Nonadiabatic and Time-Resolved Photoelectron Spectroscopy for Molecular Systems

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ABSTRACT: We quantify the nonadiabatic contributions to the vibronic sidebands of equilibrium and explicitly time-resolved nonequilibrium photoelectron spectra for a vibronic model system of trans-polyacetylene. Using exact diagonalization, we directly evaluate the sum-over-states expressions for the linear-response photocurrent. We show that spurious peaks appear in the Born–Oppenheimer approximation for the vibronic spectral function, which are not present in the exact spectral function of the system. The effect can be traced back to the factorized nature of the Born–Oppenheimer initial and final photoemission states and also persists when either only initial or final states are replaced by correlated vibronic states. Only when correlated initial and final vibronic states are taken into account are the spurious spectral weights of the Born–Oppenheimer approximation suppressed. In the nonequilibrium case, we illustrate for an initial Franck–Condon excitation and an explicit pump–pulse excitation how the vibronic wavepacket motion of the system can be traced in the time-resolved photoelectron spectra as a function of the pump–probe delay.

1. INTRODUCTION

Photoelectron spectroscopy is a well-established experimental method to probe the structure of atoms, molecules, and solids.¹,² In comparison to other spectroscopic methods such as optical absorption spectroscopy, photoelectron spectroscopy is based on non-neutral transitions between many-body states, such as initial and final states, which have vanishing matrix elements for charge-neutral transitions and might have nonvanishing matrix elements for non-neutral transitions. Hence, photoelectron spectroscopy allows us to observe dipole or quadrupole forbidden transitions, which would otherwise not be accessible in optical absorption spectroscopy.

With the appearance of femtosecond laser pulses,³ which lead to the Nobel prize in chemistry awarded to A. H. Zewail, the field of time-resolved photoelectron spectroscopy has seen tremendous developments in recent years. Femtosecond laser pulses are routinely used to probe a large variety of intramolecular effects. By combining short pulses with the new possibilities of time-resolved photoelectron spectroscopy, experimentalists are now able to realize femtosecond pump–probe photoelectron spectroscopy. Here, two independent laser pulses are employed to eject photoelectrons. The first pulse is used to excite the sample, followed by a second laser pulse after a finite delay time. The second laser pulse photoexcites the system to emit a photoelectron. The energy and angle-resolved distribution of photoelectrons can then be detected by the measurement apparatus. Tuning the delay time allows us to monitor dynamical processes in the system.

These novel techniques for time-resolved pump–probe photoelectron spectroscopy have already been used to experimentally study and characterize ultrafast photochemical dynamic processes in liquid jets,⁴ to follow ultrafast electronic relaxation, hydrogen-bond-formation, and dissociation dynamics,⁵ to probe unimolecular and bimolecular reactions in real-time,⁶ and to investigate multidimensional time-resolved dynamics near conical intersections,⁷ to mention a few, all on a femtosecond time scale.

Driven by such novel experimental possibilities, there is an ongoing demand to extend and refine the existing theory to allow for a first-principles description of time-dependent pump–probe photoelectron spectroscopy and to treat the electronic and ionic responses on an equal footing. Along these lines, first steps have already been taken, which focus on photoelectron spectroscopy in real-time. On the level of time-dependent density functional theory (TDDFT), the first pioneering work to describe photoelectron spectra was based on the momentum distribution of the Kohn–Sham orbitals recorded at a reference point far from the system.⁸ More recently, a mask technique has been developed⁹ and extended to attosecond pump–probe spectroscopy.¹⁰ This approach captures the time-dependent case intrinsically and allows, for example, to directly simulate the photoemission process for given delay times and shapes of pump and probe pulses. However, in this approach, the description is so far limited to

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classical nuclei. As a result, the vibrational sidebands in time-resolved photoelectron spectra and photoabsorption are not fully captured. Other approaches in a similar direction have been realized combining TDDFT and ab initio molecular dynamics\textsuperscript{11} or by using a reduced density matrix description, which also relies on the Born–Oppenheimer approximation.\textsuperscript{12}

On the other hand, approaches based on the Born–Oppenheimer approximation allow for a detailed analysis of angular resolved photoelectron spectra and for a reconstruction of molecular orbital densities directly from the spectra.\textsuperscript{13,14} Standard quantum chemical approaches for photoelectron spectroscopy as, for example, the double-harmonic approximation (DHA)\textsuperscript{15} allow us to capture vibrational sidebands through Franck–Condon factors.\textsuperscript{16} Here, the vibronic nature of the involved initial and final states is taken into account within a harmonic approximation of the corresponding Born–Oppenheimer surfaces. Although the vibrational sidebands of photoelectron spectra can be approximately captured in the DHA, such an approach lacks the possibility to describe time-resolved pump–probe experiments explicitly.

In this work, we attempt to compare and validate existing computational tools for photoelectron spectroscopy of vibronically coupled systems. We present an approach for time-resolved photoelectron spectra, which explicitly includes the vibronic nature of the involved states and allows us to follow the photoemission process in real-time. For a realistic model system of small trans-polyacetylene oligomer chains, we investigate time-resolved photoelectron spectra and compare it to approaches such as the double-harmonic approximation. Our study is based on exact diagonalization of vibronic Hamiltonians and real-time propagations of the time-dependent Schrödinger equation in the combined electronic and vibrational Fock space of the system. This procedure gives us access to the exact correlated electron–nuclear eigenstates and time-evolved states of the system and enables us to test different levels of approximations against our correlated reference calculations. In particular, we focus on nonadiabatic effects beyond the Born–Oppenheimer approximation.

The paper is organized as follows: In Section 2, we introduce our employed model for trans-polyacetylene oligomers and provide a comparison of the Born–Oppenheimer states of the model to the exact correlated energy eigenstates of the Hamiltonian. Different levels of approximations for the photocurrent are introduced in Section 3, and the relation of photoelectron spectra to the one-body spectral function is discussed. In addition, we focus on time-resolved pump–probe photoemission spectroscopy. In Section 4, we apply the theoretical tools of Section 3 to our model system for trans-polyacetylene and discuss spurious peaks, which appear in the Born–Oppenheimer approximation for the spectra. To illustrate our approach for explicitly time-resolved spectroscopy, we numerically simulate two pump–probe photoelectron experiments. In the first example, we consider a Franck–Condon transition as the excitation mechanism, and in the second example, we explicitly propagate the system in the presence of a pump pulse. In both cases, photoelectron spectra are recorded, and we highlight the underlying nuclear wavepacket motion and the differences to the equilibrium spectra. Finally, in Section 5, we summarize our findings and give an outlook for future work.

2. MODEL SYSTEM

2.1. Su–Schrieffer–Heeger Hamiltonian and Exact Eigenvalues and Eigenfunctions. In this section, we briefly review the Su–Schrieffer–Heeger (SSH) model\textsuperscript{17,18} for trans-polyacetylene (PA) and the exact diagonalization approach.

To model PA oligomer chains (Figure 1(a)), we employ the SSH Hamiltonian to describe π-electrons in a polymer chain.

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Hamiltonian \( \hat{H}_\text{ph} \) which models electron hopping of π-electrons within a tight binding scheme, (2) the nuclear Hamiltonian \( \hat{H}_\text{ph} \) which describes all nuclei as a chain of coupled quantum harmonic oscillators, and (3) the interaction part in the Hamiltonian \( \hat{H}_\text{ph} \) which takes the coupling of electrons and nuclei up to first-order in the nuclear displacement into account. The electron–phonon coupling \( \hat{H}_\text{e-ph} \) may be combined with the kinetic term \( \hat{H}_k \) of the π electrons. The hopping parameter \(-T\) is then replaced by \(-T + \alpha (\hat{u}_{n+1} - \hat{u}_n)\).

Physically speaking, it is more likely for electrons to hop when two nuclear positions approach each other or conversely the effective hopping parameter is decreased when the nuclei are moving apart.

To gain access to all eigenvalues and eigenstates of the system, we employ an exact diagonalization technique. In the combined electron–nuclear Fock space, we explicitly construct matrix representations for all operators present in eq 1. For the photoelectron spectra, we choose to work in Fock space containing up to eight electrons has the size \( M_{\text{tot}} = 35 \times 35 \times 35 = 313,600 \). The four (three) electron Hilbert space has a size of \( M_{\text{el}}^{(3)} = 70 \times 35 \times 35 = 85,750 \) basis functions.

The Hamiltonian in eq 1 commutes with the spin operators \( \hat{S}_x, \hat{S}_y, \hat{S}_z \), particle number \( \hat{N} \), and parity \( \hat{P} \). By exploiting all these symmetries, we first block-diagonalize the Hamiltonian by ordering basis states according to tuples of eigenvalues of all symmetry operators that commute with the Hamiltonian. All remaining blocks in the Hamiltonian are then diagonalized with a dense eigenvalue solver. In contrast to standard sparse diagonalization approaches for exact diagonalization, this procedure gives us access to the full spectrum of all \( M_{\text{tot}} \) eigenvalues and eigenvectors of the static Schrödinger equation

\[
\hat{H}_{\text{sh}} |\Psi_j^{(N)}\rangle = E_j^{(N)} |\Psi_j^{(N)}\rangle
\]

Here, the eigenstates \( |\Psi_j^{(N)}\rangle \) and eigenvalues \( E_j^{(N)} \) of the SSH polyacetylene chain refer to the exact correlated stationary states of the combined system of electrons and nuclei in Fock space. To simplify the following discussion of electron removal, we always indicate the number of electrons explicitly with a superscript \( N \).

To solve for the time evolution of arbitrary initial states \( |\Phi_j^{(N)}(0)\rangle \) in the presence of pump and probe pulses, we explicitly propagate the time-dependent Schrödinger equation

\[
\frac{\partial}{\partial t} |\Phi_j^{(N)}(t)\rangle = \hat{H}_{\text{sh}} |\Phi_j^{(N)}(t)\rangle
\]

with a Lanczos propagation scheme. In the following, the exact diagonalization of the static Schrödinger equation and the time-evolved states of the correlated system serve as exact references to test the quality and validity of approximate schemes for photoelectron spectra. Due to the exponential scaling of the Fock space size, the calculation of exact eigenstates and time-evolved wave functions is limited to small SSH chains (maximum of four oligomers in the present case). Although the exact numerical solutions are only available for small SSH chains, they serve as valuable references to test approximate schemes, which then can be employed for larger systems.

2.2. Born–Oppenheimer Approximation for the Su–Schrieffer–Heeger Hamiltonian. To introduce the required notation for the following sections and to illustrate the exact potential energy surfaces, we briefly discuss the Born–Oppenheimer approximation for the SSH model. By setting the nuclear kinetic energy in eq 1 to zero, the nuclear displacements become classical parameters and we arrive at the electronic Born–Oppenheimer Hamiltonian for the SSH chain

\[
\hat{H}_{\text{sh},\text{el}} = \hat{H}_\text{e,el} + \hat{H}_\text{ph,el}
\]

\[
\hat{H}_\text{e,el} = -T \sum_{n,\sigma} \hat{c}^\dagger_{n+1,\sigma} \hat{c}_{n,\sigma} + \hat{c}^\dagger_{n,\sigma} \hat{c}^\dagger_{n+1,\sigma} + \hat{c}_{n,\sigma} \hat{c}_{n+1,\sigma}
\]

\[
\hat{H}_\text{ph,el} = \sum_{n,\sigma} \alpha (u_{n+1} - u_n) (\hat{c}^\dagger_{n+1,\sigma} \hat{c}_{n,\sigma} + \hat{c}^\dagger_{n,\sigma} \hat{c}^\dagger_{n+1,\sigma})
\]

with the corresponding eigenvalue problem

\[
\hat{H}_{\text{sh,el}} (|u_j\rangle) |\phi_j^{(N)}(|u_j\rangle)\rangle = E_j^{(N)} (|u_j\rangle) |\phi_j^{(N)}(|u_j\rangle)\rangle
\]

The eigenvalues \( E_j^{(N)} \) and \( |\phi_j^{(N)}(u_j)\rangle \) as functions of the classical coordinates \( u_j \) denote the Born–Oppenheimer surfaces of the system. Similar to the case of the exact correlated eigenstates \( |\Psi_j^{(N)}\rangle \), we here use for electronic Born–Oppenheimer states \( |\phi_j^{(N)}\rangle \) a superscript \( N \) to distinguish between the Hilbert spaces of different electron numbers. In analogy to the exact diagonalization approach for the full Hamiltonian, as discussed in the previous section, we employ here a dense exact diagonalization scheme for the electronic Born–Oppenheimer Hamiltonian. This procedure gives us access to all exact Born–Oppenheimer surfaces and corresponding Born–Oppenheimer states of the SSH chain. In addition to the exact surfaces, we compute the Hessian of the electronic energies with respect to the displacements. By diagonalizing the Hessian, we arrive at the harmonic approximation for the Born–Oppenheimer surfaces. In Figure 2, we illustrate the exact potential energy surfaces (dashed lines) along the axis of the optical normal mode of the chain and compare to the harmonic approximation of the surfaces (solid lines). As shown from the figure, for the Hamiltonian in eq 1, the harmonic approximation is very close to the exact surfaces. Overall, the model behaves rather harmonic, and only small anharmonicities are present. We emphasize, that the almost harmonic nature of the exact potential energy surfaces originates from the quadratic interaction term in the phonon Hamiltonian \( \hat{H}_\text{ph} \) because already in the exact model Hamiltonian only quadratic nuclear interaction terms are included. The only source of anharmonicity and nonadiabaticity is the electron–phonon coupling term \( \hat{H}_\text{e-ph} \) in eq 1, which introduces only small anharmonicities and nonadiabatic couplings between different electronic surfaces.

For each fixed set of nuclear coordinates \( u_j \), the electronic eigenstates \( |\phi_j^{(N)}(u_j)\rangle \) form a complete set in the many-particle Hilbert space of the electrons. For a given set of nuclear displacement coordinates \( u_j \), we can expand the exact many-body wave function in terms of the electronic eigenstates \( |\phi_j^{(N)}\rangle \) and the nuclear eigenstates \( |\phi_j\rangle \) in terms of the Born–Huang expansion.
Figure 2. Exact Born–Oppenheimer surfaces and harmonic approximation for a trans-polyacetylene oligomer. In the panel on the left-hand side, a one-dimensional cut along the optical axis of the exact potential energy surfaces is shown in dashed lines. The harmonic approximations to the exact BO surfaces are shown in solid lines, where black lines refer to $N - 1$ electron states and the red line corresponds to the $N$-electron ground state. In the panel on the right-hand side, the corresponding photoelectron spectrum in double-harmonic approximation is shown.

\[ h_{\text{ph},j} = \hat{H}_{\text{ih},j} + \epsilon_{d,j}(\{u_n\}) \]

(7)

Here, the electronic eigenstates $\{\psi^{(N)}(\{u\})\}$ depend parametrically on the oligomer displacements $\{u\}$, and the nuclear eigenstates $\{\nu_{ij}(\{u\})\}$ are functions of $\{u\}$. We solve for the states $\{\nu_{ij}(\{u\})\}$ by diagonalizing the corresponding nuclear Born–Oppenheimer Hamiltonian.

The above form of Fermi’s Golden Rule, we can formulate the exact expression for the photocurrent \( J_k(\omega) \) in first-order perturbation theory\(^2,28,32\) as

\[ J_k(\omega) = \frac{2\pi}{\hbar} \sum_{j} \langle \psi_{k}^{(N)} | \Delta \psi_{j}^{(N)} \rangle^2 \delta(E_k - E_j - \hbar\omega) \]

(8)

Here, $\{|\psi_{k}^{(N)}\rangle\}$ denotes the final state, where the emitted photoelectron with momentum \( k \) and energy \( E_k \) is typically assumed to be in a scattering state (distorted plane wave or time-inverted scattering/LEED state, see ref 1 and references therein). The remaining part of the system is left in the excited state \( j \) with energy \( E_j \) carrying \( N - 1 \) electrons. In both subsystems, the emitted electron and remaining photofragment are in general still correlated in the combined state $\{|\psi_{k}^{(N)}\rangle\}$. The wave function $\{|\psi_{j}^{(N)}\rangle\}$ represents an initial state of the many-body system from where the photoelectron will be emitted. The above form of Fermi’s Golden Rule is strictly valid only for pure states as initial and final states. In these cases, usually the $N$-electron ground state is considered, but also excited eigenstates or superposition states are allowed. For many experimental setups, it is not justified to consider the ground state as initial state for the photoemission process. In particular, in pump–probe photoelectron spectroscopy, the system is

Table 1. Exact Correlated Energies $E_{\text{exact}}^{\text{exact}}$ and $E_{\text{BO}}^{\text{exact}}$, and Overlap between Exact and BO States\(^a\)

<table>
<thead>
<tr>
<th>state no.</th>
<th>$E_{\text{exact}}^{\text{exact}}$</th>
<th>$E_{\text{BO}}^{\text{exact}}$</th>
<th>(e,o,a)</th>
<th>overlap</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$-11.3414$</td>
<td>$-11.3419$</td>
<td>1,0,0</td>
<td>0.9986</td>
</tr>
<tr>
<td>2</td>
<td>$-11.2166$</td>
<td>$-11.2171$</td>
<td>1,0,1</td>
<td>0.9986</td>
</tr>
<tr>
<td>3</td>
<td>$-11.1583$</td>
<td>$-11.1588$</td>
<td>1,1,0</td>
<td>0.9953</td>
</tr>
<tr>
<td>4</td>
<td>$-11.0918$</td>
<td>$-11.0924$</td>
<td>1,0,2</td>
<td>0.9986</td>
</tr>
<tr>
<td>5</td>
<td>$-11.0336$</td>
<td>$-11.0341$</td>
<td>1,1,1</td>
<td>0.9955</td>
</tr>
<tr>
<td>86</td>
<td>$-9.5155$</td>
<td>$-9.5157$</td>
<td>1,10,0</td>
<td>0.9676</td>
</tr>
<tr>
<td>87</td>
<td>$-9.5076$</td>
<td>$-9.5078$</td>
<td>1,8,3</td>
<td>0.9740</td>
</tr>
</tbody>
</table>

\(^a\)All energies are given in eV. The label (e,o,a) refers to the BO quantum number of the state (electronic state, optical phonon mode, acoustical phonon mode). Note, that the exact BO energies $E_{\text{BO}}^{\text{exact}}$ provide a lower bound to the exact correlated energies $E_{\text{exact}}$. The BO photoelectron spectra acquire spurious peak amplitudes that are not present in the exact correlated spectrum.

3. THEORY OF STATIC AND TIME-DEPENDENT PHOTOELECTRON SPECTROSCOPY

In this section, we briefly review the connection between photoelectron spectra and the one-body spectral function known from literature\(^28-31\) and extend the discussion to vibronic states. For later purposes, we discuss the equilibrium and nonequilibrium spectral functions. Because our emphasis in the present work is on pump–probe photoelectron experiments for vibronic systems, we keep an explicit focus on the reference-state dependence of the vibronic one-body spectral function, and we discuss how the photocurrent can be expressed in a real-time evolution.

In terms of Fermi’s Golden Rule, we can formulate the exact expression for the photocurrent \( J_k(\omega) \) in first-order perturbation theory\(^2,28,32\) as

\[ J_k(\omega) = \frac{2\pi}{\hbar} \sum_{j} \langle \psi_{k}^{(N)} | \Delta \psi_{j}^{(N)} \rangle^2 \delta(E_k - E_j - \hbar\omega) \]
typically not in the ground state when the photoelectron is removed from the system. We illustrate the effect of different initial states for the photoemission process in detail in Section 4 and later in this section.

Generally, the coupling element $\Delta$ between initial and final states can be written in second quantization as

$$\hat{\Delta} = \sum_{l,m,\sigma} \Delta_{l,m,\sigma} \hat{c}_{l,m,\sigma} \hat{c}_{l,m,\sigma}^\dagger$$

(9)

where $\Delta_{l,m,\sigma} = \langle \phi_{l}^{(1)} | \hat{O} | \phi_{m}^{(1)} \rangle$. The coupling to the pump and probe laser pulses is usually considered in the dipole approximation with length gauge $\hat{O} \propto \vec{r} \times \mathbf{E}(r,t)$ or velocity gauge $\hat{O} \propto \dot{\vec{r}} \times \mathbf{A}(r,t)$ (without treating multiphoton processes). $\mathbf{E}$ and $\mathbf{A}$ refer to the electric field and electromagnetic vector potential, respectively. In the framework of second quantization, the wave functions $\langle \phi_{l}^{(1)} | \hat{O} | \phi_{m}^{(1)} \rangle$ form a complete set of one-body states. With no external magnetic field applied to the system, the matrix element is diagonal in spin because the operator does not act on the spin part of the wave function. The sudden approximation\(^{28,30}\) allows the final state to decouple.

$$\langle \psi_{l,k}^{(N)} | \hat{\Delta} | \psi_{j}^{(N-1)} \rangle \approx \delta_{kj} \langle \psi_{l,k}^{(N)} | \psi_{j}^{(N-1)} \rangle$$

(10)

This approximation implies that the final state is a product state between a plane wave-like state for the emitted electron and the remaining $N - 1$ electron many-body state $\langle \psi_{j}^{(N-1)} |$. At this point, we emphasize that the original matrix elements, which contribute to the photocurrent in eq 8, only contain states with fixed electron number $N$. Therefore, photoemission has to be regarded as a charge neutral excitation process induced by the presence of a laser field. Only when the excited photoelectron is starting to spatially separate from the remaining photofragment, the system is left in a charged $N - 1$ electron state. This spatial separation is also the basis of the mask approach of ref 9. In general, the emitted photoelectron can still be entangled with the remaining photofragment, for instance, in strong-coupling situations. However, for a weak-coupling situation encountered in the range of the validity of the Fermi’s Golden Rule expression for the photocurrent in eq 8, this entanglement is often neglected. If no entanglement of the emitted photoelectron and the remaining photofragment remains when the charges separate, then the state $\langle \psi_{l,k}^{(N)} |$ can be factorized. This is the basic assumption of the sudden approximation in eq 10. As a result, in the matrix elements of the photocurrent in eq 8, the neutral $N$ electron state can be replaced by an ionic $N - 1$ electron state. Only in this sense, we can talk about non-neutral excitations in a photoemission experiment, albeit initially a neutral excitation has taken place.

In terms of the usual fermionic anticommutation relation, we can write

$$\{\hat{c}_{l,k}, \hat{c}_{l,m}^\dagger\} = \delta_{kl} \delta_{lm} = \delta_{kl} \delta_{lm} - \delta_{lm} \delta_{kl}$$

(11)

and because the state $| k \rangle$ of the ejected photoelectron is an energetically high-lying state, virtual fluctuations in the reference state can be neglected. Following this argument, the last term in eq 11 can be set to zero\(^{28}\) and this allows us to write approximately

$$\hat{c}_{l,m}^\dagger \hat{c}_{l,m} \approx \delta_{kl} \hat{c}_{l,k}$$

(12)

Using eqs 9 and 10 and the approximation in eq 12 to evaluate the matrix elements in eq 8, we arrive at

$$\langle \psi_{l,k}^{(N)} | \hat{\Delta} | \psi_{j}^{(N-1)} \rangle = \sum_{l,m,\sigma} \langle \psi_{l,k}^{(N)} | \Delta_{l,m,\sigma} \hat{c}_{l,m,\sigma} \hat{c}_{l,m,\sigma}^\dagger | \psi_{j}^{(N-1)} \rangle \approx \sum_{m,\sigma} \Delta_{l,m,\sigma} \langle \psi_{l,k}^{(N)} | \hat{c}_{m,\sigma} \hat{c}_{m,\sigma}^\dagger | \psi_{j}^{(N-1)} \rangle$$

(13)

In practical applications, the matrix element $\Delta_{l,m,\sigma}$ in eq 9 is often regarded to be constant over the investigated energy range.\(^{1,2}\) This assumption is only perfectly justified in the high-energy limit (X-ray spectroscopy). In this limit, the photoelectron spectrum is directly proportional to the spectral function. The sum-over-states expression for the photocurrent in the sudden approximation (SA) is then found to take the form

$$J_{k}^{SA}(\omega) = \frac{2\pi}{\hbar} \sum_{l,m,\sigma} \Delta_{l,m,\sigma} \langle \psi_{l,k}^{(N)} | \hat{c}_{m,\sigma} \hat{c}_{m,\sigma}^\dagger | \psi_{j}^{(N-1)} \rangle (E_{k} - \hbar \omega)$$

(14)

where we have introduced the one-body spectral function $\Delta_{l,m,\sigma}(E_{k} - \hbar \omega)$. In the following, we discuss this quantity for equilibrium and nonequilibrium situations.

3.1. Spectral Function: Sum-Over-States and Time-Domain Formulation. In this section, we state and define the equilibrium and nonequilibrium spectral function. The derivation of these quantities is given in more detail in the Appendix.

3.1.1. Equilibrium Spectral Function. For the present study, it is important to distinguish between equilibrium and nonequilibrium situations. In equilibrium, $\langle \psi_{l,k}^{(N)} |$ is a eigenstate of the full vibronic Hamiltonian. The time evolution according to the time-dependent Schrödinger equation in eq 3 is in these cases trivial because eigenstates are time-invariant up to a phase. In ground-state photoemission spectroscopy, one encounters this situation if the sample is in its ground-state before it is hit by the photoemission pulse. The equilibrium spectral function is defined as

$$A_{l,m,\sigma}^{SA}(\omega) = \sum_{f} \langle \psi_{f}^{(N)} | \hat{c}_{m,\sigma}^\dagger | \psi_{l,k}^{(N-1)} \rangle \times \langle \psi_{l,k}^{(N-1)} | \hat{c}_{m,\sigma} | \psi_{f}^{(N)} \rangle \delta(\omega - E_{f})$$

(15)

Further, in equilibrium situations only diagonal terms of the spectral function ($l = m$) need to be considered.\(^{33}\)

Equation 15 can also be formulated in terms of overlaps of time-evolved states. Using this approach, the calculation of the spectral function does not rely on a sum-over-states expression. Rather, it can be computed from an explicit time propagation

$$A_{l,m,\sigma}^{SA}(t) \propto \langle \psi_{-l}^{(N-1)}(t) | \hat{c}_{m,\sigma}^\dagger | \psi_{0}^{(N)}(t) \rangle$$

(16)

with the kicked initial state $| \psi_{-l}^{(N-1)}(t_{0}) \rangle = \hat{c}_{l,m,\sigma}^\dagger | \psi_{0}^{(N)}(t_{0}) \rangle$. Depending on the size of the Hilbert space, either eq 15 or eq 16 are more efficient to evaluate. In our case, we choose to directly evaluate eq 15 using all eigenstates from our exact diagonalization procedure. However, for larger systems, where a direct diagonalization of the system Hamiltonian is computationally not feasible anymore, eq 16 provides an alternative scheme to obtain the spectral function.

A useful relation is the sum rule\(^{34}\) that is obeyed by the equilibrium spectral function

$$S = \sum_{l} \int d\omega A_{l,m,\sigma}^{SA}(\omega) = \sum_{l} \langle \psi_{l,k}^{(N)} | \hat{c}_{m,\sigma}^\dagger | \psi_{0}^{(N)} \rangle$$

(17)
where the value $S$ gives the total number of electrons $N$ in the state $|\Psi_0^{(N)}\rangle$. When computing an explicit sum-over-states summation, the limit $S = N$ is only reached if a complete set of states with a full resolution of the identity $\sum_m |\Psi_m^{(N-1)}\rangle\langle\Psi_m^{(N-1)}| = 1$ is inserted in eq 17. For an incomplete basis of states, the sum rule deviates from $N^{\text{SA}}$. Depending on the orthogonality and completeness of the employed states, $S$ can then be lower or higher than the total number of electrons in the state $|\Psi_0^{(N)}\rangle$. Therefore, in practical calculations, this sum rule can be exploited to test convergence and the completeness of the employed basis set.

3.1.2. Nonequilibrium Spectral Function. From the expression for the photocurrent in sudden approximation, eq 14, the dependence of the photoelectron spectrum on the reference state $|\Psi_0^{(N)}\rangle$ becomes apparent. As mentioned before, in most cases, the system is assumed to be in the ground state. However, in a pump–probe experiment, this assumption is not justified anymore. As we demonstrate in Section 4, quite sizable changes arise in the photoelectron spectrum when the photoelectron is ejected from a time-evolving state $|\Psi_0^{(N)}(t)\rangle$ (see discussion in next section). Ultimately peaks, which were dark for the ground state as a reference state, might become bright transitions during time evolution of a vibronic wave packet and can eventually contribute to the photoelectron spectrum.

In nonequilibrium situations, Fermi’s Golden Rule has to be extended to also allow for arbitrary states as reference states in eq 8. This can be done straightforwardly in terms of the spectral function and is explicitly calculated in the Appendix. Here, we only state the result for the nonequilibrium spectral function

$$A_{\text{SA}}^{\text{SA}}(t, \omega) = \sum_j \langle \Psi_j^{(N-1)} | \hat{c}_{\alpha}^{\dagger} \Psi_0^{(N)}(t) \rangle \langle \Psi_j^{(N)} | \hat{c}_{\alpha} \Psi_0^{(N-1)} | \delta(h\omega - E_j) \rangle$$

In nonequilibrium situations, the time-evolution of the initial state is nontrivial. Hence, the spectral function expression is not time-invariant and explicitly depends on both the time $t$ and the frequency $\omega$. Physically, we interpret time $t$ as the delay time between the pump and the probe pulse. As compared to eq 15, we now additionally allow for time-propagated reference states $|\Psi_0^{(N)}(t)\rangle$.

The nonequilibrium spectral function can also be formulated in the time-domain

$$A_{\text{SA}}^{\text{SA}}(t, \tau) = \langle \Psi_j^{(N-1)} | \hat{c}_{\alpha}^{\dagger} \Psi_0^{(N)}(t + \tau) \rangle \langle \Psi_j^{(N)} | \hat{c}_{\alpha} \Psi_0^{(N-1)} | \delta(h\omega - E_j) \rangle$$

with the kicked initial state $|\Psi_j^{(N-1)}(t)\rangle = \hat{c}_{\alpha} |\Psi_0^{(N)}(t)\rangle$, where the kick with the operator $\hat{c}_{\alpha}$ acts at time $t$ on the state $|\Psi_0^{(N)}\rangle$.

The sum rule of eq 17 also applies in nonequilibrium situations.

One point we have to mention is the neglect of the $n$-dependence in the delta function of eq 32 in the Appendix. This approximation allows us to fix peak positions to $N - 1$ electron states. Further, the approximation gives the delta peak position a clear interpretation, rather than the usage of absolute relative energies. Considering the full $n$-dependence in the delta function shifts high energy peaks to lower energy.

3.2. Approximations for Vibronic Systems. The expression for the one-body spectral function in eq 15 is still formulated in terms of correlated energy eigenstates of the full vibronic Hamiltonian. Therefore, the direct evaluation of this expression is usually a formidable task. For vibronic systems, the most straightforward approximation is to replace the correlated vibronic initial and final states by factorized Born–Oppenheimer states.

For the following discussion in Section 4, we define as a single-harmonic approximation (SHA) the case where only the initial state is replaced by the corresponding factorized Born–Oppenheimer state in harmonic approximation $|\chi(\omega_0)\phi_0^{(N)}\rangle$, and all final states are retained as correlated vibronic $N - 1$ electron states. In this case, the spectral function takes the form

$$A_{\text{SA,SHA}}(\omega) = \sum_j \langle \chi_{\omega_0} \phi_0^{(N)} | \hat{c}_{\alpha}^{\dagger} \Psi_j^{(N)} \rangle \langle \Psi_j^{(N-1)} | \hat{c}_{\alpha} \chi_{\omega_0} \phi_0^{(N)} | \delta(h\omega - E_j) \rangle$$

Interestingly, because the Born–Oppenheimer ground state is by construction not an eigenstate of the full many-body
Hamiltonian, already at the level of the SHA, the expression in eq 18 has to be applied.

As further simplification, we can consider a harmonic approximation for both the involved initial and final potential energy surfaces and replace the remaining $N-1$ electron states by Born–Oppenheimer states in harmonic approximation. This leads to the double-harmonic approximation (DHA) for the spectral function

$$A_{\text{f},m,n}^{\text{SA,DHA}}(\omega) = \sum_{n,j} \left| \langle \chi_{m,n} | \chi_{\Omega,0} \rangle \right|^2 \langle \phi_{j,n}^{(N-1)} | \hat{c}_{j,m}^{\dagger} \phi_{0}^{(N)} \rangle$$

$$\times \langle \phi_{0}^{(N)} | \hat{c}_{j,m}^{\dagger} | \phi_{0}^{(N-1)} \rangle \delta(\hbar \omega - \varepsilon_j)$$

(21)

In particular, the expression for the DHA shows that the peak heights in the photoelectron spectrum are modulated by Franck–Condon factors. Although this simplifies practical computations considerably, we show in Section 4 that spurious peaks appear in the DHA of the spectral function, which are not present in the exact spectral function, eq 15.

4. RESULTS

4.1. Comparison of BO and Exact Ground-State Photoelectron Spectra. In this section, we illustrate the different theory levels that we introduced in the previous section for the calculation of vibronic photoelectron spectra. Due to the dense diagonalization that we can perform for our model system of trans-polyacetylene, we have all correlated states and all required Born–Oppenheimer states available to perform the explicit sum-over-states that arise in the definition of the different spectral functions in eqs 15, 20, and 21. In the following, we restrict ourselves only to the ground state as the initial state for the photoemission process. We term these spectra ground-state photoelectron spectra. Later, we lift this restriction to also consider pump pulses and time-evolving reference states explicitly.

In Figure 3, we illustrate spectral functions of the SSH chain for three different cases. In case (a), the spectral function has been calculated in the double-harmonic approximation using eq 21. In case (b), the spectrum shows the spectrum calculated in the single-harmonic approximation, where eq 20 has been employed, and in case (c), the spectrum shows the exact correlated ground-state spectrum computed from eq 15. In the figure, the different peaks are labeled according to their corresponding quantum numbers (quantum numbers of electronic state, optical mode, and acoustical mode are shown). In the experiment, spectra are typically plotted as functions of the positive binding energy (see for example, Figure 4 in ref 2). To connect the plots of the present work to this convention, the absolute value of x-axis has to be considered to arrive at positive values for the binding energy. Furthermore, for all spectra, a Lorentzian broadening of the form

$$f(E; E_0, \gamma) = \frac{1}{\pi} \frac{\gamma}{(E - E_0)^2 + \gamma^2}$$

(22)

with $\gamma = 0.002$ eV has been used. Note, that a conventional broadening of 0.1 eV that is employed frequently for purely electronic Green’s functions would completely wash out the vibrational sidebands. To resolve here the vibrational sidebands of the photoelectron spectrum, a much smaller broadening of 0.002 eV has to be employed. Note that this broadening is also about an order of magnitude smaller than 1/40 eV, which gives a typical energy scale for vibronic motion at room temperature. In the experiment, the vibrational sidebands are hence only clearly visible in a low temperature limit.

In comparison to the exact spectrum in case (c), we conclude from Figure 3 that DHA and SHA both accurately predict the peak positions corresponding to the optical phonon mode in the energy range of the spectrum from −10 to −5 eV, but the spectra reveal clear differences in the energy range from −5 up to 0 eV. The accurate location of the peaks is in accord with the quality of the approximate energy values shown in Table 1. On the other hand, peak heights in the DHA are not accurate. Peaks that correspond to the optical phonon mode are most dominant in the spectrum, and their broadening overlaps and even hides peaks, which correspond to mixed or acoustical phonon modes.

As additional information, we also show in Figure 3 the sum rule calculated with eq 17 for each spectrum. The DHA spectrum violates the sum rule due to the noncompleteness of the approximation, as discussed in Section 3. In all three spectra, most of the spectral amplitude is located in energy areas below −5 eV, while only less than two percent of the spectral weight is located in the energy range above −5 eV in the DHA and SHA spectra. The most prominent feature between the different spectra is that in the DHA and SHA spectra spurious peaks appear above −5 eV that are not present in the exact correlated ground-state spectrum. We discuss the origin of this artifact of the DHA and SHA in detail in the next section.

4.2. Nonadiabaticity in Ground-State Photoelectron Spectroscopy. The prominent differences in the energy range from −5 to 0 eV between the DHA spectrum and the exact correlated spectrum shown in Figure 3 have two equally important contributions. As indicated by the name double-harmonic approximation, one performs two harmonic approximations in the DHA. It turns out that both harmonic approximations contribute independently to the spurious peaks in the spectrum. We can isolate the effect of each of the two harmonic approximations by comparing to the single-harmonic approximation. Because of our definition in eq 20, where we use correlated final states in the SHA, the only remaining approximation in the SHA is the factorized and harmonic Born–Oppenheimer initial state. Comparing the sum rules for the spectral function of the DHA with the spectral function of the SHA in Figure 3, the upper part of the spectrum (−5 to 0 eV) shows that the spectral weight of the spurious peaks is reduced from 1.6% in DHA to 0.2% in SHA. The remaining spurious amplitude, and hence the differences between the SHA spectrum in Figure 3(b) and the exact spectrum in Figure 3(c), is caused by the factorized Born–Oppenheimer initial state in the SHA.

To illustrate this further, we expand the Born–Oppenheimer ground state in the complete set of correlated eigenstates of the full many-body Hamiltonian from eq 2

$$| \phi_0^{(N)} \rangle = \sum_{n=1}^{N_{\text{max}}} \langle \psi_n^{(N)} | \phi_0^{(N)} \rangle | \psi_n^{(N)} \rangle$$

$$= \sum_{n=1}^{N_{\text{max}}} a_n | \psi_n^{(N)} \rangle$$

(23)

The magnitude for the different expansion coefficients $a_n$ is shown in Figure 4(a) in logarithmic scale. As expected, the
highest overlap is found between the Born–Oppenheimer ground state and the exact correlated ground state. For the present system, this overlap is equal to 0.9986 (see Table 1) and is marked as a black dot in the graph. The following corrections are orders of magnitudes smaller. In Figure 4(a), we illustrate with different colors the overlaps $a_n$ for $n_{\text{max}} \geq 1$ with magnitude larger than $10^{-15}$. The overlaps can be grouped in different sets, which allow us to identify different potential energy surfaces in terms of Figure 2. In Figure 4(b), we show the SHA spectral function for different upper limits of summation $n_{\text{max}}$ in the expansion of the Born–Oppenheimer initial state. If only the coefficient $a_1 = 0.9986$ with the highest overlap is included, we recover the exact correlated ground-state spectrum. This is shown in the upper spectrum in Figure 4(b). Note that we are not renormalizing the state in eq 23 after truncation, so that the sum rule corresponds for $n_{\text{max}} = 1$ to $S = 4 \times a_1^2 = 3.9888$. When more and more expansion coefficients $a_n$ with $n > 1$ are included in the expansion, the artificial peaks shown in Figure 3 in the range from $-5$ to $0$ eV start to emerge. This is illustrated in the sequence of spectra in Figure 4(b). When the expansion of eq 23 is inserted in eq 20, the spurious peaks arise due to additional cross and diagonal terms in the spectral function, which involve excited correlated eigenstates. Hence, we conclude that the artificial peaks are due to the factorized nature of the Born–Oppenheimer ground state. We emphasize, that the spurious spectral weight appears both for the Born–Oppenheimer ground state in harmonic approximation as well as for the exact Born–Oppenheimer ground state without the harmonic approximation. In both cases, the expansion in eq 23 in terms of correlated vibronic eigenstates has in general more than one term ($n_{\text{max}} > 1$), and hence, additional cross and diagonal terms in the spectral function necessarily appear. As we have demonstrated in Table 1, for the present model of trans-polyacetylene, the overlap between the exact Born–Oppenheimer, harmonic Born–Oppenheimer, and exact correlated ground state is very high due to the rather harmonic nature of the Su–Schrieffer–Heeger model. Nevertheless, the spurious spectral weights already have a magnitude of about 1.6% in DHA. For any molecular system, which is less harmonic than our model, a larger contribution to the spurious spectral peaks is expected because in the expansion more terms with a larger weight of expansion coefficients $a_n$ contribute. In this sense, the present system can be regarded as best-case scenario, and in general, the spurious spectral peaks are more pronounced. However, in the limit of large nuclear masses, the Born–Oppenheimer approximation becomes more accurate. In this limit, the Born–Oppenheimer ground state of the system becomes identical to the correlated ground state, hence leading to identical spectra.

One way to correctly incorporate nonadiabatic effects could be the inclusion of nonadiabatic couplings in the Born–Huang expansion (eq 6). Other alternatives could rely on an explicitly correlated ansatz for the combined electron-nuclear wave function, as for example, in an electron–nuclear coupled cluster approach or in a multicomponent density functional theory approach for electrons and nuclei.

4.3. Time-Resolved Pump–Probe Photoelectron Spectra. So far, we have considered the ground state as the reference state for the calculation of the spectral function. In this section, we turn our attention to explicitly time-resolved vibronic photoelectron spectra. All calculations for the remaining part of the paper are done with the exact Hamiltonian and are based on the exact time-evolution of the correlated time-dependent Schrödinger equation. To illustrate pump–probe photoelectron spectra for vibronic systems, we consider two different examples: In example 1, we initially excite our system with a Franck–Condon transition, while in example 2, we explicitly include a short femtosecond laser pulse with Gaussian envelope in our real-time propagations to simulate the pump pulse. We start in the present section with example 1.

4.3.1. Time-Resolved Photoelectron Spectra with Initial Franck–Condon Excitation. In our first example, we excite the SSH chain from the Born–Oppenheimer ground state to the first excited charge neutral $N$ electron state $\Phi^{(N)}$, while the vibrational state remains in the ground state configuration $\chi_{00}$. After excitation, the initial state for the time propagation is still a factorized Born–Oppenheimer state of the form

$$|\Psi^{(N)}(t = t_0)\rangle = |\chi_{00}\Phi^{(N)}_t\rangle$$ (24)
This type of Franck–Condon transition takes here the role of the pump pulse and is illustrated in Figure 5(a) in the left panel. The initial state in eq 24 is then propagated in real-time with the full correlated Hamiltonian in the combined electronic and vibrational Fock space of the model. Because the factorized excited Franck–Condon state is not an eigenstate of the correlated many-body Hamiltonian, a wave packet propagation is launched with this initial state, which resembles predominantly the motion of a Born–Oppenheimer state in the first excited potential energy surface. We propagate from \( t_0 = 0 \) fs to a final time of \( t_f = 110 \) fs, which corresponds to about 9/4 of the oscillation period of the nuclear wavepacket in the excited state. The oscillation spread of the center of the nuclear wavepacket is indicated by a yellow background in Figure 5(a).

After a certain delay time \( \tau \), we simulate a probe pulse by recording the photoelectron spectrum in terms of the spectral function. This amounts to replacing the reference state \( |\Psi^{(N)}(\tau)\rangle \) in eq 15 with the time-evolved state \( \Psi_j^{(N)}(\tau) \) at the pump–probe delay time \( \tau \)

\[
A^\text{SA}_{\text{in},\sigma}(\omega, \tau) = \sum_j \langle \Psi_j^{(N)}(\tau) | e^{i \omega \tau} | \Psi_j^{(N-1)} \rangle \\
\times \langle \Psi_j^{(N-1)} | e^{i \sigma \omega \tau} | \Psi_j^{(N)}(\tau) \rangle \delta(\hbar \omega - E_j)
\] (25)

In Figure 5(b), the corresponding photoelectron spectra for two different delay times of \( \tau = t_1 = 0 \) fs and \( \tau = t_2 = 110 \) fs are shown in green. The spectra after different pump–probe delays show that several peaks gain spectral amplitude, which were dark in the ground-state spectrum, and conversely, other peaks lose amplitude, which were bright before. A more complete picture of the underlying wavepacket dynamics can be obtained by plotting the spectral function \( A^\text{SA}_{\text{in},\sigma}(\omega, \tau) \) as a continuous function of the delay time \( \tau \). This is shown in Figure 5(c). Here, every slice of the 2D plot at fixed \( \tau \) corresponds to one recorded spectrum. The color code indicates the intensity of the peaks, with high photoelectron amplitude in red and lower amplitude in blue. The spacing between neighboring peaks corresponds to different vibronic states in the same potential energy surface. Besides the spectral function, we also plot with a dashed line in Figure 5(c) the center of the nuclear wavepacket (first moment) as function of the delay time \( \tau \). The oscillation time \( T_0 \) is in this case \( T_0 = 48.94 \) fs. The 2D plot of the spectral function nicely illustrates that the gain and loss of the spectral amplitude as functions of the pump–probe delay time \( \tau \) is directly linked to the underlying nuclear wavepacket motion. This is similar to optical pump–probe spectroscopy, which provides a stroboscopic picture of the nuclear dynamics of the system. The notable difference here is that outgoing photoelectrons and therefore states, which have vanishing optical matrix elements with the initial state, can also be monitored.

Typically, in nontime-resolved pump–probe photoelectron experiments, time-averages of spectra are recorded. We therefore include in Figure 5(b) a time-averaged spectrum in blue that is computed according to

\[
A^\text{SA}_{\text{in},\sigma}(\omega) = \frac{1}{t_f - t_i} \int_{t_i}^{t_f} A^\text{SA}_{\text{in},\sigma}(\tau, \omega) d\tau
\] (26)

and that can be viewed as an average of the 2D contour data of Figure 5(c) along the axis of the delay time \( \tau \). The average spectrum is useful in determining in which spectral regions the emitted photoelectrons can be found over certain oscillation periods.

### 4.3.2. Time-Resolved Photoelectron Spectra with Explicit Pump Pulse

In example 2, we investigate an explicit simulation of a pump–probe experiment in real-time. Compared to the Franck–Condon excitation, which was based only on the selection of a specific excited initial state, a more appropriate description of the excitation of the system can be realized by explicitly including the pump pulse into the time propagation. For the following discussion, we therefore add a dipole-coupling term to the Hamiltonian in eq 1

\[
\text{Ham} \rightarrow \text{Ham} + \sum_n \sum_m \frac{\langle \psi_n | \vec{d}_n \cdot \vec{r}_m | \psi_m \rangle}{\epsilon_n} \frac{\partial}{\partial \tau} \langle \psi_n | \vec{d}_n \cdot \vec{r}_m | \psi_m \rangle
\]
an exponential midpoint scheme \(37\) to account for the time-use as before a Lanczos propagator, but in addition, we employ \(\mathfrak{g}\) (the pulse. The pulse starts at \(l_{\text{ff}}\) as given in eq \(9\).

The pulse couples to both, to the dipole moment of the electrons, and to the nuclear dipole. For the electric field of the pump pulse \(E(t)\), we use a Gaussian envelope with midpoint \(t_{0} = -6\) fs, maximum envelope \(E_{0} = 0.85 \text{ V/Å}\), and variance \(\sigma = 1.5\) fs. As carrier wave, we choose a sine function with frequency \(\omega_{0} = \Delta E/\hbar = 6.20 \text{ fs}^{-1}\). The frequency of the laser pulse is chosen to be resonant for a Franck–Condon like transition with \(\Delta E\) that corresponds to the example in Section 4.3.1. For the time-propagation with the time-dependent Hamiltonian \(\hat{H}_{\text{ssh}}(t)\), we use as before a Lanczos propagator, but in addition, we employ an exponential midpoint scheme \(57\) to account for the time-dependence of the Hamiltonian. For the propagation, we choose the exact correlated vibronic ground state as initial state. This state is then propagated with the fully correlated many-body Hamiltonian including the dipole coupling to the pump laser as given in eq \(9\).

In Figure 6(a), we show the amplitude of the external laser pulse. The pulse starts at \(t_{0} = -10\) fs and is switched off at \(t_{1} = 0\) fs. As the state evolves, we compute the overlaps with the Born–Oppenheimer states as function of time. In Figure 6(b) we show which states are populated during the propagation. While at the initial time, almost the full population is in the BO ground state \(|\psi_{\text{BO}}^{(N)}\rangle\), the population moves within the first 10 fs to the state \(|\psi_{\text{CO}}^{(N)}\rangle\), i.e., the state that corresponds to a Franck–Condon transition out of state \(|\psi_{\text{BO}}^{(N)}\rangle\). After the first initial population of \(|\psi_{\text{CO}}^{(N)}\rangle\), the populations indicate two competing processes. First, the pump laser pulse transfers population from \(|\psi_{\text{CO}}^{(N)}\rangle\) to \(|\psi_{\text{BO}}^{(N)}\rangle\) because this transition is resonant. Second, once population occurs in \(|\psi_{\text{CO}}^{(N)}\rangle\), this population induces a wavepacket motion on the first excited potential energy surface, as shown in Section 4.3.1. Therefore, the system never reaches a situation where only two states are present in the system, \(|\psi_{\text{CO}}^{(N)}\rangle\) and \(|\psi_{\text{BO}}^{(N)}\rangle\), as it would be in a Franck–Condon picture of the excitation process. After the end of the pump pulse at \(t = 0\) fs, the projection of the time-evolving state on the correlated vibronic ground state is constant in time, and the ground state maintains a population of about 50%. In contrast, the projection of the correlated time-evolving state on the BO ground state exhibits small oscillations, which is shown in the inset of Figure 6(b) and which arise due to the small deviations between the BO ground state and the exact correlated ground state. As in the example before, we record photoelectron spectra as functions of pump and probe delay. The photoelectron spectrum at time \(t = t_{0} = -10\) fs is given in red in Figure 6(c). This spectrum is identical to the spectrum also shown in Figure 3(c). The spectra after the pulse has been switched off are shown in green and correspond to delay times of \(t = 0\) fs and \(t = 110\) fs. As before, we also show the time-averaged spectrum in blue. In contrast to the Franck–Condon excitation in example 1, the photoelectron spectra in this case show a pronounced large peak at about \(-7.5\) eV. This peak arises due to the remaining population of the Born–Oppenheimer ground state \(|\psi_{\text{BO}}^{(N)}\rangle\). In Figure 6(d), we show similar to Figure 5(c) the spectral function \(A_{\text{SA}}^{\text{m}}(\tau, \omega)\) as a function of the pump–probe delay \(\tau\). As before in the Franck–Condon case, also with an explicit pump pulse, we can trace the nuclear wavepacket dynamics in the photoelectron spectrum. Overall, the time-evolution of the Franck–Condon excitation captures large parts of the exact vibronic spectrum of example 2. The notable differences in the explicitly time-dependent and fully correlated vibronic case in Figure 5(c) are the additional population of the Born–Oppenheimer ground state that maintains a large spectral weight at about \(-7.5\) eV also for...
different delay times and some small nonadiabatic contributions to the spectrum in the energy range from $-4$ to $-2$ eV.

5. CONCLUSION

In summary, we have analyzed and quantified nonadiabatic contributions to the equilibrium and nonequilibrium photoelectron spectra in a model system for trans-polycyctene. We find that for low-lying states the harmonic Born–Oppenheimer photoelectron spectrum acquires, in comparison to the exact photoelectron spectrum, spurious spectral weight, which also persists when either the initial state of the photoemission process or the final state is replaced by correlated vibronic states. The origin of this behavior can be traced back to the factorized nature of the involved initial or final Born–Oppenheimer states. Only when both initial and final photoemission states are taken as correlated vibronic states are the spurious spectral peaks suppressed. We analyze this in detail by expanding the Born–Oppenheimer ground state in the complete set of correlated vibronic eigenstates of the full Hamiltonian. Inserting this expansion into the equilibrium form of the spectral function shows that additional cross and diagonal terms, which involve excited correlated eigenstates, are responsible for the spurious spectral weight.

For the example of an initial Franck–Condon transition and for an explicit pump–pulse excitation, we have demonstrated with explicit real-time propagations of the coupled polyacetylene chain how the vibronic wavepacket evolution can be traced in the photoelectron spectrum as function of pump–probe delay.

Prospects for future work include the study of temperature and pressure dependence of the photoelectron spectra as well as an extension of the present femtosecond laser excitation to ultrafast photoelectron spectroscopy with attosecond laser pulses in real nanostructured and extended systems. Another line of research is linked to the development of xc functionals for TDDFT capturing the effects discussed in this work, for example, based on electron–nuclear multicomponent density functional theory.36

# APPENDIX

A.1. Spectral Function

The one-body spectral function is defined as follows33

$$A_0(t,t') = \langle \Psi_0 | \hat{c}_i^\dagger(t) \hat{c}_j(t') | \Psi_0 \rangle$$

$$A_0(t,t') = \langle \Psi_0 | \hat{c}_i^\dagger(t') \hat{c}_j(t) + \hat{c}_j^\dagger(t') \hat{c}_i(t) | \Psi_0 \rangle$$

$$A_0(t,t') = A_0^\dagger(t,t') + A_0(t',t)$$  \hspace{1cm} (28)

with $t' > t$. The operators $\hat{c}$ and $\hat{c}^\dagger$ are here written in the Heisenberg picture. The index refers to a combined index $i = (n, \sigma)$ and $j = (m, \sigma')$, where $n$ and $m$ refer to the site number, and $\sigma$ and $\sigma'$ refer to spin up or spin down.

In this work, we only consider the first part of the commutator $A_0^\dagger(t,t')$ because we are only interested in photoemission spectra. The second term $A_0(t,t')$ leads to inverse photoemission spectra.34 In the following discussion, we distinguish two cases: (1) If $|\Psi_0\rangle$ is an eigenstate of the system Hamiltonian, we work in an equilibrium framework. (2) If $|\Psi_0\rangle$ is not an eigenstate of the system Hamiltonian, we have to work in a nonequilibrium framework.

A.2. Equilibrium Spectral Function

The equilibrium spectral function applies for situations where $|\Psi_0\rangle$ is an eigenstate of the corresponding many-body Hamiltonian $H$ of the system. Hence, we can write eq 28 in terms of a time-correlation function

$$A_0^\dagger(t,t') = \langle \Psi_0 | \hat{c}_i^\dagger(t') \hat{c}_j(t) | \Psi_0 \rangle$$

$$= \langle \Psi_0 | e^{iHt'/\hbar} \hat{c}_j^\dagger e^{-iHt/\hbar} e^{iHt'/\hbar} e^{-iHt/\hbar} | \Psi_0 \rangle$$

$$= \langle \Psi_0 | e^{-i(\Omega(t)-\Omega(t'))/\hbar} c_j^\dagger c_i e^{-i(\Omega(t)-\Omega(t'))/\hbar} | \Psi_0 \rangle$$

$$= \langle \Psi_0^\dagger(t') \hat{c}_i^\dagger | \bar{\Psi}(\tau) \rangle$$  \hspace{1cm} (29)

with $\tau = t' - t$ and the initial condition $\bar{\Psi}(\tau = 0) = \hat{c}|\Psi_0\rangle$.

Equation 28 can be reformulated to get a sum-over-states expression. This is accomplished by the insertion of an complete set of states $\sum_{m} |\Psi_m\rangle \langle \Psi_m | = 1$ and a Fourier transform with respect to the time difference $\tau = t' - t$

$$A_0^\dagger(t,t') = \langle \Psi_0^\dagger(t') \hat{c}_i^\dagger | \bar{\Psi}(\tau) \rangle$$

$$= \langle \Psi_0^\dagger e^{iHt'/\hbar} \hat{c}_j^\dagger e^{-iHt/\hbar} e^{iHt'/\hbar} e^{-iHt/\hbar} | \Psi_0 \rangle$$

$$= \sum_m \langle \Psi_0^\dagger \hat{c}_i^\dagger | \Psi_m \rangle \langle \Psi_m | \hat{c}_j | \Psi_0 \rangle e^{i(E_m - E_0)(t' - t)/\hbar}$$

$$A_0^\dagger(\omega) = \int_{-\infty}^{\omega} \sum_m \langle \Psi_0^\dagger \hat{c}_i^\dagger | \Psi_m \rangle \langle \Psi_m | \hat{c}_j | \Psi_0 \rangle e^{i(E_m - E_0 - \hbar\omega)}$$

$$A_0^\dagger(\omega) = \sum_m \langle \Psi_0^\dagger \hat{c}_i^\dagger | \Psi_m \rangle^2 \delta(E_0 - E_m - \hbar\omega)$$  \hspace{1cm} (30)

A.3. Nonequilibrium Spectral Function

In nonequilibrium situations, $|\Psi_0\rangle$ is not an eigenstate of the many-body Hamiltonian $H$. Nevertheless, it is also possible to formulate the spectral function in eq 28 as time-correlation function involving propagated states

$$A_0^\dagger(t,t') = \langle \Psi_0^\dagger(t') \hat{c}_i^\dagger(t) | \Psi_0 \rangle$$

$$= \langle \Psi_0^\dagger e^{iHt'/\hbar} \hat{c}_j^\dagger e^{-iHt/\hbar} e^{iHt'/\hbar} e^{-iHt/\hbar} | \Psi_0 \rangle$$

$$= \langle \Psi_0^\dagger(t') | e^{-i(\Omega(t)-\Omega(t'))/\hbar} c_j^\dagger c_i e^{-i(\Omega(t)-\Omega(t'))/\hbar} | \Psi_0 \rangle$$

$$A_0^\dagger(t,\tau) = \langle \Psi_0^\dagger(t + \tau) \hat{c}_i^\dagger | \bar{\Psi}(\tau + t) \rangle$$  \hspace{1cm} (31)

We introduce the relative time $\tau = t' - t$, as in Appendix A.2, while $t$ keeps its initial denotation. The state $|\bar{\Psi}(t + \tau)\rangle$ is defined as $|\bar{\Psi}(t + \tau)\rangle = e^{-i\Omega(t'/\hbar)}|\bar{\Psi}(0)\rangle$, meaning the kick $\hat{c}$ on the wavefunction acts at time $t$ during the time propagation. A Fourier transform with respect to the relative time $\tau$ yields the general expression for the sum-over-states expression.
\[ A^{\psi}_{ij}(t,t') = \langle \Psi_0 | \psi_j(t') | \psi_i(t) \rangle |\psi_i(t)\rangle |\psi_j(t')\rangle \]
\[ A^{\psi}_{ii}(t,t') = \langle \psi_j(t') | \psi_i(t) \rangle |\psi_i(t)\rangle |\psi_j(t')\rangle \]
\[ A^{\psi}_{ii}(t,t') = \sum_{n,n',m} e^{i t/\hbar (E_{n'} - E_n)} \times \langle \psi_j | \psi_i | \psi_n | \psi_m \rangle \langle \psi_n | \psi_m | \psi_{n'} | \psi_i \rangle \]
\[ A^{\psi}_{ii}(t,t') = \int_{-\infty}^{\infty} \frac{d\tau}{2\pi} \sum_{n,n',m} e^{i \tau / \hbar (E_{n'} - E_n - i\omega) + i t / \hbar (E_{n'} - E_n)} \times \langle \psi_j | \psi_i | \psi_n | \psi_m \rangle \langle \psi_n | \psi_m | \psi_{n'} | \psi_i \rangle \]
\[ A^{\psi}_{ii}(t,t') = \sum_{n,n',m} e^{i t/\hbar (E_{n'} - E_n - i\omega)} \times \langle \psi_i | \psi_{n'} | \psi_m | \psi_j \rangle |\psi_i\rangle |\psi_{n'}\rangle |\psi_m\rangle |\psi_j\rangle \]
\[ A^{\psi}_{ii}(t,t') = \sum_{n,m} \delta(E_n - E_m - i\omega) \]
\[ A^{\psi}_{ii}(t,t') = \sum_{n,m} \delta(E_n - E_m - i\omega) \]

\[ (32) \]

In our simulations, we neglect the energy dependence of the delta function in the last equation. Hence, we replace the term \( E_n \) by the energy \( E_0 \) of the state \( \Psi_0 \). This leads to

\[ A^{\psi}_{ii}(t,t') = \sum_{n,n',m} e^{i t/\hbar (E_{n'} - E_n)} \times \langle \psi_j | \psi_i | \psi_n | \psi_m \rangle \langle \psi_n | \psi_m | \psi_{n'} | \psi_i \rangle \]
\[ A^{\psi}_{ii}(t,t') = \sum_{n,m} \delta(E_n - E_m - i\omega) \]

\[ \text{REFERENCES} \]