Effect of Molecular Side Groups and Local Nanoenvironment on Photodegradation and Its Reversibility

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Effect of molecular side groups and local nanoenvironment on photodegradation and its reversibility

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ABSTRACT

Degradation of organic semiconductors in the presence of oxygen is one of the bottlenecks preventing their wide-spread use in optoelectronic devices. The first step towards such degradation in functionalized pentacene (Pn) derivatives is formation of endoperoxide (EPO), which can either revert back to the parent molecule or proceed to molecule decomposition. We present the study of reversibility of EPO formation through probing the photophysical properties of functionalized fluorinated pentacene (Pn-R-F8) derivatives. Experiments are done in solutions and in films both at the single molecule level and in the bulk. In solutions, degradation of optical absorption and its partial recovery after thermolysis were quantified for various derivatives depending on the solvent. At the single molecule level, low concentrations of each type of molecules were imaged in a variety of polymer matrices at 633 nm excitation at room temperature in air using wide-field fluorescence microscopy. Fluorescence time trajectories were collected and statistically analyzed to quantify blinking due to reversible EPO formation depending on the host matrix. To understand the physical changes of the molecular system, a Monte Carlo method was used to create a multi-level simulation, which enabled us to relate the change in the molecular transition rates to the experimentally measured parameters. At the bulk level, photoluminescence decay due to photobleaching and recovery due to EPO reconversion were measured for the same derivatives incorporated into various matrices. These studies provide insight into the synergistic effect of the local nanoenvironment and molecular side groups on the oxygen-related degradation and subsequent recovery which is important for development of organic electronic devices.

Keywords: Organic semiconductors, photodegradation, thermolysis, single molecule fluorescence spectroscopy, Monte Carlo, fluorescence intermittency

1. INTRODUCTION

The use of organic materials in low-cost (opto)electronic and photonic devices is of substantial current interest.\textsuperscript{1} Regardless of the application, successful commercialization of devices is hindered by the degradation of the organic materials due to reactions with oxygen, water, and so on. Previous studies have indicated that exposure to oxygen is detrimental to the lifetime of promising organic molecules exemplified by acene derivatives, in which the dominant oxygen-induced degradation mechanism is the formation of endoperoxides (EPO).\textsuperscript{2,3} However, EPO formation rate is environment dependent, and can be manipulated with the change in the film composition.\textsuperscript{3} To better understand which environmental changes are most effective in inhibiting EPO formation and in prolonging the lifetime of devices, we study these changes on both the nanoscale with single molecule spectroscopy\textsuperscript{3,4} and on the semi-bulk scale with absorption and photoluminescence (PL) measurements of dilute solutions and films containing low concentrations of molecules of interest incorporated into polymer matrices.\textsuperscript{5,6} Our studies probe the photophysics of technologically relevant solution-processable high-mobility functionalized acene derivatives which have been previously utilized in thin-film-transistors and other (opto)electronic applications.\textsuperscript{1,7}

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On the single-molecule level, it has been established that fluorescence intermittency (blinking) observed in Pn-R-F8 molecules incorporated in a polymer matrix is related to reversible EPO formation. However, understanding of energy and charge transfer processes between the photoexcited molecule and other molecules in the surrounding environment during the molecule degradation and recovery is difficult due to complexity of the nanoscale environment. To aid in the interpretation of the data, Monte Carlo simulations have been utilized to model fluorescence time trajectories that are affected by these processes. In particular, we seek to further refine the model and to determine the accuracy of the single molecule analysis used in experimental data processing. Through both experimental and computational approaches, we aim to better understand the requirements for molecular structure and local environment that enable stable organic (opto)electronic devices less susceptible to photodegradation.

Figure 1. Functionalized pentacene (Pn-R-F8) and indenofluorene (IF-R) derivatives used in our studies. Structures of the side groups R are also included.

2. EXPERIMENTAL

2.1 Materials

In order to investigate the effects of the side groups and host matrix on photodegradation, we chose three fluorinated pentacene (Pn) derivatives (Pn-R-F8) with functionalized side groups R of TCHS (tricyclohexylsilyl)ethynyl), NODIPS (n-octyldiisopropylsilyl)ethynyl), and TIPS (tris(isopropyl)silyl)ethynyl) shown in Figure 1. These derivatives share similar absorption and PL spectra in isolated molecules, but their intermolecular interactions, as well as interactions with the environment, depend on the side group R. These molecules have been previously utilized as acceptors in donor-acceptor heterojunctions, exhibiting strong dependence of exciton and charge carrier dynamics on the size of the R group (TCHS>NODIPS>TIPS). Additionally, reversible EPO formation has been observed on the single-molecule level in Pn-TCHS-F8 molecules incorporated in poly(methyl)methacrylate (PMMA). Moreover, this process could be controlled by the addition of a different type of molecule, such as an indenofluorene derivative (IF-TIPS in Figure 1) to the PMMA matrix.

For studies in solution, Pn-R-F8 molecules were dissolved either in benzene or in chlorobenzene (CB) at a concentration of $2 \times 10^{-5}$ M. The semi-bulk thin film preparation involved combining 30 mM Pn-R-F8 solutions in toluene with a 13 wt% PMMA solution in toluene at appropriate ratios to yield average Pn-Pn intermolecular spacings in the 3-11 nm range. In selected cases, IF-TIPS molecules (1 mM solution in toluene) were added to the mixture at concentrations that would achieve the average IF-IF intermolecular spacing of 5 nm in the film. After a 20 minute sonication of mixtures of solutions, the films (either Pn-R-F8:PMMA or Pn-R-F8:IF-TIPS:PMMA)
were deposited by spin-casting at 3000 rpm for 50 seconds from 60 µL of solution. For studies at the single-molecule level, ultra-low (10⁻¹⁰ M) concentrations of Pn-TCHS-F8 molecules were incorporated in PMMA (Mw 75,000 g/mol), polystyrene (PS, Mw 280,000 g/mol), or poly(9-vinyl)carbazole (PVK, Mw 1,100,000 g/mol) matrices as described in our previous publications.3,4

2.2 Measurements of optical absorption and photoluminescence

The propensity of Pn-R-F8 for photodegradation in solutions was measured by monitoring optical absorption of solutions irradiated by a UV lamp (300-400 nm) at 0.5 mW/cm² in air for up to 100 hours, depending on the solvent. Absorption spectra were taken using a xenon lamp illumination of the solution through a 200 µm optical fiber and collecting the transmitted light through a 50 µm optical fiber which was processed by the Ocean Optics USB2000 spectrometer. Spectrum of the solvent without Pn-R-F8 molecules served as reference.

Once the solution was completely photobleached so that no absorption in the 500-700 nm region (Fig. 2(a)) could be detected, the sample went through thermolysis. For this, the vial was sealed, wrapped in foil and submerged in water at 80°C for two sets of three hours, with absorption data collected at the end of each three hour set (Fig. 2(a)).

To measure the PL decay due to photobleaching in semi-bulk films, an Olympus IX-71 inverted microscope was used with a 10x magnification objective lens, which focused a 633 nm HeNe laser with power ranging between 300 and 700 µW onto a prepared sample. The PL was collected through a 600 µm optical fiber and processed by an Ocean Optics USB2000-FLG spectrometer. Six hundred consecutive spectra were collected with an integration time of 100 milliseconds and 5 scans to average. A dark spectrum was also taken as a background reference, and this was subtracted from the data. Each spectrum was integrated over the 500-700 nm region of interest, which provided a value for each respective time. For studies of photobleaching reversibility, after the first experimental run, the sample was kept in the dark for 20 minutes at room temperature in air, after which the experiment was repeated (run 2) on the same sample area.

Single molecule fluorescence spectroscopy was performed under a 633 nm excitation, using a 100x objective and Olympus IX-71 inverted microscope, in the wide-field geometry with an EMCCD detector, as described in detail in our previous publications.3,4

3. RESULTS AND DISCUSSION

3.1 Photodegradation and recovery in solution

Figure 2 shows (a) how photodegradation of Pn-R-F8 molecules manifests into optical absorption spectra and (b) how the polarity of the solvent affects the rate of the degradation. Upon prolonged UV irradiation, the 500-700 nm absorption band (corresponding to the S₀-S₁ transition in Pn-R-F8 molecules) disappears, and an enhanced absorption in the 350-450 nm region is observed. Similar effect was previously observed upon degradation of non-fluorinated functionalized Pn derivatives (such as Pn-TIPS) and attributed to the photodimerization and EPO formation.3 In the fluorinated derivatives, Pn-R-F8, used in our studies, both processes should be considerably less pronounced, as indeed can be seen from the comparison of the Pn-R-F8 versus Pn-TIPS data in benzene in Fig. 2(b). In particular, in benzene the half-lifetime (defined as the time over which the PL has decayed to the 50% level as compared to that in the fresh sample at t = 0) of Pn-TIPS-F8 derivative is about a factor of 4 higher than that of Pn-TIPS, consistent with our previous observations of higher stability of Pn-TIPS-F8 as compared to Pn-TIPS when incorporated into PMMA.4

The Pn-R-F8 derivatives exhibit more than an order of magnitude (a factor of 15-40, depending on the derivative) higher stability in benzene as compared to CB (Fig. 2(b)). A strong dependence of photodegradation on the solvent polarity, and the case of benzene versus CB in particular, was also observed for Pn-TIPS and attributed to the π–π interactions of the Pn molecule with the surrounding benzene molecules which prevented Pn-Pn dimerization. Similar considerations can be applicable to the case of fluorinated Pn derivatives, although quantitative assessment of relative contributions of photodimerization and EPO formation into degradation of Pn-R-F8 in solution requires further studies.
Figure 2. (a) Degradation under UV illumination over time $t$ and partial recovery of the optical absorption spectra after 6 hours of thermolysis of Pn-NODIPS-F8 solution in chlorobenzene (CB). No absorption in the 500-700 nm region was observed after 7 hours of UV illumination at which point the thermolysis was initiated. (b) Absorbance at 640 nm normalized by its value at $t = 0$ (fresh solution) at various UV irradiation times obtained in Pn-R-F8 solutions in benzene (open) and CB (solid). Data for Pn-TIPS in benzene are also included.

The effect of side groups $R$ on the photobleaching of Pn-R-F8 in solution was more pronounced in CB as compared to benzene (Fig. 2(b)), with TCHS and NODIPS derivatives exhibiting about a factor of 2 longer half-lifetimes than the TIPS derivative. This is consistent with our previous observations of higher stability of NODIPS and TCHS derivatives as compared to TIPS, in Pn-R-F8:PMMA films, indicative of a protective role of large side groups against photo-induced degradation.

In both solvents, and for all derivatives, the 3-6 hour thermolysis recovered about 6% of the original 500-700 nm absorption strength observed in fresh solutions. This recovery is most likely due to the EPO converting to the parent Pn molecules, as reversal of photodimerization would not be expected under low-temperature thermolysis conditions used in our experiments. The percentage of recovery did not considerably change between 3 and 6 hours of thermolysis; however, the absorption spectra obtained after 6 hours of thermolysis had an increased 350-450 nm spectral component as compared to those after 3 hours, which indicates that degradation processes continue during thermolysis.

3.2 Photodegradation and recovery in films

Figure 3 shows photobleaching of Pn-TCHS-F8 molecules incorporated in PMMA, measured via monitoring PL under continuous 633 nm illumination over $>600$ sec, depending on the Pn concentration (a) and on the local environment (b-c). In particular, samples with higher concentrations of Pn-TCHS-F8 exhibited better photostability, most likely due to formation of nanoaggregates with reduced free volume, which decreased the oxygen permeability resulting in inhibited reactions of Pn-TCHS-F8 molecule with oxygen. After photobleaching the Pn-TCHS-F8:PMMA sample for 10 min, about a 5% recovery in PL was observed after the sample was kept in the dark for 20 min (Fig. 3(b)) due to the EPO conversion to the parent Pn-TCHS-F8 molecule. The percentage of recovery under the same conditions doubled when IF-TIPS molecules were introduced in the film (i.e. in Pn-TCHS-F8:IF-TIPS:PMMA films, Fig. 3(c)) due to IF-TIPS molecules acting as quenchers of the highly reactive singlet oxygen produced upon the conversion of the EPO to the parent Pn-TCHS-F8 molecule, thus preventing an immediate attack of the newly recovered Pn-TCHS-F8 molecule. This effect was also observed at the single-molecule level.3
3.3 Monte Carlo simulations of single-molecule fluorescence time trajectories

3.3.1 The model

In preparation for single molecule fluorescence spectroscopy, we refined the Monte Carlo simulation, which models the fluorescence time trajectories of individual molecules. The fluorescence counts obtained from each single molecule as a function of time are processed to produce time traces similar to those of Fig. 5(a), from which “on” and “off” times can be extracted and statistically analyzed as described in our previous publication. The complementary cumulative distribution functions (CCDFs) are then created for these “on” and “off” times (e.g. Fig. 5(b)) and fit to known distributions such as Weibull, exponential and lognormal using maximum likelihood estimation (MLE). The Monte Carlo simulation creates similar fluorescence time traces (Fig. 5(a)), which can be processed identically to the experimental data. By manipulating the parameters of the simulation we can mimic the experimental changes which provides key insights into the photophysical processes under study.

Previous research has modeled the charge carrier dynamics as observed in single-molecule fluorescence experiments by using a three-level system, with a ground state, excited state and dark state. Our previous work has used a four-level system, which includes the states mentioned above and, additionally, an irreversible “photobleached” state, as well as a non-radiative recombination rate ($k_{21n}$) between the excited state and the ground state. In this model, the states 1 and 2 are the ground and excited states of the Pn-R-F8 molecule, whereas dark state 3 is the EPO obtained as a result of the reaction of Pn-R-F8 molecule with singlet.
oxygen. When the EPO reverts to the parent Pn-R-F8 molecule, the molecule can be re-excited again (“blinking”). The state 4 is a decomposed molecule. Using known experimental parameters, such as light intensity and fluorescence quantum yield of Pn-R-F8 molecules under study, we calculated realistic rates for $k_{12}$, $k_{21}$, and $k_{21n}$. In choosing realistic values for $k_{23}$, $k_{31}$ and $k_{3out}$, we focused on rates that produce numerically simulated data similar to the experimental data and that result in a range of “on” and “off” times that produce fits with the highest accuracy. For time traces of 100 sec duration typically used in our experiments and thus used in simulations, it is important that the average “on” and “off” times are considerably shorter than the 100 sec time window and yet long enough to be experimentally realistic. Modifying these values and incorporating distribution rates for the transitions provide significant insight into the accuracy of the data processing for the experimental data.

Figure 5. The Monte Carlo Simulation produces time traces (a) as outputs for the data which are converted to CCDFs (b) and fit to a variety of distributions. (c) and (d) Weibull fit values of “off” times compared to the variation in the $k_{31orig}$ rate, with $A = 1$ or $A = 1.5$ and $k_{12} = 2.9 \times 10^4 s^{-1}$, $k_{21} = 7 \times 10^7 s^{-1}$, $k_{23} = 222 s^{-1}$, and $k_{21n} \approx 1.5 \times 10^7 s^{-1}$. The $A$ fit parameters lie close to their true values and the fits for the $\lambda$ values are linear with respect to $k_{31orig}$ with the slope close to 1, as expected. In particular, $\lambda = 0.94k_{31orig} + 0.01$ for $A = 1$ and $\lambda = 0.96k_{31orig} + 0.02$ for $A = 1.5$.

The initial results from the simulation show that the “on” times for the simulation are affected by the $k_{23}$ rate and the “off” times are affected by both the $k_{31}$ and the $k_{3out}$ rates. It was also seen that when the rates were fixed, the data fit an exponential distribution. Since the experimental data are typically described by a more complicated distribution such as Weibull distribution, this indicates a variation in the transition rates with respect to time. In particular, the Weibull-distributed process is characterized by a time-dependent rate
\[ k(t) = \frac{A}{\lambda} \left( \frac{t}{\lambda} \right)^{A-1}, \text{ where } A \text{ and } \lambda \text{ are Weibull fit parameters.} \]

(If \( A = 1 \), the time-independent rate \( k = 1/\lambda \), characteristic of the single-exponential distribution, is recovered.)

To investigate how such time-dependent rate manifests into the CCDF so that such rate can be successfully recovered from CCDF fits of the experimental data, we performed Monte Carlo simulations as follows. The model was simplified down to three states (1-3). The rates \( k_{12}, k_{21}, \) and \( k_{23} \) were fixed at \( 2.9 \times 10^4, 7 \times 10^7, \) and \( 2.22 \times 10^7 \), respectively. The rate \( k_{21n} \) was fixed such that \( k_{21}/(k_{21} + k_{21n} + k_{23}) = 0.82 \), where 0.82 is the quantum yield of Pn-R-F8 molecules under study, resulting in \( k_{21n} \) approximately equal to \( 1.5 \times 10^7 \). The Weibull rate \( k(t) \) was applied to the \( k_{31} \) rate so that \( k_{31} = (A * k_{31\text{orig}})(k_{31\text{orig}} * t)^{A-1} \), where \( k_{31\text{orig}} \) is the original rate, which was varied from 0.025 – 2, \( t \) is the time the molecule has spent in the dark state (state 3), and \( A \) is the Weibull distribution parameter, which for our experiments varied from 1.2-1.5. From these rates the predicted average “on” time can be calculated using \( \tau_{on} = (k_{21} + k_{21n} + k_{23})/(k_{23} * k_{12}) \), which is 13.15 seconds. The predicted average “off” time is calculated using \( \tau_{off} = (1/k_{31\text{orig}}) * \Gamma(1 + 1/A) \), where \( \Gamma \) is the Gamma function. For the range of \( k_{31\text{orig}} \) used in our simulations the predicted average “off” times range from 0.45 – 36.1 seconds for \( A = 1.5 \) and 1.4 – 40 for \( A = 1 \). Even at the high end, the sum of the predicted average “on” times and “off” time is lower than the 100 sec limit, which should result in a sufficiently wide distribution to obtain reliable fits.

Figures 5(c) and (d) show results from the simulated data (e.g. Fig. 5(a)) for twelve sets of 100 time traces each for \( A = 1.5 \) and nine sets of 100 time traces for \( A = 1 \), where each set had a different \( k_{31\text{orig}} \) value. The CCDFs were created from these data sets and fit to various distributions (Fig. 5(b)) using MLE. As expected, the Weibull distribution produced the best fit, and the returned fit values were within error of the predicted values. In particular, when using the Weibull rate \( k(t) \) as a transition rate, the returned \( A \) values are very close to the expected value, and the fit of the \( \lambda \) vs \( k_{31\text{orig}} \) values results in a linear relationship with a slope close to one. This was accurate for both values of \( A \) used in simulations. However, the accuracy in the returned \( A \) values decreased as the \( k_{31\text{orig}} \) increased (Fig. 5(d)), which is likely due to the non-zero integration time (100 ms) dictated by our experimental settings and thus used in the simulation. When \( k_{31\text{orig}} \) is large, the predicted “off” times become small, introducing rounding errors due to integration time. This skews the distribution and reduces accuracy in the fit values.

![Figure 6. The complementary cumulative distribution functions (CCDFs) obtained from experimental data for Pn-TCHS-F8 in three different polymer host matrices: PMMA, PVK, and PS. Weibull fits are also shown.](image-url)
3.4 Single molecule spectroscopy: experiments

The CCDFs for the “on” times obtained from single-molecule fluorescence time trajectories compiled for small ensembles (85-105 molecules) of Pn-TCHS-F8 incorporated at ultra-low concentrations in PMMA, PVK, and PS host matrices are shown in Fig. 6. In all matrices, the data are not single-exponential, indicative of distributions in the transition rates $k_{23}$. The Weibull fits to these data produced $\lambda(A)$ values of 47.2 sec (1.67), 35.8 sec (1.27), and 31 sec (1.29) in PMMA, PVK, and PS, respectively. The corresponding average “on” times are then 42 sec, 35 sec, and 29 sec, respectively, which are consistent with differences in oxygen permeability and diffusion coefficients in these matrices, with, for example, PMMA having these parameters lower than PS by more than an order of magnitude, which renders it a much better matrix for single-molecule fluorescence spectroscopy. Additionally, the percentage of molecules that exhibit blinking (i.e. reversible as opposed to irreversible degradation at the 100 sec time scale of the experiment) increased from 13% in PMMA to 20% in PS. This suggests that even though the reaction of the Pn-TCHS-F8 molecule with singlet oxygen is more probable in PS (as manifested through shorter average “on” times) as compared to PMMA, the EPO conversion back to the parent Pn-TCHS-F8 molecule is also more probable in PS, possibly due to enhanced oxygen diffusion in PS that could promote singlet oxygen quenching. Similar effect was observed when IF-TIPS molecules were added to the PMMA, serving as singlet oxygen quenchers.

4. CONCLUSION

In summary, we observed dependence of photodegradation of functionalized fluorinated Pn derivatives Pn-R-F8 on the side groups R and local environment. More than an order of magnitude enhancement in photostability was observed in Pn-R-F8 solutions in benzene as compared to chlorobenzene; in both solutions, partial reversal of photodegradation could be achieved by thermolysis. In Pn-R-F8:PMMA films, addition of IF-TIPS molecules which served as quenchers for singlet oxygen enhanced the reversal, which was observed at room temperature in air. On the single molecule level, photostability of individual Pn-R-F8 molecules depended on the polymer matrix, with highest (lowest) photostability observed in PMMA (PS), in accordance with their oxygen permeability and diffusion coefficient characteristics. Computational modeling has provided insight into the experimental design for future single molecule fluorescence spectroscopy experiments so that the fits to the experimental data yield reliable values for transition rates involved in the model. Our experiments indicate that the choice of the side group R can enhance photostability by at least a factor of 2, with the additional enhancement of more than an order of magnitude provided by the suitable choice of the surrounding environment. Several routes for reversibility of photodegradation, including addition of singlet oxygen quenchers, are also available. These fundings inform us of the molecular design considerations and conditions needed to prevent degradation of organic materials for the use in future (opto)electronic devices.

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