Effect of dimerization on dynamics of spin-charge separation in Pariser-Parr-Pople model: A time-dependent density matrix renormalization group study

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Abstract

We investigate the effect of static electron-phonon coupling, on real-time dynamics of spin and charge transport in $\pi$-conjugated polyene chains. The polyene chain is modeled by the Pariser-Parr-Pople Hamiltonian with dimerized nearest-neighbor parameter $t_0(1 + \delta)$ for short bonds and $t_0(1 - \delta)$ for long bonds, and long-range electron-electron interactions. We follow the time evolution of the spin and charge using time-dependent density matrix renormalization group technique, when a hole is injected at one end of the chain in its ground state. We find that spin and charge dynamics followed through spin and charge velocities, depend both on chain length and extent of dimerization, $\delta$. Analysis of the results requires focusing on physical quantities such as average spin and charge polarizations, particularly in the large dimerization limit. In the dimerization range $0.0 \leq \delta \leq 0.15$, spin-charge dynamics is found to have a well defined behavior, with spin-charge separation (measured as the ratio of charge velocity to spin velocity) as well as, the total amount of charge and spin transported in a given time, along the chain, decreasing as dimerization increases. However, in the range $0.3 \leq \delta \leq 0.5$, it is observed that the dynamics of spin and charge transport becomes complicated. It is observed that for large $\delta$ values, spin-charge separation is suppressed and the injected hole fails to travel the entire length of the chain.

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I. INTRODUCTION

With vast advancements in technology, low-dimensional $\pi$-conjugated organic systems in recent times, have found use in single-molecule electronic and spintronic devices.\textsuperscript{1–7} Until now, these materials have been used in devices such as organic light-emitting diodes (OLEDs) and organic thin-film transistors.\textsuperscript{8–11} The $\pi$-conjugated organic materials form an interesting class of strongly correlated systems in which there exists long-range electron correlations. Therefore, the low-energy physics of these systems is different from low-dimensional strongly correlated materials described by the Hubbard model. In order to propose and design $\pi$-conjugated organic systems as components of electronic (spintronic) circuits, a proper theoretical understanding of the mechanism of charge and spin transport in these systems is thus necessary. Theoretical understanding of transport in many-particle systems with strong correlations requires appropriate techniques and formulations, mainly because transport is essentially an \textit{out-of-equilibrium} phenomena. The advent of time-dependent density matrix renormalization group (td-DMRG) technique has vastly helped in addressing this issue.\textsuperscript{12–16} Even so, spin and charge transport in $\pi$-conjugated systems have not been addressed until recently using the td-DMRG method due to the fact that, most of the existing td-DMRG algorithms are chiefly structured to handle short-range electron-electron interactions. Furthermore, those few that are capable of handling long-range interactions suffer from drawbacks of large computational resources for their study. The \textit{double time window targeting} (DTWT) technique proposed by us,\textsuperscript{17} has been employed to address the issue of dynamics of spin and charge transport in $\pi$-conjugated systems.

The issue of coupling of the electronic and lattice degrees of freedom on the stability of the metallic state in one-dimensional conductors, has been of interest for a long time.\textsuperscript{18–21} One-dimensional systems such as polyenes, typified by trans-polyacetylene ($t$-PA), belong to the class of $\pi$-conjugated molecular materials having linear (chain) topology. These materials are the simplest $\pi$-conjugated systems that have been studied extensively both experimentally and theoretically.\textsuperscript{22,23} In these systems, electronic structure is strongly affected by electron-phonon interactions leading to dimerization, which is stabilized and enhanced by electron-electron correlations. As a result, these systems have dimerized ground state and their low-energy excitations are gapped, the extent of which depends both on the degree
of dimerization and electron-electron interactions. Thus, it is to be expected that electron-phonon coupling will also influence the *out-of-equilibrium* dynamics of spin-charge separation in particular, and transport in general. Hence, an understanding of the role of dimerization on the dynamics of spin and charge transport in these systems is of considerable interest. Like most studies, we have also treated the electron-phonon coupling in the adiabatic limit (or Born-Oppenheimer approximation), i.e., the phonon degrees of freedom are considered as slow, classical variables. In this limit, the sole effect of electron-lattice coupling in polyenes manifests as bond alternation, the extent of which is dictated by the alternation (or dimerization) parameter $\delta$. A non-zero value of $\delta$ implies that the nearest-neighbor hopping integrals alternate as $t_0(1 \pm \delta)$. The object of this study is to investigate the effect of this bond alternation on the dynamics of spin and charge transport in one-dimensional systems typified by polyenes. The organization of the paper is as follows: In Sec. II, we discuss in detail, the model Hamiltonian and computational strategy used. Section III presents the results of our study along with discussions. In Sec. IV, we present our conclusions.

II. MODEL AND COMPUTATIONAL METHODOLOGY

The Pariser-Parr-Pople (PPP) model\(^\text{24,25}\) is appropriate for investigating the effect of dimerization on the dynamics of spin and charge transport in polyenes. In the second quantized representation, the PPP Hamiltonian reads as

$\hat{H}_{\text{PPP}} = \sum_{i=1}^{L-1} \sum_{\sigma} t_0 [1 - (-1)^i \delta] (\hat{c}_{i,\sigma}^{\dagger} \hat{c}_{i+1,\sigma} + \hat{c}_{i+1,\sigma}^{\dagger} \hat{c}_{i,\sigma})$

\begin{equation}
+ \sum_{i=1}^{L} \frac{U_i}{2} \hat{n}_i (\hat{n}_i - 1) + \sum_{j>i} V_{ij} (\hat{n}_i - z_i) (\hat{n}_j - z_j).
\end{equation}

Here, $L$ denotes the number of carbon atoms in the polyene chain with open boundary condition, $\hat{c}_{i,\sigma}^{\dagger}$ ($\hat{c}_{i,\sigma}$) creates (annihilates) an electron with spin orientation $\sigma$ on the $i^{th}$ carbon atom, $t_0$ is the average transfer integral without dimerization and, $0 \leq \delta \leq 1$ is the bond alternation or dimerization parameter. The strength of on-site Coulomb repulsion between two electrons of opposite spins on site $i$ is $U_i$ and $\hat{n}_i$ is the electron number operator for the same site. The term $V_{ij}$ represents the inter-site Coulomb repulsion between sites $(i,j)$, with $z_i$ being the on-site chemical potential of the $i^{th}$ carbon atom. Polyenes being
homogeneous $sp^2$ carbon systems, $U_i = U$ at all sites, and to maintain charge neutrality when a site is singly occupied, we also set $z_i = 1$ for all $i$. The inter-site interaction between electrons on sites $i$ and $j$, $V_{ij}$, is interpolated between $U$ for $r_{ij} = 0$ and $\frac{e^2}{r_{ij}}$ for $r_{ij} \to \infty$ by Ohno interpolation\textsuperscript{26} given by,

$$V_{ij} = 14.397(1.6348 + r_{ij}^2)^{-1/2}.$$ \hspace{1cm} (2)

where $V_{ij}$s are in eV and $r_{ij}$s are in Å. Although there is an algebraic fall off in the inter-site potential $V_{ij}$, all (equilibrium) properties such as optical gap, two photon gap and spin gap can be extrapolated to the thermodynamic limit. This is because the transfer part of the Hamiltonian is short-ranged and the effective correlation strength, $V_{\text{eff}} = (U - V_{12})$ is smaller than the band-width of the one-particle spectrum. Aside from this, the $\pi$-coherence length (which is the length of the $\pi$-system beyond which intensive properties such as excitation gaps, saturate) in such system is only about 20 sites long. Hence we expect that (non-equilibrium) properties such as spin and charge velocities, discussed in the subsequent section, also approach the thermodynamic limit. In this study we deal with polyene chains of 20, 30 and 40 carbon atoms, $\delta$ is set to 0.0, 0.05, 0.07, 0.15, 0.3 and 0.5, and the rest of the parameters assume standard values for the PPP model for $t$-PA and polyenes\textsuperscript{27–30}: $t_0 = -2.4$ eV, $U = 11.26$ eV, and $2\pi/3$ bond angle between successive bonds. The PPP Hamiltonian possesses charge-conjugation and inversion symmetries, and also conserves total spin. Dimerization affects the transfer term and the distance-dependent electron-electron repulsions ($V_{ij}$) only. It does not influence the on-site Coulomb repulsion ($U$) between electrons.

As $\delta$ increases from 0.0 to 0.5, hopping integrals for partial double bonds get enhanced from $-2.4$ to $-3.6$ eV, while those for partial single bonds reduce from $-2.4$ to $-1.2$ eV. As a result, the 1-norm of the Hamiltonian matrix increases, implying that $E_c = \max[|E_{\text{max}}|, |E_{\text{min}}|]$ also increases. The dimensionless time-step $\alpha = E_c \Delta t$ of a numerical scheme for solving the time-dependent Schrödinger equation, defines its stability region, with $\Delta t$ being the time-step of evolution\textsuperscript{31}. The value of $\alpha$ is constant for a given ordinary differential equation (ODE) solver, and hence as $E_c$ increases, $\Delta t$ has to be decreased. Thus, with increase in dimerization, the time-step for propagating the Schrödinger equation forward in time, has to be reduced for numerical stability. This is the scenario with commonly used ODE solvers such as the Runge-Kutta (RK) schemes, the Crank-Nicholson (CN)
method, and the multi-step differencing (MSD) techniques. Since the DTWT technique\textsuperscript{12} uses the MSD2 scheme\textsuperscript{31,32} for updating the Hilbert basis and the fourth-order RK technique for time evolution, with increase in dimerization the DTWT procedure becomes computationally time consuming. Hence, we modified the DTWT algorithm by replacing these two time evolution methods with the Chebyshev-polynomial-based expansion of the time evolution operator, \( \hat{U}(\Delta t) = \exp(-i\hat{H}\Delta t) \).\textsuperscript{33,34} The Chebyshev-polynomial-based scheme has the advantage that the expansion of \( \hat{U}(\Delta t) \) can be evaluated up to machine accuracy and is free from any time-step constraint.

The Chebyshev-polynomial-based time evolution involves propagating the state \( |\psi(t)\rangle \) by time-step \( \Delta t \) by approximating the discrete time evolution operator in terms of Chebyshev polynomials as

\[
e^{-i\hat{H}\Delta t} \approx \sum_{m=0}^{P} a_m T_m(\hat{H}),
\]

where, \( T_m(\hat{H}) \) is the \( m \)th Chebyshev polynomial of the first kind, \( \hat{H} \) represents the scaled Hamiltonian with eigenvalues ranging from \([-1.0, 1.0]\), and the coefficients \( a_m \) are given by

\[
a_m = (2 - \delta_{m0})e^{-i\Delta t\gamma}(-i)^m J_m(\Delta t\beta),
\]

where, \( \gamma = (E_{\text{max}} + E_{\text{min}})/2 \) and \( \beta = (E_{\text{max}} - E_{\text{min}})/2 \); \( E_{\text{max}}, E_{\text{min}} \) are the maximum and minimum eigenvalues of the PPP Hamiltonian, and \( J_m \) is the \( m \)th order Bessel function of the first kind. The necessity of scaling \( \hat{H}_{\text{PPP}} \) to \( \hat{H} \) arises from the argument domain of the Chebyshev polynomials of the first kind \( T_m(x); x \in [-1, 1] \). The Chebyshev polynomials can be generated using the following recursion relation\textsuperscript{35,36},

\[
T_{m+1}(\hat{H}) | \psi(t)\rangle = \left[ 2\hat{H}T_m(\hat{H}) - T_{m-1}(\hat{H}) \right] | \psi(t)\rangle,
\]

with the initial conditions, \( T_0(\hat{H}) | \psi\rangle = | \psi\rangle \), and \( T_1(\hat{H}) | \psi\rangle = \hat{H} | \psi\rangle \). However, since the coefficients \( a_m \) are known in advance [Eq. (4)], instead of using this forward recursion scheme, we use the “reverse” recursion algorithm proposed by Clenshaw\textsuperscript{37,38}, which is more stable. The Clenshaw recursion requires \( P \) sparse-matrix vector multiplications (SMVMs) of the Hamiltonian \( \hat{H} \) with the state vector \( | \psi(t)\rangle \). When \( P > \frac{1}{2} \Delta t (E_{\text{max}} - E_{\text{min}}) \), the error decays almost exponentially\textsuperscript{39}. In case of increase of 1-norm of the Hamiltonian matrix, one needs to either retain a higher \( P \) or reduce the magnitude of time step \( \Delta t \). We increase the value of \( P \) with enhancement in dimerization.
The basic DTWT algorithm remains unchanged when ODE solvers such as the fourth-order RK and MSD2 schemes are replaced by the Chebyshev-polynomial-based expansion of the time-evolution operator. However, for a given number of retained density matrix eigenvectors (DMEVs) \( m \) and a given time step \( \Delta t \), results of the Chebyshev-polynomial-based DTWT td-DMRG algorithm are more accurate than the ODE-based version of the algorithm. This is because the ODE solvers have truncation errors associated with them; the truncation errors associated with the MSD2 and the fourth-order RK procedures are \( O[(\Delta t)^3] \) and \( O[(\Delta t)^5] \), respectively.\(^{17,31,32}\) The Chebyshev-polynomial-based expansion of the time evolution operator, on the other hand, is free from such truncation errors and hence, can be evaluated up to machine accuracy by keeping \( P \) (number of SMVMs) \( > \frac{1}{2}\Delta t(E_{\text{max}} - E_{\text{min}}) \).\(^{39}\) However, the increase in accuracy is significant only at longer times; in the initial stages of time evolution, the accuracy is similar for the ODE and the Chebyshev-polynomial-based DTWT algorithms.

The number of SMVMs associated with the basic MSD2 and fourth-order RK steps are 1 and 4, respectively. For a system with \( N \) sites, in the finite-system DMRG algorithm, for a single full sweep we have \( 4(N/2 - 2) \) basic steps. Thus, total SMVMs for a single full sweep are \( 4(N/2 - 2) + 4 \). Since each single time window \( \Delta t \) is subdivided into \( p \) time slices of width \( \Delta \tau \), for propagating the wave packet by a single window, the total number of SMVMs is \( 8p(N/2 - 2) + 4p \). The first term corresponds to the MSD2 scheme for a full finite sweep of the \( N \) site system over two time windows and the second corresponds to propagation of the wave packet by a single time window. The total SMVMs per single time window are \( 4p(N - 3) \).\(^{17}\) In case of the Chebyshev-polynomial method, the total number of SMVMs per single time window \( \Delta t \) is \( 4P(N/2 - 2) \) or \( 2P(N - 4) \), \( P \) is dependent on \( \Delta t \), and usually \( P > p \). However, \( \Delta \tau \) in ODE methods should be very small (\( \sim 10^{-4} \) fs) when the 1-norm of the Hamiltonian matrix is large, for reasonable accuracy, while in the Chebyshev-based method, \( \Delta \tau \sim 10^{-1} \) fs can be employed for similar accuracy. Thus, the number of time steps in ODE based methods \( (p_{\text{ODE}}) \) is larger than in Chebyshev polynomial methods \( (p_{\text{CP}}) \). This leads to \( p_{\text{CP}} \times P \sim p_{\text{ODE}} \) and the two approaches require approximately the same computational time.

In order to investigate the effect of dimerization on the dynamics of spin and charge transport, an up spin electron is annihilated from the first carbon site of the polyene chains.
of $L$ sites in its half-filled ground state $|\phi_{gs}^0\rangle$, thereby leading to an initial wave packet $|\psi(0)\rangle$

$$|\psi(0)\rangle = \hat{c}_{1,\uparrow} |\phi_{gs}^0\rangle.$$  

(6)

This wave packet is propagated in time by solving the time-dependent Schrödinger equation numerically, using the Chebyshev polynomial-based expansion of $\hat{U}(\Delta t)$. Using the time evolved wave packets $|\psi(t)\rangle$, site charge density $[\langle \hat{n}_i(t) \rangle]$, and site spin density $[\langle \hat{s}_z^i(t) \rangle]$, at time $t$ are computed as,

$$\langle \hat{n}_i(t) \rangle = \langle \psi(t) | (\hat{n}_{i,\sigma} + \hat{n}_{i,-\sigma}) | \psi(t) \rangle,$$  

(7)

$$\langle \hat{s}_z^i(t) \rangle = \frac{1}{2} \langle \psi(t) | (\hat{n}_{i,\sigma} - \hat{n}_{i,-\sigma}) | \psi(t) \rangle.$$  

(8)

These quantities give the dynamics of the injected hole in terms of its spin and charge degrees of freedom. The real-time dynamics of the initial wave packet is studied using a modified DTWT scheme, wherein adaptation of the Hilbert space as well as time evolution, are performed with the Chebyshev polynomial-based expansion of time evolution operator. The other parameters used in our study are: DMEVs retained, $m = 300$; time step for evolution $\Delta \tau = 0.066$ fs; total evolution time $T = 33.0$ fs. However, for the purpose of this study we focus only on the initial 15.0 fs, as this is adequate for our purpose. In the modified DTWT procedure, weights of the reduced density matrices of all the time-dependent wave packets are kept the same, unlike in the originally formulated scheme, since we are using a large step size of 0.066 fs. Although time-dependent charge and spin densities are computed at all the sites, the quantities $\langle \hat{n}_{L_1}(t) \rangle$ and $\langle \hat{n}_{L_L}(t) \rangle$, and $\langle \hat{s}_z^1(t) \rangle$ and $\langle \hat{s}_z^L(t) \rangle$ are sufficient for investigating the effect of dimerization on the dynamics of the hole in terms of its spin and charge degrees of freedom; $1$ and $L$ correspond to the first and last sites of the $\pi$-conjugated chain with $L$ sites.

III. RESULTS AND DISCUSSION

The temporal variation in site charge density and site spin density for the terminal sites of a PPP chain of 40 sites are shown in Figs. 1 and 2, for various dimerization strengths $\delta$. The first significant dip in $\langle \hat{n}_{L_1}(t) \rangle$ and $\langle \hat{s}_z^L(t) \rangle$ correspond to the times $\tau_{L_1}^b$ and $\tau_{L_1}^s$, taken by the charge and spin of a hole injected at the first site, to reach the end of the chain,
FIG. 1: Temporal variation in charge densities, $\langle n_1(t) \rangle$ and $\langle n_{40}(t) \rangle$, at sites 1 and 40 in the PPP model, for different dimerizations, $\delta$, for a chain of 40 sites. In each box, $\langle n_1(t) \rangle$ is shown by thick black curve and $\langle n_{40}(t) \rangle$, by thin black curve. For $0.0 \leq \delta \leq 0.15$, $\tau_{h}^L$ is indicated by arrow.

respectively. The second significant dip in the time evolution plots of $\langle \hat{n}_1(t) \rangle$ and $\langle \hat{s}_z(t) \rangle$ occurs when the charge and spin degrees of freedom of the hole, get reflected back from the chain end and reach the injection site. These two dips correspond to times $\tau_{h}^{2L}$ and $\tau_{s}^{2L}$, respectively. This is supported by the fact that $\tau_{h}^{2L}$ and $\tau_{s}^{2L}$ are very close to twice the values of $\tau_{h}^{L}$ and $\tau_{s}^{L}$, respectively. From these times, the charge velocity ($\vartheta_{h}^{L}$) and spin velocity ($\vartheta_{s}^{L}$) are calculated in a straightforward way: $\psi_{h/s}^{L} = \left( L / \tau_{h/s}^{L} \right)$. From Figs. 1 and 2 it is observed that locating $\tau_{L}^{h/s}$ is easy for dimerizations $\delta$ up to 0.15. For dimerizations $\delta = 0.3$ and 0.5, it is however very hard to locate these minima unambiguously. This may be due to the fact that the weak bonds, when very weak (large $\delta$), do not easily transmit the charge or spin resulting in interference of the wave traveling forward with the reflected wave. This effect is seen more in the longer chains as the charge or spin needs to travel through many weak bonds.

Normally, charge and spin velocities are discussed in the bulk limit. From our earlier studies on the one-dimensional Hubbard model,[41] we have noticed that it is not possible to relate the analytically obtained charge and spin velocities $\vartheta_{\rho/\sigma}$, with $\vartheta_{L/2L}^{h/s}$ obtained for
finite small systems. This is because, the analytic velocities are obtained from the exact excitation spectrum as group velocities in the limit of momentum $\vec{q} \to 0$ while in our case $\vartheta^h/s_L$ pertain to a state which is not an eigenstate of the Hamiltonian governing the dynamics of the system. Furthermore, dimerization introduces a gap in the excitation spectrum and there is no low energy theory which completely separates the spin and charge degrees of freedom. This is also true for the one-dimensional PPP model [Eq. 1] for which however, no analytical expressions for the spin and charge velocities are available in the literature. Hence, unlike $\vartheta_\rho/\sigma$, $\vartheta^h/s_L$ depends on system size. This dependence is more pronounced in the Hückel and Hubbard models, compared to the PPP model, due to the absence of long-range interactions. The ratio of the velocities, $\vartheta^h/s_L$, due to finite size effects is system size dependent. We need to go to longer chains (longer $L$) for this ratio to be size independent. For the PPP model however, this dependence is weak (see Fig. 3).

In Table I we present the times taken by the charge and spin of the hole to propagate from one end of the chain to the other end, as well as, the charge and spin velocities, and their ratios. The times taken by the spin and charge to reach the end of the chain approximately
scale with chain length. However, the velocities of both charge and spin increase slightly with increasing chain length, which is due to weak finite-size effects. In Table I, we also present the spin and charge velocities extrapolated to infinite system size for different dimerization strengths ($\delta = 0.0, 0.05, 0.07$ and $0.15$), as well as, the ratio $\vartheta^h_\infty / \vartheta^s_\infty$. We see that the ratio $\vartheta^h_\infty / \vartheta^s_\infty$ decreases with an increase in $\delta$. In Fig. 3 is shown the dependence of the ratio of charge to spin velocities as a function of dimerization, for the chain lengths studied. We find from the plot that the ratio decreases as $\delta$ increases. Indeed, within the error of resolution of $\tau^h/s_L$, even for $\delta = 0.3$ and $0.5$, it appears that this trend continues and the ratio of the velocities approaches the non-interacting value of 1.0. This can be contrasted with the push-pull polyene systems studied by us, where the ratio of the spin and charge velocities is independent of the strength of the push-pull groups. The above feature, namely, dependence of $\vartheta^h_L / \vartheta^s_L$ on $\delta$, cannot be attributed to change in interactions brought about by geometry changes as a consequence of increased dimerization. This is because these changes are small and it has been shown in earlier studies that the contributions to the energy gaps between states due to changes in interaction parameters, $V_{ij}$, accompanying small bond length changes (due to change in $\delta$), is rather small. Therefore it appears that in dimerized models, the change in transfer integrals has a stronger role to play than the long-range interactions.

In order to investigate in detail the issue of the ratio $\vartheta^h / \vartheta^s$ tending to the non-interacting value of 1.0 as $\delta \to 1.0$, we have focused on the time dependence of the charge (spin) polarization defined as, $P_{c(s)}(t)$. These are calculated from the normalized site charge (spin) density $\rho^c_i(t)$, with background correction. Employing these renormalized quantities suppresses the large fluctuations seen in $\langle n_i(t) \rangle$ and $\langle s_i^z(t) \rangle$ allowing one to focus on the essential behavior of spin and charge transport. The quantities $\rho^c_i(t)$ and $\rho^s_i(t)$ are defined as

$$\rho^c_i(t) = \frac{\langle n_i(t) \rangle - C}{\sum_i (\langle n_i(0) \rangle - C)}; \quad \rho^s_i(t) = \frac{\langle s_i^z(t) \rangle - S}{\sum_i (\langle s_i^z(0) \rangle - S)},$$

(9)

where $C$ and $S$ are the average background charge and spin densities respectively, and are given by,

$$C = \frac{1}{N} \sum_j \langle \phi^0_{gs} | n_j | \phi^0_{gs} \rangle; \quad S = \frac{1}{N} \sum_j \langle \phi^0_{gs} | s_j^z | \phi^0_{gs} \rangle.$$

(10)
FIG. 3: Variation in the ratio of charge and spin velocities \( \left( \frac{\vartheta^h_L}{\vartheta^s_L} \right) \) with dimerization \( 0.0 \leq \delta \leq 0.15 \), in the PPP model, for different chain lengths.

For a half-filled ground state belonging to the \( S_{tot}^z = 0 \) subspace, \( C = 1.0 \) and \( S = 0.0 \). Using these observables, we define time-dependent charge and spin polarizations as, \( \vec{P}_c^{(s)}(t) = \sum_j \rho_j^{c(s)}(t) \vec{r}_j \). The time-dependence of the charge and spin polarizations qualitatively reflect the center-of-mass movement of the charge and spin peaks. However, these polarizations are gauge dependent as the system is having a net charge. Hence we have computed these by choosing as origin, the central site of a \( N \) site chain \( (N = 20, 30, 40) \).

In Fig. 4, we have plotted the variations in charge and spin polarizations with time, for a polyene chain of 40 sites. Motion of the charge (spin) across the two ends of the chain leads to oscillations in the temporal variation of \( \vec{P}_c^{(s)}(t) \). The larger the amplitude of the oscillations, the higher is the probability of charge (spin) traveling between the chain ends. It is observed from both the curves that, with an increase in dimerization, the oscillation amplitude decreases. This indicates that electron-phonon coupling reduces the probability of both the charge and spin degrees of freedom to travel between the two ends of the chain. This observation indeed supports our earlier observation that it is difficult to locate \( \tau^h/s_L \) for \( \delta = 0.3 \) and 0.5 merely because, weak bonds fail to transmit the charge or spin easily. In other words, the charge and spin for large values of \( \delta \), might not even reach the opposite end of the
TABLE I: Variation in the times $\tau^h_L$, $\tau^s_L$ (fs), and velocities $\vartheta^h_L$, $\vartheta^s_L$ (Å/fs), and ratio of velocity of charge to velocity of spin $\vartheta^h_L/\vartheta^s_L$, in the PPP model with dimerization, for different chain lengths, $L = (N - 1)r_0 - \delta$; $N$ is the number of sites and $r_0 = 1.397$ Å is the $C = C$ bond length in a regular polyene chain.

| $N$ | $\delta$ | $\tau^h_L$ | $\tau^s_L$ | $\vartheta^h_L$ | $\vartheta^s_L$ | $(\vartheta^h_L/\vartheta^s_L)$ |
|-----|----------|-------------|-------------|----------------|----------------|--------------------------|
| 20  | 0.0      | 1.81        | 3.77        | 14.66          | 7.04           | 2.08                     |
|     | 0.05     | 1.76        | 3.54        | 15.05          | 7.48           | 2.01                     |
|     | 0.07     | 1.75        | 3.45        | 15.13          | 7.67           | 1.97                     |
|     | 0.15     | 1.72        | 3.12        | 15.34          | 8.46           | 1.81                     |
| 30  | 0.0      | 2.59        | 5.44        | 15.64          | 7.45           | 2.10                     |
|     | 0.05     | 2.49        | 5.15        | 16.25          | 7.87           | 2.06                     |
|     | 0.07     | 2.45        | 4.98        | 16.51          | 8.12           | 2.03                     |
|     | 0.15     | 2.24        | 4.22        | 18.02          | 9.56           | 1.88                     |
| 40  | 0.0      | 3.44        | 6.97        | 15.84          | 7.82           | 2.03                     |
|     | 0.05     | 3.29        | 6.39        | 16.54          | 8.52           | 1.94                     |
|     | 0.07     | 3.23        | 6.17        | 16.85          | 8.82           | 1.91                     |
|     | 0.15     | 2.84        | 5.15        | 19.13          | 10.55          | 1.81                     |
| $\infty$ | 0.0      | -          | -           | 17.14          | 8.52           | 2.01                     |
|     | 0.05     | -          | -           | 18.15          | 9.35           | 1.94                     |
|     | 0.07     | -          | -           | 18.71          | 9.75           | 1.92                     |
|     | 0.15     | -          | -           | 22.99          | 12.43          | 1.85                     |

chain, especially for long chains. We also notice that an increase in dimerization suppresses spin-charge separation, which is manifested by the fact that $\vec{P}_c(t)$ and $\vec{P}_s(t)$ closely follow each other. Thus, indeed the ratio of $\vartheta^h_L/\vartheta^s_L \to 1.0$ as $\delta \to 1.0$. In other words, with an increase in electron-phonon coupling, spin-charge separation is suppressed in the PPP model.

However, the question about fate of the charge and spin of the injected hole for large
values of $\delta$, still remains. In order to investigate this issue, we have computed
\begin{align}
\frac{dN_{RL}(t)}{dt} &= \frac{N_R(t) - N_L(t)}{\Delta t}, \quad (11) \\
\frac{dS^z_{RL}(t)}{dt} &= \frac{S^z_R(t) - S^z_L(t)}{\Delta t}. \quad (12)
\end{align}
Here, $N_{R/L}(t) = \sum_{j \in R/L} \langle n_j(t) \rangle$ and $S^z_{R/L}(t) = \sum_{j \in R/L} \langle s^z_j(t) \rangle$; $R \in [N/2 + 1, N]$ and $L \in [1, N/2]$. These quantities provide information about the amount of charge and spin of the hole that is transported from the left-half to the right-half of the chain. In the initial state of the system, both the charge and spin of the hole reside solely in the left-half of the system. Hence, $\frac{dN_{RL}(t)}{dt}$ and $\frac{dS^z_{RL}(t)}{dt}$ at time $t = 0$ have the values 1.0 and 0.5, respectively. With time, as the charge and spin travel from the first to the last site of the chain, these quantities change sign and depending on the strength of electron correlations, attain values close to $-1.0$ and $-0.5$. When the charge and spin travel back from the last to the first site, these quantities again change sign and reach values close to $+1.0$ and $+0.5$. This thus indicates that, $\frac{dN_{RL}(t)}{dt}$ and $\frac{dS^z_{RL}(t)}{dt}$ oscillate between $\pm 1.0$ and $\pm 0.5$, respectively. The amplitude of oscillation of these quantities with time gives a measure of the amount of charge and spin that has been transported from the left-half to the right-half of the system. If under some
FIG. 5: Variation in $\frac{dN_{RL}(t)}{dt}$ (dark curves) and $\frac{dS_{z,RL}(t)}{dt}$ (light curves) with time, for a 40 site PPP chain, for different dimerization strengths: $\delta = 0.0, 0.05, 0.07, 0.15, 0.3$, and 0.5.

circumstance the charge (spin) fails to travel between the ends of the chain, time evolution of these two quantities does not not show any sign change.

Figure 5 depicts the time evolution of $\frac{dN_{RL}(t)}{dt}$ and $\frac{dS_{z,RL}(t)}{dt}$ for a polyene chain of 40 sites, for different dimerization strengths. It is observed that these quantities indeed oscillate between $\pm 1.0$ and $\pm 0.5$, although the amplitude of oscillations decreases with time, as $\delta$ increases. For small dimerizations ($\delta = 0.05$ and 0.07), it is observed that the temporal variations in $\frac{dN_{RL}(t)}{dt}$ and $\frac{dS_{z,RL}(t)}{dt}$ are similar to those in the undimerized case, and the number of times $\frac{dN_{RL}(t)}{dt}$ and $\frac{dS_{z,RL}(t)}{dt}$ changes sign is the same for all the three cases. However, the number of “sign changes” in the time propagation of $\frac{dN_{RL}(t)}{dt}$ is more than that of $\frac{dS_{z,RL}(t)}{dt}$, indicating that the charge degree of freedom moves faster compared to the spin degree of freedom which is merely a manifestation of spin-charge separation. Thus, for small values of electron-phonon coupling the motion of the charge and spin of the hole do not get hindered due to interferences.

However for $\delta = 0.15, 0.3$ and 0.5, the time evolution profiles of $\frac{dN_{RL}(t)}{dt}$ and $\frac{dS_{z,RL}(t)}{dt}$ are completely different from those obtained with small values of $\delta$. It is observed that the amplitude of oscillations diminishes with time indicating that the amount of charge and
spin transported across the chain decreases as electron-phonon coupling increases. This supports our observation that an increase in \( \delta \) leads to a decrease in the height of the minima, in the time evolution profiles of \( \langle n_L(t) \rangle \) and \( \langle s_z^L(t) \rangle \). Also, the number of times that \( \frac{dN_{RL}(t)}{dt} \) and \( \frac{dS_z^{RL}(t)}{dt} \) changes sign decreases, which reflects that strong electron-phonon coupling significantly reduces the probability of to-and-fro motion of the charge and spin along the chain. For \( \delta = 0.3 \) and 0.5, however, \( \frac{dS_z^{RL}(t)}{dt} \) does not change sign within the time scale of our studies. This depicts that, for these values of \( \delta \), only the charge degree of the hole can travel across the length of the chain, albeit slowly and in a very small amount. The spin of the hole fails to reach the end of the chain and “gets trapped” within the left-half of the chain. This is the reason why we find that locating \( \tau_{h/s}^L \) for large values of \( \delta \) is difficult.

Bosonization studies on one-dimensional spinful Hubbard and Luttinger liquid models, at half-filling\(^ {45-47} \), indicate that \( \delta \neq 0 \) introduces a backward-scattering operator into the bosonized Hamiltonian, which is associated with momentum transfer \( q \sim 4k_F \). Because of this umklapp scattering term, as dimerization increases, propagation of the charge and spin degrees of freedom of the injected hole between the ends of the chain become hindered, and for large values of \( \delta \) (0.3 and 0.5), fail to reach the chain end \( (L) \). Furthermore, periodic perturbation of the lattice such as Peierls distortion, which introduces a gap in the excitation spectrum, has been shown to couple the charge and spin sectors of the one-dimensional (bosonized) Hubbard Hamiltonian.\(^ {47} \) These two results, obtained with respect to spinful fermion models with short-range electron-electron interactions, seem to support both our observations in the context of the PPP model, namely, an increase in dimerization suppresses spin-charge decoupling, and inhibits the oscillatory to-and-fro motion of the charge and spin of the injected hole.

IV. CONCLUSION

To conclude, we have shown that dimerization modifies the dynamics of spin and charge transport in the PPP model. For small dimerizations the system shows a smooth decrease in the ratio of charge to spin velocities, although both the velocities decrease slightly. For large values of \( \delta \), the situation is different. It is difficult to obtain the charge and spin velocities as strong interference effects smear out their motion. However, a more careful
analysis based on net charge transport across the chain shows that both the charge and spin stay almost localized for the duration of our time evolution. Thus, although dimerization of polyenes predominantly changes the one-electron part of the Hamiltonian, it strongly affects spin-charge separation. Large values of dimerization reduce the amount of charge and spin transported across the system as well as, suppress spin-charge separation, as evidenced by the decrease in $\hat{\vartheta}^h_L / \hat{\vartheta}^s_L$ ratio and by closely following time evolution profiles of spin and charge polarizations.

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