Photoinduced In-Gap States in the One-Dimensional Extended Hubbard Model

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We investigate time evolution of optical conductivity in the half-filled one-dimensional extended Hubbard model driven by a transient laser pulse, by using the time-dependent Lanczos method. Photoinduced in-gap excitations exhibit qualitatively different structure in the spin-density wave (SDW) in comparison to the charge-density wave (CDW) phase. In the SDW, the origin of a low-energy in-gap state is attributed to even-odd parity of photoexcited states, while in the CDW an in-gap state is due to confined photo-generated carriers. The signature of the in-gap states can be identified as a characteristic oscillation in the time evolution of physical quantities.

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Time-resolved spectroscopy can provide new insights into the dynamical properties of condensed matters. Among many time-resolved spectroscopies, pump-probe optical measurements have been performed extensively to investigate strongly correlated electron systems. Pumping pulse applied to one-dimensional (1D) Mott insulators with spin-density-wave (SDW) correlation creates not only photoinduced metallic state [1, 2] but also a photoinduced charge-density-wave (CDW) state [3]. In two dimensions, the photoexcitation of underdoped cuprates induces a signature of enhanced superconductivity [4, 5]. Even in a conventional superconductor, an observation of amplitude mode [6] is under intensive investigation within the field of nonequilibrium and nonlinear phenomena induced by transient electric field.

The time evolution of optical conductivity in the ultrafast process has theoretically been examined [7–11], since this quantity contains information on photoinduced charge dynamics and initial relaxation through internal degrees of freedom of a given closed system. In Mott insulators, the photoexcitation is expected to induce new excitation channel inside the Mott gap. The nature of those states has not been fully understood from theoretical viewpoint even for 1D Mott insulators where, due to spin charge separation, the spin degrees of freedom affect the charge dynamics much less than in the 2D Mott insulator [12, 13].

In this Letter we address the physical properties of photoinduced states, their positions within the Mott gap, and their influence on the time evolution of the nonequilibrium optical properties. For this purpose, we calculate the time-dependent optical conductivity of a 1D extended Hubbard model driven by a transient laser pulse. In this model, one can change the ground state from SDW phase to CDW phase by changing interaction parameters and tune the laser pulse so as to enhance charge correlation even within the SDW phase [13]. We find that photoinduced in-gap excitations appear in a different manner, depending on the nature of the initial ground state, i.e., SDW or CDW state. In the SDW, a dominant in-gap state emerges at low-energy region reflecting the energy difference between even-odd parity of photoexcited states. In the CDW, in-gap states appear in the middle of the Mott gap and their origin can be attributed to photo-generated carriers. We also find that the signature of the in-gap states emerges in the time evolution of physical quantities such as kinetic energy and peak intensity of the in-gap states as a characteristic oscillation.

We investigate the extended one-dimensional (1D) Hubbard model at half filling,

\[ H = -\frac{\hbar^2}{2} \sum_{i,\sigma} \left( c_{i+1,\sigma}^\dagger c_{i,\sigma} + H.c. \right) + U \sum_i \left( n_{i,\uparrow} - \frac{1}{2} \right) \times \left( n_{i,\downarrow} - \frac{1}{2} \right) + V \sum_i (n_i - 1)(n_{i+1} - 1), \]

where \( c_{i,\sigma}^\dagger c_{i,\sigma} \) is the creation (annihilation) operator of an electron with spin \( \sigma \) at site \( i \), \( n_i = n_{i,\uparrow} + n_{i,\downarrow} \), \( t_h \) is the hopping constant, \( U \) and \( V \) are on-site and nearest-neighbor repulsions, respectively.

The application of a spatially uniform and time-dependent electric field expressed by a vector potential \( A(t) \) is incorporated into the hopping terms via the Peierls substitution:

\[ c_{i,\sigma}^\dagger c_{i+1,\sigma} + H.c. \rightarrow e^{iA(t)} c_{i,\sigma}^\dagger c_{i+1,\sigma} + H.c., \]

leading to \( H \rightarrow H(t) \). We assume that \( A(t) \), representing the pumping pulse, is expressed in terms of the temporal gauge

\[ A_{\text{pump}}(t) = A_0 e^{-(t-t_0)^2/2\Delta^2} \cos [\omega_0 (t - t_0)], \]

where \( \omega_0 \) is the frequency of the laser pulse.
where the temporal distribution of $A_{\text{pump}}(t)$ is Gaussian centered around $t_0$, while the width is controlled by the parameter $\sigma$. $\omega_0$ is the central frequency. The average incoming number of photons per lattice site during the pump is estimated to be $\propto A_{\text{pump}}^2 \omega_0 t_d$.

In our calculation, the system's temporal evolution is evaluated by the time-dependent Lanczos method, where, within the reach of the exact diagonalization, the time-dependent wave function $\psi(t)$ can be obtained exactly. Based on the information of $\psi(t)$, we calculate the time-dependent current density $j(t) = \langle \psi(t) | \hat{j} | \psi(t) \rangle$, where $\hat{j}$ is the current operator.

Motivated by the experimental pump-probe setup, we use the following strategy to obtain the time-dependent optical conductivity after pumping. For a given pump, we trace the temporal evolution of the system twice: first, without the probe and the second time with the applied probe. The current induced by the probing pulse $j_{\text{pr}}(t)$ is obtained from the current difference between these two consecutive simulations. The subsequent Fourier transformations on $j_{\text{pr}}(t)$ and $A_{\text{p}}(t)$ produce the time-dependent optical conductivity $\sigma(\omega, \Delta \omega)$ as

$$\sigma(\omega, \Delta \omega) \approx j_{\text{pr}}(z)/izA_{\text{pr}}(z),$$

where $\Delta \omega$ is the time difference between the probing pulse and the pumping pulse and $z = \omega + i\delta$ with a small positive number $\delta$. Introduction of $\delta$ is necessary for obtaining spectral weight that is due to the Drude peak in the limit of $\omega \to 0$. We have checked that $\sigma(\omega, \Delta \omega)$ gives a good agreement with that obtained by a nonequilibrium linear-response theory as long as the strength and the width of the probing pulse are small enough. We note that the result is also insensitive to the details of the probing pulse, regardless of the shape of the vector potential and the central frequency.

In our numerical calculations, we consider a half-filled periodic lattice with $L$ sites. We set $t_h = 1$ as a unit of energy and $h = 1$. Throughout this work, the value of $U$ is chosen to be $U = 10$, which is in the strong coupling regime. The central frequency of the pumping pulse $\omega_p$ is always set to match the absorption-peak energy of a given system. The small positive number $\delta$ for the calculation of $\sigma(\omega, \Delta \omega)$ is taken to be $\delta = 1/L$.

Let us start with the case where the initial ground state is the SDW state. Figure 1(a) shows the real part of the time-dependent optical conductivity, $\text{Re}(\sigma(\omega, \Delta \omega))$, for a $L = 14$ ring with $V = 3$. Before pumping, there is a single absorption peak at $\omega = 6.1$, which is due to the exciton. With pumping, the exciton absorption peak loses its weight, while a low-energy peak at $\omega = 0.2$ emerges in the Mott gap. The low-energy peak persists throughout the time evolution. Figure 1(b) shows $A_0$ dependence at $\Delta \omega = 6$ for $V = 4.5$. As expected, with increasing $A_0$, the exciton peak at $\omega = 4$ loses its weight continuously and the low-energy in-gap peak increases.

We now discuss the origin of the low-energy peak seen in Fig. 1(a) and 1(b). Since the peak appears only after the excitation with the pump pulse, its origin should be related to photoexcited states. The photoexcitation excites carriers into the state that gives rise to the absorption peak in the equilibrium, which is an optically allowed odd-parity state. The probe pulse couples in part to the photoinduced carriers in the odd-parity state, resulting in an excitation from the optically allowed state to an optically forbidden even-parity state. In Mott insulators, it is known that the optically forbidden excitonic state is located slightly above the optically allowed excitonic state. Therefore, nonequilibrium optical conductivity should detect such an excitation. This is the origin of the low-energy peak. Its energy thus corresponds to the energy separation between the optically allowed state (exciton peak) and the forbidden state. In order to confirm this, we perform the same calculation on a smaller system size. The optically forbidden state is known to be lower in energy than the allowed one if the small system has periodic boundary conditions. A $L = 10$ lattice with $V = 4.5$ is such a case. If the optically forbidden state is lower in energy than optically allowed one, a negative-energy excitation appears after pumping, leading to negative weight at positive-energy side since $\omega \sigma(\omega)$ is odd in $\omega$. This actually occurs in the $L = 10$ lattice as shown in Figure 1(d), where $\text{Re}(\sigma(\omega, \Delta \omega))$ is compared with that of the $L = 14$ lattice. The data for $L = 10$ exhibits a peak at $\omega = 0.4$ with a negative weight in contrast to the $L = 14$ case with a positive weight. This confirms our hypothesis that low-energy in-gap state comes from the excitation from the optically allowed to forbidden state.

It is vital to test whether the photoinduced low-energy in-gap excitation demonstrated in Fig. 1(a) and 1(b) can be detected by pump-probe optical experiments. A re-
cent time-dependent optical reflectivity measurement on ET-F$_2$TCNQ has indicated the presence of photoinduced low-energy states located around $\sim$100 meV by analyzing the intensity oscillation near an exciton peak [2]. This energy is well below the energy of the exciton peak, 675 meV. The situation looks similar to Fig. 1(a) and (b). In order to explore the possibility to detect the in-gap excitation, we calculate $\text{Re} \sigma(\omega, \Delta t)$ and make a Fourier analysis as is the case of the experiment. Figure 2(a) shows the contour plot of $\text{Re} \sigma(\omega, \Delta t)$ ($1 \leq \omega \leq 10$) for a $L = 10$ ring with $U = 10$ and $V = 3$. We note that in this system and parameter set the photoinduced in-gap state emerges with positive-spectral weight at $\omega = 0.2$. We can find an oscillating behavior of a photoinduced state below the exciton peak at $\omega = 6.3$ as a function of $\Delta t$. The oscillation has a period of $\Delta \sim 16$. The Fourier amplitudes of the maximum intensity near $\omega = 6$ and the corresponding position are shown in Fig. 2(b) together with $\text{Re} \sigma(\omega, \Delta t = 6)$. We observe a dominant peak at $\omega \sim 2\pi/16 \sim 0.4$ in both the position and the intensity. The peak position of the Fourier amplitude roughly matches the position of the photoinduced in-gap state at $\omega = 0.2$, but close to each other. We also find that such a low-frequency oscillation close to the energy of the in-gap state does not occur in the cases with the negative-weight peak, e.g., $V = 2$ at $L = 10$ (not shown). Therefore, we consider that the oscillation appearing in the exciton peak reflects the energy of the in-gap state.

We next consider the case of a large pump width $t_d = 5$ for $V = 4.5$, where a strong enhancement of short-range charge correlation after pumping has been reported in Ref. [18]. Large width of the pump pulse $t_d$ leads to well-tuned energy transfer between the pump pulse and the system. After the pump pulse we thus expect an enhanced population of excited states around $\omega_0$, which is tuned to the position of the absorption peak. Such a non-thermal population of states around $\omega_0$ naturally leads to the appearance of the negative-weight of the absorption peak as shown in Fig. 3(c). In addition to the negative weight, we find a pronounced structure around $\omega = 2$ in Fig. 3(c) as comparing with the case of small $t_d$ [Fig. 3(b)] where additional structure around $\omega = 1$ exists. Since the pumping pulses with $t_d = 0.5$ used in Fig. 3(b) also induce an enhancement of charge correlation but with less extent than that in Fig. 3(c) [18], the pronounced structure around $\omega = 2$ in Fig. 3(c) may be attributed to a signature of enhanced charge correlation. In fact, the persistence of such an in-gap state is similar to the case of CDW phase as will be discussed below.

The CDW state is stable for $V > 5$. Figure 3(a) shows the $\Delta t$ dependence of $\text{Re} \sigma(\omega, \Delta t)$ for $V = 6$. The weight

FIG. 2. (Color Online) (a) Contour plot of $\text{Re} \sigma(\omega, \Delta t)$ ($1 \leq \omega \leq 10$) for the SDW phase of the $L = 10$ extended Hubbard model with $U = 10$ and $V = 3$. The amplitude is normalized by the maximum value of the equilibrium absorption peak. The conditions of pumping pulse are $A_0 = 0.3$ and $t_d = 0.5$. (b) The amplitudes of Fourier components of the maximum intensity (red broken line) near $\omega = 6$ and the corresponding position (black solid line) in (a). The amplitude is normalized by the maximum value. The blue thin line represents $\text{Re} \sigma(\omega, \Delta t = 6)$ for the same system.

FIG. 3. (Color Online) $\text{Re} \sigma(\omega, \Delta t)$ for the CDW phase of the extended Hubbard model with $U = 10$. (a) $\Delta t$ dependence for $L = 14$ and $V = 6$. $A_0 = 0.3$ and $t_d = 0.5$. (b) Different parameter sets for $L = 14$ and $V = 7$. $A_0 = 0.3(1)$, $t_d = 1(0.1)$, and $\Delta t = 20(2)$ for solid (broken) line. The in-gap peak with positive weight at $\omega = 5$ is insensitive to the parameter sets. (c) Equilibrium optical conductivity $\text{Re} \sigma(\omega)$ for the same parameter set as (a) but away from half filling (7 up-spin electrons and 6 down-spin electrons). (d) The same as (c) but the same parameter set as (b).
of the absorption peak at $\omega = 6.2$ substantially decreases with pumping. A low-energy structure with both positive and negative weights around $\omega = 0.3$ emerges and exhibits a strong $\Delta t$ dependence, which has not been seen in Fig. 1. Because of strong time dependence, we consider that the low-energy excitations is not a signature of photoinduced states. On the other hand, a strong in-gap peak appears at $\omega = 2.7$ with weak $\Delta t$ dependence. We regard this as a relevant photoinduced state. The $\text{Re}(\omega, \Delta t)$ for $V = 7$ also shows similar in-gap excitation that is independent of the pumping condition and $\Delta t$ as shown in Fig. 3(b). Therefore, the in-gap state is expected as a characteristic of CDW phase.

We now discuss the origin of the in-gap state in the CDW phase. Again, we should notice that the pumping creates photo-carriers in the system. However, in the CDW phase, a metallic or low-energy excitation is unstable as seen in Fig. 3. This is expected to be related to the charge distribution of the CDW ground state with the characteristic arrangement of alternating doubly occupied and empty sites. The photo-carriers are created by transforming pairs of doubly occupied and empty sites to singly occupied ones, but the movement of electrons is still blocked by the presence of neighboring doubly occupied sites. This may lead to a finite-energy excitation inside the original CDW gap. This is confirmed by calculating the optical conductivity for the extended Hubbard model away from half filling with single hole. Figure 3(c) and 3(d) exhibit the equilibrium $\text{Re}(\omega)$ for $U = 6$ and $U = 7$ in the CDW phase, respectively, with 7 up-spin electrons and 6 down-spin electron in the $L = 14$ lattice. In-gap peaks appear at $\omega = 2.8$ and $\omega = 4.6$ for $U = 6$ and $U = 7$, respectively. Their energies are close to that of the photoinduced in-gap excitation in Fig. 3(e) and 3(b), indicating that the presence of carriers is indeed the origin of the in-gap states.

The intensity of the in-gap state at $\omega_{\text{in-gap}} = 4.45$ oscillates with $\Delta t$ as shown in Fig. 4(a). After Fourier analysis, we find characteristic frequencies at $\omega = 4$ and $\omega = 14$ in Fig. 4(b) (black solid line). The former is very close to the energy of the in-gap state seen in $\text{Re}(\omega, \Delta t) = 6)$. We find that not only the intensity of the in-gap state but also the kinetic energy, which is an average value of the kinetic terms in $H$ (1), displays pronounced weight in the Fourier spectrum around $\omega_{\text{in-gap}}$ as seen in Fig. 4(b) (red broken line). Similar to the case of SDW, this characteristic oscillation related to the in-gap state is also recognized from the analysis on the remaining of the main absorption peak (not shown here). This means that the energy of the in-gap state may be detected by measuring oscillations of physical quantities. The response at $\omega = 14$ is due to high-energy excitations that are due to symmetry restrictions not picked up by the optical conductivity measurement. These excitations are determined by $U$ and $V$, leading to oscillations with short period as seen in Fig. 4(a).

In summary, we have analyzed physical properties of photoinduced states generated by a short laser pulse and their influence on the time evolution of various physical properties in the 1D the extended Hubbard model at half filling. We have calculated the time-dependent optical conductivity driven by a transient laser pulse and clarified the difference in the nonequilibrium optical response between the SDW and the CDW phases. The photoinduced in-gap excitations exhibit qualitatively different structure in the SDW or CDW ground state. In the SDW, a dominant in-gap state emerges at low-energy region reflecting the energy difference of even and odd parity of photoexcited states, where the even (odd) parity state corresponds to optically forbidden (allowed) state. In the CDW, an in-gap state appears in the middle of the Mott gap and its origin can be attributed to photogenerated carriers. The signature of the in-gap states in both cases emerges in the time evolution of physical quantities as a characteristic oscillation with a characteristic frequency roughly matching the energy of the in-gap states. Such a phenomenon has been reported in the time-dependent optical reflectivity measurement on ET-$F_2$TCNQ, a prototype 1D Mott insulator [2]. Observation of this characteristic frequency represents an experimental probe to identify photoinduced in-gap states.

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[1] H. Okamoto, H. Matsuzaki, T. Wakabayashi, Y. Takahashi, and T. Hasegawa, Phys. Rev. Lett. 98, 037401 (2007).

[2] S. Wall, D. Brida, S. R. Clark, H. P. Ehrke, D. Jaksch, A. Ardavan, S. Bonora, H. Uemura, Y. Takahashi, T. Hasegawa, H. Okamoto, G. Cerullo, and A. Cavalleri, Nat. Phys. 7, 114 (2011).

[3] H. Matsuzaki, M. Iwata, T. Miyamoto, T. Terashige, K. Iwano, S. Takahashi, M. Takamura, S. Kumagai, M. Yamashita, R. Takahashi, Y. Wakabayashi, and H. Okamoto, Phys. Rev. Lett. 113, 096403 (2014).

[4] D. Fausti, R. I. Tobey, N. Dean, S. Kaiser, A. Dienst, M. C. Hoffmann, S. Pyon, T. Takayama, H. Takagi, A. Cavalleri, Science 331, 189 (2011).

[5] D. Nicoletti, E. Casandruc, Y. Laplace, V. Khanna, C. R. Hunt, S. Kaiser, S. S. Dhesi, G. D. Gu, J. P. Hill, and A. Cavalleri, Phys. Rev. B 90, 100503(R) (2014).

[6] W. Hu, S. Kaiser, D. Nicoletti, C. R. Hunt, I. Gierz, M. C. Hoffmann, M. Le Tacon, T. Loew, B. Keimer, and A. Cavalleri, Nat. Mat. 13, 705 (2014).

[7] R. Matsunaga, Y. I. Hamada, K. Makise, Y. Uzawa, H. Terai, Z. Wang, and R. Shimano, Phys. Rev. Lett. 111, 057002 (2013).

[8] T. Papenfort, V. M. Axt, and T. Kuhn, Phys. Rev. B 76, 224522 (2007).

[9] J. Untherinninghofen, D. Manske, and A. Knorr, Phys. Rev. B 77, 180509(R) (2008).

[10] M. Eckstein and M. Kollar, Phys. Rev. B 78, 205119 (2008).

[11] Y. Kanamori, H. Matsueda, and S. Ishihara, Phys. Rev. B 82, 115101 (2010).

[12] M. Eckstein, M. Kollar, and P. Werner, Phys. Rev. B 81, 115131 (2010).

[13] G. De Filippis, V. Cataudella, E. A. Nowadnick, T. P. Devereaux, A. S. Mishchenko, and N. Nagaosa, Phys. Rev. Lett. 109, 176402 (2012).

[14] E. Iyoda and S. Ishihara, Phys. Rev. B 89, 125126 (2014).

[15] Z. Lenarčič, D. Golež, J. Bonča, and P. Prelovšek, Phys. Rev. B 89, 125123 (2014).

[16] H. Okamoto, T. Miyagoe, K. Kobayashi, H. Uemura, H. Nishioka, H. Matsuzaki, A. Sawa, and Y. Tokura, Phys. Rev. B 83, 125102 (2011).

[17] Z. Lenarčič and P. Prelovšek, Phys. Rev. Lett. 111, 016401 (2013).

[18] H. Lu, S. Sota, H. Matsueda, J. Bonca, and T. Tohyama, Phys. Rev. Lett. 109, 197401 (2012).

[19] P. Prelovšek and J. Bonča, in Strongly Correlated Systems - Numerical Methods, Springer Series in Solid-State Sciences, Vol. 176, edited by A. Avella and F. Mancini (Springer Berlin Heidelberg, 2013) pp. 1-30.

[20] Y. Mizuno, K. Tsutsui, T. Tohyama, and S. Maekawa, Phys. Rev. B 62, R4769 (2000).

[21] M. Takahashi, T. Tohyama, and S. Maekawa, J. Phys. Chem. Solids 63, 1599 (2002).