Vibrational Stark Effect Mapping of Polaron Delocalization in Chemically Doped Conjugated Polymers

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ABSTRACT: The nitrile vibrational modes of the 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane (F4TCNQ) anion occur across a range of different frequencies in doped conjugated polymers. We show that these shifts can be understood as resulting from the vibrational Stark effect. F4TCNQ− serves as a sensitive vibrational Stark probe, and its nitrile stretches provide a direct read-out of the locally experienced electric fields from the nearby polaron on the doped polymer backbone. The shift of the F4TCNQ− B2u mode and broadening of the B2u mode in doped poly(3-hexylthiophene-2,5-diyl) (P3HT) confirm that the dopant anion’s long axis is oriented perpendicular to the P3HT backbone. The magnitude of the vibrational shifts prove that the anion−polaron Coulombic interaction. This work provides the first direct experimental measurement of polaron delocalization in doped conjugated polymer films and verifies theoretical models relating polaron coherence to the shape of the doped polymer’s mid-IR electronic absorption spectrum.

1. INTRODUCTION

Chemical doping of conjugated polymers has emerged as a viable method for tuning the electronic properties of organic electronic devices such as thermoelectrics and photovoltaics.1−3 In the simplest case, p-type doping of polymer-based semiconductors can be achieved through the introduction of strong oxidizing agents whose LUMO levels are energetically lower than the HOMO levels of the semiconducting polymer. This leads to doping via integer charge transfer (ICT), where each dopant molecule removes an electron from the polymer π-system, producing both a cationic hole that can delocalize along the conjugated backbone (also termed a polaron) and a counterion from the reduced dopant.

A variety of recent work has suggested that the ICT-separate hole and dopant counterion Coulombic interaction plays a fundamental role in determining the carrier behavior in doped conjugated polymers: the stronger the Coulombic interaction, the lower the polaron delocalization and carrier mobility.5−11 The experimental evidence for charge carrier−counterion interactions, however, is indirect, based mostly on features observed in the polaron mid-IR absorption spectrum. One of the goals of this paper is to provide a means by which to directly measure the local electric fields experienced by polarons and dopant counterions in doped conjugated polymers.

The approach we take is based on the fact that the strength of the Coulombic interaction between polarons on the conjugated polymer backbone and dopant counterions is actually recorded in the vibrational spectra of certain dopant anions. Our first hint that this might be possible comes from previous work by both our group15 and others16−25 who have used the nitrile stretching vibrational spectrum of the 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane (F4TCNQ) dopant as a diagnostic tool for tracking the doping mechanism in conjugated polymers, which takes place either via ICT or by the formation of charge transfer complexes (CTCs). Nitrile stretching modes have the advantage that they occur in a mid-IR spectral “window” where no other molecular vibrations appear, so that they are relatively easy to measure and characterize. In the context of studying CTCs, the nitrile stretching modes of the neutral F4TCNQ molecule undergo a predictable softening in their resonant vibrational energy that is roughly proportional to the oxidation state of the molecule. As a result, vibrational absorption at intermediate energies between the neutral and anionically charged species is understood to represent a fractional charge transfer state for the dopant molecule.

The sensitivity of dopant vibrations to the local polymer environment also can be seen in work from Hase et al.,26 who
recently studied the nitrile stretching vibrational spectrum of the F4TCNQ radical anion inside blend-doped films of poly(3-hexylthiophene-2,5-diyl) (P3HT). These workers showed that the amplitude of certain F4TCNQ− vibrational features can change quite dramatically (and even disappear) when the doped films are subjected to thermal annealing cycles. This demonstrates that the precise vibrational spectrum of the F4TCNQ dopant anion is in fact dependent on the local structural morphology. As a result, it has become increasingly clear that the F4TCNQ vibrational spectrum can change in subtle ways as a result of the charge state of the molecule and the local structural morphology in which it resides.

In this paper, we provide a comprehensive picture of the Coulombic interactions in F4TCNQ-doped P3HT films that links the behavior of the F4TCNQ nitrile vibrational spectrum to the degree of polaron delocalization in the doped polymer, the local structural disorder, and the polaron-dopant Coulombic interactions that ultimately control the charge carrier mobility. We show that the counterion vibrational spectrum is directly influenced by the local electric field emanating from the nearby hole and thus can serve as an exquisite reporter of the degree of polaron delocalization on the nearby conjugated polymer.

Figure 1. (a) Normalized mid-IR electronic absorption spectra for P3HT films sequentially doped with F4TCNQ (1 mg mL−1) from solvent blends composed of CF and DCM. (b) Normalized mid-IR electronic absorption spectra of doped P3HT films where the initial films are created to have a range of crystallinities and then subsequently doped with F4TCNQ (1 mg mL−1) from DCM. For the blue and dark green curves, commercially available P3HT with a regioregularity of 91−94% was used, and the relative film crystallinity was controlled by varying the drying time of the casting solvent. A specially synthesized batch of P3HT with virtually 100% regioregularity was cast from ODCB to obtain the most ordered conditions shown in light green (see the Supporting Information for X-ray-based structural characterization). The A/B spectral ratio (see text) for both panels a and b can be estimated on the vertical axis by following the horizontal dashed lines. (c) F4TCNQ nitrile stretch vibrational spectra of the doped films in panel a (also directly visible in panel a as tiny peaks near 0.27 eV). The shaded Gaussians correspond to peak fits for the conditions using 100% CF as the doping solvent, reduced to 0.3 intensity for ease of viewing (see the Supporting Information for fitting details). (d) F4TCNQ nitrile stretching vibrational spectra of the doped films in panel b. (e) Scatter plot displaying the experimentally measured A/B ratio for the samples in panels a and b plotted against the vibrational energy of the F4TCNQ− B1u mode. (f) Scatter plot displaying the A/B ratio for all the samples in panels a and b plotted against the peak width of the F4TCNQ− B2u vibrational mode.
We then argue that the behavior of the F$_4$TCNQ$^-$ nitride stretching modes can be quantitatively described by using the framework of the vibrational Stark effect (VSE). There is a rich existing literature using the vibrational Stark shifts for a variety of nitride, carbonyl, and other pendant groups to directly measure the local electric fields experienced in biological systems.$^{7−9}$ This method also allows for tuning the degree of crystallinity in doped conjugated polymer films. For example, doped P3HT crystallinity can be tuned either by changing the processing conditions of the precast polymer film$^{14}$ or by careful choice of the solvent used in the subsequent doping step.$^{15}$

Figure 1a shows a series of mid-IR absorption spectra for F$_4$TCNQ-doped P3HT films where the solvent used to introduce the dopant is composed of a mixture of chloroform (CF) and dichloromethane (DCM). By varying the blend ratio of these two solvents, we can exert a high degree of control over the resultant morphology. In previous work, we used grazing incidence wide-angle X-ray scattering (GIWAXS) to characterize the crystallinity of films produced this way and found that although the morphology and even the doping mechanism both become associated with increased amorphous regions as CF content increases in the doping solvent, the ordered regions that remain become more highly crystalline.$^{15}$

This is due to the strong dissipating power of the CF solvent, which allows for recrystallization of the ordered fractions of the precast P3HT film that remain following doping. Thus, the crystalline domains that do survive have a much higher average polaron coherence length than crystallites in films prepared with doping solvents where redissolution does not occur.$^{15}$

The crystallinity of F$_4$TCNQ-doped P3HT films also can be manipulated during the initial polymer spin-coating step, as shown in Figure 1b. In this case, we exploit both the polymer regioregularity and the polymer casting solvent drying kinetics to control the polymer order. We previously have used GIWAXS to show that when a commercial P3HT film is cast from CF, a fast-drying solvent, the resultant polymer film is highly amorphous with relatively little crystallinity.$^{6}$ In contrast, when commercially available P3HT is cast from the more slowly drying solvent o-dichlorobenzene (ODCB), the films have more time to order and become more crystalline. Finally, by using a specially synthesized batch of P3HT with virtually 100% regioregularity,$^{5,6}$ also cast from ODCB, we can obtain highly crystalline P3HT films (see the Supporting Information for experimental details). We also showed that when the film crystallinity is tuned this way, the crystalline regions of the polymer dope first, and the amorphous regions only become doped at very high doping concentrations after the crystalline regions are filled.$^{6}$

3. MID-IR SPECTROSCOPY OF F$_4$TCNQ-DOPED P3HT FILMS

We have previously shown that sequential processing, casting the polymer film first and then infiltrating the dopant in a second solution- or gas-phase step, is a versatile technique capable of producing high-quality doped semiconducting polymer films.$^{6,12−14}$ This method also allows for tuning the degree of crystallinity in doped conjugated polymer films. For example, doped P3HT crystallinity can be tuned either by changing the processing conditions of the precast polymer film$^{15}$ or by careful choice of the solvent used in the subsequent doping step.$^{15}$

3.1. Spectral Signatures of P3HT Polaron Coherence Following Doping with F$_4$TCNQ. Spano and co-workers have developed a theory for doped conjugated polymers that describes how changes in the energetic and structural disorder of a polymer affect polaron coherence, that is, the degree to which the polaron is delocalized along the P3HT backbone and/or between adjacent π-stacked P3HT chains.$^{6,10}$ The theory indicates that as the polaron coherence changes, there is a corresponding shift in both the position and shape of the mid-IR polaron electronic absorption spectrum. In this model, one of the primary features that affects the polaron coherence length is the strength of the Coulombic interaction between the positively charged, delocalized hole on the polymer and the corresponding countercharge located on the dopant molecule. The model can reproduce experiment remarkably well for a variety of conjugated polymer–dopant pairs.

In the present work, the most crystalline of the F$_4$TCNQ-doped P3HT samples (Figure 1 purple curve in panel a and
light green curve in panel b) show the characteristic P3HT polaron absorption from highly ordered environments. Because the hole is better able to delocalize in such environments, there is less Coulombic attraction between the hole on the polymer and dopant counterion, as discussed further in section 3.1. This weakened Coulombic interaction and larger coherence lengths are manifest spectroscopically by both increased relative absorption intensity for peak “A” (the rough feature centered near ~0.16 eV), and a shift to lower energies for peak “B” (the main, smoother feature whose maximum falls between 0.35 and 0.55 eV depending on the sample).

In contrast, the curves in maroon (panel a) and blue (panel b), which are for the most disordered doped P3HT films, show mid-IR absorption spectra characterized by a relatively low absorption intensity of the A peak and a shift of the B peak to higher energies. This is a hallmark of poor polaron coherence.7−10 Because the amplitude of peak A decreases relative to peak B with increasing hole localization, the amplitude ratio of peaks A/B becomes a simple metric for tracking changes in polaron delocalization in doped conjugated polymers.7−10

For the method of controlling morphology by adjusting the doping solvent (panel a), the A/B ratio increases from A/B = 0.55 when casting F4TCNQ from pure DCM, up to A/B = 0.81 when casting the dopant from pure CF. When altering the morphology during the polymer casting step (panel b), the change in the A/B ratio is nearly identical, with A/B = 0.56 when the polymer is cast from CF and A/B = 0.81 for slow-dried films fabricated from 100% regioregular P3HT. Because of the relatively small physical size of the F4TCNQ anion, the Coulombic interaction between the anion and the hole polaron has enough strength that increasing the local order of the polymer can only tune the degree of polaron delocalization over a somewhat limited range compared to what is achievable with larger dopants designed to minimize the polaron–dopant Coulombic interaction.13,14

The theory developed by Spano and co-workers provides a direct connection between polaron coherence lengths and spectral A/B ratios.9 In this work, we experimentally measure the change in polaron coherence length by estimating the strength of the electric field between the F4TCNQ+ counterion and the hole polaron on P3HT. Our approach is possible because the theoretical relationship between the A/B spectral ratio and polaron coherence is predicted to be roughly linear across a wide range of coherence length values.9 For our doped P3HT films with different degrees of order, the measured A/B ratios range from ~0.55 to 0.81, which should correspond to an ~25% change in the average polaron coherence length.9 In the sections that follow, we will show how shifts in the energy of the F4TCNQ+ vibrational modes allow us to map out the local electrical fields in doped P3HT films to precisely determine the change in polaron coherence length.

3.2. F4TCNQ+ Nitrile Vibrational Spectrum in Doped P3HT Films. F4TCNQ+ has two IR-active vibrational modes with significant oscillator strength in the so-called transparent window between 1800 and 2500 cm−1; these are C≡N stretching modes with B1u and B2u symmetry, illustrated in Figure 2a, which in doped P3HT films occur near 2190 and 2170 cm−1, respectively. An example IR spectrum of F4TCNQ+ doped P3HT produced by sequential processing is shown in Figure 2b. Meneghetti and Pecile have noted that the location and geometry of the nitrile substituents on this molecule make these two vibrational modes uniquely susceptible to their local environment.15 In fact, this susceptibility to local environment should make these modes relatively poor indicators for quantitatively measuring the degree of charge transfer, a subject of much recent interest.15,17

The literature on the vibrational spectroscopy of the F4TCNQ+ radical anion inside sequentially solution-doped P3HT (with the dopant cast from 1 mg mL−1 n-butyl acetate) and inside blend-doped P3HT samples taken from various literature references, in descending order on the legend: Pingel et al. (ref 16), Ghani et al. (ref 18), Mendez et al. (ref 58), Jacobs et al. (ref 21), Hase et al. (ref 26), Neelamraju et al. (ref 20), and Watts et al. (ref 22).

Figure 2. (a) Atomic displacement vectors for the two F4TCNQ C≡N vibrational modes with B1u and B2u symmetry. Compression and expansion of the C≡N bonds is indicated red and blue arrows, respectively. The difference dipoles (green arrows) for these two modes point along the long axis of the molecule for the B1u mode and across the short molecular axis for the B2u mode. (b) FTIR spectrum of the F4TCNQ+ radical anion inside sequentially solution-doped P3HT (with the dopant cast from 1 mg mL−1 n-butyl acetate) and (c) inside blend-doped P3HT samples taken from various literature references, in descending order on the legend: Pingel et al. (ref 16), Ghani et al. (ref 18), Mendez et al. (ref 58), Jacobs et al. (ref 21), Hase et al. (ref 26), Neelamraju et al. (ref 20), and Watts et al. (ref 22).
the $B_{1u}$ and $B_{2u}$ modes usually appear near 2194 and 2170 cm$^{-1}$, respectively. However, the spectra in Figure 2c show an additional vibrational peak between these two modes, centered near 2185 cm$^{-1}$. The specific origin of this peak has been a topic of debate.$^{16,18,20–22,58}$ In early work, this 2185 cm$^{-1}$ peak was tentatively assigned to a C=CN stretching mode of $A_2$ symmetry,$^{16,17}$ which would require a break in planarity of the $F_4TCNQ$ anion. Such a nonplanar geometry might occur if $F_4TCNQ$-doped P3HT in a $\alpha$-stacking arrangement.$^{59}$ The more recent consensus, however, is that $F_4TCNQ$ resides in the lamellar side-chain regions of the doped P3HT crystallites,$^{6,11,12,24,26,53,60}$ so that the 2185 cm$^{-1}$ mode must result from some other phenomenon.$^{22}$

One important clue to the assignment of the 2185 cm$^{-1}$ peak comes from Hase et al.$^{65}$ who showed that upon exposing $F_4TCNQ$-doped P3HT films to extended thermal annealing cycles, the $B_{1u}$ absorption feature near 2194 cm$^{-1}$ becomes reduced in amplitude, leaving behind the peak near 2185 cm$^{-1}$ and the $B_{1u}$ mode near 2170 cm$^{-1}$. Watts et al. subsequently noted that if the unknown peak near 2185 cm$^{-1}$ did belong to an $A_2$ mode, then the two features near 2185 and 2194 cm$^{-1}$ should show similar behavior upon annealing.$^{22}$ The fact that annealing causes a loss in amplitude only for the 2194 cm$^{-1}$ mode suggests that this vibrational feature is associated with a local structure that can change upon annealing. Thus, it seems far more likely that the two features near 2185 and 2194 cm$^{-1}$ both belong to the $B_{1u}$ mode, where the specific vibrational energies are reporting something about variations in the local environment.

With the idea that the $F_4TCNQ^-$ nitrile stretching modes shift with their local environment, we now examine how the doped film processing method, which controls P3HT crystallinity and polaron delocalization, affects the resulting dopant nitrile vibrational spectrum. Figure 2b shows the intensity-normalized nitrile vibrational spectrum of the $F_4TCNQ$ radical anion inside sequentially doped P3HT using the relatively noninteracting solvent n-butylic acid (NBA) to infiltrate the dopant at a concentration of 1 mg mL$^{-1}$. It is clear from comparing panels b and c in Figure 2 that the dopant anion vibrational spectrum for sequentially doped films is considerably different from those obtained from blend doping. Rather than seeing two peaks for the $B_{1u}$ mode distributed between 2194 and 2185 cm$^{-1}$, the sequentially doped film shows only a single feature centered at 2187 cm$^{-1}$. This suggests that for sequential processing, which generally produces more ordered doped P3HT films, the dopant molecules have a singular well-defined geometry in the polymer crystallites. The higher-energy $B_{1u}$ peak near 2194 cm$^{-1}$, in contrast, must belong to anions that Coulombically trap the hole in the morphologically disordered polymer regions that predominate in blend-doped films.

3.3. Correlation between Polaron Coherence and $F_4TCNQ^-$ Vibrational Spectrum. Now that we have established a basis for understanding how the doped polymer structural environment should affect the vibrational spectrum of $F_4TCNQ^-$, we can consider precisely how the local degree of order and polaron coherence affect the nitrile stretching frequencies. Figure 1c shows the vibrational spectrum of the $F_4TCNQ$ radical anion inside P3HT films sequentially doped with $F_4TCNQ$ cast from CF/DCM solvent blends. We see that when the dopant is infiltrated from pure CF, the principal $B_{1u}$ nitrile absorption feature occurs with $\nu_{\text{max}} = 2185$ cm$^{-1}$ (purple curve). As the polaron becomes increasingly localized by increasing the fraction of DCM in the doping solvent, the $B_{1u}$ vibrational peak monotonically shifts to $\nu_{\text{max}} = 2189$ cm$^{-1}$ (maroon curve). It is worth noting that even in these high-DCM processed P3HT films, which are the most disordered made by sequential doping, the $B_{1u}$ mode is still five wavenumbers lower in energy than the 2194 cm$^{-1}$ peak seen in the even more disordered blend-doped P3HT films.

When we analyze the principal vibrational features in the sequentially doped films seen in Figure 1c in more detail, we see that they are actually best fit to two Gaussians: a dominant feature that shifts between 2188 and 2185 cm$^{-1}$ (shaded purple curve reduced to 0.3 intensity) and a smaller shoulder (shaded maroon curve) located near 2194 cm$^{-1}$ (see the Supporting Information for fitting details). As the polarons in the crystalline regions become more delocalized by increasing the fraction of CF, the intensity of this small peak should near 2194 cm$^{-1}$ monotonically decreases and disappears entirely when the dopant is cast from 100% CF. The fact that the peak near 2194 cm$^{-1}$ decreases with a higher fraction of CF solvent and thus increasing local crystalline order fits well with the notion that this higher energy peak is associated with highly localized polarons, which are more likely to exist in more disordered polymer regions. This also helps to explain why the peak near 2194 cm$^{-1}$ presents so prominently in films produced by blend doping, since blend-cast films are expected to have a broader distribution of morphological states.

A similar trend for the $F_4TCNQ^-$ $B_{1u}$ vibrational mode also occurs when the doped polymer crystallinity is tuned by changing the how the polymer film is cast (with the dopant solvent held constant), as shown in Figure 1d. When the underlying P3HT film is 100% regioregular and cast from ODCB, the $F_4TCNQ^-$ $B_{1u}$ mode has $\nu_{\text{max}} = 2186$ cm$^{-1}$. As the polaron becomes more localized when the initial P3HT film is cast from CF, we see the $B_{1u}$ mode shift up to $\nu_{\text{max}} = 2191$ cm$^{-1}$.

For both sets of doped P3HT films, we see a remarkably strong correlation between the P3HT polaron coherence, as measured by the A/B spectral intensity ratio, and the position of the $F_4TCNQ^-$ $B_{1u}$ nitrile stretch, as summarized in Figure 1e. This correlation is one of the principal results of this work: the position of the $B_{1u}$ stretching peak of the dopant provides a local measure of polaron coherence on the polymer. The $F_4TCNQ^-$ $B_{1u}$ nitrile stretching spectra seen in Figure 2c behave in a fundamentally predictable way if the degree of polaron coherence is known. Thus, the variations measured by different groups directly reflect differences in polaron coherence resulting from different methods of fabricating the doped films.

As an aside, we note that in both sets of doped P3HT films studied in Figure 1, we see an increased amplitude for the $F_4TCNQ$ CTC vibrational peak near 2200 cm$^{-1}$ with increasing polymer disorder.$^{6}$ The reason the CTC peak becomes more prominent when the doping solvent has an increasing fraction of CF is because the use of CF as a sequential doping solvent causes significant disruption of the P3HT crystal lattice. This is what both increases the total fraction of disordered regions and also increases the order of the crystalline regions that remain.$^{65}$ When using more common and milder processing solvents, however, it is clear that CTC formation is mostly limited by the relative amount of disordered polymer regions.

For the subset of our $F_4TCNQ$-doped P3HT samples where the $F_4TCNQ^-$ $B_{2u}$ mode did not have a neighboring low
energy shoulder (which has been previously attributed to the B_{3g} mode near \nu = 2160 cm^{-1})^{15,66} or simply remained an unassigned mode\[^{20,22}\] we can cleanly fit both the position and width of the B_{2u} peak. We find that the vibrational energy of this mode does not shift in different P3HT environments, always remaining within a 2 cm\(^{-1}\) range. However, we do see that the full width at half-maximum (FWHM) of the F_4TCNQ\(^-\) B_{2u} feature broadens from 8.1 cm\(^{-1}\) for doped P3HT films with the most localized polaron (lowest A/B ratio) up to 12.5 cm\(^{-1}\) for doped films with the most delocalized polaron (highest A/B ratio), a correlation summarized in Figure 1f. The fact that the peak width of the B_{3g} mode correlates with polaron coherence, not the peak vibrational energy, provides another crucial piece of information that will allow us to quantify both the orientation of the F_4TCNQ anion relative to the P3HT polaron and the degree of polaron delocalization.

Although the two series of doped P3HT films we have studied in Figure 1 modify the local polymer crystalline order during different processing steps, they show a similar range of polaron coherence, as evidenced by the A/B spectral ratio, and a similar behavior of the B_{1u} and B_{2u} vibrational modes, summarized in Figures 1e and 1f. For the most locally ordered P3HT films from each set (100% RR precast P3HT film and commercial P3HT with the dopant cast from 100% CF), the mid-IR spectra have an identical A/B ratio of 0.81 and correspondingly similar B_{1u} vibrational peak positions of 2185–2186 cm\(^{-1}\) as well as similarly broad B_{2u} vibrational peak widths of ~12.2–12.5 cm\(^{-1}\). This represents the highest A/B ratio, smallest B_{1u} vibrational frequency, and broadest B_{2u} peak width that we have observed for this polymer–dopant combination. Thus, it is likely these values are approaching the fundamental upper limit for polaron delocalization when F_4TCNQ is used to dope highly ordered P3HT.

Why would there be a fundamental upper limit to polaron delocalization? We expect polaron delocalization to be determined by two primary factors: (i) the inherent energetic disorder due to structural imperfections in the polymer crystallites and (ii) the strength of the Coulombic interaction between the hole and dopant counteranion.\[^7\]–\[^10\] Because the dopant preferentially occupies crystalline regions of P3HT films,\[^6,53\] the distance between the anion and the polymer backbone, \(d_{\text{anion}}\), depends mainly on the physical size of the anion.\[^13,14\] For the doped P3HT samples whose properties are shown in Figure 1, \(d_{\text{anion}}\) is essentially fixed, so the increases we see in polaron coherence and the corresponding changes in the vibrational modes must result from increases in the local structural order of the polymer.

All of this means that the extent of polaron delocalization in the most-ordered polymer regions is limited by the minimum possible strength for the hole–counterion Coulombic interaction. F_4TCNQ-doped P3HT has a lamellar spacing of \(~17\text{ Å},\[^6,15\] so the maximum possible dopant–polaron distance is \(~7–8\text{ Å}.\[^1\] Because of this relatively short distance, the Coulombic interaction from F_4TCNQ\(^-\) always causes some localization of the polaron beyond what is intrinsically allowed by a highly ordered polymer structure.\[^6,9\] This explains why the F_4TCNQ-doped P3HT system has a relatively limited doping efficiency and carrier delocalization\[^67\] and why the use of physically larger dopants that screen the Coulombic interaction can produce polaronics with even greater delocalization in P3HT (A/B ratios greater than unity).\[^13,14\] The idea that dopant anions can only reside in a limited set of places also helps to explain why F_4TCNQ-doped P3HT films always display some absorption of the most weakly shifted B_{1u} nitrile peak near 2185 cm\(^{-1}\); even in blend-cast films with a relatively poor overall morphology, there are always a few doped P3HT crystallites with a reasonable degree of order.

The strength of the Coulombic interactions between the polaronic charge carriers and their dopant counterions is also expected to impact the measured electrical conductivity of doped conjugated polymer films. In theory, smaller Stark shifts indicate lower Coulombic binding between the polaron and counterion, which should result in higher electrical conductivity. For the series of samples where the polymer crystallinity is controlled in the polymer casting step, this is precisely what we see, as shown in Table S9. However, for the series of samples where the doping solvent properties were altered, a significant number of F_4TCNQ dopants begin to dope via CTC formation. CTC doping results in carriers that are several orders of magnitude less mobile than those created by ICT doping, so the presence of a significant number of CTCs leads to a decrease in conductivity despite the diminished Stark shift of the counterions in the crystalline regions. We have explored the nature of the CTC doping in samples like these and the effects on charge transport characteristics in previous work.\[^12\]

4. F_4TCNQ AND THE VIBRATIONAL STARK EFFECT

Now that we have established a clear correlation between the behavior of the C≡N vibrational stretches of F_4TCNQ\(^-\) and the degree of polaron coherence in doped P3HT films, we can turn our attention to considering the mechanisms that cause these changes to take place.

The central thesis of this paper is that the blue-shift of the B_{1u} mode and broadening of the B_{2u} mode with increasing A/B ratio occur as a direct result of the vibrational Stark effect (VSE). The VSE arises from the interaction between a molecule’s vibrational mode and an externally applied electric field. If vibrational excitation of a particular mode changes the dipole of the molecule, then the difference dipole vector between the ground and excited vibrational states can interact with the aligned component of an external electric field, causing a shift in the resonant energy for that vibrational mode. Because external fields are usually weak relative to the internal fields that hold a molecule together, the VSE can be well described by first-order perturbation theory, and the shift of the vibrational frequency is roughly linear with the local field strength.

The VSE is well studied in the literature, and its primary use has been to understand the electric fields present in biological systems.\[^27–39\] The VSE also has been recently used to directly study ionic interactions in solution, allowing researchers to determine if dissolved salts exist as contact ion pairs, solvent-separated ion pairs, or truly free solvated ions.\[^63–67\] In many applications, VSE probes are simple alkylnitrile substituents, for which the difference dipole is aligned along the internuclear axis of the C≡N triple bond. The F_4TCNQ anion, in contrast, has two pairs of nitride groups. Figure 2a shows the atomic displacement vectors for the F_4TCNQ\(^-\) B_{1u} and B_{2u} nitride stretching modes with bond compression indicated in red, bond expansion in blue, and the difference dipole vectors in green. The difference dipole vector for the B_{1u} mode points along the long axis of the molecule, while that of the B_{2u} mode points transversely across the short axis. So although F_4TCNQ was not specifically designed for this purpose, it is clear that

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this dopant molecule contains all of the essential machinery required to operate as a vibrational Stark probe.

4.1. Coulombic Interactions and the Vibrational Stark Effect in F4TCNQ-Doped P3HT Films. The fact that the difference dipoles of the B1u and B2u modes point in perpendicular directions allows us to directly infer the orientation of the dopant anion relative to the polaron on the polymer backbone that generates the local electric field. As mentioned above, X-ray diffraction work from several groups has determined that F4TCNQ\(^-\) sits in the lamellar side-chain regions of the P3HT crystallites.\(^{3,4,11,12,24,36,33,35,36,60,61,63,64}\) In addition, Brinkmann and co-workers, who studied rub-parallel aligned F4TCNQ-doped P3HT films by using polarized absorption spectroscopy, have shown that the long axis of the F4TCNQ\(^-\) anion sits perpendicular to the polymer backbone.\(^{62,63,64}\) This means that the difference dipole vector of the B1u mode is oriented parallel to the electric field lines emanating from the hole on the polymer backbone (Figure 3), while that of the B2u mode lies perpendicular to the local field.

Figures 3a,c illustrate how the VSE shifts the resonant energy of the F4TCNQ\(^-\)\(B_{1u}\) mode in direct response to changes in the magnitude of the P3HT hole-generated electric field. At fixed \(d_{\text{anion}}\), the strength of the field will decrease as the polaron becomes increasingly delocalized. This provides a simple explanation as to why the frequency of the F4TCNQ\(^-\)\(B_{1u}\) mode, whose difference dipole is oriented parallel with the field lines from the polaron, exhibits a strong vibrational Stark shift in response to changes in polaron coherence length (i.e., with A/B spectral ratio, as seen in Figure 1e).

In contrast, the peak vibrational energy of the F4TCNQ\(^-\)\(B_{2u}\) mode does not change with polaron coherence length; it only broadens. This is because the orientation of the B2u mode difference dipole is mostly orthogonal to the hole-generated electric field. This means that any Coulombic interaction between the polaron and the difference dipole of this mode must originate from an off-axis electric field component. In the simplest case where the polaron is centered on the polymer-dopant axis (Figures 3a,c), there is no net off-axis field component that might be aligned with the B2u mode, explaining why this peak does not shift in response to changes in polaron coherence.

However, there is no guarantee that a polaron will always be centered along the dopant–polaron axis; in fact, we expect some degree of along-the-chain disorder of the dopant location relative to the polaron position in F4TCNQ-doped P3HT films. This means that the magnitude of the polaron \(E_{\text{field}}\) component aligned with the B2u mode dipole will increase not only with the amount of off-center displacement but also with the degree of polaron delocalization, as illustrated in Figures 3b,d. If the total lateral disorder is modest, the result will be a net broadening of the B2u mode rather than a shift, exactly as observed in Figure 1f. This is because, on average, the polaron is still centered relative to the location of the anion, so the off-axis component is relatively weak compared to

Figure 3. Cartoons illustrating how the electric field from a variety of polaron geometries influences the vibrations of the F4TCNQ anion. (a) A localized P3HT polaron centered along the dopant–polymer axis produces a strong electric field aligned with the F4TCNQ\(^-\)\(B_{1u}\) difference dipole, resulting in a VSE shift to a higher vibrational energy. The centered geometry has no electric field component aligned along the \(B_{2u}\) difference dipole, so there is no VSE shift of that mode. (b) A localized P3HT polaron with a slightly off-axis geometry slightly lowers the electric field component experienced by the \(B_{1u}\) vibration and slightly increases that experienced by the \(B_{2u}\) vibration, leading to a slight broadening of both modes. (c) A more delocalized P3HT polaron centered along the dopant–polymer axis exerts a lower field strength on the F4TCNQ anion than the localized polaron case in panel a, resulting in a smaller VSE shift of the \(B_{1u}\) mode, which appears at a lower vibrational energy. (d) A more delocalized P3HT polaron with an off-axis geometry has little effect on the \(B_{1u}\) mode, but the increased electric field component along the \(B_{2u}\) mode leads to an increase in VSE broadening.
the field strengths experienced along the $B_{1u}$ difference dipole direction.

It makes sense that there should be a relatively strong correlation between the width of the $F_4TCNQ^-$ $B_{2u}$ nitrile stretch and the degree of polaron delocalization, but it is also possible that broadening of the $B_{2u}$ mode might occur if the dopant anions had significant orientational disorder relative to the direction of the electric field from the polaron. If there were orientational disorder, however, the $B_{2u}$ peak would be expected to narrow as the degree of local polymer crystalline order increased. Instead, we observe the opposite: the $B_{2u}$ peak broadens as the polymer film becomes more ordered. This verifies that the observed correlation of the anion vibrational spectrum with the delocalization of the polaron indeed comes from the most ordered regions in the doped films and is not representative of orientationally disordered areas of the film.

Finally, it is conceivable that at sufficiently high doping levels a second polaron on a neighboring chain could exert a field on the $F_4TCNQ$ molecule that would work to counteract the field from the first polaron, reducing the observed Stark shift. However, Figure S13a shows that the magnitude of the Stark shift does not change over a wide range of doping concentrations. This confirms that the doping levels we used to analyze the Stark shifts in this work are not high enough to encounter this effect, though it might be observed at higher dopant to monomer ratios.

4.2. Stark Tuning Rate: Estimating Local Electric Field Strengths with Vibrational Mode Shifts. The above evidence strongly indicates that the VSE can explain all of the observed changes in the $F_4TCNQ^-$ nitrile vibrations as resulting from changes in polaron delocalization in doped P3HT films. In the next sections, we work to quantify the strength of these interactions to experimentally extract the anion–hole distance and degree of polaron delocalization.

It is well established that there is a linear relationship between the shift of a vibrational mode and the strength of the applied field. The magnitude of the vibrational shift per unit applied field is known as the Stark tuning rate (STR) parameter. To the best of our knowledge, the STR parameters have only been experimentally measured for a very limited number of conjugated molecules. For Stark probe molecules with multiple nitrile groups, similar to $F_4TCNQ$, the STRs tend to fall within the range of $1–3 \text{ cm}^{-1}/(\text{MV cm}^{-1})$. This is considerably higher than the STR’s of isolated nitrile groups, which typically are a factor of 2 smaller.31,32,42 Suydam and Boxer experimentally measured the STRs for a variety of mono- and disubstituted nitrile containing conjugated molecules.30 These workers reported a STR of $2.9 \text{ cm}^{-1}/(\text{MV cm}^{-1})$ for the $B_{2u}$ mode and $1.4 \text{ cm}^{-1}/(\text{MV cm}^{-1})$ for the $B_{1u}$ mode of a dicyanomethylene–polyene compound of similar size to $F_4TCNQ$. We will assume that these values also represent along-the-chain disorder, along the $F_4TCNQ$ $B_{2u}$ direction. The magnitude of this component is an order of magnitude smaller than that along the $B_{1u}$ direction.

determine how the polaron coherence length changes across the experimentally observed range of $A/B$ values. We begin by considering the predicted magnitudes of the fields experienced by the $B_{1u}$ and $B_{2u}$ modes as a function of polaron delocalization and the anion–hole separation distance, $d_{anion}$. For this calculation, we consider the field experienced by the anion at a single point located at $d_{anion}$ away from the center of the P3HT chain. We then approximate polaron delocalization as resulting from a single positive charge spread out uniformly along a one-dimensional line of length $L$, given in units of the number of P3HT monomers. We then model changes in polaron coherence as $L$ increases from 1 to 9 (Figure 4a), so that the one-dimensional polaronic charge density goes as $1/L$. Our model assumes a dielectric constant of unity, as the short-range interactions between the polaron and anion likely do not experience the dielectric value of the bulk material—an assumption which is consistent with what others have chosen.
for similar inquiries. We note that our model ignores interchain polaron coherence—a choice made because work from Ghosh and co-workers predicts that the way in which the coherence length changes with the spectral A/B ratio is roughly the same regardless of whether the delocalization is spread over one chain or between multiple chains. Because of this, we do not expect delocalization between neighboring chains to significantly affect the electric fields experienced by a nearby F4TCNQ anion. Finally, even though we expect some triangles), the relative electric field is an interaction described by Coulomb's law.

The main difficulty associated with quantitatively determining the change in P3HT polaron coherence length from the VSE of the F4TCNQ nitrile stretching modes is that we only have access to the experimentally measured vibrational shifts that occur within a range of externally applied fields: we do not know the absolute magnitudes of these fields because we do not know the position of the B1u stretching mode inside a P3HT film in the absence of any external fields. Thus, the best we can do is to use the measured shift in the B1u peak vibrational energy, along with the STR, to determine the change in electric field associated with the ~25% increase in polaron coherence length measured by the spectral A/B ratio in Figure 1. Once we have calculated the change in electric field experienced by the B1u mode, we can then use the measured change in width of the B2u mode, which experiences a field from the same delocalized polaron, to experimentally determine both the value of dni, and the physical polaron coherence length increase that can explain the vibrational Stark features of both modes.

Figure 4b shows how the strength of the electric field aligned with the B1u difference dipole is affected as polaron delocalization is increased from 1 to 9 monomer units. As argued above, the maximum possible dopant–polaron distance in a P3HT crystal is half the lamellar spacing, or only ~7–8 Å. Thus, by examining a range of possible anion–hole distances up to 8 Å, we can narrow down the possible range of fields exerted by a P3HT polaron on the nitrile stretching modes of F4TCNQ-. In Figure 4b, the dark blue squares corresponding to dni = 8 Å show that increasing the degree of polaron delocalization diminishes the strength of the electric field experienced by the anion to ~39% of its original value. Because this is the largest possible anion–hole distance, it also represents the minimum possible decrease in the relative field strength for a 1 to 9 unit change in polaron delocalization. For shorter anion–hole distances, such as dni = 5 Å (red triangles), the relative electric field strength drops to ~26% of its starting value as the polaron delocalizes from 1 to 9 monomer units. We do not expect more sophisticated models to deviate significantly from these trends, since fundamentally this is an interaction described by Coulomb's law.

For the B2u mode, our simplified model does not predict any net electric field alignment with the difference dipole. However, we can approximate the effects of along-the-chain disorder (cf. Figures 3b and 3d) by considering the field contribution from just half of the delocalized polaron along the positive axis. Figure 4c shows the magnitude of the electric field component from half of the delocalized charge that is aligned parallel with the B2u mode as a function of both polaron coherence length and dni. In this case, we see that the electric field does not vary monotonically with polaron delocalization because of competing effects. At shorter polaron coherence lengths, additional delocalization creates additional

off-axis field alignment, which raises the overall magnitude of the field component along the B2u difference dipole. However, at longer coherence lengths, additional delocalization lowers the line charge density, which contributes to a net reduction in the total magnitude of the field.

In our simple model, the predicted magnitudes of the fields experienced by the F4TCNQ- B2u mode are nearly an order of magnitude lower compared to those experienced by the B1u mode. This is indeed consistent with the fact that the B2u mode only slightly broadens and does not shift with changing P3HT polaron delocalization, as measured by the A/B spectral ratio, while the B1u mode shows a definite Stark shift as polaron coherence changes with the local polymer order.

As mentioned above, theoretical work based on Spano’s model indicates that our range of measured A/B ratios corresponds to an ~25% increase in P3HT polaron coherence length; this change in coherence is the same whether or not the delocalization is along or between the polymer chains. With an estimated B2u STR of ~2.9 cm−1/(MV cm−1) and the observed (single edge) vibrational peak broadening of 2.2 cm−1, the observed 25% change in polaron delocalization corresponds to a change in electric field of 0.76 MV cm−1. Note that because the Efield alignment does not change monotonically for this mode, we cannot determine the sign of this change or the distance from which the delocalized polaron exerts the field.

However, the same delocalized polaron electric fields that change the width of the B2u mode also cause the Stark shift of B1u mode. Pairing our estimated STR parameter for the B1u mode as ~1.2 cm−1/(MV cm−1) with the experimentally measured vibrational shift of 5.79 cm−1, the electric field experienced by this mode decreases by 4.9 MV cm−1 over this range of A/B ratios. Figures 4b,c show that the only way for F4TCNQ- to experience both of these field components for an ~25% change in polaron coherence at a single distance is when the intrachain coherence length increases from 6 to 7.5 monomer units with an anion–hole separation distance of ~6 Å. A set of tables summarizing the possible changes in field strength for different coherence lengths and separation distances can be found in the Supporting Information. Thus, by use of the VSE, it is possible to experimentally link the spectral A/B ratio to a specific degree of intrachain polaron delocalization.

Rather than assuming the ~25% change in delocalization length from theory, we can also do the same analysis by first assuming a fixed ~6 Å distance for dni based on X-ray diffraction. The measured field-induced changes of the two F4TCNQ- modes are then only consistent with a coherence length change of ~25% from 6 to 7.5 monomer units, so that no matter what our starting assumptions, there is only a single internally consistent value for the experimentally determined intrachain polaron coherence lengths.

Thus, by using the two orthogonally oriented nitrile vibrational modes of F4TCNQ- as independent measurements of the electric field, we can build a comprehensive picture of the charge carrier–dopant interactions in F4TCNQ-doped P3HT films. The experimentally determined change in intrachain polaron coherence length from 6 to 7.5 monomer units is consistent with what has been put forward previously for this system. The Stark-determined dni, of 6 Å also fits well with the X-ray scattering determined lamellar stacking distances (see Table S3) as well as previously modeled anion distances based on the profile shape of the mid-IR polaron absorbance of F4TCNQ-doped P3HT system.
5. CONCLUSIONS

In summary, we have shown that the nitrile stretches of the commonly used dopant \( \text{F}_4\text{TCNQ} \) make an exquisite vibrational Stark probe of the local electric field in doped conjugated polymer films, allowing us to uncover detailed information about P3HT polaron coherence. The fact that we observe one mode shift with polaron coherence while the other only broadens directly demonstrates that the dopant anion has a preferred orientation with respect to the polymer backbone. This is consistent with the idea that dopants prefer to reside in the lamellar regions of polymer crystallites, reinforcing conclusions in the literature reached by X-ray scattering \( ^{6,11,12,24,26,53,60,61,63,64} \) and by polarized spectroscopy of rub-aligned films. \( ^{52,65,68–70} \)

More importantly, we have provided the first experimental evidence that shows it is the degree of polaron delocalization that is directly responsible for the commonly seen changes in the \( \text{F}_4\text{TCNQ} \) nitrile vibrational spectrum in doped P3HT films with varying degrees of crystalline order. This verifies theories that argue the polaron A/B spectral ratio (and position of the B polaron absorption band) are indeed direct experimental signatures of polaron coherence. With a simple model, we can use the magnitudes of the observed vibrational spectral shifts and broadening to pin down precisely how the local electric field changes with polaron delocalization in different P3HT environments. The field magnitudes experimentally prove that the anion–polaron distance in \( \text{F}_4\text{TCNQ} \)-doped P3HT films is \( \sim 6 \, \text{Å} \) and that intrachain polaron coherence lengths of roughly \( 6–7.5 \, \text{monomer repeat units} \) can be produced by varying the degree of local order through the film processing conditions.

Our results also allow us to explain the previous uncertainty over how to assign the vibrational spectrum of \( \text{F}_4\text{TCNQ} \) in doped P3HT films. The various vibrational lines that have been observed occur as a direct result of the unique preparation conditions and dopant/polymer blend ratios which produce a range of structural morphologies and polaron delocalization lengths. This in turn causes the dopant anions to experience a range of local electric fields, ultimately shifting the resonant vibrational energies. This also means that using the positions of the \( \text{F}_4\text{TCNQ} \) anion nitrile vibrations to determine the precise degree of charge transfer in charge-transfer complexes, usually associated with a feature near \( 2200 \, \text{cm}^{-1} \), is potentially fraught with difficulty as electric fields from nearby integer charge-transfered polaron may also be responsible for shifts of this mode.

For the \( \text{F}_4\text{TCNQ} \)-P3HT materials combination studied here, the relatively modest intrachain polaron coherence lengths we measure are a direct result of the fact that the \( \text{F}_4\text{TCNQ} \) anion always resides within \( 1 \, \text{nm} \) of the polaron, so that polaron coherence is always limited by the local Coulombic interaction with the anion, independent of how well ordered the P3HT crystallites are. Thus, although \( \text{F}_4\text{TCNQ} \) is certainly a useful molecular dopant for studying polaron behavior in conjugated polymers, the relatively strong Coulombic interaction it partakes in puts a fundamental limit on the degree of polaron coherence for this polymer–dopant combination. Fortunately, these issues can be avoided with the use of other dopants that better screen the polaron–counterion Coulombic interaction. \( ^{13,14} \)

We close by noting that in many ways measuring the electric field landscape in doped semiconducting materials is precisely the kind of application that vibrational Stark probes were designed to explore. The nitrile groups that make this kind of analysis possible are commonly added to p-type dopants to make them stronger oxidizers. Thus, application of dopants as VSE probes is a natural use for these molecules. Although the current generation of nitrile-containing dopants like \( \text{F}_4\text{TCNQ} \) were not designed to be used this way, it is exciting to imagine an entirely new class of p-type dopants that could be specifically engineered to fulfill the dual roles of being strong oxidizers that also map the electric fields from polarons in these materials. For these reasons, we are optimistic that the VSE will become a more generally applied principle in this area for probing the Coulombic interactions in doped conjugated polymers.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.chemmater.1c02934. Detailed information about materials used, fabrication and doping of polymer films, FTIR measurements, peak fitting of the \( \text{F}_4\text{TCNQ} \), C–N vibrational spectra, tabulated fitting parameters, dopant–polaron Coulombic modeling, and GIWAXS measurements of doped polymer films (PDF)

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Notes
The authors declare no competing financial interest.

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