Low-lying excitations of polyydiacetylene

A. Race, W. Barford, and R. J. Bursill

1Department of Physics and Astronomy, The University of Sheffield, Sheffield S3 7RH, United Kingdom
2School of Physics, The University of New South Wales, Sydney NSW 2052, Australia

(Received 1 September 2000; revised manuscript received 12 December 2000; published 27 June 2001)

The Pariser-Parr-Pople Hamiltonian is used to calculate and identify the nature of the low-lying vertical transition energies of polyydiacetylene. The model is solved using the density matrix renormalization group method for a fixed acetylenic geometry for chains of up to 102 atoms. The nonlinear optical properties of polyydiacetylene are considered, which are determined by the third-order susceptibility. The experimental $1B_u$ data of Giesa and Schultz are used as the geometric model for the calculation. For short chains, the calculated $E(1B_u)$ agrees with the experimental value, within solvation effects ($\sim 0.3$ eV). The charge gap is used to characterize bound and unbound states. The $nB_u$ state is above the charge gap and hence a continuum state; the $1B_u$, $2A_g$, and $mA_g$ states are not and hence are bound excitons. For large chain lengths, the $nB_u$ state tends towards the charge gap as expected, strongly suggesting that the $nB_u$ state is the conduction band edge. The conduction band edge for polyydiacetylene is agreed in the literature to be $\sim 3.0$ eV. Accounting for the strong polarization effects of the medium and polaron formation gives our calculated $E_s(nB_u) \sim 3.6$ eV, with an exciton binding energy of $\sim 1.0$ eV. The $2A_g$ state is found to be above the $1B_u$ state, which does not agree with relaxed transition experimental data. However, this could be resolved by including explicit lattice relaxation in the Pariser-Parr-Pople-Peierls model. Particle-hole separation data further suggest that the $1B_u$, $2A_g$, and $mA_g$ states are bound excitons, and that the $nB_u$ is an unbound exciton.

DOI: 10.1103/PhysRevB.64.035208 PACS number(s): 71.20.Rv, 71.35.–y, 71.45.Lr, 71.10.Fd

I. INTRODUCTION

Polymers, and other molecular materials, that exhibit nonlinear optical properties and electroluminescence have attracted much interest among theorists and experimentalists, owing to their possible use in organic technology. Poly(diacetylene)s (PDA’s) with the general formula shown in Fig. 1(a) are the $\pi$-conjugated organic polymers studied here: they exhibit nonlinear optical properties, near-perfect crystal structure, and doping-dependent transport properties. These properties make them ideally suited for use in the theorist’s calculation. A full characterization of the PDA electronic properties would mean a better understanding of $\pi$-conjugated polymers in general. In PDA’s with a very low polymer content ($X_p < 10^{-5}$ in weight), interchain interaction is minimal and thus it can be considered an ideal one-dimensional model system.

There have been many calculations performed on PDA with varying degrees of success. Parry conducted a self-consistent field (SCF) calculation and found an energy gap that was too low. Boudreaux’s SCF-X$\alpha$ calculation gives an energy gap of 2.21 eV, which agrees well with the long-chain experimental extrapolations for the energy gap of 2.11 eV and 2.25 eV (yellow phase). A further SCF-X$\alpha$ calculation by Boudreaux accounts for charge density waves by the same geometric Huük model of Whangbo et al. The ab initio crystal orbital calculations of Kertesz et al. give energy gaps that are too high. Predicted bond lengths show impressive agreement with x-ray data in Karpfen’s ab initio SCF calculation. He performed a total energy minimization to calculate bond lengths, and showed that the energy difference between butatriene and acetylene structure per unit cell is 0.52 eV. This agrees with previous calculations and experiment. A similar calculation yielding similar results was performed by Brédas et al.

Wilson, Cojan et al., and Cade and Movaghar performed Hückel band structure calculations for the infinitely long PDA chain. Cojan et al. and Wilson did not include bond alteration in their calculation, and thus modeled a compound with one distinct chemical bond. The energy gap was found to be too low. Cade and Movaghar used two degrees of freedom in their calculation as a bond alteration parameter.

FIG. 1. Chemical structure of a diacetylene trimer molecule. (a) Schematic representation of the orbitals giving rise to the different structures. The shaded orbitals give rise to $\sigma$ bonds, while the unshaded orbitals contain $\pi$ electrons in $2p$ orbitals. The $2p_z$ orbitals all overlap along the length. The molecular plane is in the $x$-$y$ plane. (b) Acetylene structure. (c) Butatriene structure.
Extended Hückel calculations have been performed by Parry\textsuperscript{21} and Wangbo \textit{et al.}\textsuperscript{12} The work of Parry is restricted to one distinct chemical structure and, as with other parameterizations, yields a value for the energy gap that is too low. The model employed by Wangbo has a hopping term of the form

$$t = t_0 (1 \pm \delta)$$

with $0 < \delta < 1$. This produces results that are, again, quantitatively too low; however, it does generate an energy difference of 0.48 eV between the butatriene and acetylene structures, in excellent agreement with other calculations and experiment.\textsuperscript{15,16} In summary, therefore, all these models are seen to give good qualitative predictions for bond alteration, but the data are seen to be consistently redshifted owing to their single-particle picture.

The importance of screening in PDA is still considered controversial by many authors, because it determines the role of electron-electron correlations. A simple way to account for screening is to adjust the single-chain electron-electron interaction potential $V(r)$, a method suggested by many authors. For example, Abe lowered the Hubbard parameter $U$ from 11 to 5 eV and used a large long-range dielectric constant in an Ohno potential.\textsuperscript{22} However, this again causes redshifted energy gaps and results in unphysical atomic cohesion energies. We emphasize that there is clear and indisputable evidence that electron-electron interactions are of paramount importance in the electronic structure calculations of PDA. First, in polymeric crystal form, the linear absorption spectrum is seen to be symmetric and peaks at $\sim 1.8$–$1.9$ eV. However, the onset of photoconduction occurs not at this value but at $\sim 2.3$–$3.0$ eV (corresponding to uncorrelated $\pi-\pi^*$ transitions).\textsuperscript{23} In addition, comparison of the linear absorption with the electroabsorption spectrum shows an EA feature at $\sim 2.0$ eV corresponding to the first derivative of the absorption: this is attributed to the Stark shift of the exciton. There is also an oscillating Franz-Keldysh band-edge structure at $\sim 2.5$–$3.0$ eV, with a line shape that deviates from the first derivative of the absorption. The oscillations are due to interband transitions and coincide with the onset of photoconduction. These data independently suggest that the exciton binding energy is $\sim 0.5$ eV.\textsuperscript{24–26} We note that these values show a great deal of variability owing to the variety of phases and the disorder present in PDA. Hence, despite the apparent success of the Su-Schrieffer-Heeger model\textsuperscript{27,28} neutral excitons with large binding energies are the low-lying excitations in PDA.\textsuperscript{24,29,30} This is borne out by the Pariser-Parr-Pople calculation presented in this work, which incorporates electron-electron interactions.

## II. MODELING POLYDIACETYLENE

### A. Electronic structure

The atomic orbital model used in the calculation is described as follows. Carbon has the atomic configuration $1s^22s^22p^2$. Since carbon has four valence electrons, there is an alternating structure of two $sp^2$ and two $sp$ hybridized carbon atoms that can account for the idealized PDA chemical composition. (See Fig. 1.)

Since PDA is spatially centrosymmetric, and thus shows $C_2$ symmetry, the wave functions possess mirror plane and centrosymmetry symmetries. The group notation for mirror plane symmetries is $A$ for the symmetric and $B$ for the antisymmetric case. Inversion symmetries are labeled $g$ for symmetric and $u$ for antisymmetric. The ground state is therefore labeled $1A_g$ and the first optically active dipole from the ground state has to be the $1B_u$ state. The wave functions are either even ($A_g$) or odd ($B_u$) under an inversion.

### B. Geometric structure

Although there is a myriad of literature on the optical properties of PDA, there is still little about their structural and conformational properties, which prevents extensive theoretical investigation. The work of Giesa and Schultz\textsuperscript{10} follows that of Wudl and Bitler\textsuperscript{31} describes the synthesis and thorough characterization of the series of alternating all-trans-polyenes without substituents at the vinylic bonds that are studied in this paper. PDA exists in three distinctly colored phases: when in a good solvent (e.g., chloroform) it is yellow. On addition of a nonsolvent (e.g., hexane) PDA changes to a blue or red form. In the yellow phase, PDA single chains exist in coil conformations. Adding a nonsolvent causes PDA to form aggregates or microgels, becoming rodlike, fully conjugated, and semicrystalline. Hence, we propose that \textit{in vacuo} calculations be compared with PDA in a nonsolvent to ensure an ideal system. The work of Giesa and Schultz is on PDA in a hexane solution.

To maintain consistency with the literature, the unsubstituted model compounds are named as follows:

$$C_n^1[N],$$

where the number of unsaturated carbon atoms $n$ specifies each compound, and the number of formal monomer units $N$ are added in the square brackets for clarity. The relation between $n$ and $N$ is

$$n = 2(2N + 1).$$

These structures are shown in Fig. 2. In contrast with the simplest conjugated polymer represented by trans-polyacetylene, the carbon backbone of PDA contains two, additional, localized $\pi$ electrons in every unit cell. Thus, there are two possibilities of bond alternation in this system that lead to nonequivalent structures with nondegenerate ground states, as shown in Figs. 1(b) and 1(c). Optical absorption measurements of Gross and Sixl\textsuperscript{9} have found that in the ground state of long-chain PDA molecules the acetylene structure is energetically more favorable than the butatriene structure. Several authors have confirmed this theoretically and experimentally. According to the electron spin resonance (ESR) measurements of Neumann and Sixl\textsuperscript{12} and Huber \textit{et al.},\textsuperscript{33} PDA has a diradical character for chain lengths $2 \leq N \leq 4$. In addition, Kollmar and Sixl found that a
phase change occurs from the butatriene to acetylene form for $N<4$. However, work on the molecular geometries of $C_{14}$ and $C_{22}$ by x-ray structure analysis has determined bond lengths and angles that represent a typical acetylene structure (see Table I) and no evidence for a butatriene form is found; this contradicts the work of Refs. 32–34. In addition, Giesa and Schultz shows that substitution does not alter the polymer structure significantly, further adding doubt to the findings of Kollmar and Sixl. Increasing the size of the polymer does elongate double and triple bonds, while reducing the single bond; however, this is not pronounced. Hence, overall, these findings justify the use of the model acetylene structure employed in this paper, because an increase in conjugation length does not show a substantial effect on bond lengths and angles.

To gain insight into the influence of conjugation length on the electronic properties of PDA, ultraviolet-visible spectroscopy, Raman scattering, $^{13}$C NMR shifts, and bond lengths in these compounds have been studied by Giesa and Schultz and are compared in this paper with the theoretical calculation. The extrapolation of these data to the infinite chain length provides yet another method of testing theory. This can be used to investigate the effects of side groups and cis bonds on the delocalized system. Hence, owing to PDA’s simple electronic and geometric ground-state structure, it can be easily modeled by the Pariser-Parr-Pople Hamiltonian.

III. THE PARISER-PARR-POPLE HAMILTONIAN

The Pariser-Parr-Pople Hamiltonian is

$$\hat{H}_{ppp} = \sum_{\langle ij \rangle} t_{ij} (c_{i \sigma}^\dagger c_{j \sigma} + c_{j \sigma}^\dagger c_{i \sigma}) + U \sum_i (n_i - \frac{1}{2})(n_i - \frac{1}{2})$$

$$+ \sum_{\langle i \sigma \alpha \sigma' \rangle} V_{ij}(n_{i \sigma} - 1)(n_{j \sigma'} - 1),$$

where the operator $c_{i \sigma}^\dagger (c_{i \sigma})$ creates (annihilates) a $2p_z$ electron of spin $\sigma$ at site $i$ (the $l$th carbon atom), $t_{ij} (>0)$ is the transfer integral between the nearest neighbor atomic orbitals. The first term in the Hamiltonian allows the electrons to ‘‘hop’’ from one atom to another, and represents the kinetic energy gained from delocalizing an electron from its atomic site. $n_{i \sigma} = c_{i \sigma}^\dagger c_{i \sigma}$ is the number density operator, and $\langle ij \rangle$ denotes nearest neighbors. $U$ is the Pariser-Parr-Pople Coulomb repulsion between two electrons occupying the same $2p_z$ orbital and $U=10.06$ eV. $V_{ij}$ is the long-range Coulomb repulsion and, in this work, is the Ohno function, $V_{ij} = U/\sqrt{1 + \beta r_{ij}^2}$, where $\beta = (U/14.397)^2$ and $r_{ij}$ is the interatomic distances measured in Å.

To find quantitative values for the resonance integrals, a relationship between the bond lengths known from the x-ray data in Table I and the resonance integrals is needed. This is found by linearizing the exponential law found in the literature. The general expression for the resonance integrals is given by

$$t_{ij} = t_0 + \alpha (r_{ij} - r_0),$$

where $\alpha$ is the coupling constant of the electron-phonon interactions, $r_{ij}$ is the length of the bond between carbon atoms $i$ and $j$, and $r_0$ is the average bond length of the reference system. This is an expansion around the interchain transfer

![Diagram](Image)

FIG. 2. The idealized chemical structure of the series of PDA’s used in this work. Me$_3$ represents three methyl structures. $C_n[N]$ unambiguously represents the structures, where $n$ is the number of carbon atoms and $N$ is the number of monomer units.

### Table I. Bond lengths for various forms of PDA. All values are in Å. (Data are taken from Ref. 10.)

<table>
<thead>
<tr>
<th></th>
<th>R</th>
<th>$d$ (C=(\equiv)C)</th>
<th>$D$ (C=C)</th>
<th>$D$ (C=(\equiv)C)</th>
<th>$sp^2$ angle$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDAs</td>
<td>P-THD</td>
<td>1.205</td>
<td>1.426</td>
<td>1.359</td>
<td>119.1</td>
</tr>
<tr>
<td></td>
<td>P-MBS</td>
<td>1.195</td>
<td>1.364</td>
<td>1.424</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PTS-6</td>
<td>1.191</td>
<td>1.356</td>
<td>1.428</td>
<td>121.9</td>
</tr>
<tr>
<td>PDAs used in</td>
<td>$C_{14}[3]$</td>
<td>H</td>
<td>1.202</td>
<td>1.329</td>
<td>142.7</td>
</tr>
<tr>
<td>this paper</td>
<td>$C_{22}[5]$</td>
<td>H</td>
<td>1.208</td>
<td>1.360</td>
<td>141.2</td>
</tr>
</tbody>
</table>

$^a$Bond angles in degrees between an $sp^2$ carbon atom and the next $sp$ atom.

$^b$R = -CH$_3$-OTos at 295 K.

$^c$R = -CH$_3$SO$_3$-Ph-p OMe at 295 K.

$^d$R = -CH$_3$N(Ph)$_2$ at 295 K.
The parameters in expression (4) are derived using polyacetylene as a reference system. First, it is pertinent to consider the gradient of the resonance integral with respect to bond length. It has been shown that the difference $\delta t$ of the resonance integral is $0.2162$ eV and the corresponding average change in bond lengths $\delta l$ is $0.0471$. Thus, the electron-phonon coupling constant is

$$\alpha = \frac{\delta l}{\delta t} = 4.593 \text{ eV } \AA^{-1}. \tag{7}$$

A full list of Hamiltonian parameters derived from polyacetylene used in this work is shown in Table II, and a schematic of the unit cell is found in Fig. 3.

It is instructive to consider the terms of the Hamiltonian (4) separately: the hopping part of the Hamiltonian for the unit cell is

$$\hat{H}_{\text{unit cell}} = -[t_s(c_{1j}^{\dagger}c_{2j} + \text{H.c}) + t_d(c_{3j}^{\dagger}c_{4j} + \text{H.c}) + t_t(c_{3j}^{\dagger}c_{4j} + \text{H.c}) + t_d(c_{4j}^{\dagger}c_{1j} + \text{H.c})]. \tag{8}$$

where $t_s$, $t_d$, and $t_t$ are the hopping terms associated with the single, double, and triple bonds, respectively. By introducing Bloch states the following equation for the single-particle dispersion is found:

$$E_w = 2zt_0, \tag{6}$$

where $z$ is the number of nearest neighbors, and thus for linear polymers $z = 2$.

FIG. 3. Resonance integrals used for PDA with bond alteration in the $i$th unit cell. $t_s$ is the resonance integral for the single, $t_t$ for the triple, and $t_d$ is for the double bond. The periodicity of polydiacetylene is given by $4a$, where $a$ is the undimerized bond length.

FIG. 4. Schematic energy band structure of polydiacetylene in the reduced Brillouin zone for the acetylene structure. $E'_g$ is the energy gap within the valence bands. $E_g$ is the energy gap between the upper valence band and the conduction band. Numerical estimates are from the parameters of Table II (see text).

$$E_k = \pm \left\{ \frac{1}{2}(2t_s^2 + t_t^2 + t_d^2) \pm \frac{1}{4}(t_s^2 - t_t^2) + t_d^2(t_s^2 + t_d^2) + 2t_d^2t_s\cos 4ka \right\}^{1/2}. \tag{9}$$

It is apparent that two valence bands and two conduction bands exist, and they are labeled as $E_{c1}$, $E_{c2}$, $E_{v1}$, and $E_{v2}$. The energy gaps and total bandwidth are given by

$$E_g = E_{c1}(0) - E_{c2}(0) = |2t_s - (t_d + t_t)| = 1.33 \text{ eV}, \tag{10}$$

$$E'_g = E_{c2}\left( \pm \frac{\pi}{4a} \right) - E_{c1}\left( \pm \frac{\pi}{4a} \right) = t_t - t_d = 0.641 \text{ eV},$$

$$E_w = E_{c2}(0) - E_{v1}(0) = 2t_s + t_d + t_t = 11.127 \text{ eV}.$$
Essential States

\[ \text{Band of States} \]

\[ \begin{align*}
  mA_g & \\
  nB_u & \\
  1Ag & \\
  1Bu & \\
  1Ag & 
\end{align*} \]

FIG. 5. The four essential states.

Conduction band signifies the noninteracting limit, excitons are energetically situated below the conduction band.

IV. RESULTS AND DISCUSSION

A. Essential states

Nonlinear optical experimental data can be used to characterize the low-lying states of π-conjugated polymers. The relative ordering of these states can elucidate much of the physics of such systems.\(^{30,39,40}\) Nonlinear optical processes in polymers with \(C_2\) symmetry are determined by the third-order susceptibility, \(\chi^3(-\omega_1 - \omega_2 - \omega_3; \omega_1, \omega_2, \omega_3)\), which is calculated by the sum over all available states. This makes calculating \(\chi^3\) in principle difficult, owing to the large number of paths included. However, it is found that there are certain essential states that are important when considering the \(\chi^3\) of π-conjugated polymers, as only the essential states make a significant contribution to it. The four essential states are the ground state, the first odd-parity exciton state, \(1Bu\), the charge transfer state, \(mA_g\), and the \(nB_u\) state (see Fig. 5). Another important state is the \(2Ag\); this is a two-photon state, whose position relative to the \(1Bu\) state determines whether a polymer exhibits photoluminescence. The importance of these states is the result of the strong and dominant dipole couplings amongst them.

TABLE III. Dipole moments and corresponding energies of the essential states defined in the text for chains of up to 26 atoms (6 monomer units).

<table>
<thead>
<tr>
<th>Chain length</th>
<th>Dipole moment (\mu_{ij} = \langle \hat{\mu} \rangle )</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n)</td>
<td>(N)</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>10</td>
<td>2</td>
</tr>
<tr>
<td>14</td>
<td>3</td>
</tr>
<tr>
<td>18</td>
<td>4</td>
</tr>
<tr>
<td>22</td>
<td>5</td>
</tr>
<tr>
<td>26</td>
<td>5</td>
</tr>
</tbody>
</table>
to higher-order corrections, and the fit required cannot be used to characterize states. In this work, therefore, a polynomial fit is preferred for extrapolating the data.

The calculated vertical excitation energies for the $1B_u$, $2A_g$, $mA_g$, and $nB_u$ state and charge gap, $\Delta$, are plotted in Fig. 6 against $1/n$. In addition, the polynomial extrapolations for the long-chain limit are shown in Table IV for some of the states. The charge gap is a useful criterion for characterizing excitonic and unbound states, and is given as follows:

$$\Delta = E(n+1) + E(n-1) - 2E(n). \quad (12)$$

Here $E(n)$ is the ground-state energy of the $n$-electron system. The charge gap signifies the lowest-energy excitation of an electron from the valence to conduction band. Thus states above $\Delta$ are unbound excitons; those below it are bound. Of the states plotted in Fig. 6 only the $nB_u$ is positioned above the charge gap, suggesting that the $nB_u$ is a continuum state. Conversely, the $1B_u$, $2A_g$, and $mA_g$ states are below it and are thus bound excitons. This is shown to be true in Sec. IV C by examining the particle-hole separation of these states.

The experimental data obtained by Giesa and Schultz are also plotted in Fig. 6. A polynomial extrapolation of the experimental data yields $E_x \sim 2.5$ eV; however, this is considered unreliable as there are so few points used in the fit. For short chains the calculated and experimental $1B_u$ energies are remarkably within solvation (polarization) effects of the experimental medium ($\sim 0.3$ eV). This solvation value is derived from the work of Moore et al. who predicts 0.3 eV for the $1B_u$ state. In addition, the Bursill and Barford calculation on polyenes found $E(1B_u)$ to be 0.3 eV above the experimental value. Their recent calculations on poly(phenylene) (PPP) yields similar results. Our calculated $E_x(1B_u)$ for PDA is $\sim 3.0$ eV, and hence correcting for solvation effects gives our calculated $E_x(1B_u) \sim 2.7$ eV.

The energy of the $nB_u$ state is seen to tend to that of the charge gap for $n \rightarrow \infty$, as seen in Fig. 6. Hence, the extrapolated long-chain $nB_u$ energy is found to be $\sim 5.7$ eV, and this state is predicted to be the conduction band threshold. However, the onset of photoconduction (continuum limit) is $\sim 3.0$ eV. The large polarization effects of the surrounding medium can correct for this: these are estimated by Moore et al. to be $\sim 1.5$ eV. In addition, polaron formation means that a further $\sim 2 \times 0.3$ eV can be subtracted from 5.7 eV. Including all these effects gives $E_x(nB_u) \sim 3.6$ eV, which is in reasonable agreement with the experimental conduction band threshold quoted earlier. Therefore, the exciton binding energy is $E_b = E_x(nB_u) - E_x(1B_u)$, and from our calculation is found to be $3.6 \pm 0.9$ eV. This is in reasonable agreement with binding energies found in the literature.

Having identified the important nonlinear optical states in PDA from the dipole moments, and calculated the corresponding excitation energies, an energy level diagram can be drawn. This is shown for 26 atoms in Fig. 7. As expected there is a large $1A_g \rightarrow 1B_u$ dipole moment, corresponding to the one-photon absorption, while the band threshold $(nB_u)$ has a weak coupling to the ground state. It is seen that the first excited $A_g$ state is above the $1B_u$ state, which would suggest that PDA shows photoluminescence. (There is a direct $1A_g \rightarrow 1B_u$ transition.) However, experiments have shown that PDA is not photoluminescent: measurements of

TABLE IV. Extrapolated long-chain energies for the essential states and the charge gap.

<table>
<thead>
<tr>
<th>State</th>
<th>$E_x$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$1B_u$ Theory</td>
<td>3.021</td>
</tr>
<tr>
<td>Exp.</td>
<td>2.5113</td>
</tr>
<tr>
<td>Charge gap, $\Delta$</td>
<td>5.7351</td>
</tr>
</tbody>
</table>
In order to help characterize the low-lying $A_g$ and $B_u$ states the particle-hole separation is calculated and gives an indication of the spatial extent of a given state: an explanation of how it is calculated is given in the Appendix. Suffice it here to say that we use the one-particle singlet excitation correlation function, which directly relates a hole in the valence band to an electron in the conduction band. In essence, if a particular state’s particle-hole separation is seen to increase with system size, the electron and hole are unbound. However, if this quantity reaches a maximum, the Coulomb attraction between the electron and hole is strong enough to bind them together. In Fig. 8 the particle-hole separations are given as a function of system size in units of the average C–C bond length for the $1B_u$, $2A_g$, and $mA_g$ states. It is clear that the $1B_u$, $2A_g$, and $mA_g$ states are all bound excitons because the electron-hole separation reaches a maximum value with increasing chain length. These signify the composite particles mentioned in Sec. IV B. However, the particle-hole separation of the $nB_u$ state increases linearly with system size: this suggests that the $nB_u$ state is either very weakly bound or unbound. We conclude therefore that for PDA molecules of up to 26 carbon atoms the electron-hole continuum (i.e., the band edge) is expected to be the $nB_u$ state, further strengthening the arguments of Sec. IV B.

### C. Particle-hole separation

In order to help characterize the low-lying $A_g$ and $B_u$ states the particle-hole separation is calculated and gives an indication of the spatial extent of a given state: an explanation of how it is calculated is given in the Appendix. Suffice it here to say that we use the one-particle singlet excitation correlation function, which directly relates a hole in the valence band to an electron in the conduction band. In essence, if a particular state’s particle-hole separation is seen to increase with system size, the electron and hole are unbound. However, if this quantity reaches a maximum, the Coulomb attraction between the electron and hole is strong enough to bind them together. In Fig. 8 the particle-hole separations are given as a function of system size in units of the average C–C bond length for the $1B_u$, $2A_g$, and $mA_g$ states. It is clear that the $1B_u$, $2A_g$, and $mA_g$ states are all bound excitons because the electron-hole separation reaches a maximum value with increasing chain length. These signify the composite particles mentioned in Sec. IV B. However, the particle-hole separation of the $nB_u$ state increases linearly with system size: this suggests that the $nB_u$ state is either very weakly bound or unbound. We conclude therefore that for PDA molecules of up to 26 carbon atoms the electron-hole continuum (i.e., the band edge) is expected to be the $nB_u$ state, further strengthening the arguments of Sec. IV B.

### V. CONCLUSIONS

The DMRG calculation of a suitably parametrized Pariser-Parr-Pople Hamiltonian for a rigid geometry can be used to describe the electronic structure of short- or long-chain polydiacetylene polymers. The role of electron-electron interactions is fundamental in the low-lying excitations, and this is borne out in our calculation. We have found that the frequency dependence of third-harmonic generation in Langmuir-Blodgett films of PDA indicate that an $A_g$ singlet state is $\sim 0.11$ eV below the lowest energy $B_u$ state. In addition, the recent work of Kohler and Schilke has shown an $A_g$ symmetry state $\sim 0.21$ eV lower than the lowest $B_u$ symmetry state. Our model calculates vertical transitions only and is seen to agree with many theoretical vertical-energy calculations of PDA. Singlet exciton relaxation studies on isolated polydiacetylene chains by subpicosecond pump-probe experiments have suggested that the nonradiative singlet $B_u$ exciton relaxation involves two $A_g$ states in series. As it is well known that electron-phonon coupling is strong in PDA, we propose that lattice relaxation effects produce these $A_g$ states. It is hoped that this will be resolved by studies of lattice relaxation explicitly within the Pariser-Parr-Pople-Peierls Hamiltonian, which are now in progress. Similar studies by Barford and co-workers on polyenes have found that the relaxation energies of the $1B_u$ and $2A_g$ states can be reduced by as much as 0.3 and 1.0 eV, respectively. Thus, including lattice relaxation in the Pariser-Parr-Pople-Peierls model is expected to bring the $2A_g$ state below the $1B_u$ state in agreement with experiment.

![FIG. 8. Particle-hole separation for the $1B_u$, $2A_g$, $mA_g$, and $nB_u$ states in units of the average C-C distance as a function of the number of carbon atoms, $n$.](image-url)
promotes an electron from where a dimer is a double or triple bond. The local molecular orbitals correspond to the bonding (|1⟩) and antibonding (|2⟩) combinations of the atomic orbitals on each dimer, where a dimer is a double or triple bond. The local molecular orbitals delocalize via the single bonds. Then, the creation operator,

\[ S_{ij}^\dagger = \frac{1}{\sqrt{2}} (a_{i2}^\dagger a_{j1} + a_{i1}^\dagger a_{j2}) \]  

promotes an electron from |1⟩ on dimer j to |2⟩ on dimer i. However, since the local molecular orbitals are not exact Bloch transforms of the band molecular orbital it is also necessary to include the Hermitian conjugate of \( S_{ij} \). The exact Bloch transforms of the band molecular orbitals are Wannier molecular orbitals. The Wannier orbitals of the valence band are predominantly |1⟩, with a small admixture of |2⟩ from neighboring dimers.

We now define the exciton correlation function with respect to the ground state, as

\[ C_{ij}(|n\rangle) = \langle n| (S_{ij}^\dagger S_{ij}) |1A_e^+\rangle, \]  

and the mean square of the particle-hole separation is

\[ \langle (i-j)^2 \rangle = \frac{\sum_{ij} |C_{ij}|^2}{\sum_{ij} C_{ij}^2}. \]  

In practice, we do not consider all combinations of i and j, but restrict ourselves to i and j symmetrically spaced around the middle dimer.


