Effects of thermal disorder on the electronic properties of ordered polymers

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The effects of thermal disorder on the electronic properties of crystalline polymers were investigated. Atomic configurations of the material were obtained using classical Monte Carlo simulations at room temperature, while electronic structure calculations were performed using the density functional theory based charge patching method and the overlapping fragment method. We investigated two different stable configurations of crystalline poly(3-hexylthiophene) (P3HT) and calculated the density of electronic states and the wave function localisation. We found that the effect of disorder in side chains is more pronounced in the more stable configuration of P3HT than in the other one due to the larger conformational freedom of side chains. The results show that disorder in main chains has a strong effect on the electronic structure and leads to the localisation of the wave functions of the highest states in the valence band, similar to localisation that occurs in amorphous polymers. The presence of such states is one possible origin of thermally activated electrical transport in ordered polymers at room temperature.

1 Introduction

Materials based on conjugated polymers attract a lot of interest due to their applications in electronic devices such as organic field-effect transistors, organic light-emitting diodes and organic solar cells.\textsuperscript{1-5} These materials contain both crystalline and amorphous regions and consequently their electronic properties strongly depend on their morphology. In crystalline regions of the material, thiophene rings are connected into main (backbone) polymer chains, which then form two dimensional lamellar structures separated by insulating side chains.

Crystalline polymer regions exhibit better transport characteristics than amorphous regions. At first, one may expect that the electronic states in crystalline regions are fully delocalised due to the effect of periodicity. On the other hand, wave functions of electronic states in amorphous regions are well localised due to the effects of disorder. Calculations of the electronic structure of amorphous polymers show that wave functions of the highest states in the valence band are localised on few rings only.\textsuperscript{6,7} The electronic states in crystalline regions may also exhibit localisation due to the effects of thermal disorder; at finite temperature the atoms are displaced from their equilibrium positions in a random manner and the atomic positions no longer exhibit periodicity. The importance of the effects of thermal (or dynamic, which is an alternative expression often used in the literature) disorder in small molecule based organic crystals is now widely recognized.\textsuperscript{8-14} On the other hand, the effects of thermal disorder in polymers are less well understood. Currently available simulation results\textsuperscript{15-17} suggest that the highest valence band states in ordered polymer materials at room temperature are localised. However, most of these calculations either consider a single polymer chain or do not include the effects of side chains.

The aim of this work is to investigate in detail the effects of thermal disorder on the electronic properties of ordered polymer materials and to identify relative importance of various sources of thermal disorder. For concreteness, we choose the poly(3-hexylthiophene) (P3HT) polymer for our study. We calculate the electronic density of states (DOS) and the localisation of hole wave functions. Atomic configurations at finite temperature are obtained using Monte Carlo (MC)\textsuperscript{18} simulations, while the charge patching method (CPM)\textsuperscript{6,7,19,20} and the overlapping fragment method (OFM)\textsuperscript{7,21} are used for electronic structure calculations.

The disorder in the structure at finite temperature comes both from the disorder in the shape of the flexible alkyl side chains (that will be referred to as side chain disorder from now on) and from the disorder due to variations of torsion angles between thiophene rings in the main chains and the position of main chains (that will be referred to as main chain disorder from now on).\textsuperscript{22} One should note that disorder of main or side chains does not imply an amorphous material. While it is well known that the wave functions in conjugated polymers are localised on the main chain, side chains with a disordered

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shape still affect the electronic structure as they create a disordered electrostatic potential on the main chain. Disorder in the shape of the main chain mostly affects the electronic structure through variations in electronic coupling between the rings. To isolate the effects of main and side chain disorders we investigate three types of structures: (1) the structures with straight main chains and disordered alkyl side chains; (2) the structures with disordered main chains in the absence of side chains; (3) the structures with both main and side chains disordered.

We find that the effects of disorder are least pronounced in the structures with disordered side chains. Strong wave function localisation of highest valence band states occurs in structures with main chains disordered, both with and without side chains. Such a localisation is one possible origin of thermally activated transport observed in all reported P3HT mobility measurements.

2 Methods for atomic and electronic structure calculations

The method used for electronic structure calculations is schematically described in Fig. 1. The atomic structure is obtained using in-house developed code for MC simulations. It is subsequently used to calculate the electronic structure, using the CPM and the OFM.

2.1 Lattice constants

The initial structure for MC simulations is the ideal crystalline structure of P3HT. The atomic structure of crystalline P3HT was extensively studied and different possible configurations were found. Two different stable crystalline structures are simulated at 300 K: aligned and shifted, which are shown in Fig. 2. In the aligned structure, thiophene rings from two adjacent main chains in the \( \pi-\pi \) stacking direction are aligned. On the other hand, in the shifted structure, thiophene rings from two adjacent chains in the \( \pi-\pi \) stacking direction are mutually shifted by the half of the unit cell in the main chain direction. Parameters of the unit cells are found using NPT (constant pressure and temperature) MC simulation at 300 K and 101.325 kPa. During the simulation, the size of the box in the main chain direction is kept constant, while two other dimensions are changed. The energy of the system is modelled as a sum of the long-range (van der Waals and Coulomb) interactions between atoms from different chains. Expression and parameters for the long-range interactions are taken from the OPLS parameters set, which was previously successfully applied for the simulations of the same material. Cutoff for these interactions is taken to be 12 Å. The unit cell parameters are determined as the parameters obtained when the system reaches thermal equilibrium. For the aligned structure these are: \( a/2 = 15.7 \, \text{Å}, \quad b = 8.2 \, \text{Å} \) and \( c = 7.77 \, \text{Å} \), while for the shifted structure these are: \( a/2 = 15.7 \, \text{Å}, \quad b = 8.1 \, \text{Å} \) and \( c = 7.77 \, \text{Å} \). All unit cell angles are taken to be 90°. These parameters are in very good agreement with previous computational results for the same structures of P3HT. For example, the unit cell parameters obtained for the aligned structure in ref. 24 are: \( a/2 = 16 \, \text{Å}, \quad b = 8.2 \, \text{Å} \) and \( c = 7.81 \, \text{Å} \), while for the shifted these are: \( a/2 = 16 \, \text{Å}, \quad b = 7.85 \, \text{Å} \) and \( c = 7.81 \, \text{Å} \). On the other hand, experimental results based on the X-ray diffraction measurements suggest somewhat higher value for the side chain stacking direction of 16.8 Å. This difference might originate from the assumption of the ideal crystal structure without disorder made in the calculations, which is not the case in reality. Interdigitation between side chains from different lamellas is weaker in more disordered structures, and consequently, the unit cell parameter for the side chain stacking direction is higher. We find that the shifted structure is more stable at 300 K, since its potential energy per number of rings is 0.38 eV lower than the corresponding energy in the aligned structure, confirming the
results given in ref. 24 that the shifted structure is more energetically favourable.

2.2 Atomic structure

With the initial structure at hand, MC simulations are performed to obtain the snapshots of the atomic structure at room temperature. During the MC simulations bond lengths and bond angles are kept constant, while some or all torsion angles are changed. Variations of torsion angles affect electronic coupling between orbitals more strongly than variations of bond lengths and bond angles. For example, the thermal energy at room temperature $k_B T = 25 \text{ meV}$ leads to displacement of an atom due to bond stretching on the order of $0.02 \AA$, while the same energy leads to interring torsion angle change on the order of $45^\circ$, which yields the atomic displacements of more than $1 \AA$. For this reason, it is reasonable to keep bond angles and bond lengths constant. In each step of the MC simulation a new configuration is generated by changing the torsion angles and moving the whole polymer chain. The new configuration is accepted if it satisfies the Metropolis condition.\textsuperscript{18} The energy of the system is calculated as a sum of torsion potentials and long range van der Waals and Coulomb interactions. Thiophene–thiophene torsion potential is taken from ref. 6, while thiophene-side chain torsion potential is taken from ref. 24. Long-range interactions are modelled in the same way as for lattice constants calculations. An OPLS parameter set is also used for the torsion potentials within side chains. Periodic boundary conditions are applied in each direction of the unit cell for the systems with disorder in side chains. Boundary conditions are open in the main chain direction for the systems with disorder in main chains. The final atomic structure is taken after the system is thermally equilibrated at 300 K, which is evidenced from the saturation in the dependence of potential energy on the number of MC steps.

2.3 Electronic structure

The final atomic configuration obtained from MC simulations is used as input for electronic structure calculations. For the system containing more than thousand atoms density functional theory (DFT) calculations would be too computationally demanding. Therefore, the CPM is applied instead of DFT. The main idea of the CPM is to calculate the electronic charge density directly, instead of obtaining it from self-consistent solution of Kohn–Sham DFT equations. The charge density is calculated as a sum of contributions (called motifs) of all atoms in the system. Motifs are extracted from the DFT calculations of a small prototype system containing 3 neighbouring rings of P3HT and they are the same as used in ref. 6. The CPM was previously applied to calculate the electronic structure of amorphous P3HT.\textsuperscript{5,7} The details of the CPM are presented in ref. 19.

When electronic charge density is obtained, the single-particle Kohn–Sham Hamiltonian is constructed by solving the Poisson equation for the Hartree potential and using the local density approximation (LDA) for the exchange–correlation potential.\textsuperscript{19} The single particle Hamiltonian is then diagonalized using the OFM. This method uses the eigenstates of small fragments of the system as the basis set. In this case, fragments are systems of 3 neighbouring rings of P3HT (trimers). In previous work we demonstrated that a good basis set is obtained when the fragments mutually overlap.\textsuperscript{7} To calculate the electronic states in the desired spectral region (within 0.5 eV below the top of the valence band) with good accuracy, only one eigenstate (the HOMO state) of each fragment is needed. When the basis set is obtained, transfer and overlap integrals between basis states are calculated and subsequently, the Hamiltonian generalized eigenvalue problem is solved. These calculations can be done using parallel computer architectures, which saves a lot of computational time. A detailed description of the OFM can be found in ref. 21.

In this work, the effect of polarons was not included. Previous DFT calculations of long straight polythiophene chains at zero temperature indicate that polaron binding energy is on the order of few meVs only and that it can be ignored.\textsuperscript{35,36} However, it is more difficult to assess the role of polarons at finite temperature when thermal disorder is present in the material. Since the main goal of this work is to understand the effect of thermal disorder on wave function localisation and DOS, as well as the contributions from main and side chains to thermal disorder, polaronic effects were not considered.

Next, we discuss the appropriateness of LDA for the description of the localisation effects. The localisation effects that we observe essentially come from two effects: (i) variations of on-site energies of rings; (ii) variations in electronic coupling between the rings. Effect (i) comes mainly from disordered long range electrostatic potential that side chains, the rest of the main chain and other main chains produce on a certain ring. Within DFT, electrostatic potential is taken into account through the Hartree term in the Kohn–Sham equation, which is an exact term. Therefore, for effect (i), the use of LDA in our calculation is not an issue. To check if LDA gives reliable values of electronic coupling (effect (ii)), we have performed the calculation of electronic energy levels of a ten unit long straight thiophene oligomer using either the LDA or B3LYP functional for the same atomic configuration. These calculations have been performed using the NWChem code\textsuperscript{7,34} and the 6-31G basis set was used to represent the wave functions. We find that the spacing between energy levels calculated using these two functionals differs typically by 15%. Since the spacing between energy levels of straight oligomers is proportional to electronic coupling between the rings, we conclude that possible uncertainties in electronic coupling calculated using LDA are on the order of 10–20% and such uncertainties are not expected to significantly affect the localisation lengths.

3 Results and discussion

The effects of thermal disorder in crystalline P3HT are investigated by examining 3 different types of structures: the structures with disorder in side chains, the structures with disorder in main chains and the structures with disorder in both side and main chains. In the first case, main chains are kept rigid during the MC simulation, while side chains are allowed to
move freely. In the second case, side chains are removed (more precisely, replaced with a hydrogen atom) before the electronic structure calculations start, in order to isolate the effects of main chain disorder. In the case of structures with the presence of disorder in both main and side chains, the same atomic configurations are used as for the second case, but side chains are not removed in this case. To get sufficiently large statistics, for each of the investigated cases and for both aligned and shifted structures, 100 different realizations are generated. Each configuration contains 10 polymer chains stacked in the π–π direction, while each chain contains 10 rings. The total number of atoms in each configuration is 2520. For subsequent analysis only 10 highest states from each configuration are taken into account, since they cover a spectral range of about 0.5 eV below the HOMO level, which is the range of interest for the electrical transport properties. To analyse the effects of disorder, we calculate the DOS and the localisation length of hole states.

### 3.1 Density of electronic states

Densities of states obtained from the calculations are shown in Fig. 3. In the case of disorder in side chains in the aligned structure, DOS is nearly discrete, composed of several peaks (Fig. 3a). These peaks correspond to the peaks in the DOS of the ideal crystalline structure without any disorder. In the shifted structure, peaks are broader and overlap more than in the aligned structure, making DOS continuous (Fig. 3b). Thus, the effects of side chain disorder on DOS are more pronounced in the shifted structure than in the aligned. The difference in DOS for shifted and aligned structures can be explained by the difference in the spatial distribution of side chains in these two structures. In the aligned structure, side chains connected to the aligned thiophene rings from neighbouring chains in the π–π direction are at the same side of the main chains (chains denoted by 1 and 2 in Fig. 2a). On the other hand, in the shifted structure these side chains are at the opposite sides. The distance between nearest side chains in the shifted structure (chains denoted by 1 and 2 in Fig. 2b) is greater than in the aligned. Therefore, side chains in the shifted structure have more conformational freedom than in the aligned. This is evidenced by the distributions of the thiophene-side chain torsion angle for both structures, given in Fig. 4a and b. The distribution of thiophene-side chain torsion angles is significantly wider in the shifted structure, which results in a higher degree of side chain disorder. This difference in morphology leads to the difference in the electronic structure.

In the structures with disordered main chains, DOS is continuous (Fig. 3c and d). In the case of aligned structures, distribution of energies is significantly wider than in the shifted structure. Distributions of thiophene–thiophene torsion angles for the aligned and shifted structures are similar (Fig. 4c and d) and agree well with the results of ref. 16. Therefore, the difference in the distribution of the energies is not caused by the shape of the chains. Electronic coupling between different chains is 0.11 eV in the ideal aligned structure and 0.07 eV in the ideal shifted structure. This substantial difference leads to the wider distribution of the energies in the aligned structure. When both main and side chains are disordered, the DOS is continuous (Fig. 3e and f) without any apparent difference between structures. This similarity in DOS can be explained by the effect of compensation between wider energy distribution in the shifted structure when only side chains are disordered and wider energy distribution in the aligned structure when only main chains are disordered.

A suitable measure of disorder in the system is the distribution of diagonal Hamiltonian elements given by $H_{ii} = \langle i | H | i \rangle$, where $|i\rangle$ are wave functions of trimers and $H$ is the Hamiltonian of the system. Wave functions of the trimers are localised on the main chain and on the carbon atoms in the side chains closest to the thiophene rings. Consequently, if main chains of trimers are rigid, their wave functions and energies of HOMO levels will be equal. When only disorder in side chains is applied, the difference between diagonal Hamiltonian elements arises only
from variations in $H$, due to the variations in the electrostatic potential caused by side chain disorder. On the other hand, when disorder in main chains is present, variations in $H_{ii}$ arises both from $H$ and $|i\rangle$, since wave functions of trimers now differ significantly. Distributions of diagonal elements of the Hamiltonian are given in Fig. 5. As expected, distributions are widest in the case when both disorders are present (Fig. 5e and f). Having in mind the results for DOS presented above, which suggest that disorder in main chains has more impact on the electronic structure of P3HT than disorder in side chains, one may find unexpected that distributions given in Fig. 5c and d are similar to the distributions given in Fig. 5a and b. Side chains have more conformational freedom than main chains, especially in the shifted structure. Therefore, their disorder affects the electrostatic potential more than disorder in main chains. When effects of disorder in main chains are isolated, side chains are removed and, consequently, variations of electrostatic potential are weaker than in the case of disorder in side chains, which leads to the weaker variations in $H$.

### 3.2 Wave function localisation

Wave functions of HOMO levels for 6 different cases are shown in Fig. 6. In the ideal crystalline structure the wave function of the HOMO level (and any other level) is completely delocalised, as Bloch theory predicts. When the side chain disorder is partially applied, wave functions remain delocalised (Fig. 6a and b). They are not delocalised along the whole structure, as in the ideal structure. Delocalisation is broken both in the $\pi-\pi$ stacking direction and in the main chain direction. On the other hand, wave functions of HOMO levels in the case of disorder in main chains are localised, both with and without side chains included (Fig. 6c–f). They are localised on 5–15 rings, usually on 2 neighbouring chains (as in Fig. 6c and e). Therefore, thermal disorder in the crystalline P3HT localises the wave function of the HOMO level, as in the amorphous phase. 6,7

To investigate the effects of disorder on the wave function localisation more precisely, we calculate two localisation lengths for each state: localisation in the $\pi-\pi$ stacking direction $L_\pi$ and localisation in the main chain direction $L_c$. If the wave functions are represented in the orthonormal basis set of well localised orbitals, the localisation length can be generally defined as

$$ L = 1 \left/ \sum_m |d_m|^4 \right., $$

where $d_m$ is the expansion coefficient of the wave functions in the orthonormal basis $|m\rangle$. The basis set used in the OFM calculations is not orthonormal. The orthonormal basis set is constructed by transformation $|m\rangle = \sum_m T_{m\ell} |\ell\rangle$ using transformation matrix $T = (S^{-1/2})^\ast$, where $S$ is the original overlap matrix and $|\ell\rangle$ are original basis wave functions. Expansion coefficients of the orthonormal basis set are related to the original coefficients $c_i$ via $d_m = \sum_\ell (S^{1/2})_{m\ell} c_i$. For the orthonormal basis set the condition $\sum_m |d_m|^2 = 1$ is satisfied. In our case, this sum can be divided into two sums, one over different chains and other over rings in one chain: $\sum_{j=1}^{N_c} \sum_{j=1}^{N_r} |\hat{\mu}_j|^2 = 1$, where $\hat{\mu}_j$ is the localisation length in the $j$-th ring of the $i$-th chain.
where \( N_c \) and \( N_m \) are the number of chains and the number of rings within one chain, respectively. Following the general definition of the localisation length, \( L_b \) is defined as \( L_b = 1 / \sum_{i} |\beta|^2 \), where

\[
|\beta|^2 = \sum_{j=1}^{N_m} |\beta_j|^2
\]

Similarly, \( L_c \) is defined as \( L_c = 1 / \sum_{j} |\beta''|^4 \), where

\[
|\beta''|^4 = \sum_{i=1}^{N_c} |\beta_i|^2
\]

Plots of the dependence of \( L_b \) on the energy of the electronic states are shown in Fig. 7. In the ideal crystalline structure (both aligned and shifted), \( L_b \) of the HOMO level is equal to the number of chains, which is 10 in this case. Lower states in the ideal structure have an \( L_b \) of either 6.67 or 10. In the case of the aligned structure with disordered side chains (Fig. 7a), the values of \( L_b \) for the HOMO level vary from 4 to 10. Other states have \( L_b \) which is around the value of \( L_b \) in the ideal structure. In the case of the shifted structure, distribution of the energies of states is wider than in the aligned. Consequently, it is difficult to isolate the values of \( L_b \) for HOMO levels from Fig. 7b. Looking into the range of 0.2 eV below the highest energy, the value of \( L_b \) is between 3 and 6, which is lower than the values of \( L_b \) in aligned structures. Therefore, wave functions of HOMO levels are more localised in the shifted structure than in the aligned. In all remaining cases (Fig. 7c–f) we get qualitatively similar results. Values of \( L_b \) for the highest occupied states are low, they take values from 2 to 4 chains. States with lower energies have wider distributions of \( L_b \), suggesting that delocalised states exist. The shape of the plot of \( L_b \) is similar to the plot of the hole localisation length of the amorphous P3HT, given in ref. 7.

Results for \( L_c \) are similar to the results for \( L_b \). The minimal value for \( L_c \) when both side and main chains are disordered is 3. Electronic coupling is stronger in the main chain direction than in the \( \pi-\pi \) stacking direction and therefore localisation in the \( \pi-\pi \) stacking direction is stronger than in the chain direction.

It is interesting to note that the highest states in the valence band are typically localised on 2 neighbouring chains when main chains are disordered. The state will be localised on two chains if the electronic coupling \( t_{mn} = \langle m | H | n \rangle \) between orbitals \( m \) and \( n \) from different chains is greater than variations of the diagonal Hamiltonian elements. The distributions of the quantity \( d = t_{mn} - |H_{mn} - H_{nn}| \) are given in Fig. 8. Since \( d \) takes positive values as well, existence of strong coupling between chains is confirmed, which explains localisation on two neighbouring chains.

The total localisation length can be found using the general definition previously given. A plot of its dependence on the energy of the states is given in Fig. 9. For the highest states, when main chains are disordered, it takes values from 5 to 15 rings (Fig. 9c–f). These values are slightly higher than the values of the localisation length of highest states in the valence band of amorphous P3HT, which is around 5. This difference is expected, since crystalline P3HT, despite the high degree of disorder, is still more ordered than amorphous. In the sense of the hole localisation length, effects of disorder in crystalline and amorphous P3HT are similar. Delocalised states (localised on more than 10 rings) start to appear only few hundreds of meV below the top of the valence band. This is in agreement with the findings presented below.
in ref. 17 where DOS and localization length in PBTTT are calculated. Results for the carrier localization orbital density of HOMO levels of crystalline P3HT at 300 K, given in ref. 15, indicate the presence of both localised (4–10 rings) and weakly (more than 10 rings) localised states. These calculations were performed without alkyl side chains and for isolated main chains. We obtain qualitatively the same results for $L_c$, since we find states localised on few rings within a chain and states that are extended over the entire chain. To conclude, our findings agree with previous findings that wave functions of the highest states in the valence band are localised and that delocalised (or weakly localised) states also exist below these states.

Results for total localization length and $L_q$ are qualitatively similar for the structures with isolated disorder in main chains and with both disorder in main and side chains. Nevertheless, side chains have a significant quantitative effect on electronic properties. This can be seen by comparing the DOS (Fig. 3c vs. Fig. 3e and Fig. 3d vs. Fig. 3f), localization length (Fig. 9c vs. Fig. 9e and Fig. 9d vs. Fig. 9f) and on-site Hamiltonian elements (Fig. 5c vs. Fig. 5e and Fig. 5d vs. Fig. 5f). By comparing the figures one can also see that the effect of side chain disorder is stronger in the shifted than in the aligned structure, which is due to the larger conformational freedom of the shifted structure, as discussed in Section 3.1. Therefore, to obtain reliable results, side chains should be included in the calculations.

3.3 Consequences for electrical transport

We now discuss the consequences of our findings about thermal disorder on electrical properties of the material. In small molecule-based organic crystals (SMOCs) the effects of thermal disorder were used to explain the temperature dependence of the mobility where the mobility that decreases with increasing temperature is typically observed. On the other hand, all mobility measurements of P3HT, even for highest quality ordered samples, yield a thermally activated temperature dependence.

Our results suggest that in ordered P3HT there is a spectral region within first 200 meV below the top of the valence band with electronic states localised to just a few rings. In a combined molecular dynamics–electronic structure study in ref. 15 such states were found to be persistently localised in the sense that their position does not vary over time on the order of few nanoseconds. Below the spectral region with localised states, there is a region where both localised and delocalised states exist. It is well understood that the spatial and energetic distribution of electronic states that we obtained leads to thermally-activated transport; at low temperature most carriers populate localised states at the top of the valence band, which yield low mobility, while at higher temperatures less localised or delocalised states become more populated and the transport is much better then.

However, what is the main difference between polymers and SMOCs where a different temperature dependence of mobility is observed? In SMOCs, thermal disorder leads to localised electronic states, as well. However, the spectral region where these states exist is much narrower. For example, in ref. 13 the spectral region with strongly localised states has the width of approximately $0.2t$, where $t$ is the electronic coupling transfer integral between two neighbouring molecules (which is typically on the order of 100 meV in SMOCs). In ref. 10 this range is equally narrow and is comparable to or even smaller than thermal energy $k_B T$ at room temperature. For this reason, thermally activated behaviour is not observed in SMOCs.

The comparison between the effects of thermal disorder in ordered polymers and SMOCs illustrates the dual role of temperature when thermal disorder and transport properties are concerned. The temperature acts on the one hand to create well localised states and on the other hand to promote the carriers from such localised states to delocalised states with better transport. In SMOCs higher temperatures lead to better localization of the states and consequently to a smaller mobility. On the other hand, in ordered P3HT polymers, in the range of temperatures from 100 K to 300 K, the temperature has a weak effect on the DOS and on the dependence of localization length on energy. This conclusion was obtained from the comparison of these two quantities at 100 K and 300 K, presented in Fig. 10. A weak effect of temperature on the degree of localization is observed? In SMOCs, thermal activation of carriers from localised states to delocalised or less localised states with better transport. While we show that the effects of thermal disorder alone lead to thermally activated mobility, this does not necessarily imply that they are the dominant cause of thermally activated mobility observed in realistic samples. Various imperfections of chemical or structural nature produce traps where carriers can be localised and these can also lead to thermally activated mobility.

4 Conclusion

In this work the effects of thermal disorder on the electronic structure of crystalline P3HT were investigated. The influence of side chains and main chains on the thermal disorder was investigated separately for the first time. The main conclusions
from the obtained results can be summarized as follows. The disorder in side chains has a significant effect on the electronic structure of P3HT. The effect is more pronounced in the shifted structure than in the aligned, due to higher conformational freedom of side chains. The disorder in main chains has a strong effect on the electronic structure, leading to the localization of HOMO level wave functions to few rings only. Such a degree of localization is similar to the localisation in amorphous P3HT and it is a possible cause of thermally activated mobility that is typically observed in ordered polymers.

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