1. INTRODUCTION

The quantum transport in molecular junctions has seen great progress in recent years. Single molecular junction has been widely studied to use nonequilibrium many-body quantum phenomena on the nanoscale. Application has been demonstrated successfully in molecular electronics/molecular devices and scanning tunneling microscopy (STM). Interests phenomena have emerged such as Coulomb blockade, Kondo effect, chiral-induced spin selectivity, phonon blockade, hysteresis, etc. The charge occupation number at the molecular bridge site often changes with the molecular geometry. In fact, vibrational contributions have been observed in the experimental measurement of currents or current–voltage curves. By and large, organic molecules are flexible, naturally leading to large vibrational coupling.

A number of theoretical methods have been developed to tackle those problems of quantum transport in molecular junctions with interactions, such as nonequilibrium Green function (NEGF) methods, scattering theory, and master equation methods. NEGF and master equation approaches usually rely on perturbation treatments. We intend to develop a nearly exact, robust, and systematic numerical method to study quantum transport problems with consideration of many-body effects.

Numerical methods have been developed to achieve this goal, such as quantum Monte Carlo, dynamical mean-field theory (DMFT), numerical renormalization group, hierarchical equations of motion (HEOM), and the multilayer multiconfiguration time-dependent Hartree (ML-MCTDH) method. In this paper, we propose to use the time-dependent density matrix renormalization group (TDMRG) method, in close comparison with ML-MCTDH. DMRG is a specific structure of a tensor network which is nearly exact for one-dimensional quantum many particle systems. It was established by White in 1992. Since then, many new algorithms have been developed, such as dynamical DMRG, finite temperature, and time-evolving methods. Recently, we have implemented the GPU acceleration algorithms and proposed a general automatic construction method for the matrix product operators. We further presented a general picture for carrier transport in organic semiconductors combining DMRG and the Kubo formula with the Holstein–Peierls model featuring both inter- and intramolecular vibrational couplings and applied its algorithm to calculate the thermoelectric power factor. There have been theoretical works to calculate electron transport of molecular junction using DMRG without considering the electron phonon interactions. In this work, we develop an algorithm to study a general model of vibrationally coupled electron transport in molecular junctions, based on time-dependent DMRG, at both zero and finite temperature.

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2. THEORETICAL MODEL AND METHODOLOGICAL APPROACH

2.1. Model Hamiltonian. The vibrationally coupled electron transport model in the molecule junction has been proposed previously.\(^\text{13}\) The simplest model consists of a single molecular site, harmonic oscillators coupled to the molecule, and the left and right leads. The Hamiltonian reads:

\[
H = H_{\text{el}} + H_{\text{nuc}} + H_{\text{el-nuc}}
\]

\[
H_{\text{el}} = E_d d^+ d + \sum_{k_l} E_k c_{k_l}^+ c_{k_l} + \sum_{k_k} E_{k_k} c_{k_k}^+ c_{k_k} + \sum_{k_k} V_{d k_k} (c_{k_k}^+ d + d^+ c_{k_k}) + \sum_{k_k} V_{d k_k} (c_{k_k}^+ d + d^+ c_{k_k}) \tag{1}
\]

\[
H_{\text{nuc}} = \frac{1}{2} \sum_j \left( P_j^2 + \alpha_j^2 Q_j^2 \right) = \sum_j \alpha_j \left( a_j^+ a_j + \frac{1}{2} \right) \tag{2}
\]

\[
H_{\text{el-nuc}} = d^+ d \sum_j c_j Q_j - d^+ d \sum_j \frac{c_j}{\sqrt{2\alpha_j}} \left( a_j^+ + a_j \right) \tag{3}
\]

which describes pure electronic terms of the system and leads, vibration of the system, and electron vibration couplings, respectively. The bath Hamiltonian is diagonal; namely, there is no coupling between bath sites. \(d^+ (d)\) is the electron creation (annihilation) operator of the molecular bridge. \(c\)'s are from the left and right leads, respectively. \(a_j^+\) and \(a_j\) are phonon creation and annihilation operators, while \(P_j/Q_j\) are the momentum and displacement operators of normal modes. \(E_d\) is the onsite energy of the molecular bridge, and \(E_{k_k}\) and \(E_{k_k}\) are the onsite energies of the left and right electrodes, respectively. \(V_{d k_k}\) and \(V_{d k_k}\) are coupling strengths between the molecular bridge and leads. These are calculated by the following width functions and the corresponding continuous functions:

\[
\Gamma(E) = \left\{ \begin{array}{ll}
2\pi \sum_{k_k} |V_{d k_k}|^2 \delta(E - E_{k_k}) & |E| \leq 2|\beta| \\
0 & |E| \geq 2|\beta| 
\end{array} \right.
\]

\[
\Gamma(E) = \left\{ \begin{array}{ll}
\frac{\alpha_j^2}{\beta_j^2} \sqrt{4\beta_j^2 - E^2} & |E| \leq 2|\beta_j| \\
0 & |E| \geq 2|\beta_j| 
\end{array} \right.
\]

\[
\Gamma_l(E) = \Gamma(E - \mu_L) \\
\Gamma_R(E) = \Gamma(E - \mu_R) \tag{5}
\]

The applied bias voltage changes the chemical potentials of left and right leads with a Fermi surface at 0 eV.

\[
\mu_{L/R} = \pm V/2 \tag{6}
\]

The electronic continua are discretized evenly, i.e., with equal separation in the interval \([-2\beta, 2\beta]\), though different schemes had been tried.\(^\text{45}\) \(\omega_j\) is the frequency of normal modes. \(c_j\) is the electron–nuclear coupling constant, or \(c_j / (\alpha_j \sqrt{2\omega_j})\) is the dimensionless electron–phonon coupling constant. These can be calculated by the following spectral density function and the corresponding continuous functions is the Ohmic spectral density function:

\[
J(\omega) = \frac{\pi}{2} \sum_j \frac{c_j^2}{\omega_j} \delta(\omega - \omega_j) \tag{7}
\]

\[
I_{\text{O}}(\omega) = \frac{\pi}{2} a_0 e^{-\omega \omega_j / \alpha_j} \tag{8}
\]

The discretization is different from that of the fermion bath, which is not evenly distributed, and \(\omega_j\) and \(c_j\) can be acquired from the integral of \(\rho(\omega)\):

\[
\rho(\omega) = \frac{N + 1}{\alpha_j} e^{-\omega / \alpha_j} \tag{9}
\]

\[
\int_0^{\omega_j} \text{d} \omega \rho(\omega) = j, \quad j = 1, 2, ..., N \tag{10}
\]

\[
\omega_j = -\omega_j \ln \left( 1 - \frac{j}{N + 1} \right) \tag{11}
\]

\[
c_j = \frac{2}{\pi} \int_0^{\omega_j} \frac{I_{\text{O}}(\omega)}{\rho(\omega)} \text{d} \omega = \sqrt{\frac{a_0^2 \omega_j}{N + 1}} \tag{12}
\]

Due to the role of leads or chemical potential, a grand canonical ensemble should be considered. At finite temperature, the initial density matrix is

\[
\rho = \rho_0^0 e^{-\beta (H_0 - \mu_L N_L - \mu_R N_R)} \tag{13}
\]

in which

\[
H_0 = \sum_{k_k} E_{k_k} c_{k_k}^+ c_{k_k} + \sum_{k_k} E_{k_k} c_{k_k}^+ c_{k_k} + H_{\text{nuc}}^0 \tag{14}
\]

and \(\rho_0^0\) is the initial density matrix of the molecular bridge. \(N_\xi = \sum_{k_k} c_{k_k}^+ c_{k_k}(\xi = L, R)\) represents the occupation number in each lead. Eq 15 represents a grand canonical ensemble description. \(H_{\text{nuc}}^0\) is the equilibrium initial phonon bath Hamiltonian. It equals \(H_{\text{nuc}}\) if the bridge state is unoccupied initially, and it becomes \(H_{\text{nuc}} + \sum_j c_j Q_j\) if the bridge state is occupied initially, namely, a displaced harmonic oscillator. The nonequilibrium initial phonon bath has been discussed in the literature.\(^\text{46,47}\) At zero temperature, the density matrix becomes a pure state, which should be the ground state of \(H_0 - \mu_L N_L - \mu_R N_R\).

With the time-evolving Hamiltonian and initial density matrix, we are interested in the electric current as a time derivative of bath occupation \((h = e = 1)\), and the current operator can be evaluated by the following commutator:

\[
I_L(t) = -\frac{\text{d} N_L(t)}{\text{d} t} = \frac{-\text{tr}[\rho \text{e}^{iHt} [H, N_L] \text{e}^{-iHt}]}{\text{tr}[\rho]} \tag{15}
\]

\[
I_R(t) = \frac{\text{d} N_R(t)}{\text{d} t} = \frac{\text{tr}[\rho \text{e}^{iHt} [H, N_R] \text{e}^{-iHt}]}{\text{tr}[\rho]} \tag{16}
\]

\[
I = \text{tr}[H, N_\xi] = i \sum_{k_k} V_{d k_k} (-c_{k_k}^+ d + d^+ c_{k_k}) \tag{17}
\]

In this paper, we evaluate current as the average of left and right currents, and another important physical quantity is the population of the system, following Wang et al.\(^\text{10,13}\)
2.2. TD-DMRG Approach. 2.2.1. MPS and MPO. In order to treat the time-dependent transport problems in a rigorous way, we employ time-dependent density matrix renormalization group (TD-DMRG) algorithms. Below, we will introduce DMRG briefly in the language of matrix product state (MPS) and matrix product operator (MPO) and extend the algorithms to finite temperatures and identical particles.

The wave function is described as a chain product of matrices in the language of MPS:

$$\Psi = \sum_{|\sigma_1, \sigma_2, \ldots|} A[1]^{\sigma_1} A[2]^{\sigma_2} \ldots A[n]^{\sigma_n} |\sigma_2 \ldots \sigma_n\rangle,$$

in which $a_i$ denotes the dimension of the $i$th virtual bond, and $\sigma$ is the physical bond index as shown in Figure 1a.

![Figure 1. Schematic diagram of (a) MPS and (b) MPO. \{\sigma, \sigma'\} are the physical bonds, and \{a\} is the virtual bonds.](image)

In principle, if the bond dimensions are infinite, then TD-DMRG is able to solve the time-dependent Schrödinger equation exactly. In practice, they have to be truncated. Therefore, extending the virtual bond dimensions can systematically improve the accuracy of the method. The physical bond dimensions denote the basis of each site. Similarly, the operator can be constructed as the following chain of matrices:

$$\hat{O} = \sum_{|\sigma_1, \sigma_2, \ldots|} W[1]^{\sigma_1} W[2]^{\sigma_2} \ldots W[n]^{\sigma_n} |\sigma_2 \ldots \sigma_n\rangle$$

in which $a_i$ denotes the dimension of the $i$th virtual bond, and $\sigma$ and $\sigma'$ are the physical bond indexes as shown in Figure 1b.

2.2.2. Time Evolution Algorithm. To solve the time-dependent Schrödinger equation, we use the time-dependent variational principle (TDVP) method for time evolution. The Dirac–Frenkel TDVP is expressed as

$$\frac{\partial}{\partial t} \langle \Psi | \hat{H} | \Psi \rangle = 0$$

Using the tangent space of $\langle \Psi |$, the equation can be written as

$$\frac{\partial |\Psi\rangle}{\partial t} = -i \hat{H} |\Psi\rangle$$

The expectation value of any operator $\hat{F}$ in $P$ space at 0 K is

$$\langle \hat{F}(t) \rangle = \langle \Psi(0) | e^{-i\hat{H}t/2} \hat{F} e^{-i\hat{H}t/2} |\Psi(0)\rangle$$

in which the time evolution is done as in eq 31, step by step numerically.

2.2.3. Finite Temperature and Imaginary Evolution Algorithm. At a finite temperature, the initial state is not a pure quantum state. It cannot be described by a single MPS. We then apply the thermo field dynamics approach (or “ancilla” or “purification” approach). For any density matrix $\rho_p$ of a quantum system P we can find a pure state $|\Psi_{PQ}\rangle$ in space $P$ and its auxiliary space $Q$ such that $\rho_p = Tr_Q(|\Psi_{PQ}\rangle\langle\Psi_{PQ}|)$. If $\rho_p$ has orthonormal decomposition,

$$\sum_{i} p_i |i_p\rangle \langle i_p|,$$

then one way to define the pure state is

$$|\Psi_{PQ}\rangle = \sum_{i} \sqrt{p_i} |i_p\rangle |i_q\rangle,$$

which can be easily proved to meet $\rho_p = Tr_Q(|\Psi_{PQ}\rangle\langle\Psi_{PQ}|)$.

Then, the thermal initial density matrix is

$$\frac{e^{-\beta H}}{Tr(e^{-\beta H})}$$

So we can define $\langle \Psi(\beta) |$ and $\langle \Psi(\beta) | P |\Psi(\beta)\rangle$ which is the imaginary time evolution of $\sum |i_p\rangle \langle i_p|$, and normalize it at each step. The expectation value of any operator $\hat{F}$ in $P$ space is

$$\frac{Tr\langle \hat{F} e^{-i\hat{H}t/2} | e^{-i\hat{H}t/2} \hat{F} e^{-i\hat{H}t/2} |\Psi(\beta)\rangle}{Tr\langle e^{-i\hat{H}t/2} | e^{-i\hat{H}t/2} \hat{F} e^{-i\hat{H}t/2} |\Psi(\beta)\rangle}$$

In eq 25, $P$ is the projection operator made up of orthonormal vectors belonging to the tangent space. $Tr_{PQ}$ can be regarded as an orthogonal projection onto the current tangent space of $|\Psi(\beta)\rangle$. Here,

$$\hat{P}[1: i] = \sum_{a_i} |a_i [1: i]\rangle S[1: i] |a_i [1: i]\rangle$$

$$\hat{P}[i + 1: n] = \sum_{a_i, a'_{i+1}} |a_i [i + 1: n]\rangle S[i + 1: n] |a_i [i + 1: n]\rangle$$

$$\hat{I}_i = \sum_{a_i} |a_i\rangle \langle a_i|$$

$$\hat{P}[1: 0] = \hat{P}[n + 1: n] = 1$$

Here, $S$ is the overlap matrix due to the nonorthogonality of the renormalized basis. Then, after a tiny time $\Delta t$ propagation, the wave function of eq 25 can be solved as

$$\langle \Psi(t + \Delta t) | = e^{-\beta \Delta t} |\Psi(t)\rangle$$

The expectation value of any operator $\hat{F}$ at 0 K is

$$\langle \hat{F}(t) \rangle = \langle \Psi(0) | e^{-i\hat{H}t/2} \hat{F} e^{-i\hat{H}t/2} |\Psi(0)\rangle$$

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Actually, the initial thermal density matrix (eq 15) is a grand canonical ensemble (as indicated in Supporting Information 5), which can be obtained by replacing $H$ with $H - \mu N$ in eqs 33 and 34. Since it is diagonal, we can calculate $e^{\beta(H-\mu N)/2} \sum |i\rangle \langle i|/Z^{1/2}$ directly, and there is no need to use TD-DMRG to calculate the imaginary time evolution.

However, we have seen in eq 15 that the initial thermal distribution is the direct product of the system, and all baths and the baths are free fermions and bosons. So, we can use the thermal Bogoliubov transformation. Several articles have already considered the transformation for phonons, but not for fermions. The derivation and equations and the results and comparison between the transformation and imaginary time evolution are given in Supporting Information 4.

2.2.4. Jordan-Wigner Transformation. To deal with the many-particle problem for fermions, the antisymmetry property of operators is non-negligible. We employ the Jordan–Wigner transformation to express the Fermion operators into Pauli matrices:

$$
\begin{align*}
\hat{a}_i^+ &= \prod_{l=1}^{k-1} (-\sigma_l^z)\hat{c}_l^+ \\
\hat{a}_i^- &= \prod_{l=1}^{k-1} (-\sigma_l^z)\hat{c}_l^-
\end{align*}
$$

$$
\begin{align*}
\hat{a}_i^+\hat{a}_m^-\sigma_i^+\sigma_m^- &= \sigma_i^+\sigma_m^- \\
\hat{a}_i^+\hat{a}_m^- &= \sigma_i^+\sigma_m^- (n < m)
\end{align*}
$$

Now, the electronic Hamiltonian and the current operator become:

$$
\begin{align*}
H_{el} &= \sum_k E_k c_k^+ c_k + \left( \sum_k V_{dk}\sigma_k^+\sigma_d^- \prod_{l=(k,d)} (-\sigma_l^z) \right) + H_c \\
I &= \frac{i}{2} \left( \sum_k V_{dk}\sigma_k^+\sigma_d^- \prod_{l=(k,d)} (-\sigma_l^z) - H_c \right) \\
&\quad - \frac{i}{2} \left( \sum_k V_{dk}\sigma_k^+\sigma_d^- \prod_{l=(k,d)} (-\sigma_l^z) - H_c \right)
\end{align*}
$$

in which $k$ can denote the molecular bridge and left and right lead sites, and $(k,d)$ means the total product is acquired from all the sites between site $k$ and site $d$.

2.3. Orbital Ordering and Entanglement Entropy. The orderings of the bath sites (electrons of the leads and phonons) in MPS can make a difference to numerical accuracy. Zwolak et al. discussed the purely electronic model and gave a quite reasonable ordering for the star geometry. They ranked the lead electronic sites by energy. Thus, the left and right lead sites could be mixed up. Ranking sites by energy and mixing the left and right lead sites are to localize the entanglement in the bias window and between nearly iso-energetic sites. Different ordering schemes have also been suggested for the chain geometry. However, electron–phonon couplings have not been considered.

In this paper, we extend the appropriate ordering described above to the electron–phonon model. Except for the left and right lead baths, there is a phonon bath. Since they are all only entangled with the system bath, we will not mix the phonon sites and lead electron sites. Therefore, there are only four relative orderings of the phonon sites, and they are given in Figure 2. The white and black circles are lead sites, corresponding to sites above and below the Fermi surface, respectively, i.e., unoccupied and occupied initially. The gray circle is the system site. The blue circles are the phonon sites, and they can be placed on four different relative positions. The relative orderings of the lead sites are kept the same as in ref 39. Since entropy is easy to get in the language of MPS, we will calculate the effective entropy and bond entropy of different orderings and compare and discuss the four orderings in section 3.4.

3. RESULTS AND DISCUSSION

We employed the TD-DMRG method to investigate the electron transport in a vibrationally coupled molecular junction. The parameters utilized in the model, as detailed in section 2.1, were taken from the literature, $\beta = 1$ eV, $\alpha_e = 0.2$ eV, and $\omega_c = 500$ cm$^{-1} = 0.062$ eV, and remained constant throughout this study. For convergence tests and comparison with ML-MCTDH results, we included the discretization scheme and TD-DMRG parameters in Supporting Information 2. Figure 3 displays the additional parameters we varied to explore vibrationally coupled electron transport at 0 K. One of the

![Figure 2](image_url)

**Figure 2.** Four orderings of all the sites at 0 K. a and b mean that the phonon sites are placed closely next to the system site, from right or left, respectively. c and d mean that the phonon sites are placed far from the system site. The frames mean that ordering of the phonon or electron sites in the frames is fixed.

![Figure 3](image_url)

**Figure 3.** Diagrammatic presentation of the vibrationally coupled molecular junction electron transport at 0 K. The onsite energy $E_d$ of the molecular bridge, initial occupancy of the bridge state, reorganization energy $\lambda$, and voltage are tuned to explore the transport. The blue areas are occupied by electrons, and the white areas are empty. The orange circles are phonons.
these parameters is the on-site energy of the molecular bridge relative to the Fermi surface of the leads, denoted as $E_d$. Its value can be adjusted using gate voltage. Additionally, we set the initial occupancy of the molecular bridge to either zero or one. This choice leads to different phonon initial states and can significantly impact electron transport. Another crucial parameter is the strength of electron phonon coupling, determined by $\alpha$ in eq 14 or by the reorganization energy, $\lambda$. Last, the voltage or chemical potential of the leads in eq 8 serve as another parameter, for which we provide current−voltage curves in Supporting Information 2. The cases involving finite temperature are discussed in Supporting Information 4.

3.1. Steady Currents, Oscillation Periods, and Peak Values. We then analyze the steady currents, oscillation periods, and peak values of the currents for different reorganization energies. The whole dynamics can be found in Supporting Information 3. We focus our discussion on three initial states: $E_d = 0.5$ eV and initially unoccupied, $E_d = -0.5$ eV and initially occupied, and $E_d = -0.5$ eV and initially unoccupied. In Figure 4a, we observe that for $E_d = 0.5$ eV, the steady current increases as the electron phonon coupling strength rises. Conversely, for $E_d = -0.5$ eV, the opposite trend is observed. Notably, the initial occupancy does not impact the steady current significantly.

In Figure 4c, we examine the peak value of the currents, which present the oscillation amplitudes. The highest peaks remain unchanged regardless of changes in the reorganization energy in initially unoccupied cases, as depicted in Figure 4b. However, in $E_d = -0.5$ eV and the initially occupied case, the maximum current decreases rapidly with increasing electron phonon couplings. On the other hand, the lowest peaks increase with larger electron phonon couplings in all three cases, resulting in a decay of the oscillation amplitudes. A more detailed analysis and visualization can be found in Supporting Information 3, and we will provide a semiquantitative explanation below.

3.2. Explanation Using Polaron Transformation and Canonical Transformation. The phenomenon in Figure 5a is just like the Marcus formula: \begin{equation}
    k_{\text{Marcus}} = \frac{V^2}{\hbar} \sqrt{\frac{\pi}{2k_B T}} \exp\left[-\frac{(\Delta G + \lambda)}{4k_B T}\right]
\end{equation}
in which increasing $\lambda$ can increase or decrease the rates depending on the sign of $\Delta G$. However, it is important to note that this formula is specifically applicable in the context of organic semiconductors and may not be directly applicable to molecular junctions. Additionally, its effectiveness might be limited when considering the temperature effects.

In the initially unoccupied cases, the observed phenomena can be explained by employing the polaron transformation. For the model in eqs 1−4, it is known that the phonon and system components can be disentangled using a straightforward polaron transformation, disregarding the lead sections.
Indeed, the electron–phonon coupling has the ability to “lower” the onsite energy of the bridge, and this reduction corresponds to the reorganization energy \( \lambda = \sum_j c_j^2/(2\omega_j^2) \). This mechanism provides an explanation for the observed changes in steady currents. In the initially unoccupied case, where \( E_d = 0.5 \) eV is far above the Fermi surface of the leads, reducing it can enhance the currents. Conversely, in \( E_d = -0.5 \) eV and initially unoccupied case, where \( E_d \) is considerably below the Fermi surface of the leads, reducing it causes the “effective” onsite energy even further away from the Fermi surface. As a result, the steady currents decrease.

In Figure 4b,c, the plots for the initially unoccupied case demonstrate that the electron phonon coupling does not influence the oscillation periods. However, the amplitudes of the oscillations are reduced due to the presence of decoherence. In contrast, the \( E_d = -0.5 \) eV and initially occupied case exhibits distinct behavior. Changing the initial occupancy significantly alters the oscillation behavior. As mentioned in the following paragraph of eq 15, different initial occupancies lead to distinct \( H^{0}_{\text{nuc}} \) terms. In the case of the initially occupied situation, the phonons do not behave as simple harmonic oscillators before time evolution, but rather as displacement harmonic oscillators:

\[
H = E_d d^+d + \sum_j \omega_j \left( a_j^+a_j + \frac{1}{2} \right)
+ \sum_j \frac{\xi_j}{\sqrt{2\omega_j}} d^+d \left( a_j^+ + a_j \right)
+ \sum_j \omega_j \left( a_j^+a_j + \frac{1}{2} \right)
\tag{41}
\]

The electron and phonon parts are disentangled in this approximation. This “new approximation”, as we will refer to it in the subsequent discussions, differs significantly from the polaron transformation. For the polaron transformation, the phonon operators are replaced by their expectation value. However, in eq 43, the electron operator \( d^+d \) is replaced by its expectation value to eliminate electron phonon coupling. Additionally, it differs from eq 40 in terms of lowering the onsite energy by \( \sum_j c_j^2/\omega_j^2 = 2\lambda \) instead of \( \lambda \).

For the case when \( d^+d \approx 1 \), the Hamiltonian described in eq 43 reduces to a purely electronic Hamiltonian. This should yield similar results for the electron phonon model and an “equivalent” purely electronic model, where the steady current can be calculated using the Landauer formula.\(^{11,16,17,56}\) Figure 5 allows us to verify this assumption. Figure 5a illustrates that the steady currents of the \( E_d = -0.5 \) eV and initially occupied electron phonon model closely match those of the corresponding purely electronic model and predictions from the Landauer formula. Similarly, Figure 5b,c demonstrate that the oscillation period and peak value of the currents exhibit remarkable similarity between the electron phonon model and corresponding purely electronic model. Thus, the behavior observed in
Figure 4 can be primarily explained as purely electronic behavior rather than as an effect of electron phonon coupling. This comparison further highlights the differences between the polaron transformation and canonical transformation. While they both lower the effective on-site energy and eliminate the electron phonon coupling, the former is primarily used for conceptual understanding and unable to eliminate electron phonon coupling. In contrast, with the latter, the many body effects can be partially eliminated by appropriately selecting the parameters.

It is verified in Figure 5 that the approximation $d^+d \approx 1$ is valid under specific conditions. It is a good approximation when the initial occupancy is unity, and the effective on-site energy is considerably below the Fermi surface. This gives rise to an interesting phenomenon. In $E_d = 0.5 \text{ eV}$ far above the Fermi surface and initially occupied case, different reorganization energies can result in distinct steady occupancies. Figure 6a demonstrates this behavior, showing that for small reorganization energies the occupation number tends to approach zero from unity, while for quite large reorganization energies the occupancy remains close to one. Comparing the $E_d = 0.5 \text{ eV}$ and initially unoccupied case with the initially occupied case, Figure 6b reveals that for small reorganization energies the steady occupancy is independent of the initial occupancy. However, for large reorganization energies, two distinct steady occupancies emerge. This phenomenon is known as bistability.

Consequently, distinct steady occupation numbers arise (the “actual” reduction in onsite energy can be a fraction depending on the steady population\(^\text{(57)}\)). For $\lambda = 0.248 \text{ eV}$, $E_d - 2\lambda$ remains above the Fermi surface, rendering the canonical transformation invalid after a certain time and resulting in the same steady occupancy as the initially unoccupied case. Thus, the polaron transformation and canonical transformation offer a simple criterion for the emergence of bistability.

3.3. Comparison with NEGF. In addition to the bistability phenomenon, the two transformations can be utilized for comparison with the results obtained by the NEGF method. An NEGF approach based on the polaron transformation has been proposed to address electron transport coupled with vibrations.\(^\text{13,58–60}\) Figure 7 illustrates the stationary current obtained from the NEGF method alongside the results from eqs 40 and 43, all compared to the ML-MCTDH method, which provides numerically exact results for different electron phonon couplings. The figure shows that the NEGF method is in good agreement with the numerically exact results only for small reorganization energies, where it aligns well with the results obtained from eq 40, as they are both based on polaron transformation. However, eq 40 only considers the polaron shift of the onsite energy, hence we refer to the results from eq 40 as “polaron shift.”\(^\text{13}\) For larger reorganization energies, the $d^+d \approx 1$ approximation (eq 43) performs significantly better, confirming the validity of our approximation (if we calculate the final occupation number numerically and multiply it with $2\lambda$, the result is even better\(^\text{(57)}\)). When dealing with large electron phonon coupling, simulating the current accurately by using the NEGF method becomes challenging. Nevertheless,
for larger reorganization energies, eq 43 provides a closer approximation, consistent with our previous comparisons of the two transformations in different regimes.

In Figure 7, where the on-site energy is set to 0 eV, we observe that for small reorganization energies, the polaron shift approximation in eq 40 provides a valid description. However, for strong electron–phonon couplings, the new approximation in eq 43 outperforms the polaron shift approximation. This is clearly demonstrated in Figure 8a. However, in the $E_d = -0.5$ eV case in Figure 8b, the polaron shift one is not as accurate across all ranges and performs even worse for large reorganization energies. On the other hand, eq 43 exhibits improved accuracy in this scenario. Overall, Figure 8 illustrates that the effective bridge energy differs among the different models used, highlighting the respective scopes of application for the two approximations. Specifically, eq 43 proves to be a quite reliable approximation in regions where the on-site energy is far below the Fermi surface and in regions with strong electron–phonon couplings. The $E_d = 0.5$ eV case is not discussed due to the presence of bistability phenomena, which significantly influence the steady current as a result of the initial occupation. More detailed information about the time-dependent current can be found in Supporting Information 2.

3.4. Ordering and Entanglement Entropy. In the context of MPS, obtaining the ordering and entanglement entropy is relatively straightforward. The effective entropy can be defined as follows:

$$S_{\text{eff}} = \ln \left( \frac{1}{L - 1} \sum_{n} e^{S_n} \right)$$

The given equation, $L$ represents the total number of sites, $S_n$ is the entropy for bipartite cuts, and $n$ runs over the entire

![Figure 9](image-url)  
Entropy curves are shown for the purely electronic model. (a) Bond entropy. The peak of the orange line is at site 79 (the initial site number is zero), and the peak of the other is at site 80. (b) Effective entropy at about 77.5 fs. $T = 0$ K, $V = 0.1$ V.

![Figure 10](image-url)  
Entropy curves are shown for the electron–phonon model. (a) Effective entropy at about 77.5 fs. (b) Bond entropy of a. c and d are the same as a and b except for the position of the phonon sites. $V = 0.2$ V.
MPS chain. Figure 9 represents the effective and bond entropies of the purely electronic model. The system consists of a total of 161 sites (Supporting Information 2). The ordering follows Zwolak’s ordering convention, where the lead sites are arranged based on increasing onsite energy, and the single system site is positioned in the middle. As a result, the left half of the system is initially occupied, while the right half remains unoccupied. Figure 9a compares the orange line corresponding to \( E_g = 0.5 \text{ eV} \) with an initial occupation of zero and the blue line representing \( E_d = -0.5 \text{ eV} \) with an initial occupation of 1. Due to the electron–hole symmetry, these lines should be identical, as verified by Figure 9. It can also be observed that the peak occurs at the interface of the zero Fermi energy. This suggests that the entropy between sites with different occupation numbers or on opposite sides of the Fermi surface is greater than the entropy between sites with similar occupation numbers.\(^{39,43}\)

In the electron–phonon model, there are 50 phonon modes (Supporting Information 2). In Figure 10a and b, the phonon sites are put on the left or right side of the electronic site within the system, between the lead sites (as shown in Figure 3, labeled as a and b). For the case of \( E_g = -0.5 \text{ eV} \), when the phonon sites are on the left-hand side, the entanglement entropy is smaller; while for the case of \( E_d = 0.5 \text{ eV} \), the ordering with phonon sites on the right-hand site results in smaller entanglement entropy. There is something interesting in Figure 10b, for phonon sites on the left or right of the system site, there is a platform or a peak plus a platform, and the location of the peak is contrary for \( E_g = -0.5 \text{ eV} \) versus \( E_d = 0.5 \text{ eV} \).

In Figure 10b, the orange line exhibits a platform from site 79 to site 129, representing the bond entropy between nearest neighbor phonon sites and the bond entropy between terminal phonon sites and either lead sites or the single system site. On the other hand, the red line shows a peak at site 79, located between the single system site and the lead sites, followed by a platform extending from site 80 to site 130, representing bond entropy between nearest neighbor phonon sites and bond entropy between terminal phonon sites and either lead sites or the single system site. This behavior can be understood by considering the combination of maximal entropy across the Fermi surface\(^{39,43}\) and the polaron transformation. Using the polaron transformation\(^{13,55}\), the system site and the phonon sites can be treated as a single entity. In the case of the red line, there is expected to be a large entropy between the almost unoccupied system site and the lead site below the Fermi surface.\(^{39,43}\) However, the entropy of the phonon site that interacts with the lead site above the Fermi surface should be small. This leads to the observed peak at site 79 and the subsequent platform between the phonon sites shown by the red line. Similarly, for the orange line, the entropy of site 79 is expected to be large due to the same reason, while the entropy of site 130 should be small. Again, there is a platform between phonon sites, resulting in the shape of the orange line. The shapes of the green and blue lines in Figure 10b can also be explained using the same reason, except that the onsite energy is now below the Fermi surface. Consequently, the peak appears at the opposite position.

In Figure 10c and d, the arrangement of phonon sites is the same as in Figure 10a and b, except that the phonon sites are put outside of all the electron sites (as shown in Figure 3, labeled as c and d). Due to this change in placement, there is no longer a high-entropy platform between the phonons, resulting in smaller effective entropy values, although the largest bond entropy remains the same. For the case of \( E_g = 0.5 \text{ eV} \), when the phonon sites are placed close to the lead sites above the Fermi surface, the entanglement entropy is smaller, as indicated by the red lines in Figure 10c and d. Conversely, for the case of \( E_g = -0.5 \text{ eV} \), the situation is reversed, with a larger entanglement entropy observed for the green lines in Figure 10c and d. When comparing Figure 10b and d, it can be observed that the left part of red and green lines and the right part of orange and blue lines remain the same, respectively.

Based on the analysis of the electron phonon model for a single molecular junction with the given parameters \( \mu = 1 \text{ eV}, \alpha_d = 0.2 \text{ eV}, \omega_d = 500 \text{ cm}^{-1} = 0.062 \text{ eV}, \lambda = 0.248 \text{ eV}, \) and \( V = 0.2 \text{ V} \), it appears that the optimal ordering is to position the phonon sites outside the electron sites. However, since the entropy values are relatively small compared to the bond dimension (as indicated in Supporting Information 2), the ordering does not significantly impact the time-dependent current.\(^{39}\) Additionally, it is worth noting that the ordering remains unchanged during time evolution. In future research, the identified orderings can serve as a good initial guess combined with on-the-fly swapping techniques.\(^{61}\) This approach can be employed to explore the optimal ordering for transport problems involving vibrationally coupled single-molecular junction as well as multimolecular junctions.

4. CONCLUSIONS

In this study, the TD-DMRG algorithms were utilized to investigate the time-dependent current and occupation number of a molecular junction with vibrational coupling. The calculations were performed for both zero- and finite temperature conditions. To gain insights into the behavior of steady currents in the regime of strong electron–phonon coupling, a novel transformation and approximation method was developed.

The proposed TD-DMRG method was extensively tested across a wide range of parameters to demonstrate its correctness and computational feasibility. The results obtained from the method were found to be in excellent agreement with those obtained from ML-MCTDH calculations as well as analytical results obtained in the purely electronic limit.\(^{3,5,6}\) This successful comparison validates the accuracy and robustness of the TD-DMRG method for general molecular transport problems involving both electrons and phonons. In the strong electron phonon coupling regime, which is challenging for the NEGF method,\(^{5,6}\) our new approximation proves particularly effective. In addition, the study also delves into the entropy of different orderings, expanding the analysis to electron–phonon model problems. These entropy variations are further understood through application of the polaron transformation. Finally, in Supporting Information 4, thermal Bogoliubov transformation of both fermions and bosons is employed to address the finite temperature problems and compared with the results from imaginary time evolution results.

In this paper, the focus was primarily on the charge transport in molecular junctions. Other important aspects are heat transport, thermoelectric properties, and spin currents.\(^{11,12,20,24}\) As we demonstrated recently, TD-DMRG can be effectively applied to evaluate the thermal electric conversion power factor,\(^{38}\) and it is expected to explore many thermal effects in molecular junctions, including the Seebeck effect observed in polymeric materials\(^{64,65}\) and spin Seebeck effect.\(^{17,66}\) In future
work, we will develop TD-DMRG algorithms for studying transport in molecular junctions. This may involve combining open quantum system methods with MPS\textsuperscript{25,26} and going beyond the Lindblad equation\textsuperscript{22} to avoid the need for discretizing of the Fermi bath. These advancements will enable more comprehensive investigations into the thermal effects, thermoelectric properties, and spin currents in molecular junctions.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jctc.3c00340.

Display of the Poincaré recurrence; convergence tests for the TD-DMRG parameters and comparison with ML-MCTDH; dynamic images of the electron phonon model and purely electronic model; derivation of thermal Bogoliubov transformation of both fermions and bosons at finite temperature; and discussions about the grand canonical ensemble in this model (PDF)

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Notes
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