decay together with a time constant, τ < ~1 ps. The band PA1 (~0.75 eV) in fact coincides with the energy difference E(mA0)−E(1Bu) extracted from the EA and TPA spectra above, and thus is assigned as 1Bu → mA0 optical transition that originates from the photogenerated singlet excitons. The energy difference E(MLCT)−E(mA0) (~0.95 eV) indicates that PA1 cannot be due to the transition MLCT → mA0; this conclusion is also consistent with the lack of EA signal from the MLCT states, as discussed above. The band PA2 is interpreted as due to the transition 1Bu → kA0 from the same singlet excitons, in analogy with other π-conjugated polymers.26 We thus conclude that the primary excitations in Pt-1 are still π-π* excitons, even that the MLCT state is the lowest singlet exciton in this polymer. This happens since the pump pulse predominantly populates the π-π* manifold, and there is little interaction between the π-π* and MLCT states during the ultrafast hot exciton thermalization process.

Figure 5(b) shows the steady state PM spectrum of Pt-1 films. The spectrum is dominated by a broad PA band that peaks at ~1.95 eV, which we interpret as due to triplet-triplet transition from the lowest π-π* triplet, T1, since it has the same dynamics as that of the PH emission band in this polymer.11 This cw PA band is very similar to the broad TA band obtained in the ps time domain; and we thus interpret the ultrafast TA band as due to triplet excitons. Consequently, the ultrafast decay of bands PA1 and PA2 into the TA band is caused by the ISC process from the singlet to the triplet π-π* manifolds. We thus conclude that the ISC process in Pt-1 is much faster than in any other π-conjugated polymer (apart from triplet formation via singlet fission). The reason for the ultrafast ISC in this Pt-polymer is the much stronger SOC that is due to the ‘heavy atom’ effect of the intrachain Pt atoms. A more rigorous comparison between a Pt-polymer and its non-Pt analog (that have a different structure than Pt-1) is shown in the Supplement Information (Fig. S-2). It shows again that the PA bands related to the π-π* singlet exciton decay much faster in the Pt-polymer compared to the non-Pt polymer analog.

Figure 5 (c, d) shows the transient PM spectrum of Pt-3 solution at various times, t. As in Pt-1 the spectrum here also contains three PA bands: PA1 at ~0.7 eV, PA2 at 1.7 eV and a relatively broad TA band at 1.8 eV that overlaps with PA2. The bands PA1 and PA2 are correlated since they decay together with τ ~ 6 ps. Similar as in Pt-1 we interpret the ultrafast PA bands in Pt-3 as due to singlet excitons in the π-π* manifold, and thus their decay reflects the ISC process dynamics in this polymer. The substantially slower obtained ISC process in Pt-3 explains the relative stronger cw FL emission of this polymer compared to that of Pt-1 (Fig. 1). The dominant PA band (T) in the cw PM spectrum at 1.7 eV is interpreted as due to T1 → Tn transitions in the triplet manifold, and thus the PA band in Fig. 5(a) reflects the transient triplet PA. We note, however, that the cw PM spectrum of Pt-3 may have some contribution from polarons since a PA band related to polarons is observed below ~0.5 eV.27

Discussion

In this work two Pt-containing polymers (namely Pt-1 and Pt-3) with different organic spacer length in between each two adjacent intrachain Pt atoms were synthesized and extensively studied using a variety of NLO spectroscopies that include electroabsorption, two-photon absorption, and ultrafast and steady state photomodulation. The NLO spectra were compared to the absorption and luminescence spectra. From quantum chemistry calculation and the NLO measurements we conclude that the lowest singlet state in Pt-1 is a Metal-to-Ligand Charge Transfer (MLCT) state, which lies below the lowest π-π* exciton; however, the order is reversed in Pt-3. Surprisingly, the primary photoexcitations in both polymers are singlet π-π* excitons, irrespective of the excited state order. We note, however that the electron in the MLCT state is localized on the Pt atom center orbital. Consequently the MLCT states have relatively large SOC and their energy is practically independent on the linker length. In contrast the energy of the π-π* transitions substantially depend on the linker length. Because of the π-electron delocalization within the π-conjugated linker, the π-π* IBu state has relatively weak SOC, which is further reduced for longer linker. This rationalizes our experimental results of the way the intrachain Pt atom influences the ISC rate in both Pt-1 and Pt-3.

From the ps transient decay of the photogenerated π-π* excitons we deduce a record ISC timescale of <~1 ps and ~6 ps in Pt-1 and
Pt, respectively. The larger ISC rate in Pt-1 is attributed to stronger SOC that originates from the localized MLCT singlet and triplet states through which the ISC transition may occur. Such energetics is attributed to a smaller spacer between adjacent intrachain Pt atoms. We conclude that the controllable spacer between the intrachain Pt atoms not only tunes the effective spin-orbit coupling, but also influences the excited state order of the MLCT and π-π* states. This implies synthetic design strategies to achieve optimal photocitation spin dynamics in metal-containing π-conjugated polymers for organic OLED and OPV technologies. In particular, Pt-polymers with larger spacer in between each adjacent Pt-atoms may have smaller ISC rate and consequently larger fluorescence emission, which may be readily used for white OLEDs.

Methods

CW spectroscopies. The absorption spectrum was measured with a Cary UV-Vis-NIR spectrometer. For the cw PL emission, an Ar+ laser beam at 3.5 eV was used to excite the Pt-polymer films that were kept at 40 K under dynamic vacuum. The PL emission was collected using a large f-number lens, and spectrally and spatially filtered to eliminate the excitation laser light. Corrections due to the system spectral response and and wavelength to photon energy transformation were readily done. The steady state PM spectrum was obtained using a standard setup[9] with pump excitation from an Ar+ laser beam at 3.5 eV. The pump beam was modulated at a frequency, f = 300 Hz by a mechanical chopper. A beam from an incandescent tungsten/halogen lamp was used as the probe. The PM spectrum in the form of (∆I/T), where ∆T is the change in the transmission, T induced by the pump beam was measured using a lock-in amplifier referenced at the laser for the mid-IR spectral range; and a high power (energy/pulse = 10 mJ) 1 kHz) laser for the near-IR/visible spectral range. In both laser systems the pump excitation was set at fpy = 3.1 eV. For the low intensity measurements we used an optical parametric oscillator (Opal, Spectral-Physics) that generates probe fpy continuously changing from 0.55 to 1.05 eV, and from 0.14 to 0.43 eV, respectively[10]. For the high intensity measurements, white light super-continuum was generated for the probe fpy ranging from 1.15 to 2.7 eV. In general, the transient PM signal, ∆I/T(τ) is negative for photoinduced absorption (PA) and positive for photo-bleaching (PB)[11]. The transient PM spectra from the two laser systems were normalized to each other in the near-IR-visible spectral range, for which the IR probe photon energy from the low power laser system was doubled.

TPA measurements. The TPA spectrum was measured using the polarized pump-probe correlation technique with the low repetition rate high-power laser system at time delays of several ps[12]. The partially polarized pump beam was set at 15° to each other, which is much below the polymer absorption band; whereas the probe beam from the white light super-continuum covered the spectral range from 1.7 to 2.8 eV. The temporal and spatial overlap between the pump and probe beams on the sample film leads to PA signal that peaks at t = 0; this transient PA has a temporal profile identical to the cross-correlation trace of the pump and probe pulses[13]. The ultrafast PA is TPA of one pump photon and one probe photon, of which spectrum yields the TPA spectrum in the spectral range between 3.25 to 4.35 eV.

EA spectroscopy. For the EA measurements we used thin polymer films spin cast on an EA substrate (template)[14]. The EA template consisted of two interdigitated sets of Au m wide gold electrodes, which were patterned on a sapphire field[15], and verified that no EA signal was observed at f = 1 kHz. The temperature of the sample was kept at 5 K and its temporal profile was identical to the cross correlation trace of the pump and probe pulses[16]. The ultrafast PA is TPA of one pump photon and one probe photon, of which spectrum yields the TPA spectrum in the spectral range between 3.25 to 4.35 eV.

Excited states were calculated using time-dependent DFT (TDDFT) technique. These are currently methods of choice for quantum-chemical modeling of ground and excited state electronic properties in molecules of intermediate size. For all simulations we used hybrid B3LYP[17] functional coupled with LANL2dz(Pt)/6-31G* (all other atoms) basis set as implemented in the Gaussian09 suite[18]. Such model chemistry has shown an excellent quantitative performance in similar organometallic compounds in a number of previous studies[19]. We lowest singlet (triplet) states have been calculated and analyzed using the natural transition orbital (NTO) approach[20] to determine their characteristic nature. A moderately polar solvent, ethanol (ε = 24.9), is included in this study via the conductor-like polarizable continuum model (CPCM) as implemented in Gaussian09 software package[21] in order to mimic the polymer’s polarizable dielectric environment. Even though this may be an overestimation of the dielectric constant, the effects of this dielectric medium throughout our study are minor and the conclusions are the same as for the simulations in the gas phase.

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Author contributions
M.T. and A.G. were responsible for the transient PM measurements in the visible and mid-IR, respectively; S.S. was responsible for the EA and TPA measurements; T.D. was responsible for the cw spectroscopies; A.G. and C.-X.S. were responsible for the experimental figures; C.-X.S. also wrote the first draft of the manuscript; S.T. was responsible for the quantum chemical calculations and manuscript second draft; Z.V.V. was responsible for the project planning, group managing, and final writing.

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