Macromolecules

Article

pubs.acs.org/Macromolecules

Quantitative Measurements of the Temperature-Dependent ² Microscopic and Macroscopic Dynamics of a Molecular Dopant in a **Conjugated Polymer**

⁴ Jun Li,[†][®] Correy Koshnick,[‡] Souleymane O. Diallo,[∥] Sophia Ackling,[#] David M. Huang,[#][®] ⁵ Ian E. Jacobs,[‡][®] Thomas Harrelson,[†][®] Kunlun Hong,[⊥][®] Guangwu Zhang,[§] Joseph Beckett,[§] ⁶ Mark Mascal,[§] and Adam J. Moulé^{*,†}

7 [†]Department of Chemical Engineering, [‡]Department of Materials Science and Engineering, and [§]Department of Chemistry,

University of California, Davis, Davis, California 95616, United States 8

^{||}Neutron Scattering Science Division and ^{\perp}Center for Nanophase Materials Sciences, Oak Ridge National Laboratory, Oak Ridge, 9 Tennessee 37831, United States 10

[#]Department of Chemistry, School of Physical Sciences, The University of Adelaide, Adelaide, SA 5005, Australia 11

Supporting Information 12

ABSTRACT: Understanding the nature of dopant dynamics in 13 the solid state is critical for improving the longevity and stability 14 of organic electronic devices and for optimizing the doping-15 induced solubility control (DISC) patterning method. In this 16 work, we use quasi-elastic neutron scattering (QENS) and 17 fluorescence quenching techniques to develop a comprehensive 18 picture of both the microscopic and macroscopic dynamics of 19 20 the soluble p-type molecular dopant tetrafluoromethyloxycarbonyltricyanoquinodimethane (F4MCTCNQ) in the 21 conductive polymer poly(3-hexylthiophene-2,5-diyl) (P3HT). 22 Specifically, fast dynamics (ps-ns) of the dopant, such as the 23 methyl and the methoxycarbonyl group rotations, are observed 24 25 in QENS experiments. From confocal fluorescence microscope 26 experiments, longer-range/slower dopant diffusion (ms-days) is 27



captured. However, in order to fit these data, it is necessary to incorporate a Langmuir isotherm equilibrium between the neutral and ionized dopant molecules. Ionized F4MCTCNQ is strongly favored by the equilibrium, but it diffuses 3 orders of magnitude 28

slower than neutral species. Moreover, the macroscopic diffusion is found to depend mostly on the minority concentration of 29 neutral dopant molecules in the film. Finally, the global diffusion coefficient of the monoester-substituted dopant F4MCTCNQ is 30

shown to be more than 1 order of magnitude smaller than that of the widely used dopant F4TCNQ. 31

INTRODUCTION 32

33 The successful development of organic electronic devices, such 34 as organic photovoltaics (OPVs),^{1,2} organic light-emitting 35 diodes (OLEDs),^{3,4} and organic field-effect transistors 36 (OFETs),^{5,6} is largely attributable to the tunability of their 37 electrical properties. Like inorganic semiconductors, organic 38 semiconductors (OSCs) can be doped to facilitate increased 39 charge transport^{7,8} as well as decreased injection or extraction 40 barriers via changes in the Fermi energy.9 However, the 41 undesired diffusion of dopants in organic electronic architec-42 tures has been known to cause serious stability and longevity 43 issues.^{10–12} Controlling dopant diffusion is also an important 44 aspect of doping-induced solubility control (DISC), a new 45 patterning method recently developed which allows the 46 solubility of a conductive polymer to be switched on and off 47 by doping and dedoping, respectively.^{13–15} It has also been 48 demonstrated that DISC is capable of using both vapor and

solution processes to achieve micrometer and submicrometer 49 resolution, respectively, in a method that is compatible with 50 roll-to-roll solution processing.^{13,16} For DISC patterning, the 51 lateral diffusion of the dopant limits the ultimate resolution of 52 the pattern. Therefore, for both the stability of completed 53 organic devices and for fabrication of these devices using 54 solution compatible methods, understanding the dynamics of 55 the dopants is extremely important to further application 56 development.

The measurement of small molecular dopant diffusion is 58 particularly challenging because dopants are present in trace 59 quantities and composed of the same light elements (C, H, O, 60 N, S) as the host OSCs. Most of these studies so far have 61

Received: March 30, 2017 **Revised:** May 31, 2017

62 focused on the diffusion of dopants from one layer to another 63 in a vertical layer stack. These samples are prepared by either 64 vacuum evaporating or laminating doped and undoped layers of 65 materials. Secondary ion mass spectrometry (SIMS) is by far 66 the most commonly used technique for diffusion studies due to 67 its high spatial and depth resolution in differentiating 68 molecules. A recent study shows that SIMS can be utilized to 69 monitor the diffusion of 2,3,5,6-tetrafluoro-7,7,8,8-tetracyano-70 quinodimethane (F4TCNQ), which is a widely used p-type 71 molecular dopant, in the hole transport layer of OLEDs.¹⁷ The 72 SIMS signal from the fluorine atoms gives a unique isotopic 73 label. Similar work using SIMS was also performed more 74 recently on the diffusion of the soluble p-dopant molybdenum 75 tris[1-(methoxycarbonyl)-2-(trifluoromethyl)-ethane-1,2-di-76 thiolene] $(Mo(tfd-CO_2Me)_3)$ into polymer and polymer blend 77 films using the Mo label.¹⁸ Likewise, excellent work was $_{78}$ conducted by Treat et al. where the diffusion of phenyl-C₆₁-79 butyric acid methyl ester (PCBM) in poly(3-hexylthiophene-80 2,5-diyl) (P3HT) was quantified using SIMS and a deuterium-81 labeled PCBM (d-PCBM).¹⁹ The same group also quantified 82 the lateral diffusion of *d*-PCBM in P3HT using dynamic SIMS, 83 yielding a one-dimensional diffusion coefficient using Fick's 84 second law.²⁰ However, SIMS is a destructive technique, and 85 additional etching steps are needed. In addition, a unique atom 86 or isotopic labeling is required in order to use SIMS. Other 87 measurement techniques, such as X-ray photoelectron spectroscopy (XPS), ultraviolet photoelectron spectroscopy (UPS), 88 89 neutron/X-ray/visible light scattering, and photoluminescence, 90 have also been used to measure the diffusion of dopants in 91 layered samples.^{21–29} Most of these studies reported whether 92 dopants diffused beyond their original layer and at which temperature threshold. Although some were able to quantify 93 94 dopant concentration at an interface or surface, only one study 95 was able to measure diffusion as a function of distance and time. 96 Thus, Fischer et al. quantified the temperature dependence of 97 the diffusion coefficient in a vertical geometry using a time-98 resolved optical technique.²⁴ Finally, a series of articles have 99 reported the observation of dopant diffusion (or lack thereof) 100 by creating bi- or trilayers and subsequently measuring changes 101 in the electronic properties of layer stack.³⁰⁻³⁴ For these 102 electronic measurements, the movement of dopants is inferred but cannot be quantified. 103

Previous studies have shown that dopant molecules can 104 105 diffuse in a small molecule or polymer OSC host. It is known 106 that larger dopant molecules diffuse more slowly but also have 107 been shown to disturb the packing of the OSC and thereby to 108 increase disorder in the local density of states.³⁵ Very few studies have quantified the diffusion rate of dopants. It has been 109 assumed that the onset of diffusion occurs as a result of heating 110 to above the glass transition temperature (T_g) of host OSC. 111 However, one study showed that F4TCNQ diffuses at room 112 113 temperature in P3HT²³ and another showed that PCBM diffuses into P3HT at 50 °C, which is below the T_{σ} of P3HT.¹⁹ 114 To the best of our knowledge, no molecular level studies have 115 116 looked into the mechanism for dopant binding or investigated 117 how the properties of the host material affect the onset of 118 dopant diffusion.

In this article, we investigate the temperature-dependent microscopic and macroscopic dynamics of the soluble p-type molecular dopant tetrafluoromethyloxycarbonyltricyanoquinodimethane (F4MCTCNQ) in P3HT. F4MCTCNQ is chosen due to its superior solubility/miscibility and thereby thereby there are a solution of the solution matrix of the solution of the solution of the solution of the solution matrix of the solution of the sol chemical structures of the materials investigated in this paper 125 are shown in Figure 1. F4MCTCNQ was also chosen because 126 f1



Figure 1. Chemical structures of F4TCNQ, F4MCTCNQ, P3HT, *d*-P3HT, and the simplified P3HT analogue (M8) investigated in this work.

its methyl group carries three identical protons that can be used 127 as a unique isotopic label in neutron scattering experiments. 128 Quasi-elastic neutron scattering (QENS) is a well-established 129 nondestructive technique to probe dynamic processes in soft 130 matter.³⁷ In the first part of this article, we use QENS to 131 measure the dynamics of F4MCTCNQ at the molecular scale 132 in a sample of fully deuterated P3HT (d-P3HT). The protons 133 on the F4MCTCNQ are the only hydrogen sites in the system, 134 and ¹H has a much larger inelastic scattering cross section than 135 any other isotope in the sample, allowing us to track the 136 temperature-dependent motions of the dopant molecule only. 137 The diffusion rate and jump distance for various microscopic 138 motions are measured, and the motions are assigned with the 139 aid of density functional theory (DFT) calculations. In the 140 second part of the article, we use confocal fluorescence 141 spectroscopy to spatially and temporally track the 142 F4MCTCNQ concentration in patterned P3HT films, which 143 allows us to quantify the macroscopic diffusion of the dopant. 144 We show that macroscopic diffusion is principally dependent 145 on the minority concentration of neutral dopant molecules in 146 the film. This multiscale investigation gives an unprecedented 147 mechanistic insight into dopant diffusion in OSCs. 148

EXPERIMENTAL SECTION

149

Materials. P3HT (regioregular >98%, $M_n = 54-75$ kDa), 150 chlorobenzene, deuterated chlorobenzene, and molybdenum oxide 151 (MoO₃) were purchased from Sigma-Aldrich. *d*-P3HT (regioregular 152 >97%) and F4MCTCNQ were synthesized using the methods 153 reported in the literature.^{36,38} All chemicals were used as received 154 unless otherwise indicated. PELCO TEM grids (1GN400, nickel) were 155 purchased from Ted Pella, Inc. 156

Sample Preparation and Measurements. *Neutron Scattering.* 157 To prepare the blend sample for QENS measurements, solutions of 158 F4MCTCNQ and *d*-P3HT in deuterated chlorobenzene were first 159 mixed to achieve a 17% mole fraction of dopant to polymer. The blend 160 solution was then drop-cast onto a clean glass slide. The sample was 161 placed into a vacuum chamber to completely evaporate the solvent. 162 The resulting blend layer was scraped from the glass slide using a clean 163 razor blade. Both blend and neat samples were ground to a power 164 using a mortar and pestle for 15 min and then weighed prior to being 165 loaded into a 0.5 mm thick flat aluminum container for further 166 analysis. 167

QENS measurements were performed on the backscattering 168 spectrometer called BASIS at the Spallation Neutron Source (SNS) 169 170 at Oak Ridge National Laboratory (ORNL) with an energy resolution 171 of 3.5 μ eV (full width at half-maximum, fwhm), a dynamic range of 172 ±170 μ eV, and a scattering wavevector Q range from 0.3 to 2.0 Å^{-1.39} 173 The elastic scans were obtained from 50 to 375 K at a heating rate of 1 174 K/min, and the high statistical QENS spectra were recorded at 273, 175 323, and 373 K. The spectra collected at 50 K were used as the 176 instrument resolution function, and all the data were corrected for 177 detector efficiency using a vanadium standard.

Confocal Spectroscopy. To prepare samples for the macroscopic 178 179 diffusion measurements, P3HT films of ~50 nm thickness were first 180 obtained by spin-coating 10 mg/mL P3HT from a chlorobenzene 181 solution onto clean glass substrates (cleaned in ultrasonic baths of 182 acetone, 5% Mucasol detergent, and deionized water, followed by 183 drying with nitrogen and exposure to UV/ozone for 30 min). A TEM grid (400 mesh; hole width 38 μ m; bar width 26 μ m) was 184 185 subsequently placed onto the P3HT film using Kapton tape. Next, a 186 2.2 nm F4MCTCNQ layer was deposited onto the P3HT film using 187 an MBraun thermal evaporator at a deposition rate ~0.1 Å/s at ~80 °C, followed by the deposition of 50 nm MoO₃ and 100 nm Ag, also 188 189 via evaporation. Prior to deposition, the evaporation chamber was pumped down to a pressure of 5×10^{-6} mbar. Finally, the samples 190 were encapsulated in an epoxy resin under a glass coverslip to ensure 191 192 complete isolation from O_2 and H_2O . All fabrication steps were 193 performed inside a nitrogen glovebox.

The confocal fluorescence (FL) microscope images were obtained on a Zeiss LSM 700 microscope equipped with a 488 nm laser and a 20× objective with 0.6 numerical aperture. These confocal FL images 197 (16-bit, 2048 × 2048 pixels of 320 μ m size) were captured at the same 198 position before and after annealing at various temperatures (298, 323, 199 348, and 373 K) for various time steps by tracking the position of the 200 marked center of the TEM grid.

Data Analysis. Neutron Data Analysis. For the data analysis, the 201 202 dynamic range was from -120 to $+120 \ \mu eV$. The relevant momentum transfer range in this study was limited to 0.5-1.3 Å⁻¹ given the signal-203 204 to-noise ratio, the flat-plate geometrical limit, and the uncertainty of 205 the background subtraction. DAVE software was used for data 206 reduction and analysis.⁴⁰ The neat *d*-P3HT spectra were subtracted 207 from the F4MCTCNQ doped d-P3HT spectra for all the QENS data 208 analyses to ensure that the signal from F4MCTCNQ could be isolated. In general, the experimentally observed scattering intensity 2.09 210 $S_{exp}(Q,\omega)$ at each Q and temperature is expressed as the convolution 211 of the model function $S_{\text{model}}(Q,\omega)$ and the resolution function $R(Q,\omega)$, 212 plus a linear background term $B(Q,\omega)$:⁴¹

$$S_{\text{exp}}(Q, \omega) = S_{\text{model}}(Q, \omega) \otimes R(Q, \omega) + B(Q, \omega)$$
(1)

214 Here, the model function can be separated into an elastic component 215 (a delta function $\delta(\omega)$) and a quasi-elastic contribution $S_{\text{oe}}(Q_i\omega)$:

$$S_{\text{model}}(Q, \omega) = \text{DWF}(Q)[A_0(Q)\delta(\omega) + (1 - A_0(Q))S_{\text{qe}}(Q, \omega)]$$
216
(2)

217 where DWF(Q) is the Debye–Waller factor due to the vibrational 218 motions and $A_0(Q)$ is the elastic incoherent structure factor (EISF) 219 defined as the fraction of elastic scattering. The proposed expression 220 for $S_{qe}(Q,\omega)$ will be addressed in detail in the Results and Discussion 221 section.

Macroscopic Diffusion Model. The location of dopants within the 222 223 P3HT films was determined by measuring the fluorescence intensity of 224 P3HT as a function of position using a confocal microscope and 488 225 nm excitation. The concentration of the dopants was determined by comparing to a calibration curve of fluorescence intensity in P3HT as a 226 227 function of known doping density and assuming that there is no 228 vertical doping gradient (reasonable for diffusion over μ m in an ~50 229 nm thick film). A two-dimensional two-species (2D2S) diffusion 230 model was developed to fit the 2D doping profiles. The two diffusing species in this model were treated using a Langmuir isotherm 231 232 equilibrium,⁴² which was precedented for P3HT/dopants in a previous 233 publication.¹⁴ In terms of detailed fits, the 2D concentration profile 234 images were initially evolved based on the above-mentioned diffusion 235 model and then evolved using a stepwise solution to Fick's second law.

The finite-difference method using a forward time-centered difference 236 scheme was used to evolve the model in time. After each profile was 237 evolved one time step into the future, the ratio of doped to undoped 238 F4MCTCNQ was re-equilibrated as a function of position and 239 concentration using the Langmuir equilibrium. In this rapid 240 equilibration, it is assumed that the reaction rate between two species 241 is much faster than the rate of diffusion. In doing so, the diffusion 242 coefficients for the two species could be treated as fitting parameters 243 and optimized based on minimization of the sum of squared residuals. 244

Computational Method. An eight-unit methyl-substituted oligo- 245 thiophene (M8) oligomer was used as a model for a P3HT chain. The 246 oligomer was optimized in isolation at the B3LYP/6-31G* level of 247 theory with Grimme D2 dispersion correction (B3LYP-D).^{43,44} In the 248 absence of a dielectric medium, two of these optimized oligomers were 249 then placed at two distinct intermolecular separations (6.30 and 6.96 250 Å), and a F4MCTCNQ molecule was placed midway between and 251 parallel with the two M8 chains. These two intermolecular separations 252 were chosen based on the optimized M8/F4TCNQ system.⁴⁵ The 253 chains were fixed while the F4MCTCNQ was free to relax during a 254 geometry optimization at the B3LYP-D/6-31G* level. A dielectric 255 constant of 3.0 was used as it has been shown to be appropriate for 256 organic semiconductors.^{46–49} The lowest energy geometry was found 257 to be that with the ring of the F4MCTCNQ stacked over the central 258 inter-thiophene bond and the ester methyl group on the opposite side 259 of the M8 chain methyl groups. The two separations were chosen as 260 twice the distance between a single M8 chain and F4MCTCNQ 261 molecule from an unconstrained geometry optimization using B3LYP 262 with and without the dispersion correction, respectively. The two M8 263 oligomers were constrained in the optimization of the full system to 264 prevent unphysical wrapping of the chains around the F4MCTCNQ 265 molecule. These two intermolecular separations were used to give an 266 estimate of possible variations in the energy barriers to rotation and 267 translation due to variations in the interchain separation. Since it was 268 found that introducing solvent into this system does not affect the 269 intermolecular separation, the values from systems without the 270 dielectric continuum were used. 271

To calculate the potential energy barrier for F4MCTCNQ 272 translation along the polymer backbone axis, atoms in the M8 chains 273 were fixed while the F4MCTCNQ was free to relax in a set of 274 optimizations, subject to a position constraint applied by fixing one 275 atom in the F4MCTCNQ. This set of translation optimizations 276 spanned up to two thiophene units.

To calculate the potential energy barrier for F4MCTCNQ methyl 278 rotation between the two M8 chains for both interchain separations, all 279 atoms in the system excluding the ester methyl hydrogens were fixed, 280 and these hydrogens were free to relax in a set of optimizations subject 281 to dihedral angle constraints spanning 120°. In addition, calculations 282 on an isolated F4MCTCNQ molecule were performed, in which no 283 atom positions were fixed. Solvent was used in these calculations. 284 Similar calculations were performed to calculate the barrier for the 285 F4CTCNQ methoxycarbonyl rotation, but only for the isolated 286 molecule and for the molecule between M8 chains separated by 6.96 287 Å. 288

RESULTS AND DISCUSSION

Part One: Microscopic Dynamics of Molecular ²⁹⁰ **Dopant.** *Elastic Incoherent Neutron Scattering.* The micro- ²⁹¹ scopic dynamics of the F4MCTCNQ molecule were ²⁹² investigated using neutron scattering. To determine the ²⁹³ temperature range at which the dynamic transitions of protons ²⁹⁴ take place and become observable on BASIS, facile elastic ²⁹⁵ incoherent neutron scattering experiments were performed ²⁹⁶ first. The elastic scans of the F4MCTCNQ doped d-P3HT ²⁹⁷ blend measured from 50 to 375 K at various Q values are ²⁹⁸ shown in Figure 2a. Generally, the elastic intensity decreases ²⁹⁹ f2 monotonically with temperature due to Debye–Waller thermal ³⁰⁰ factors, while any observed dynamics of the species lead to ³⁰¹ further reduction of the elastic intensity.⁵⁰ The divergence from ³⁰²

289



Figure 2. (a) Normalized elastic scattering intensity at various *Q* and (b) mean-square displacement (MSD) as a function of temperature for the F4MCTCNQ doped *d*-P3HT blend.

303 the monotonic decrease as a function of increasing temperature 304 indicates the onset of molecular motions for hydrogen sites in 305 this system.

To enhance conceptual understanding of this dynamic 306 process, the temperature dependence of the proton mean-307 square displacement (MSD) determined from analysis of the 308 elastic scans (Supporting Information section 2) was 309 310 investigated. From Figure 2b, the MSD values do not vary much at temperatures below 180 K because only some local 311 vibrations are expected to occur at such low temperatures. As 312 the temperature increases, two deviations (changes in slope) 313 are observed: one at about 180 K and another at 254 K. We 314 attribute the first deviation at ~180 K to the activation of 315 ³¹⁶ methyl group rotations, which are well-known to occur ³¹⁷ between 100 and 220 K.⁵¹⁻⁵³ On the other hand, the second deviation at ~254 K implies that another dynamic transition 318 319 takes place in this temperature region. This transition could not be assigned unambiguously without further analysis. 320

³²¹ *Quasi-Elastic Neutron Scattering (QENS).* To assign the ³²² dynamic transition starting at ~254 K, the high statistical ³²³ scattering data were collected at temperatures above 254 K in ³²⁴ order to capture this transition, and detailed analyses were ³²⁵ performed. As mentioned earlier, neat *d*-P3HT spectra were ³²⁶ also collected at the corresponding temperatures and subtracted ³²⁷ from the spectra of the doped samples (Supporting Information ³²⁸ section 3). In this manner, any changes of the scattering spectra ³²⁹ explicitly result from the dynamics of the hydrogen sites in the ³³⁰ dopants. Representative neutron scattering spectra of subtracted 331 doped *d*-P3HT at Q = 0.9 Å⁻¹ at three temperatures (273, 332 323, and 373 K) are shown in Figure 3. The experimental 333 f3



Figure 3. Representative BASIS spectra of subtracted doped *d*-P3HT at Q = 0.9 Å⁻¹ at 273 K (orange), 323 K (dark cyan), and 373 K (purple), along with the resolution function obtained at 50 K (gray dashed line). Both unnormalized (top) and normalized (bottom) data with respect to their intensity at $\hbar \omega = 0$ are shown.

resolution functions collected at 50 K are also plotted. In $_{334}$ general, the spectra can be described as a superposition of two $_{335}$ contributions: (1) a Gaussian distribution corresponding to the $_{336}$ elastic portion and (2) a broad shoulder in the quasielastic $_{337}$ region. The broadening of the quasi-elastic components with $_{338}$ increasing temperature seen from Figure 3 (it is more obvious $_{339}$ in their normalized spectra) provides clear evidence of the $_{340}$ occurrence of dynamic processes, which is consistent with the $_{341}$ MSD results from elastic scans. Similar broadening features are $_{342}$ also observed at other Q ranges (Supporting Information $_{343}$ Figures S2 and S3).

In order to extract diffusive information from QENS, the 345 experimental spectra were fit using a model function discussed 346 in the Data Analysis section (eqs 1 and 2). Again, since two 347 dynamic processes are observed above 254 K, two stretched 348 exponential relaxation (or Kohlrausch–Williams–Watts, 349 KWW) functions were included in the quasi-elastic term 350 $S_{qe}(Q,\omega)$ in eq 2, expressed as 351

$$S_{\rm qe}(Q, \omega) = P \int \exp\left[-\left(\frac{t}{\tau_1}\right)^{\beta_1}\right] \exp\left(i\frac{E}{\hbar}t\right) dt + (1-P) \int \exp\left[-\left(\frac{t}{\tau_2}\right)^{\beta_2}\right] \exp\left(i\frac{E}{\hbar}t\right) dt$$
(3) 352

where τ and β are the characteristic time and stretching 353 exponent, respectively, while the subscripts 1 and 2 represent 354 different dynamic components. The parameter *P* is defined as 355 the fraction of the first Fourier-transformed stretched 356 exponential relaxation term. *E* is the energy change of the 357 neutrons, and \hbar is the reduced Planck constant. Using this 358 model, we expect that the rotations of methyl group can be 359 captured by the faster stretched exponential term (set $\tau_1 < \tau_2$), 360 f4

385

4

f5



Figure 4. (a) Relative weight of the first KWW term (eq 3) as a function of Q at various temperatures. (b) Representative QENS spectrum and fit with a single KWW function of subtracted doped *d*-P3HT at a Q of 0.9 Å⁻¹ and temperature of 273 K. (c) Representative QENS spectrum and fits with two KWW functions of subtracted doped *d*-P3HT at a Q of 0.9 Å⁻¹ and temperature of 373 K.

³⁶¹ while the other motion of the hydrogen sites can be captured ³⁶² using the slower stretched exponential term.

We performed fits to all the spectra and found that a single 363 364 stretched exponential term was sufficient for satisfactory fits of 365 the 273 K data (the fit weight P values converge to unity, 366 shown in Figure 4a), but at 323 and 373 K, fitting $S_{ab}(Q,\omega)$ 367 required incorporating two KWW terms for all data except Q =368 1.3 Å⁻¹. Examples of these one and two KWW fits are 369 presented in Figures 4b and 4c, respectively. As can be seen, the 370 fits capture the experimental spectra reliably over the entire 371 energy range in both cases. We attribute the successful fit using 372 only one KWW term at 273 K to the fact that the dynamics of methyl group rotations dominate at a temperature that is just 373 374 slightly above 254 K. Reduced P values, therefore, are expected once temperature increases, which is exactly what is observed. 375 The P values follow a clear decreasing trend upon heating. At 376 377 373 K, an average of \sim 20% of the total inelastic hydrogen scattering is due to the second dynamic process. The full 378 experimental spectra and fits at all conditions are shown in 379 Supporting Information Figures S4-S6. 380

³⁸¹ The successful fits of QENS spectra using a stretched ³⁸² exponential model allow us to further explore the nature of the ³⁸³ dynamic motions. Normally, the mean relaxation time $\langle \tau_i \rangle$ of ³⁸⁴ process *i* is extracted by

$$\langle \tau_i \rangle = \frac{\tau_i}{\beta_i} \Gamma \left(\frac{1}{\beta_i} \right) \tag{4}$$

386 where τ_i and β_i are fitting parameters from the stretched 387 exponential model in eq 3, and $\Gamma(x)$ is the gamma function. 388 The Q^2 dependence of calculated values in the form of $\langle \tau_i \rangle^{-1}$ is 389 plotted in Figure 5. The comparison of absolute $\langle \tau_i \rangle^{-1}$ values 390 clearly implies that two distinct diffusive motions are observed 391 as expected. In addition, Figure 5 shows that for both fast and 392 slow processes the $\langle \tau_i \rangle^{-1}$ values exhibit a quadratic behavior at 393 low Q values and then a saturation trend at high Q values. This 394 nonlinear variation of $\langle \tau_i \rangle^{-1}$ as a function of Q^2 indicates 395 noncontinuous diffusion of hydrogen sites,⁵⁴ which is 396 particularly true for the case of methyl group rotations. 397 Therefore, a well-studied jump diffusion model (Singwi– 398 Sjölander model) was employed to describe both dynamic 399 processes, given by⁵⁵

$$\frac{1}{\langle \tau \rangle} = \frac{DQ^2}{1 + DQ^2 \tau}$$
(5)



Figure 5. Inverse of mean relaxation time (symbols) of the fast process (a) and slow process (b) obtained from KWW fits as a function of Q^2 at various temperatures. The corresponding solid lines are the best representative fits to the jump diffusion model.

where τ is the residence time between jumps of a diffusive 401 particle and *D* is the diffusion coefficient. The mean-square 402 jump length can also be determined by $\langle L^2 \rangle = 6D\tau$ in this 403 model. As can be seen from the solid lines in Figure 5, these fits 404 yield a reasonable agreement with the data, and the resulting 405 diffusion parameters at each temperature are summarized in 406 Table 1. 407 ti

By comparing the two dynamic processes, we notice that the 408 observed diffusion coefficients of the faster process are 409 approximately 1 order of magnitude greater than those of the 410 f6

t2

f6t2

Table 1. Model Parameters Obtained from Fits of the Q^2 -Dependent $\langle \tau_1 \rangle^{-1}$ and $\langle \tau_2 \rangle^{-1}$ Using a Jump Diffusion Model

	temp (K)	diffusion coeff D $(\times 10^{-6} \text{ cm}^2/\text{s})$	residence time $ au$ (ps)	mean jump length $(\langle L^2 \rangle)^{1/2}$ (Å)
fast process	273	3.13 ± 1.05	34.38 ± 8.50	2.54 ± 0.74
	323	5.90 ± 0.90	20.23 ± 1.62	2.68 ± 0.31
	373	10.17 ± 0.89	12.13 ± 0.64	2.72 ± 0.19
slow process	273			
	323	0.86 ± 0.12	100.15 ± 18.01	2.27 ± 0.26
	373	1.68 ± 0.15	46.57 ± 10.22	2.17 ± 0.34

411 slower process. Also, the residence time values are factor of 5 412 lower at the same temperature for the faster process. The mean 413 jump lengths, on the other hand, are comparable at 2–3 Å. 414 With respect to the individual process, the rotational diffusion 415 coefficient of the methyl group increases from 3.13×10^{-6} 416 cm²/s at 273 K to 10.2×10^{-6} cm²/s at 373 K. Although we are 417 unable to capture the slower motion process at 273 K, the 418 diffusion coefficient of the second dynamic process still exhibits 419 a 2-fold increase from 323 to 373 K.

Further analysis of the temperature dependence of the diffusion coefficients and residence time was performed. Here, Az2 Arrhenius behavior was assumed to fit the data (seen in Figure Az3 6) and an activation energy E_A associated with each process was



Figure 6. Diffusion coefficient (top) and residence time (bottom) as a function of 1000/T for the first dynamic (purple circles) and second dynamic (dark cyan squares). The corresponding solid lines are fits to an Arrhenius behavior.

424 determined, which is summarized in Table 2. As can be seen, 425 the activation energy of the slow process (\sim 14 kJ/mol) is 426 slightly higher than that of the fast process (\sim 10 kJ/mol).

> Table 2. Activation Energy E_A (kJ/mol) Obtained from Arrhenius Fits of Diffusion Coefficient D and Residence Time τ Associated with Dynamics of Hydrogen Atoms^{*a*}

	$E_{\rm A}$ from diffusion coeff (kJ/mol)	<i>E</i> _A from residence time (kJ/mol)	$E_{\rm A}$ from elastic scattering (kJ/mol)
fast process	10.4 ± 0.5	9.6 ± 0.8	7.0 ± 0.3
slow process	13.4 ± 2.8	14.9 ± 4.4	

^{*a*}The E_A value derived from elastic scattering is also shown for comparison.

Generally, the E_A of methyl group rotation is reported to be $_{427}$ 6–12 kJ/mol for small molecules and polymers. $_{51,59-61}^{51,59-61}$ $_{428}$ To more accurately determine the E_A for this specific sample, 429 we first performed density-functional theory (DFT) calcu- 430 lations on an isolated dopant molecule. The F4MCTCNQ 431 methyl hydrogens were rotated through 120° in 30° increments 432 with the rest of the atoms in the system fixed. As can be seen in 433 Figure 7b (orange curve), the energy barrier to methyl rotation 434 f7 occurs at a 60° offset from the global minimum energy 435 configuration. The calculated barrier of ~4 kJ/mol is roughly 436 consistent with the experimental barrier of ~10 kJ/mol. 437 However, since this calculated energy barrier could also be 438 affected by the P3HT chains, the barrier in the presence of fixed 439 chains was investigated, and the actual barrier is likely to be 440 between these values. We assumed here, based on the 441 published geometry of P3HT in the presence of 442 F4TCNQ^{62,63} that a F4MCTCNQ molecule is sandwiched 443 between two M8 chains as seen in Figure 7a. Given the possible 444 sensitivity to the M8 chain distance, calculations at two distinct 445 intermolecular separations (6.30 and 6.96 Å) were performed. 446 Figure 7b shows that the magnitude of the rotational barrier 447 increases slightly with increasing interchain separation. In both 448 cases, the barrier to methyl rotation was determined to be 8-10 449 kJ/mol, yielding a linear distance of 1.8 Å and arc length of 2.1 450 Å for moved hydrogen atom in this rotation. Given the typical 451 errors of this type of calculation of at least 1 kcal/mol (~4 kJ/ 452 mol), this result is consistent with the QENS experiments. The 453 value of the jump length, however, is a slightly lower than that 454 determined experimentally, which could be due to the 455 approximations made in the calculations, which were performed 456 in vacuum at zero temperature and with fixed positions for the 457 thiophene backbone. The good match between QENS 458 experiments and DFT calculations validate the assignment of 459 the fast dynamic process to the methyl rotation of 460 F4MCTCNO.

With regard to the slower observed motion, we can 462 confidently rule out based on DFT calculations that it is due 463 to the translational diffusion of F4MCTCNQ along the 464 polymer backbone. Figure 7c shows the calculated energy 465 profile for the translation of a F4MCTCNQ molecule for one 466 thiophene unit (\sim 4 Å) in each direction from its lowest energy 467 position for the 6.96 Å separation and half a thiophene unit (~ 2 468 Å) in each direction for the 6.30 Å separation. The closer 469 interchain distance between thiophene chains yields substan- 470 tially higher energy barriers to translation in both directions 471 (~60 kJ/mol). However, even calculation using a larger 472 interchain separation still yields a large energy barrier of ~ 30 473 kJ/mol. Extending the translation for the larger interchain 474 distance to one thiophene unit demonstrates local minima at 475 every half-thiophene unit translation, corresponding to 476 F4MCTCNQ ring stacking directly over a thiophene ring. In 477 any case, the calculated translational barriers are considerably 478 higher than what were measured experimentally from QENS 479 $(\sim 14 \text{ kJ/mol}).$ 480

Given the small activation energy and jump length of the 481 slower motion, it seems more reasonable that this motion is 482 associated with a localized motion that is internal to the 483 F4MCTCNQ molecule. Closer inspection of the translation 484 calculations in Figure 7c implies that low-energy vibrational 485 motion around the position minima may be responsible for the 486 slower motion. In other words, the whole F4MCTCNQ 487 molecule might move back and forth within a potential well. 488 However, further calculations using a simple harmonic 489



Figure 7. (a) Sandwich geometry comprising two M8 oligomers and a F4MCTCNQ molecule. The corresponding diffusive motions of each DFT calculations are also described schematically. (b) Rotation energy of the ester methyl hydrogens through 120° for two interchain separations and the isolated dopant molecule. Inset: calculated hydrogen displacements for the F4MCTCNQ methyl rotation. (c) Translation energy of the monoester molecule over one thiophene unit (6.30 Å interchain separation) and two thiophene units (6.96 Å interchain separation). The dashed gray line is the quadratic fit in order to yield a harmonic potential. (d) Rotation energy of ester group for the isolated dopant molecule (top) and a 6.96 Å interchain separation (bottom).

490 oscillator model, shown as the dashed gray line in Figure 7c, 491 enable us to disprove this hypothesis. The calculated energy 492 spacing between vibrational energy levels (~2.5 meV, 493 Supporting Information section 6) is significantly higher than 494 the energy spread of the QENS measurements (on the order of $_{495}$ μ eV), indicating that it is impossible to capture this dynamic 496 motion in the energy window on BASIS. Other possible sources of the slower motion are the localized rotation of methoxy-497 carbonyl group $(-COOCH_3)$ or the rotation of methoxy group 498 $(-OCH_3)$ in F4MCTCNQ. We can safely rule out the 499 500 methoxy rotation assignment because of its high rotation barrier (>50 kJ/mol) in the literature calculations even for an 501 isolated molecule.^{64,65} On the other hand, DFT calculations of 502 the methoxycarbonyl rotation of an isolated F4MCTCNQ 503 ₅₀₄ molecule (Figure 7d) yields a barrier of ~ 16 kJ/mol, which is 505 consistent with the QENS results. Although the calculation on 506 the sandwich geometry (6.96 Å interchain separation) produces 507 a significantly higher barrier, a lower value wound be expected if 508 the relaxation of the chains were allowed. In addition, given the $_{509}$ fact that the inferred $E_{\rm A}$ values of the slower motion are in good 510 agreement with reported values for methoxycarbonyl rota-511 tions,⁶⁵ we tentatively assign the slower motion to this rotation 512 of the dopant.

513 Additional insight, particularly into the geometry of the 514 motion, can also be obtained by investigating the elastic 515 incoherent structure factors (EISF). The Q-dependent EISF 516 values at three investigated temperatures are plotted in Figure 517 8. A 3-fold jump model has been widely used to describe 518 methyl group rotations:^{66,67}

f8

f8



Figure 8. Elastic incoherent structure factor (EISF) obtained for subtracted doped *d*-P3HT as a function of Q at 273 K (orange), 323 K (dark cyan), and 373 K (purple). The solid lines are product fits to a rotational jump diffusion on a circle model (set N = 30 in eq 7) and a methyl group 3-fold jump model. The 3-fold jump model of the methyl group is also shown for comparison. Inset: obtained EISF fit parameters using the product diffusion model in form of eq 7.

$$\text{EISF}_{\text{methyl}}(Q) = f_1 + (1 - f_1) \left(\frac{1}{3} [1 + 2j_0(\sqrt{3} Qr)] \right)$$
(6) 519

where j_0 is the spherical zeroth-order Bessel function, f_1 is the s20 fraction of immobile protons, and *r* is the radius, which was set s21 to 1.03 Å based on the known geometry of methyl group.⁶⁸ s22 However, the fit of methyl group 3-fold jumps (dashed line) s23 shows significant deviation from the EISF values and is also s24



Figure 9. Schematics of (a) sample preparation and (b) resulting sample geometry and confocal experimental setup. (c) The confocal fluorescence images of patterned films before (top row) and after (bottom row) annealing at various temperatures (298, 323, 348, and 373 K) for time *t*.

⁵²⁵ greater in all cases. This clearly indicates that the methyl group ⁵²⁶ rotation is not the only component that contributes to the ⁵²⁷ QENS signal even at 273 K and that the slower motion must ⁵²⁸ also be taken into account, which again is in good agreement ⁵²⁹ with our observations from the elasitic QENS data. Therefore, ⁵³⁰ we rewrite eq 6 by including a well-established rotational ⁵³¹ diffusion model, where it is assumed that the scattering center ⁵³² jumps between N equivalent points on a circle: ^{69,70}

$$EISF(Q) = \left[f_1 + (1 - f_1) \left(\frac{1}{3} [1 + 2j_0(\sqrt{3} Qr)] \right) \right] \\ \times \left[f_2 + (1 - f_2) \left(\frac{1}{N} \sum_{i=1}^N j_0 \left(2QR \sin \frac{\pi i}{N} \right) \right) \right]$$
(7)

5

534 where f_2 is the fraction of immobile protons involved in the s35 methoxycarbonyl rotation, R is the circle radius, and r is still set 536 to 1.03 Å. The product form means that the whole methyl 537 group moves during the methoxylcarbonyl rotation. The 538 detailed derivation is shown in Supporting Information section 1. For $N \to \infty$, the limiting case of continuous rotational 539 diffusion on a circle is reached. By means of this improved 540 541 model, the EISF behavior is able to be captured over the whole 542 investigated Q range as seen the solid lines in Figure 8 where N543 is set to 30 (a satisfactory fit is found for $N \ge 2$). In all cases, 544 the parameter f_1 always converges to 0, revealing that all 545 hydrogen sites are involved in the methyl rotations at any given 546 temperature. The parameter f_2 and R, on the other hand, are 547 found to vary slightly and converge with increasing N (Figure 548 S7), the corresponding converged values of which are 549 summarized in the inset of Figure 8. As can be seen, the 550 values of f_2 follow a clear decreasing trend from 0.86 to 0.29 551 upon heating, which is consistent with results from Figure 4 552 that increased scattering from the slower motion is observed as 553 temperature increases. Moreover, the diffusive motion of the slower motion is restricted on a circle of $\sim 2.0-3.7$ Å radius, 554 which is also consistent with the geometry of the methoxy- 555 carbonyl rotation, where the distance from the center of methyl 556 hydrogen sites to the rotation axis is estimated to be ~ 1.7 Å. 557

Part Two: Macroscopic Dynamics of Molecular 558 **Dopants.** The previous section demonstrated that two 559 molecular-scale motions of F4MCTCNQ in P3HT can be 560 observed and assigned from QENS experiments. Unfortunately, 561 the energy window of the BASIS spectrometer was insufficient 562 to observe rarer and longer-range diffusive processes that 563 contribute to macroscopic diffusion. Therefore, laser scanning 564 confocal microscopy (LSCM) was used to quantify the 565 macroscopic diffusion of F4MCTCNQ in P3HT. 566

In order to make this macroscopic measurement possible, a 567 specific sample architecture was required, which was derived 568 from our recently developed dopant-induced solubility control 569 (DISC) patterning method.¹³ Schematics of the sample 570 preparation and resulting sample geometry are shown in 571 Figures 9a and 9b, respectively, and a detailed sample 572 f9 description is provided in the Experimental Section. It is 573 worth emphasizing that it is very important to insert a MoO₃ 574 (or similarly performing oxide) layer between the patterned 575 film and the Ag layer in this experiment because the high work 576 function MoO_3 (WF = 6.8 eV) can act as an insulator to 577 prevent dopant-metal reactions.⁷¹ It is also well-known that 578 the fluorescence (FL) of P3HT is quenched by the presence of 579 dopants or any electron-accepting molecules.⁷²⁻⁷⁵ By mapping 580 out the FL intensity from LSCM and combining with a 581 calibration curve (Figure S8), we were able to track the dopant 582 concentration in the P3HT film spatially and temporally. We 583 assumed here that there was no vertical concentration gradient 584 of dopants, and thus all diffusion occurs within the two- 585 dimensional plane of the P3HT film. This assumption is valid 586 considering that the P3HT film is only 50 nm thick and 587 fluorescence is measured spatially over $\sim 1-10 \ \mu m$. Figure 9c 588



Figure 10. (a) LSCM fluorescence images of patterned F4TCNQ/P3HT films before (left) and after (right) annealing at 348 K for 50 min. (b) The 2D doping concentration maps converted from LSCM fluorescence images using a calibration curve at t = 0 (left) and t = 50 min (right). (c) The 2D doping concentration maps for a two-dimensional one-species (2D1S) model fit (left) and its absolute residuals (right). (d) Cross-sectional doping concentration profiles through images in (b) and (c) (indicated by white dashed lines) using a 2D1S diffusion model. (e) The 2D doping concentration profiles through images in (b) and (c) (indicated by white dashed lines) using a 2D1S diffusion model. (e) The 2D doping concentration profiles through images in (b) and (e) (indicated by white dashed lines) using a 2D2S diffusion model.

show examples of confocal FL images before (top) and after 589 (bottom) annealing for a time *t* at four different temperatures. 590 The narrowing gaps in the FL intensity clearly indicate that the 591 F4MCTCNQ diffuses from the doped (dark) regions to the 592 undoped (bright) regions in the sample over time, as expected. 593 Preliminary analysis based upon the annealing time at the 594 different temperatures suggests that the macroscopic diffusion 595 rate of the dopant is at least 1 order of magnitude higher for 596 every 25 K temperature increase from 298 to 373 K. 597

We quantified the macroscopic diffusion rate and mechanism by by fitting the dopant concentration as a function of lateral oposition and diffusion time. The F4TCNQ/P3HT system was used to validate our models. Initially, we attempted to fit the dot aby using a one-dimensional (1D) Fickian diffusion model that has been used previously to determine the diffusion co4 coefficient of PCBM in a P3HT film,²⁰ the details of which are shown in Figure S9. This 1D model dose not successfully co6 capture the F4TCNQ diffusion behavior. Therefore, we

extended this 1D model to a two-dimensional, one-species 607 (2D1S) diffusion model, which appeared to be more realistic. 608 Images of 60 \times 60 μ m were used for all the fits in order to save 609 computational time. The experimental confocal FL images 610 before and after annealing at 348 K for 50 min and converted to 611 2D doping concentration maps are presented in Figures 10a 612 flo and 10b, respectively, both of which clearly show F4TCNO 613 diffusion. For this 2D1S model, only one diffusive species was 614 taken into account. The model fit and corresponding residuals 615 are shown in Figure 10c. A direct comparison of cross-sectional 616 profiles from these experimental and fitting images (indicated 617 by the white dashed lines in Figures 10b and 10c, respectively) $_{618}$ is also shown in Figure 10d. Again, a single diffusing species 619 model yields an unsatisfactory fit for dopants concentration 620 versus temperature and time, which can be seen from the 621 discrepancy between measured (green line) and fit (red line) 622 doping levels. 623

Macromolecules



Figure 11. (a) A representative F4MCTCNQ diffusion fit at 348 K using a two-dimensional two-species (2D2S) model. 2D doping concentration maps were obtained experimentally from LSCM at t = 0 (top left) and t = 560 min (top right), 2D2S fit (bottom left), and absolute residuals (bottom right). (b) Cross-sectional doping concentration profiles through images in (a) (indicated by white dashed lines) using a 2D2S diffusion model. (c) Cross-sectional doping concentration profiles using a two-dimensional one-species (2D1S) diffusion model. (d) Determined two diffusion coefficients from 2D2S fits as a function of annealing temperatures for F4MCTCNQ (solid symbols) and F4TCNQ (open symbols) as well as corresponding Arrhenius fits. (e) 2D mole ratio profiles of slower (ionized) species to faster (neutral) species at t = 560 min at 348 K for F4MCTCNQ diffusion. (f) Schematic of F4MCTCNQ dopant diffusion derived from 2D2S diffusion model analysis.

We recently showed that an equilibrium between neutral 624 625 F4TCNQ and ionized F4TCNQ exists in sequentially processed doped P3HT films, which can be described by 626 $627 \text{ F4TCNQ} + \text{P3HT} \Rightarrow \text{F4TCNQ}^- + \text{P3HT}^{+.14}$ In the cited 628 study, F4TCNQ was added into the P3HT film using an orthogonal solvent. We showed that the orthogonal solvent 629 could not dissolve or deposit F4TCNQ into the crystalline 630 domains of P3HT, but instead only doped the amorphous 631 domains of the polymer. In addition, we quantified the site 632 density versus doping level for doping of the amorphous P3HT 633 634 using a Langmuir isotherm model. We posit that the same 635 model is very likely to explain the equilibrium for the F4TCNQ 636 dopant in evaporation patterned P3HT films. The Langmuir 637 isotherm model can be expressed as

$$\theta = \frac{C_{\text{ionized}}}{C_{\text{sat}}} = \frac{K_{\text{eq}}C_{\text{neutral}}}{1 + K_{\text{eq}}C_{\text{neutral}}}$$
(8)

638

where θ is the fraction of occupied doping sites in the 639 amorphous domains of the film, C_{ionized} is the film doping 640 concentration, C_{sat} is the saturated film doping concentration, 641 C_{neutral} is the concentration of neutral dopant molecules, and 642 K_{eq} is the equilibrium constant. 643

Given that both neutral and ionized dopants coexist in this 644 sample, we developed a two-dimensional two-species (2D2S) 645 diffusion model assuming (1) that the neutral and ionized 646 species obey the aforementioned Langmuir isotherm equili-647 brium and (2) that the rate of equilibration between the two 648 species is much faster than the rate of diffusion. Since increasing 649 the temperature would shift the equilibrium toward a higher 650 population of neutral dopants, the temperature-dependent K_{eq} 651 $= \exp(-\Delta G^{\circ}/(k_{\rm B}T))$, where ΔG° is taken to be the HOMO- 652 LUMO difference between P3HT and the dopant (i.e., roughly 653 the change in energy of the charge transfer process; 0.23 eV for 654 F4TCNQ and 0.14 eV for F4MCTCNQ³⁶), $k_{\rm B}$ is the constant, 655 656 and T is the temperature. The 2D mapped doping 657 concentration profiles were then fit with a 2D solution to 658 Fick's second law, yielding a diffusion coefficient for each 659 diffusive species (denoted as D_{neutral} and D_{ionized} in order to 660 differentiate them from the diffusion coefficients obtained from 661 QENS). The performed 2D2S fit for the F4TCNQ diffusion at 662 348 K is shown in Figure 10e. By comparing the experimental 663 and fit doping concentration profiles and inspecting the low 664 residual between the data and fit in Figures 10b and 10e, it is 665 clear that this 2D2S model is able to reproduce the lateral diffusion rates and mechanism in a consistent manner. Cross-666 sectional profiles through these images (indicated by the white 667 dashed lines) are shown in Figure 10f and directly compared 668 with the 2D2S fit. A full comparison between 2D1S and 2D2S 669 model fits for F4TCNQ diffusion at other investigated 670 temperatures can be found in Figures S10 and S11. 671

This 2D2S diffusion model was applied to describe the 672 673 diffusion of F4MCTCNQ in the P3HT films. A representative example of the 2D2S fit for F4MCTCNQ at 348 K is shown in 675 Figure 11a, and the cross-sectional doping concentration profiles through these images are provided in Figure 11b. The fits at other investigated temperatures can be found in 677 Figure S12. Similar to the case of F4TCNQ, satisfactory fits can 678 be produced. For consistency, the 2D1S model fits on 679 F4MCTCNQ were also performed, and the resulting cross-680 sectional profiles at the same position are shown in Figure 11c 681 (corresponding 2D images are presented in Figure S13). The 682 683 discrepancy between measured and fit doping levels, especially 684 in the doped region, using the 2D1S model again shows that a 685 two species diffusion model is necessary to fit the diffusion data 686 with sufficient accuracy. Additionally, since the highest doping 687 ratio of ~4.3 mol % in the feature centers is much lower than 688 the reported saturated doping level (>17 mol %),^{36,62} it is 689 reasonable to assume that ionized dopants are the majority of 690 the diffusive species in the system. Also, since the highest 691 doping ratio is below the $C_{\rm sat}$ determined for the amorphous 692 domains of P3HT in the previous study,¹⁴ it is likely that most 693 or all of the dopants are located in and transported through the amorphous domains of P3HT. 694

The resulting two diffusion coefficients of the faster and 695 696 slower species obtained from the 2D2S diffusion model are 697 plotted as a function of the inverse of the annealing temperatures (1000/T) in Figure 11d. As can be seen, two 698 699 distinct diffusion coefficients are observed. The diffusion 700 coefficient of the faster species D_{neutral} is approximately 2–3 701 orders of magnitude greater than that of the slower species $_{702}$ D_{ionized} at the same temperature for both the F4MCTCNQ and 703 F4TCNQ dopants in P3HT. Since the Langmuir isotherm 704 model tracks the number of available doping sites in the P3HT, it is reasonable that the ratio of ionized to neutral dopants 705 706 increases with reduced doping density. The mole ratio of the 707 ionized (slower) to neutral (faster) dopant species is therefore tracked in this diffusion model. A representative 2D map of the 708 ionized/neutral doping ratio for F4MCTCNQ/P3HT at 348 K 709 710 is shown in Figure 11e. It is clear that the concentration of 711 neutral species is orders of magnitude lower than the ionized 712 species (also seen in Figure S14) and that this ratio is position-713 dependent, reflecting the local equilibrium between unfilled 714 P3HT sites and dopants. Additionally, the ratio of ionized to 715 neutral molecules increases as a function of diffusion distance, 716 from \sim 100 in the initially doped region to \sim 500 in the initially 717 undoped region (Figure S14). This increased ratio implies that 718 the neutral dopant can diffuse for longer times in the highly doped part of the sample, where there are fewer free undoped 719 sites on the P3HT. As the dopant leaves the highly doped 720 region, the neutral dopant is presented with a large number of 721 free doping sites and so readily binds. The two-species diffusion 722 processes is illustrated schematically in Figure 11f. 723

Since the temperature-dependent diffusion coefficients 724 follow, within error, Arrhenius behavior (the straight lines in 725 Figure 11d), we are able to determine the activation energy E_A 726 associated with the diffusion process for each species. In our 727 calculations, the $D_{neutral}$ values at 298 K are excluded due to 728 their uncertainties, seen from the large error bars in Figure 11d. 729 The determined E_A values for translational diffusion are listed 730 in Table 3. Interestingly, the E_A values for both neutral and 731 t3

Table 3. Activation Energy E_A (kJ/mol) for Translational Diffusion Obtained from Arrhenius Fits of Diffusion Coefficients ($D_{neutral}$ and $D_{ionized}$) of Neutral and Ionized Dopant Molecules, Respectively^{*a*}

	$E_{A_{neutral}}$ (kJ/mol)	$E_{A_ionized}$ (kJ/mol)
F4MCTCNQ	53.8 ± 1.9	55.5 ± 2.4
F4TCNQ	74.2 ± 7.3	55.9 ± 2.6
^{<i>a</i>} E, values of F4TCN() diffusions are also sh	own for comparison.

ionized F4MCTCNQ (~55 kJ/mol) are consistent with the 732 translational barriers from DFT calculations in Figure 7c 733 (especially for the smaller interchain separation). We note that 734 the jump distance for an ionized dopant depends strongly on 735 the energy landscape. As can be seen from Figure 7c, an open 736 site (no other dopant in the path) and sufficient energy to 737 surmount a 50-60 kJ/mol barrier for hopping to the next site 738 are required for the ionized species to jump into the nearby 739 potential well that is one bond. In addition, Kramer's theory 740 was applied to estimate the diffusion coefficient of the dopant 741 based on the DFT potential energy curves (Supporting 742 Information section 11). The calculation yields a diffusion 743 coefficient for ionized F4MCTCNQ of 1.4 \times 10 $^{-14}$ cm $^2/s$ at 744 298 K, which again falls into the range measured in the confocal 745 experiments. However, the geometry of the DFT calculations 746 depicted a highly simplified ordered P3HT domain, which is 747 likely to different from the amorphous P3HT domain in which 748 the dopant is expected to reside at the doping density used in 749 the experiment. Given this discrepancy between the simulated 750 geometry and likely sample geometry, we speculate that the 751 measured $E_{A \text{ ionized}}$ likely represents an average value and that 752 the actual hopping barriers in amorphous domains will be more 753 heterogeneous than those estimated in the DFT calculations. 754

In addition, we can infer from the diffusion model that the 755 jump distance for the neutral dopant is considerably larger than 756 for the ionized dopant, even though their activation energies are 757 similar. We know that the neutral dopant is not Coulombically 758 bonded to the P3HT. We also know from the comparison of 759 the 2D1S and 2D2S models that the neutral species is needed 760 to explain the bulk diffusion profile as a function of time and 761 temperature. It can be seen that the jump attempt frequency for 762 the neutral and ionized molecules are similar, since since this is 763 related to the size of the molecule and the curvature of the 764 potential energy surface, which are likely similar of the two 765 species. However, the jump distance is not determined from the 766 Arrhenius fit but rather by the availability of binding sites. For 767 an ionized dopant, the next binding site is one thiophene unit 768 away, and thus the jump distance is smaller. For the neutral 769 dopant, the next binding site is on average a much larger 770 771 distance so although the neutral species is a minority species 772 and has a similar jump attempt frequency to ionized 773 F4MCTCNQ, the macroscopic diffusion is largely driven by 774 the diffusive motion of neutral dopant molecules.

Finally, comparing the diffusion coefficients of F4TCNQ to 775 776 F4MCTCNQ, we note that F4TCNQ and F4TCNQ⁻ 777 respectively diffuse approximately 10-100 times faster than 778 F4MCTCNQ and F4MCTCNQ⁻ at investigated temperatures 779 (Figure 11d). However, the $E_{A neutral}$ of F4TCNQ is shown to 780 be higher than that of F4MCTCNQ. The reduced global 781 diffusion of F4MCTCNQ, therefore, indicates that the methyl 782 ester side chain on the F4MCTCNQ somehow reduces its 783 jump distance. To pin down the reason for the reduced diffusion rate of the substituted dopant, we will present detailed 784 comparisons of temperature-dependent as well as time-785 786 dependent diffusion coefficients between F4TCNQ and its soluble analogues in a separate and upcoming paper. In 787 addition, examining reported diffusion studies on PCBM in the 788 789 P3HT matrix, we can also conclude that the diffusions of 790 molecular dopants F4TCNQ and F4MCTCNQ are faster than that of PCBM in the P3HT $(D = \sim 10^{-14} \text{ cm}^2/\text{s}^{76-78} \text{ and } D =$ 791 $\sim 10^{-11} \text{ cm}^2/\text{s}^{79}$ at around 413 K). 792

The results and model presented here represent the first detailed mechanistic study of a specific dopant diffusing in a specific semicrystalline semiconducting polymer. The results model could be applicable to a wider and more general model for dopant diffusion in a wide variety of organic semiconductors. For this model to be generally applicable, a plarge number of different dopants would need to be nivestigated in a large number of different semiconductors. Also, a general means to determine the temperature-dependent binding energies for dopants and the energy landscape for SOSCs is needed.

804 CONCLUSION

805 In conclusion, we have unveiled the diffusion dynamics of the 806 p-type dopant, F4MCTCNQ, in the semiconducting polymer 807 P3HT for both microscopic and macroscopic time/distance 808 scales using two different techniques. From QENS experiments, 809 two local motions (<4 Å) are observed: methyl group rotations sin with $D \approx 10^{-6} - 10^{-5}$ cm²/s and methoxycarbonyl group sin rotations with smaller $D \approx 10^{-7} - 10^{-6}$ cm²/s, both of which s12 exhibit small E_A values (<15 kJ/mol) that are confirmed by 813 DFT calculations. The macroscopic dynamics, on the other 814 hand, are captured by an LSCM imaging technique. A two-815 dimensional two-species model is developed to interpret this 816 long-range diffusion behavior, which includes a Langmuir 817 isotherm equilibrium between neutral and ionized dopant 818 molecules. As the dominant species, the ionized F4MCTCNQ 819 molecules exhibit slower diffusion than the neutral molecules 820 due to a much shorter jump distance. Both the activation 821 energy for diffusion and the jump distance of the ionized 822 species are in good agreement with the estimates from DFT 823 calculations associated with the translational motions of 824 F4MCTCNQ along the P3HT backbone (E_A of ~50 kJ/mol s25 and jump distance of ~ 2 Å). In contrast, the neutral species has 826 a significantly greater jump distance, which makes the neutral 827 species the dominant diffusing species at the macro scale. 828 Lastly, by comparing the diffusion coefficients between 829 F4MCTCNQ and F4TCNQ, we can also conclude that methyl 830 ester substitution in F4MCTCNQ not only increases 831 solubility/miscibility and thus doping effectiveness but also 832 helps to anchor dopants into position postdeposition due to

reduced diffusion. In theory, the two-dimensional one-/two- 833 species diffusion developed in this paper could be generalized 834 to determine the diffusion coefficient of any fluorescence 835 quenching species that can be evaporated into a polymer. 836 Studies of the dopant dynamics at both microscopic and 837 macroscopic levels are important for understanding the 838 degradation of organic devices as well as improving DISC 839 patterning methodology. 840

ASSOCIATED CONTENT	841
S Supporting Information	842
The Supporting Information is available free of charge on the	843
ACS Publications website at DOI: 10.1021/acs.macro-	844
mol.7b00672.	845
Figures S1–S14 (PDF)	846
AUTHOR INFORMATION	847
Corresponding Author	848
*E-mail: amoule@ucdavis.edu (A.J.M.).	849
ORCID [©]	850
Jun Li: 0000-0001-7850-3722	851
David M. Huang: 0000-0003-2048-4500	852
Ian E. Jacobs: 0000-0002-1535-4608	853
Thomas Harrelson: 0000-0002-8689-4273	854
Kunlun Hong: 0000-0002-2852-5111	855
Notes	856
The authors declare no competing financial interest.	857

ACKNOWLEDGMENTS

This project was carried out with funding from the U.S. 859 Department of Energy, Office of Basic Energy Sciences, 860 Division of Materials Sciences and Engineering, under Award 861 DE-SC0010419. The DFT work was undertaken with the 862 assistance of resources and services from the National 863 Computational Infrastructure (NCI), which is supported by 864 the Australian Government. Computational resources provided 865 by the University of Adelaide's Phoenix High Performance 866 Computing service are also gratefully acknowledged. Deuter- 867 ated P3HT was synthesized at the Center for Nanophase 868 Materials Sciences, a DOE Office of Science User Facility. A 869 portion of this research was performed at Spallation Neutron 870 Source, a DOE Office of Sciences facility. We thank Prof. Greg 871 Miller (UC Davis) for advice on the development of the 872 diffusion model. 873

874

858

(1) Taima, T.; Sakai, J.; Yamanari, T.; Saito, K. Doping Effects for 875 Organic Photovoltaic Cells Based on Small-molecular-weight Semi- 876 conductors. Sol. Energy Mater. Sol. Cells **2009**, 93, 742–745.

(2) Hains, A. W.; Liang, Z.; Woodhouse, M. A.; Gregg, B. A. 878 Molecular Semiconductors in Organic Photovoltaic Cells. *Chem. Rev.* 879 **2010**, *110*, 6689–6735. 880

(3) Zhou, X.; Blochwitz, J.; Pfeiffer, M.; Nollau, A.; Fritz, T.; Leo, K. 881 Enhanced Hole Injection into Amorphous Hole-Transport Layers of 882 Organic Light-Emitting Diodes Using Controlled p-Type Doping. *Adv.* 883 *Funct. Mater.* **2001**, *11*, 310–314. 884

(4) Reineke, S.; Thomschke, M.; Lüssem, B.; Leo, K. White Organic 885 Light-emitting Diodes: Status and Perspective. *Rev. Mod. Phys.* **2013**, 886 85, 1245–1293. 887

(5) Braga, D.; Horowitz, G. High-Performance Organic Field-Effect 888 Transistors. *Adv. Mater.* **2009**, *21*, 1473–1486. 889

(6) Lu, G.; Blakesley, J.; Himmelberger, S.; Pingel, P.; Frisch, J.; 890 Lieberwirth, I.; Salzmann, I.; Oehzelt, M.; Di Pietro, R.; Salleo, A.; 891 892 Koch, N.; Neher, D. Moderate Doping Leads to High Performance of
893 Semiconductor/insulator Polymer Blend Transistors. *Nat. Commun.*894 2013, 4, 1588.

895 (7) Heeger, A. J. Semiconducting and Metallic Polymers: The Fourth 896 Generation of Polymeric Materials (Nobel Lecture). *Angew. Chem., Int.*

897 Ed. 2001, 40, 2591–2611. 898 (8) Lüssem, B.: Riede, M.: Leo, K. Doping of Organic S

898 (8) Lüssem, B.; Riede, M.; Leo, K. Doping of Organic Semi-899 conductors. *Phys. Status Solidi A* **2013**, *210*, 9–43.

900 (9) Zhao, X.; Zhan, X. Electron Transporting Semiconducting 901 Polymers in Organic Electronics. *Chem. Soc. Rev.* **2011**, 40, 3728– 902 3743.

903 (10) Walzer, K.; Maennig, B.; Pfeiffer, M.; Leo, K. Highly Efficient
904 Organic Devices Based on Electrically Doped Transport Layers. *Chem.*905 *Rev.* 2007, 107, 1233–1271.

906 (11) Olthof, S.; Tress, W.; Meerheim, R.; Lüssem, B.; Leo, K. 907 Photoelectron Spectroscopy Study of Systematically Varied Doping 908 Concentrations in An Organic Semiconductor Layer Using A 909 Molecular P-Dopant. *J. Appl. Phys.* **2009**, *106*, 103711.

910 (12) Lee, J. H.; Lee, J.; Kim, Y. H.; Yun, C.; Lüssem, B.; Leo, K. 911 Effect of Trap States on the Electrical Doping of Organic 912 Semiconductors. *Org. Electron.* **2014**, *15*, 16–21.

913 (13) Jacobs, I. E.; Li, J.; Burg, S. L.; Bilsky, D. J.; Rotondo, B. T.; 914 Augustine, M. P.; Stroeve, P.; Moulé, A. J. Reversible Optical Control 915 of Conjugated Polymer Solubility with Sub-micrometer Resolution. 916 ACS Nano **2015**, *9*, 1905–1912.

917 (14) Jacobs, I. E.; Aasen, E. W.; Oliveira, J. L.; Fonseca, T. N.;
918 Roehling, J. D.; Li, J.; Zhang, G.; Augustine, M. P.; Mascal, M.; Moule,
919 A. J. Comparison of Solution-mixed and Sequentially Processed
920 P3HT:F4TCNQ Films: Effect of Doping-induced Aggregation on
921 Film Morphology. J. Mater. Chem. C 2016, 4, 3454–3466.

(15) Jacobs, I. E.; Wang, F.; Hafezi, N.; Medina-Plaza, C.; Harrelson,
T. F.; Li, J.; Augustine, M. P.; Mascal, M.; Moulé, A. J. Quantitative
Pedoping of Conductive Polymers. *Chem. Mater.* 2017, *29*, 832–841.
(16) Jacobs, I. E.; Aasen, E. W.; Nowak, D.; Li, J.; Morrison, W.;
Roehling, J. D.; Augustine, M. P.; Moulé, A. J. Direct-Write Optical
Patterning of P3HT Films Beyond the Diffraction Limit. *Adv. Mater.*2017, *29*, 1603221.

929 (17) Tyagi, P.; Dalai, M. K.; Suman, C. K.; Tuli, S.; Srivastava, R. 930 Study of 2,3,5,6-tetrafluoro-7,7[prime or minute],8,8[prime or 931 minute]- tetracyano quinodimethane Diffusion in Organic Light 932 Emitting Diodes Using Secondary Ion Mass Spectroscopy. *RSC Adv.* 933 **2013**, 3, 24553–24559.

934 (18) Dai, A.; Wan, A.; Magee, C.; Zhang, Y.; Barlow, S.; Marder, S.
935 R.; Kahn, A. Investigation of P-dopant Diffusion in Polymer Films and
936 Bulk Heterojunctions: Stable Spatially-confined Doping for All937 solution Processed Solar Cells. Org. Electron. 2015, 23, 151–157.

938 (19) Treat, N. D.; Brady, M. A.; Smith, G.; Toney, M. F.; Kramer, E.
939 J.; Hawker, C. J.; Chabinyc, M. L. Interdiffusion of PCBM and P3HT
940 Reveals Miscibility in a Photovoltaically Active Blend. *Adv. Energy*941 *Mater.* 2011, *1*, 82–89.

942 (20) Treat, N. D.; Mates, T. E.; Hawker, C. J.; Kramer, E. J.; 943 Chabinyc, M. L. Temperature Dependence of the Diffusion 944 Coefficient of PCBM in Poly(3-hexylthiophene). *Macromolecules* 945 **2013**, 46, 1002–1007.

946 (21) Qi, Y.; Sajoto, T.; Kröger, M.; Kandabarow, A. M.; Park, W.;

947 Barlow, S.; Kim, E.-G.; Wielunski, L.; Feldman, L. C.; Bartynski, R. A.;
948 Brédas, J.-L.; Marder, S. R.; Kahn, A. A Molybdenum Dithiolene
949 Complex as p-Dopant for Hole-Transport Materials: A Multitechnique
950 Experimental and Theoretical Investigation. *Chem. Mater.* 2010, 22,
951 524–531.

(22) Li, J.; Rochester, C. W.; Jacobs, I. E.; Friedrich, S.; Stroeve, P.;
Riede, M.; Moulé, A. J. Measurement of Small Molecular Dopant
F4TCNQ and C60F36 Diffusion in Organic Bilayer Architectures.
ACS Appl. Mater. Interfaces 2015, 7, 28420-28428.

956 (23) Li, J.; Rochester, C. W.; Jacobs, I. E.; Aasen, E. W.; Friedrich, S.;
957 Stroeve, P.; Moulé, A. J. The Effect of Thermal Annealing on Dopant
958 Site Choice in Conjugated Polymers. *Org. Electron.* 2016, 33, 23–31.
959 (24) Fischer, F.; Hahn, T.; Bässler, H.; Bauer, I.; Strohriegl, P.;
960 Köhler, A. Measuring Reduced C60 Diffusion in Crosslinked Polymer

Films by Optical Spectroscopy. *Adv. Funct. Mater.* **2014**, *24*, 6172–961 6177. 962

(25) Tyagi, P.; Tuli, S.; Srivastava, R. Study of Fluorescence 963 Quenching due to 2, 3, 5, 6-Tetrafluoro-7, 7', 8, 8'-Tetracyano 964 Quinodimethane and Its Solid State Diffusion Analysis using 965 Photoluminescence Spectroscopy. J. Chem. Phys. 2015, 142, 054707. 966

(26) Zhang, L.; Zu, F.-S.; Deng, Y.-L.; Igbari, F.; Wang, Z.-K.; Liao, 967 L.-S. Origin of Enhanced Hole Injection in Organic Light-Emitting 968 Diodes with an Electron-Acceptor Doping Layer: p-Type Doping or 969 Interfacial Diffusion? *ACS Appl. Mater. Interfaces* **2015**, *7*, 11965–970 11971. 971

(27) Jung, M.-C.; Qi, Y. Dopant Interdiffusion Effects in n-i-p 972 Structured Spiro-OMeTAD Hole Transport Layer of Organometal 973 Halide Perovskite Solar Cells. *Org. Electron.* **2016**, *31*, 71–76. 974

(28) Gao, W.; Kahn, A. Controlled P Doping of the Hole-transport 975 Molecular Material N,N-diphenyl-N,N-bis(1-naphthyl)-1,1-biphenyl- 976 4,4-diamine with Tetrafluorotetracyanoquinodimethane. *J. Appl. Phys.* 977 **2003**, *94*, 359–366. 978

(29) Kolesov, V. A.; Fuentes-Hernandez, C.; Chou, W.-F.; Aizawa, 979 N.; Larrain, F. A.; Wang, M.; Perrotta, A.; Choi, S.; Graham, S.; Bazan, 980 G. C.; Nguyen, T.-Q.; Marder, S. R.; Kippelen, B. Solution-based 981 Electrical Doping of Semiconducting Polymer Films over a Limited 982 Depth. *Nat. Mater.* **2016**, advance online publication. 983

(30) Gao, Z. Q.; Mi, B. X.; Xu, G. Z.; Wan, Y. Q.; Gong, M. L.; 984 Cheah, K. W.; Chen, C. H. An Organic P-type Dopant with High 985 Thermal Stability for an Organic Semiconductor. *Chem. Commun.* 986 **2008**, 117–119. 987

(31) Gao, W.; Kahn, A. Controlled P-doping of Zinc Phthalocyanine 988 by Coevaporation with Tetrafluorotetracyanoquinodimethane: A 989 Direct and Inverse Photoemission Study. *Appl. Phys. Lett.* **2001**, *79*, 990 4040–4042. 991

(32) Dai, A.; Zhou, Y.; Shu, A. L.; Mohapatra, S. K.; Wang, H.; 992 Fuentes-Hernandez, C.; Zhang, Y.; Barlow, S.; Loo, Y.-L.; Marder, S. 993 R.; Kippelen, B.; Kahn, A. Enhanced Charge-carrier Injection and 994 Collection via Lamination of Doped Polymer Layers p-Doped with a 995 Solution-Processible Molybdenum Complex. *Adv. Funct. Mater.* **2014**, 996 24, 2197–2204. 997

(33) Drechsel, J.; Männig, B.; Kozlowski, F.; Pfeiffer, M.; Leo, K.; 998 Hoppe, H. Efficient Organic Solar Cells Based on a Double p-i-n 999 Architecture Using Doped Wide-gap Transport Layers. *Appl. Phys.* 1000 *Lett.* **2005**, *86*, 244102. 1001

(34) Meerheim, R.; Olthof, S.; Hermenau, M.; Scholz, S.; Petrich, A.; 1002 Tessler, N.; Solomeshch, O.; Lüssem, B.; Riede, M.; Leo, K. 1003 Investigation of C60F36 as Low-volatility P-dopant in Organic 1004 Optoelectronic Devices. J. Appl. Phys. **2011**, 109, 103102. 1005

(35) Lin, X.; Purdum, G. E.; Zhang, Y.; Barlow, S.; Marder, S. R.; 1006 Loo, Y.-L.; Kahn, A. Impact of a Low Concentration of Dopants on 1007 the Distribution of Gap States in a Molecular Semiconductor. *Chem.* 1008 *Mater.* **2016**, 28, 2677–2684. 1009

(36) Li, J.; Zhang, G.; Holm, D. M.; Jacobs, I. E.; Yin, B.; Stroeve, P.; 1010 Mascal, M.; Moulé, A. J. Introducing Solubility Control for Improved 1011 Organic P-type Dopants. *Chem. Mater.* **2015**, *27*, 5765–5774. 1012

(37) Sakai, V. G.; Arbe, A. Quasielastic Neutron Scattering in Soft 1013 Matter. Curr. Opin. Colloid Interface Sci. 2009, 14, 381–390. 1014

(38) Shao, M.; Keum, J.; Chen, J.; He, Y.; Chen, W.; Browning, J. F.; 1015 Jakowski, J.; Sumpter, B. G.; Ivanov, I. N.; Ma, Y.-Z.; Rouleau, C. M.; 1016 Smith, S. C.; Geohegan, D. B.; Hong, K.; Xiao, K. The Isotopic Effects 1017 of Deuteration on Optoelectronic Properties of Conducting Polymers. 1018 *Nat. Commun.* **2014**, *5*, 3180. 1019

(39) Mamontov, E.; Herwig, K. W. A Time-of-flight Backscattering 1020 Spectrometer at the Spallation Neutron Source, BASIS. *Rev. Sci.* 1021 *Instrum.* **2011**, *82*, 085109. 1022

(40) Azuah, R. T.; Kneller, L. R.; Qiu, Y.; Tregenna-Piggott, P. L.; 1023 Brown, C. M.; Copley, J. R.; Dimeo, R. M. DAVE: A Comprehensive 1024 Software Suite for the Reduction Visualization, and Analysis of Low 1025 Energy Neutron Spectroscopic Data. J. Res. Natl. Inst. Stand. Technol. 1026 **2009**, 114, 341. 1027

(41) Hempelmann, R. Quasielastic Neutron Scattering and Solid State 1028 Diffusion; Clarendon Press: 2000. 1029 1030 (42) Langmuir, I. The Adsorption of Gases on Plane Surfaces of 1031 Glass, Mica and Platinum. J. Am. Chem. Soc. **1918**, 40, 1361–1403.

1032 (43) Becke, A. D. Density-functional Thermochemistry. III. The Role 1033 of Exact Exchange. *J. Chem. Phys.* **1993**, *98*, 5648–5652.

1034 (44) Grimme, S. Semiempirical GGA-type Density Functional 1035 Constructed with a Long-range Dispersion Correction. *J. Comput.* 1036 *Chem.* **2006**, *27*, 1787–1799.

(45) Ackling, S. Interplay of Structural, Dynamical, and Electronic
Properties in Doped Semiconducting Polymer Systems. M.Sc. Thesis,
The University of Adelaidem, 2017.

1040 (46) Lemaur, V.; Steel, M.; Beljonne, D.; Brédas, J.-L.; Cornil, J. 1041 Photoinduced Charge Generation and Recombination Dynamics in 1042 Model Donor/Acceptor Pairs for Organic Solar Cell Applications: A 1043 Full Quantum-Chemical Treatment. *J. Am. Chem. Soc.* **2005**, *127*, 1044 6077–6086.

1045 (47) Mityashin, A.; Olivier, Y.; Van Regemorter, T.; Rolin, C.; 1046 Verlaak, S.; Martinelli, N. G.; Beljonne, D.; Cornil, J.; Genoe, J.; 1047 Heremans, P. Unraveling the Mechanism of Molecular Doping in 1048 Organic Semiconductors. *Adv. Mater.* **2012**, *24*, 1535–1539.

(48) Gregg, B. A.; Hanna, M. C. Comparing Organic to Inorganic
Photovoltaic Cells: Theory, Experiment, and Simulation. J. Appl. Phys.
1051 2003, 93, 3605–3614.

1052 (49) Madigan, C. F.; Bulović, V. Solid State Solvation in Amorphous 1053 Organic Thin Films. *Phys. Rev. Lett.* **2003**, *91*, 247403.

1054 (50) Mamontov, E.; Luo, H.; Dai, S. Proton Dynamics in N,N,N,N-

1055 Tetramethylguanidinium Bis(perfluoroethylsulfonyl)imide Protic Ionic 1056 Liquid Probed by Quasielastic Neutron Scattering. *J. Phys. Chem. B* 1057 **2009**, *113*, 159–169.

1058 (51) Frick, B.; Fetters, L. J. Methyl Group Dynamics in Glassy
1059 Polyisoprene: A Neutron Backscattering Investigation. *Macromolecules*1060 **1994**, 27, 974–980.

1061 (52) Ngai, K. L.; Capaccioli, S.; Paciaroni, A. Change of Caged 1062 Dynamics at Tg in Hydrated Proteins: Trend of Mean Squared 1063 Displacements after Correcting for the Methyl-group Rotation 1064 Contribution. *J. Chem. Phys.* **2013**, *138*, 235102.

1065 (53) Paternó, G.; Cacialli, F.; García-Sakai, V. Structural and 1066 Dynamical Characterization of P3HT/PCBM Blends. *Chem. Phys.* 1067 **2013**, 427, 142–146.

1068 (54) Gautam, S.; Liu, T.; Rother, G.; Jalarvo, N.; Mamontov, E.;
1069 Welch, S.; Sheets, J.; Droege, M.; Cole, D. R. Dynamics of Propane in
1070 Nanoporous Silica Aerogel: A Quasielastic Neutron Scattering Study. J.
1071 Phys. Chem. C 2015, 119, 18188–18195.

1072 (55) Singwi, K. S.; Sjölander, A. Diffusive Motions in Water and Cold 1073 Neutron Scattering. *Phys. Rev.* **1960**, *119*, 863–871.

1074 (56) Batchelder, L. S.; Niu, C. H.; Torchia, D. A. Methyl 1075 Reorientation in Polycrystalline Amino Acids and Peptides: A 1076 Deuteron NMR Spin-lattice Relaxation Study. *J. Am. Chem. Soc.* 1077 **1983**, 105, 2228–2231.

1078 (57) Takegoshi, K.; Imashiro, F.; Terao, T.; Saika, A. 1H and 13C 1079 NMR Study on Rotation of Congested Methyl Groups in Methyl 1080 Substituted Phenanthrenes, Fluorenes, and Fluorenones. *J. Chem. Phys.* 1081 **1984**, *80*, 1089–1094.

(58) Beckmann, P. A.; Cheung, A. M.; Fisch, E. E.; Fusco, F. A.;
1083 Herzog, R. E.; Narasimhan, M. Methyl and Tertbutyl Reorientation
1084 and Distributions of Activation Energies in Molecular Solids. A
1085 Nuclear Spinrelaxation Study in 2,4 and 2,5ditertbutylhydroxybenzene.
1086 J. Chem. Phys. 1986, 84, 1959–1968.

(59) Chahid, A.; Alegria, A.; Colmenero, J. Methyl Group Dynamics
1088 in Poly(vinyl methyl ether). A Rotation Rate Distribution Model.
1089 *Macromolecules* 1994, *27*, 3282–3288.

(60) Mukhopadhyay, R.; Alegria, A.; Colmenero, J.; Frick, B. Methyl
1091 Group Dynamics in Poly(vinyl acetate): A Neutron Scattering Study.
1092 Macromolecules 1998, 31, 3985–3993.

1093 (61) Moreno, A. J.; Alegría, A.; Colmenero, J.; Frick, B. Methyl 1094 Group Dynamics in Poly(methyl methacrylate): From Quantum 1095 Tunneling to Classical Hopping. *Macromolecules* **2001**, *34*, 4886– 1096 4896. Article

P3HT. Org. Electron. 2013, 14, 1330–1336. 1099 (63) Harrelson, T. F.; Cheng, Y. Q.; Li, J.; Jacobs, I. E.; Ramirez- 1100 Cuesta, A. J.; Faller, R.; Moulé, A. J. Identifying Atomic Scale Structure 1101 in Undoped/Doped Semicrystalline P3HT Using Inelastic Neutron 1102 Scattering. Macromolecules 2017, 50, 2424–2435. 1103

(64) Wiberg, K. B.; Laidig, K. E. Barriers to Rotation Adjacent to 1104 Double Bonds. 3. The Carbon-oxygen Barrier in Formic Acid, Methyl 1105 Formate, Acetic Acid, and Methyl Acetate. The Origin of Ester and 1106 Amide Resonance. J. Am. Chem. Soc. **1987**, 109, 5935–5943.

(65) Heijboer, J.; Baas, J.; van de Graaf, B.; Hoefnagel, M. A 1108 Molecular Mechanics Study on Rotational Motions of Side Groups in 1109 Poly(methyl methacrylate). *Polymer* **1987**, *28*, 509–513. 1110

(66) Gabrys, B.; Higgins, J. S.; Ma, K. T.; Roots, J. E. Rotational 1111 Motion of the Ester Methyl Group in Stereoregular Poly(methyl 1112 methacrylate): A Neutron Scattering Study. *Macromolecules* **1984**, *17*, 1113 560–566.

(67) Arrighi, V.; Higgins, J. S.; Burgess, A. N.; Howells, W. S. 1115 Rotation of Methyl Side Groups in Polymers: A Fourier Transform 1116 Approach to Quasielastic Neutron Scattering. 1. Homopolymers. 1117 *Macromolecules* **1995**, *28*, 2745–2753. 1118

(68) Kimura, K.; Kubo, M. Structures of Dimethyl Ether and Methyl 1119 Alcohol. J. Chem. Phys. **1959**, 30, 151–158. 1120

(69) Barnes, J. D. Inelastic Neutron Scattering Study of the Rotator 1121 Phase Transition in N-nonadecane. J. Chem. Phys. **1973**, 58, 5193–1122 5201. 1123

(70) Fujara, F.; Wefing, S.; Spiess, H. W. Dynamics of Molecular 1124 Reorientations: Analogies between Quasielastic Neutron Scattering 1125 and Deuteron NMR Spin Alignment. J. Chem. Phys. **1986**, 84, 4579–1126 4584. 1127

(71) Winkler, S.; Amsalem, P.; Frisch, J.; Oehzelt, M.; Heimel, G.; 1128 Koch, N. Probing the Energy Levels in Hole-doped Molecular 1129 Semiconductors. *Mater. Horiz.* **2015**, *2*, 427–433. 1130

(72) Yim, K. H.; Whiting, G. L.; Murphy, C. E.; Halls, J. J. M.; 1131 Burroughes, J. H.; Friend, R. H.; Kim, J.-S. Controlling Electrical 1132 Properties of Conjugated Polymers via a Solution-Based p-Type 1133 Doping. *Adv. Mater.* **2008**, *20*, 3319–3324. 1134

(73) Tsoi, W. C.; Spencer, S. J.; Yang, L.; Ballantyne, A. M.; 1135 Nicholson, P. G.; Turnbull, A.; Shard, A. G.; Murphy, C. E.; Bradley, 1136 D. D. C.; Nelson, J.; Kim, J.-S. Effect of Crystallization on the 1137 Electronic Energy Levels and Thin Film Morphology of P3HT:PCBM 1138 Blends. *Macromolecules* **2011**, *44*, 2944–2952. 1139

(74) Yu, G.; Gao, J.; Hummelen, J. C.; Wudl, F.; Heeger, A. J. 1140 Polymer Photovoltaic Cells: Enhanced Efficiencies via a Network of 1141 Internal Donor-Acceptor Heterojunctions. *Science* **1995**, 270, 1789–1142 1791. 1143

(75) Ferguson, A. J.; Kopidakis, N.; Shaheen, S. E.; Rumbles, G. 1144 Quenching of Excitons by Holes in Poly(3-hexylthiophene) Films. J. 1145 Phys. Chem. C 2008, 112, 9865–9871. 1146

(76) Watts, B.; Belcher, W. J.; Thomsen, L.; Ade, H.; Dastoor, P. C. 1147 A Quantitative Study of PCBM Diffusion during Annealing of 1148 P3HT:PCBM Blend Films. *Macromolecules* **2009**, *42*, 8392–8397. 1149

(77) Labram, J. G.; Kirkpatrick, J.; Bradley, D. D. C.; Anthopoulos, T. 1150 D. Measurement of the Diffusivity of Fullerenes in Polymers Using 1151 Bilayer Organic Field Effect Transistors. *Phys. Rev. B: Condens. Matter* 1152 *Mater. Phys.* **2011**, *84*, 075344.

(78) Labram, J. G.; Kirkpatrick, J.; Bradley, D. D. C.; Anthopoulos, T. 1154 D. Impact of Fullerene Molecular Weight on P3HT:PCBM Micro-1155 structure Studied Using Organic Thin-Film Transistors. *Adv. Energy* 1156 *Mater.* **2011**, *1*, 1176–1183. 1157

(79) Berriman, G. A.; Holdsworth, J. L.; Zhou, X.; Belcher, W. J.; 1158 Dastoor, P. C. Molecular versus Crystallite PCBM Diffusion in P3HT: 1159 PCBM Blends. *AIP Adv.* **2015**, *5*, 097220. 1160