Structural relaxations in electronically excited poly(\textit{para}-phenylene)

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Structural relaxations in electronically excited poly(\textit{para}-phenylene) are studied using many-body perturbation theory and density-functional-theory methods. A sophisticated description of the electron-hole interaction is required to describe the energies of the excitonic states, but we show that the structural relaxations associated with exciton formation can be obtained quite accurately within a constrained density-functional-theory approach. We find that the structural relaxations in the low-energy excitonic states extend over about 8 monomers, leading to an energy reduction of 0.22 eV and a Stokes shift of 0.40 eV.

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The sustained growth of interest in conjugated polymers arises from their potential as the active material in low-cost field-effect transistors, photovoltaic devices and light emitting diodes (LEDs).\textsuperscript{1, 2} Investigating electronically excited states is an important part of this effort, which has proved a great challenge for theoretical techniques. An accurate description of the excited electronic states of conjugated polymers requires the inclusion of the strong electron-hole interaction and the structural relaxations which occur in the excited state.

In its simplest form, a polymer LED consists of a layer of conjugated polymer sandwiched between two electrodes. Electrons and holes are injected into the polymer where they are attracted to one another and form bound excitons. Light is then emitted by exciton recombination. In an excited state the polymer may lower its energy by structural relaxations so that the peak optical emission frequency is lower than the peak absorption frequency. Although this “Stokes shift” can, at least in principle, be measured, the actual relaxations have not been determined experimentally. These relaxations localize excitons and are relevant for the technologically important processes of exciton migration and recombination.

We have investigated structural relaxations in the low-energy excited states of poly(\textit{para}-phenylene) (PPP). A PPP LED has been demonstrated which emits blue light in a band around 2.7 eV.\textsuperscript{3, 4} PPP is, however, insoluble and therefore difficult to process and, instead, soluble derivatives of PPP with various side groups are preferred for manufacturing LEDs. The low-energy optical properties of PPP and its derivatives are similar, although the structural relaxations in the excited states depend on the nature of the side groups. We have chosen to study PPP because it has a simpler structure than its derivatives.

As we show below, systems containing more than 100 atoms are required for studying the excited-state relaxations of PPP, which pose a significant problem to theory. Such systems can easily be treated by density-functional theory (DFT)\textsuperscript{5}, but DFT alone often has difficulties in describing excited states. Excited states can be described by many-body perturbation theory (MBPT)\textsuperscript{6}, but this is numerically demanding and cannot currently be applied to such large systems. However, we find that the forces in the excited states of PPP obtained within DFT are very similar to those within MBPT, which indicates that the relaxed structures and Stokes shift obtained within DFT are reliable.

A PPP chain (Fig. 1) consists of phenyl rings joined by single C-C bonds, with adjacent rings being rotated with respect to one another by a torsional angle \(\theta\). Our DFT structures for the ground states of the single chain and crystal are in excellent agreement with a previous DFT calculation and in accord with experiment.\textsuperscript{6} Our
DFT band gap for the single chain is 2.44 eV and that for the crystal is 1.70 eV, which is in good agreement with a previous DFT calculation.[7]

We began by using a simple model of how an optical excitation affects the bonding in PPP: we performed DFT calculations of the changes in the electronic charge distribution due to a single excitation from the highest occupied molecular orbital (HOMO) to the lowest unoccupied molecular orbital (LUMO). In both the isolated chain and crystalline forms we found an increase in the electronic charge in the A bonds and to a lesser extent in the B bonds (see Fig. 1). These changes indicate the development of double-bonding character which is expected to favor both a more planar arrangement of rings (reduction in $\theta$) and compression of the A and B bonds.

In a perfect chain or crystal an exciton would, in principle, be delocalized because localization within some region of space costs kinetic energy. However, a larger energy may be gained by structural relaxations so that an exciton may localize itself. The low-energy excitonic states of PPP may therefore be self-localized by structural relaxations, i.e., local reductions in $\theta$ and the lengths of the A and B bonds.

The structural relaxations in the lowest excited states of a single PPP chain can be investigated using a constrained DFT method. To this end we model the excited state by promoting an electron from the HOMO to the LUMO at wave vector $k = 0$ and performing full structural relaxations in unit cells of various sizes. A unit cell containing 28 monomers was found to be sufficient to converge the excited state structure. The important parameters of the minimum energy structure are shown in Fig. 2. The relaxations are largely confined to a region of approximately 8 monomers over which the torsional angle and the lengths of the A bonds are significantly reduced, and the lengths of the B bonds are somewhat reduced. The gain in energy from the relaxation is 0.22 eV, and the Stokes shift, calculated as the difference between the band gaps of the ground and excited state structures, is 0.40 eV.[8]

We have analyzed the localization of the exciton in terms of the compression of the A and B bonds and the reduction in the torsional angle, $\theta$. We performed two constrained DFT calculations for the excited state in which (1) only relaxations along the axis of the chain and (2) only relaxations in the plane perpendicular to the chain were allowed. The relaxation energies of 0.14 eV and 0.17 eV, respectively, add up to significantly more than the full relaxation energy of 0.22 eV, indicating that the modes are strongly coupled. The relaxations along the chain led to a Stokes shift of only 0.06 eV and to very weak localization of the exciton, whereas the relaxations perpendicular to the chain gave a Stokes shift of 0.16 eV and much stronger localization. The torsional relaxation is therefore very important in determining the Stokes shift, although the coupling to the bond compression is strong and both types of relaxation are important in determining the full relaxation energy and Stokes shift.

Stokes shifts in PPP derivatives have been measured in dilute solutions and solid films. A common approach is to use “laddered” PPP derivatives in which some or all of the phenyl rings are joined to their neighbors by chemical “bridges”. These bridges hinder rotation about the A bonds which, according to our picture, is expected to reduce the Stokes shift. Hertel et al. report absorption and emission measurements on a series of laddered and non-laddered polymers, which clearly show that laddering reduces the Stokes shift. As shown in Fig. 3b of Hertel et al., in dilute solution the non-laddered polymer dodecyloxy-poly(para-phenylene) (DDO-PPP) exhibits a Stokes shift of 0.6 ± 0.2 eV. Considering the presence of the solvent and different side chains in the experimental system the agreement with our single chain PPP value of 0.40 eV is reasonable. The measured Stokes shifts for the laddered polymers are smaller, in agreement with our picture that changes in $\theta$ couple strongly to the band gap. Our picture also suggests that structural relaxations for excitons in laddered PPP derivatives should only lead to a weak localization effect.

We now investigate the reliability of the constrained-
DFT results for the relaxation and Stokes shift of the exciton. The formation of a completely delocalized exciton leads to a vanishingly small change in the charge density and therefore vanishingly small forces on the atoms. If, however, a lattice distortion occurs which reduces the band gap in some region of space then the exciton will be attracted to this region. If the lowering of the energy of the exciton is greater than the energy required to form the distortion then the exciton will stabilize the distortion, leading to a self-localized exciton. The reduction in the band gap due to a lattice distortion is an effect which is described approximately within our DFT calculations. The excitonic energies themselves, however, are strongly modified by many-body effects arising from the electron-hole interaction. We now investigate the dependence of these many-body effects on the lattice distortion. If this dependence is sufficiently weak the constrained DFT approach will be reliable.

A rigorous approach to electronic many-body effects for excited electronic states is given by many-body perturbation theory (MBPT). We have used the GW approximation to describe the addition or removal of an electron and the Bethe-Salpeter equation (BSE) for the excitation of an electron including the electron-hole interaction. These techniques have recently been used to describe the excitonic states of solids, clusters, and polymers. MBPT is numerically very demanding and cannot currently be applied directly to a polymer with a self-localized exciton.

To investigate the effects of electronic correlation and validate the constrained-DFT approach we proceeded as follows. We performed DFT and MBPT calculations for a series of structures involving torsional angles and compressions of the A and B bonds which are similar to those found in the self-localized exciton of Fig. 2. We considered the relaxation of the two central monomers of Fig. 2 as indicating the maximum relaxations which occur in the exciton. We then periodically repeated this structure, obtaining a reference system which can be investigated within both constrained DFT and MBPT. We defined a structural parameter \( x \) which takes the value \( x=0 \) for the ground state structure (\( \theta=33.7^\circ, A=1.456 \text{ Å}, \) and \( B=1.380 \text{ Å} \)), and \( x=1 \) for the geometry corresponding to the central monomers of the self-localized exciton (\( \theta=9.1^\circ, A=1.427 \text{ Å}, \) and \( B=1.366 \text{ Å} \)). Intermediate values of \( x \) correspond to linearly interpolating \( \theta, A \) and \( B \) between the extremal values. Fig. 3 shows the excitation energies of this periodic structure as a function of \( x \). The four curves denote the DFT energy gap, the quasiparticle (QP) energy gap, and the transition energies of the lowest spin-singlet and spin-triplet excitons obtained from solving the BSE. The QP gaps are 2.3-3.0 eV larger than the DFT gaps due to the significant QP corrections typical of semiconducting systems. The exciton energies, on the other hand, are 1.5-2.9 eV smaller than the QP gaps due to the attractive electron-hole interaction. The

![FIG. 3: Excitation energies calculated within the DFT and GW approaches and the GW-BSE singlet and triplet energies as a function of the structural parameter \( x \) for a single PPP chain.](image)

We have analyzed the reasons for the success of the constrained DFT approach for calculating excited state geometries. The details of this analysis will be published elsewhere, but we conclude that the various approximations should work well for the chain or solid when the excitons are delocalized over many atoms and when the relaxations are relatively small and do not include reconstructive changes such as bond breaking. The constrained DFT approach would be inaccurate if the excitonic wave function contained contributions from more than one electron or hole band, but this mixing could...
be calculated from BSE calculations on high symmetry structures and then used in DFT studies of large structures. Our study suggests that the constrained DFT approach is likely to work for other conjugated polymers.

Our main calculations are for single chains, in which the electron-hole interaction is larger than in solids. Therefore our calculations are an even stronger test of the idea of neglecting the electron-hole interaction when calculating the excited state relaxations than would be encountered in the more technologically relevant solid state. In the solid, many chains are packed together at van der Waals distances, which gives rise to three main additional features: (i) electronic overlap between chains, (ii) van der Waals interactions, and, in particular, (iii) the effect of inter-chain dielectric screening on the electron-hole interaction. The first of these is well described within DFT while MBPT includes the second and third.

To study the effects of inter-chain dielectric screening we considered a 3D crystalline array of PPP chains in which the electronic overlap between chains is small and consequently the DFT results are only weakly perturbed. The calculated QP correction to the DFT gap and the electron-hole interaction are, however, both reduced by more than 1 eV as a consequence of the more effective dielectric screening in a 3D solid. The QP state, which describes an additional electron or hole on one chain, shows significant interaction with the polarizable neighboring chains, and the QP gap closes. In addition, the QP corrections also depend on $x$, indicating that the intra-chain screening changes when the geometry relaxes. The exciton energies are, however, only weakly affected by the changes in the inter- or intra-chain screening. In the solid the singlet (triplet) energy ranges from 3.66 eV (3.14 eV) at $x=0$ to 2.55 eV (2.23 eV) at $x=1$, which is only a little higher than the single-chain results. This insensitivity derives from the charge-neutral character of the exciton, which is much less influenced by electrostatic screening than charged single-particle excitations. Similar effects have also been observed for other polymers by van der Horst et al. The changes in the excitation energies as $x$ varies from 0 to 1 ($–1.11$ eV for the singlet, $–0.91$ eV for the triplet) are very similar to the change of the DFT gap ($–0.95$ eV), confirming the reliability of the constrained-DFT forces. The weak interactions between chains have to be handled with care, however, and the self-trapping of the exciton in the solid state will be the subject of a further study.

In summary, we have shown that while the electron-hole interaction is very important in determining the excitonic energies in PPP it is less important for the excited state geometries. We have demonstrated that a simple constrained DFT approach is sufficient for calculating relaxations in the low-energy excited states of PPP. In this case we find that the structural relaxations extend over about 8 monomers, leading to an energy reduction of 0.22 eV and a Stokes shift of 0.40 eV. The DFT method is tractable for studying the influence of structural disorder on the optical properties of polymers.

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[15] These results also imply that the structural relaxations and Stokes shifts in the lowest singlet and triplet states of PPP should be very similar.