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Quantum Dynamics of the Hubbard-Holstein Model in Equilibrium and Non-Equilibrium: Application to Pump-Probe Phenomena

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The spectral response and physical features of the 2D Hubbard-Holstein model are calculated both in equilibrium at zero and low chemical dopings, and after an ultra short powerful light pulse, in undoped systems. At equilibrium and at strong charge-lattice couplings, the optical conductivity reveals a 3-peak structure in agreement with experimental observations. After an ultra short pulse and at nonzero electron-phonon interaction, phonon and spin subsystems oscillate with the phonon period $T_{ph} \approx 80$ fs. The decay time of the phonon oscillations is about 150 – 200 fs, similar to the relaxation time of the charge system. We propose a criterion for observing these oscillations in high $T_c$ compounds: the time span of the pump light pulse $\tau_{pump}$ has to be shorter than the phonon oscillation period $T_{ph}$.

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It is accepted that the physics of the high critical temperature (high $T_c$) cuprate superconductors is controlled by the interplay of different kinds of elementary excitations. The Mott-Hubbard nature of these systems was quickly recognized, and highlighted the relevance of the electron-phonon interaction (EEI) [1]. Angle resolved photoemission spectroscopy (ARPES) [2] and optical conductivity (OC) [3] measurements revealed the importance of the electron-phonon interaction [4]. In spite of the general consensus that both EEI and EPI manifest themselves in the physical features of these compounds, it is still unclear which of these two interactions is responsible for particular properties. This difficulty arises because the EEI and EPI have similar spectroscopic energy scales. Ultra-fast pump-probe experiment (PPE), which probes the time dependence of the properties of a system after it was displaced from equilibrium by a laser pump pulse, is a promising method for disentangling charge, magnetic, and lattice degrees of freedom. Indeed, since laser pulses couple to charge excitations, the characteristic response times of the different subsystems are not determined solely by their own characteristic energies but are strongly influenced by the strength of their entanglement with the perturbed charge density.

Although pumping in PPEs on high $T_c$ compounds [5–11] or other materials (e.g., organic charge-transfer crystals [12–14]) is always done by an ultra-short pulse of light, there is a significant variation in the experimental setup. Conditions that can vary include fluence, photon density (total number of photons per site absorbed during the pulse), frequency of the pump photons $\omega_{pump}$, pulse duration $\tau_{pump}$, time resolution, and, of particular importance, the quantity probed after the time delay. Examples of such quantities are lattice structure and dynamics measured by ultra-fast electron diffraction [5, 7] and electron temperature derived from ARPES [6]. However, one of the most informative PPE techniques is associated with energy- and time-resolved analysis of the electrodynamic response, such as reflectivity [9, 10, 12–14], transmission [11], or full OC [8]. In particular, PPEs on high $T_c$ compounds [5–11] revealed three basic timescales. The charge immediately reacts at times < 50 fs. The phonons which are directly coupled to charges are involved in time scale of 100-200 fs, and the energy is gradually dispensed to the whole phonon subsystem (heat reservoirs) by anharmonic interactions during further 500-1000 fs.

In this letter we study the OC of the two dimensional (2D) Hubbard-Holstein model (HHM) both in equilibrium and after an ultra-short powerful light pulse. We choose this model because it reproduces the equilibrium OC of the doped compounds, where experimental studies resolve three peaks [15]: two in the infrared region around 2000 cm$^{-1}$ (250 meV) and 5000 cm$^{-1}$ (0.6 eV), and one in the visible range around 1.6 x 10$^4$ cm$^{-1}$ (2 eV). Calculations in the framework of the t-J-Holstein model reproduce the middle peak at 0.6 eV [16–21] and, in more sophisticated approximations, also can distinguish the low energy peak at 2000 cm$^{-1}$ [19–21]. However, since this model projects out charge fluctuations, the highest peak at 2 eV is simply absent. Therefore, although the t-J-Holstein model is well suited to describe quasistationary states in the presence of a constant electric field that is not too large [22], since pumping by high-energy pulses is associated with transitions into the upper Mott-Hubbard (or charge transfer) band, the HHM is the only model able to describe the time evolution of charge, magnetic,
and lattice degrees of freedom in PPEs. We are aware of theoretical studies of pump-probe physics in models similar to the HHM in one dimension (1D) with classical [23] and quantum [24] treatment of phonons, and in 2D with a classical treatment of phonons [14, 25]. Here we present the first full quantum treatment of thermodynamic equilibrium and time-resolved optical spectroscopy of an electron-lattice system described by the HHM in 2D. Such calculations became feasible due to the recently developed double phonon cloud method [26] circumventing the problem of dramatic increase of the full phonon basis with the size of the system. It has been shown [26] that such a method, by using an optimized truncated phonon basis containing phonon clouds typical of weak- and strong-coupling limits, is able to describe correctly the polaronic response in any coupling regime. The HHM Hamiltonian coupled to an external classical vector potential is \( H = H_t + H_U + H_{EPI} \), where

\[
H_t = -t \sum_{i,\mu,\sigma} (\sigma_i A(t)c_{i+\mu,\sigma}^\dagger c_{i,\sigma} + H.c.),
\]

\[
H_U = U \sum_i (n_i,\uparrow - \frac{1}{2})(n_i,\downarrow - \frac{1}{2}),
\]

\[
H_{EPI} = \omega_0 \sum_i a_i^\dagger a_i + g\omega_0 \sum_i (a_i^\dagger + a_i)(1 - n_i).
\]

Here \( t \) is the hopping amplitude, \( c_{i,\sigma}^\dagger \) is the fermionic creation operator, \( \mu \) is a unit vector along the axes of the lattice, \( a_i^\dagger \) creates a phonon at site \( i \) with frequency \( \omega_0 \), and \( n_i \) is the electron number operator. The EPI strength is defined by the dimensionless coupling constant \( \lambda = g^2\omega_0/4t \). We choose model parameters typical for high \( T_c \) materials: \( t = 0.25 \text{ eV} \approx 2000 \text{ cm}^{-1} \) and \( \omega_0 = 0.2t \approx 400 \text{ cm}^{-1} \) [27]. The value of the Hubbard repulsion \( U = 10t \) = 2.5 eV yields low-energy physics very similar to that of the t-J model with \( J = 4t^2/U = 800 \text{ cm}^{-1} \). The time-dependent potential vector is [24]:

\[
A = A_0 e^{-i\omega_0\tau^2/\tau_{\text{pump}}} \cos(\omega_{\text{pump}}(\tau - \tau_0)),
\]

and the OC, after the pulse, is given by [28]:

\[
\sigma(\omega, \tau) = \frac{1}{M\omega^2} \int_0^\infty i e^{i(\omega + i\eta)\tau} \langle j(t), j(0) \rangle |\tau\rangle dt,
\]

where \( |\tau\rangle = T e^{-i\int_0^\infty H(\tau_1)dt_1} |\tau = 0\rangle \), \( T \) is the time ordering operator, \( j(t) \) is the current operator in the Heisenberg representation along one of the lattice direction axes, \( \eta \) is a broadening factor taking into account additional dissipative processes, and \( M \) is the number of lattice sites. The \( |\tau\rangle \) state is obtained through the Lanczos time propagation method [29]. In the following we use periodic boundary conditions on 2D lattices [30] with \( M = 10 \) for equilibrium and \( M = 8 \) for pump-probe calculations. The charge and spin degrees of freedom are evaluated exactly, whereas the quantum phonons are treated with the double phonon cloud method [26].

First we study the properties of the HHM in equilibrium. The kinetic energy \( K \) and the average number of phonons \( N \) versus \( \lambda \) at HF and in one-hole subspace (one extra hole compared to HF in a system with \( M = 10 \) lattice sites). The strength of the EPI has a negligible effect at HF due to the uniform charge distribution, but in the doped system, the doped holes become lattice polarons at \( \lambda_c \approx 0.4 \) [31].

We now turn to the OC of the HHM in equilibrium. Figure 2(a) compares the OC at zero (\( \lambda = 0 \)) and strong (\( \lambda = 0.575 \)) EPI in the half-filled system; Fig. 2(b) presents the OC at zero and weak (\( \lambda = 0.1 \)) EPI in one-hole subspace. The OC for strong EPI at half filling and at low hole concentration are compared in Fig. 2(c). The OC of the HHM at half filling displays, in agreement with experiment [15], a gap with the peak of upper Hubbard band at \( \omega \approx 8t \approx 1600 \text{ cm}^{-1} \). In contrast to the t-J-Holstein model, where this upper band is absent, the HHM can describe experiments using high-energy pump photons, \( \omega_{\text{pump}} > 8t \). Similar to the ground state properties shown in Fig. 1, the OC at HF weakly depends on \( \lambda \). The high-energy edge of the gap in OC is gradually filled by phonon-dressed excitations when \( \lambda \) increases.

On the contrary, the OC of the weakly doped system depends strongly on \( \lambda \) as shown in Fig. 2(b). Upon chemical doping the gap is filled through a transfer of spectral weight towards lower energies. The most prominent feature of the OC of the weakly doped system without EPI is a low energy peak at \( \omega \approx 2J = 1600 \text{ cm}^{-1} \) (solid green line in Fig. 2(b)) which is well known from the pure t-J model. However, this narrow peak cannot explain the broad mid-infrared feature in the underdoped high \( T_c \) compounds, because its energy is too low, the peak width is too narrow, and, for most model parameters, its doping dependence is opposite that detected in experiments (see [19, 32, 33]). Besides, the OC without EPI does not display the 2-peak structure of the experimental mid-infrared response [19]. However, even a very small EPI

![FIG. 1: (Color online) (a) Kinetic energy K and (b) average number of phonons N versus \( \lambda \) at HF and in one-hole subspace.](image-url)
(λ = 0.1) induces a low-energy peak at ω ≥ ω₀ just above the phonon frequency (dashed black line in Fig. 2(b)). At the same time, the intensity of the 2J peak is renormalized. Finally, the OC at larger values of EPI (λ = 0.575) agrees well with the experimental positions of the OC peaks at energies 1500-2000 cm⁻¹ and 5000-6000 cm⁻¹ (red solid curve in Fig. 2(c)) observed in strongly underdoped high T_c compounds. The HHM model, like the t-J-Holstein model [19, 34], explains the doping dependence of the spectra in the mid-infrared region. In addition, the Holstein model [19, 34], explains the doping dependence of the chosen hopping solving the time dependent Schroedinger equation. For distinguishing between these two scenarios. Ref. [33] (see also [35]). Time-dependent PPEs can distinguish between these two scenarios.

We now turn to the study of pump-probe dynamics, by solving the time dependent Schroedinger equation. For the chosen hopping t, the real time unit is ħ/t = 2.62 fs. We fix ħωpump = 8.5t, above the peak of the charge transfer gap, and A₀ = 0.2, corresponding to the absorption of about 1/2 photon during the pulse with τpump = 2.62 fs and τ₀ = 5.24 fs. We study the dynamics in the first 250 fs because our model is missing a heat reservoir [36] where energy is dispersed in experiments during 500-1000 fs.

The time evolution of the kinetic energy K, the average value of Hubbard energy E_U = ⟨H_U⟩, the local spin-spin correlation S = ∑ᵢ,j S_i · S_j , the average number of phonons N, and the average value of the contribution describing the charge-lattice coupling E_I = ⟨τ|gω₀ ∑ᵢ (aᵢ† + aᵢ)(1 − nᵢ)|τ⟩ are shown in Fig. 3 for three different values of EPI at half filling. Quantities associated with charge and spins (K, E_U, and S) display quick changes during the first 10-20 fs (Fig. 3a, b, and c) followed by slower dynamics depending on the strength of the EPI. On the other hand, quantities associated with lattice degrees of freedom (N and E_I) show dynamics on a time scale associated with the period of phonon oscillations T_{ph} ≈ 80 fs (Fig. 3d). Furthermore, after about two phonon oscillations, both at λ = 0.1 and λ = 0.5 a relaxed state is reached within a time around 150 - 200 fs. It is interesting to note that the spin correlation function S (Fig. 3c) at λ ≠ 0 also displays oscillations at timescales similar to the phonon period T_{ph} ≈ 80 fs, highlighting the strong coupling of spin and lattice degrees of freedom. Notably, after the laser pulse, slower charge dynamics are observed only in the presence of a significant EPI (see inset Fig. 3b): at λ = 0, E_U displays oscillations[37] around an essentially constant value, whereas at λ = 0.5, a relaxation is ob-

![FIG. 2: (Color online) OC at equilibrium (in units of σ₀ = e²/ħa) for different EPI values, at HF and in one-hole subspace.](image)

![FIG. 3: (Color online) Time evolution of: (a) Kinetic energy K; (b) and inset in (b): U interaction contribution E_U = ⟨H_U⟩; (c) and inset in (c): local spin-spin correlation; (d): average number of phonons N; inset in (d): EPI energy at λ = 0 (dotted green line), λ = 0.1 (solid black line), and λ = 0.5 (dashed red line).](image)
served. The relevance of the EPI is also indicated by strong $\lambda$ dependence of $K$ after the pumping: the kinetic energy gain, that is $\lambda$ independent in the ground state, quickly reduces by increasing $\lambda$.[37].

The oscillation behavior with characteristic phonon frequencies predicted here, has not been observed in experiment [5–7, 9–11]. To our mind, the difference lies in the pump-width to phonon period ratio $\tau_{\text{pump}}/T_{\text{ph}}$ which is small $\tau_{\text{pump}}/T_{\text{ph}} \ll 1$ in our calculation and large $\tau_{\text{pump}}/T_{\text{ph}} > 1$ in experiments [5–7, 9–11]. This interpretation is supported by numerous PPEs in charge-transfer organic compounds [12–14], where the relation $\tau_{\text{pump}}/T_{\text{ph}} \ll 1$ holds, and the oscillations of physical quantities with phonon period are observed. Moreover, experiments on high $T_c$ materials with record ultra-short $\tau_{\text{pump}} = 12$ fs pulse [8] do observe oscillations with the frequencies of bond-bending and apical oxygen phonons. In addition, the wavelet transformation of time dynamics of the OC shows that the energy transfer into the lattice system is associated with a time around $\tau_tr \approx 150$ fs, which is similar to that of phonon oscillations [8]. Hence, interplay of similar periods $\tau_{tr} \approx T_{\text{ph}}$ may result in beat behavior of lattice properties as observed in our calculations at $\lambda = 0.5$ (Fig. 3d). We conclude that in the most of PPEs on high $T_c$ materials [5–7, 9–11] the phonon oscillations are self-averaged because of long pump duration whereas the shortening of the pump-width is crucial to make the pump-probe phenomena similar to those observed in charge transfer organic compounds [12–14].

Fig. 4 shows the OC after the pulse at HF with (a) zero, (b) weak, and (c) strong EPI. The gap present in the undoped system is filled after the pulse, but in a way that is strongly dependent on the EPI strength. The non-equilibrium OC without EPI shows almost time-independent Drude behavior (Fig. 4a). On the other hand, for weak EPI ($\lambda = 0.1$) as shown in Fig. 4b, the OC oscillates and contains two main peaks at the same energies as observed in the weakly doped system at weak EPI (Fig. 2b). These peaks correspond to $2J$ and $\omega_0$, which are related to charge scatterings with magnons and phonons, respectively. These two features, associated with spin and lattice degrees of freedom, become visible at very short times after the pump pulse. This result agrees with results obtained within the Holstein model [40]. Finally, the non-equilibrium OC at $\lambda = 0.5$ has no features in common with the OC at equilibrium (compare Fig. 4c and Fig. 2c): there is no dominant peak at $5000$ cm$^{-1}$ or at $2000$ cm$^{-1}$. Instead, a low energy peak with phonon sidebands is observed indicating that the lattice polaron is not yet formed. Indeed polaron formation at intermediate EPI requires long times[40], due to the high potential barrier between free and self-trapped states.

At zero (weak) EPI, a feature is clearly (tentatively) revealed in the non-equilibrium OC at $\omega \sim 3J$, due to the transition into the bimagnon state. This excitation has been observed by Perkins et al. [38] in undoped copper oxides and, theoretically, can be seen in the equilibrium OC only in the presence of EPI [39] (see inset in Fig. 4a for equilibrium OC at different EPIs). However, out of equilibrium, due to lower symmetry, such a transition is allowed even at $\lambda = 0$.

In conclusion, we studied spectral properties of under-doped high $T_c$ materials (2D Hubbard-Holstein model with quantum phonons) in equilibrium and during the first 250 fs following excitation by a laser pulse. In particular, we found phonon-governed oscillations that can be observed in pump-probe experiments provided that the pulse time $\tau_{\text{pump}}$ is shorter or at least comparable with the typical phonon times $T_{\text{ph}} = 2\pi/\omega_0$. We showed that the analysis of OC in and out of equilibrium in the mid-infrared range can reveal the EPI strength in cuprates.

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[27] The integration over the phonon degrees of freedom (J.E. Hirsch and E. Fradkin, Phys. Rev. B 27, 4302 (1983)) provides an attractive interaction between electrons that is nonlocal in time. Only in the extreme antiadiabatic regime the interaction becomes instantaneous leading to an effective Hubbard model with a reduced $U$ value. In the adiabatic regime, investigated here, the retardation effects due to the lattice displacement propagation play a crucial role.
[34] A. S. Mishchenko et al., EPL 95, 57007 (2011).
[37] In addition to physically meaningful time dynamics finite size oscillations are observed corresponding to electron motion from one edge of finite system to another with the time scale $(h/t)/(8\sqrt{2}\approx 10 fs$ [24].