Dynamical coupling and separation of multiple degrees of freedom in a photoexcited double-exchange system

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We present a theory of ultrafast photo-induced dynamics in a spin-charge coupled system, motivated by pump-probe experiments in perovskite manganites. A microscopic picture for multiple dynamics in spin and charge degrees is focused on. Real-time simulations are carried out by two complimentary methods. Our calculation demonstrates that electron motion governs a short-time scale where charge and spin dynamics are combined strongly, while, in a long-time scale controlled by spin relaxation, charge sector does not follow remarkable change in spin sector. Present results are in contrast to a conventional double-exchange picture in equilibrium states.

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Ultrafast photo-control of electronic and magnetic structures has attracted much attention for a long time from viewpoint points of fundamental physics and technological application. In correlated electron systems with multiple degrees of freedom, i.e., charge, spin, orbital and lattice, a stable equilibrium phase is determined by a subtle balance of several interactions between them [1]. In a bare stable state at vicinity of phase boundary, a gigantic change in electronic structure is triggered by an optical pump pulse. Recently developed several temporal experiments enable us to access directly to the photo-dynamics of the multiple degrees of freedom [2].

Exotic equilibrium phenomena in correlated electron systems, such as, high-Tc superconductivity, colossal magnetoresistance (CMR), multiferroics, and others, are often attributed to strong correlation between multiple degrees of freedom. It is not quite trivial whether this naive strong-coupling picture between multiple degrees is applicable to the photo-induced dynamics or not. One example is seen in perovskite manganites [3, 4, 5, 6]. A key issue is a coupling between spin and charge. The CMR effects are resulted from a subtle balance of the charge ordered (CO) insulating phase associated with the antiferromagnetic (AFM) order and the ferromagnetic metallic (FM) phase, and are addressed in strong coupling between localized spins and conductive electrons [1]. By irradiation of a femto-second laser pulse in the CO insulating phase near the boundary, a transient metallic state expected from a reflectivity change appears within 100fs. While a macroscopic magnetization suggested from the magneto-optical Kerr rotation increases with the time of few ps [7-10, 11].

These results imply that changes in spin and charge sector, which are recognized to be strongly correlated in manganites, are not always accompanied with each other in photo-excited state. This dynamical coupling and separation of multiple degrees of freedom is not only a clue for an overall understanding of the optical-control electronic phases in correlated system. This also provides a key to uncover exotic phenomena in equilibrium state; roles of the multiple degrees are able to be identified separately. In spite of the progressive searches, however, a microscopic picture of multiple dynamics in photo-excited states has not been clarified yet.

In this Letter, motivated from the pump-probe experiments in manganites, we present a theory of the photo-induced dynamics in a spin-charge coupled system described by the double-exchange (DE) model. We focus on dynamical coupling and separation between spin and charge. To attack this issue, two complimentary methods, the exact-diagonalization (ED) and the Hartree-Fock (HF) methods, are adopted. Calculations demonstrate that electron motion governs the short-time scale where charge and spin sectors change cooperatively. On the other hand, in the long-time scale controlled by spin relaxation, charge sector does not follow change in the long-range spin correlation.

Let us introduce the extended DE model,

$$\mathcal{H}_{DE} = -\alpha t \sum_{\langle ij \rangle} \left(c_{i\alpha}^\dagger c_{j\alpha} + H.c.\right) - J_H \sum_i \mathbf{s}_i \cdot \mathbf{S}_i + U \sum_i n_{i\uparrow} n_{i\downarrow} + V \sum_{\langle ij \rangle} n_{i\uparrow} n_{j\uparrow} + J_S \sum_{\langle ij \rangle} \mathbf{S}_i \cdot \mathbf{S}_j,$$ (1)

de re counting electron at site $i$ with spin $\sigma(\uparrow, \downarrow)$, and $\mathbf{S}_i$ is the operator for the localized spin. We introduce the number operator $n_{i\alpha}(\equiv \sum \epsilon_{\alpha} c_{i\alpha}^\dagger c_{i\alpha})$ and the spin one $\mathbf{s}_i = \frac{1}{2} \sum_{\alpha} c_{i\alpha}^\dagger \mathbf{\sigma}_{\alpha} c_{i\alpha}$ with the Pauli matrices $\mathbf{\sigma}$ for the conduction electrons. The first and second terms describe the electron transfer, $\alpha t$, and the Hund coupling, $J_H$, respectively. The on-site Coulomb repulsion $U$, the nearest-neighbor (NN) one $V$, and the AFM superexchange interaction $J_S$ between the localized spins are also taken into account. All energy and time parameters are given as a unit of $t$ and $t^{-1}$, respectively. A magnitude of the transfer integral is changed by changing $\alpha$ from one. For a typical value of $t$ in manganite, a unit of time, $t^{-1}$ corresponds to 0.5fs~1fs.

To analyze this model, two complimentary methods, the ED and HF ones, are utilized. In the ED method, the time-dependence of the wave function and the excitation spectra are able to be obtained exactly. On the other hand, in the HF one, simulations in a large system size and long time distance are possible, and the spin relaxations are taken into account.
We introduce the Fourier transform (FT) of the operator, Lanczos methods [12]. One-dimensional clusters of system size \(N_L = L(\leq 13)\) with the open-boundary condition are adopted. Electron number is \(N_{\text{ele}} = (L + 1)/2\) corresponding to the 1/4 filling. For simplicity, amplitude of the localized spin is set to be 1/2. We take a damped oscillator form for the vector potential of the pump photon as \(A_{\text{pump}} = A_0 e^{-i\omega_0 t - 2\gamma_0 t}\) at time \(t\) with frequency \(\omega_0\) and a damping constant \(\gamma_0\). A center of the wave packet is defined by \(t = 0\). The wave function for the one-photon absorbed state at time \(\tau (> \gamma_0^{-1})\) is derived by the first order perturbation with respect to \(A_{\text{pump}} [13]\

\[
|\Psi(\tau)\rangle = \mathcal{N} e^{-i\mathcal{H}_\text{DE} \tau} \left[ \frac{\gamma_0}{(\omega_0 - \mathcal{H}_\text{DE} + E_0)^2 + \gamma_0^2} \right] j|0\rangle , \tag{2}
\]

where \(j = i\alpha \sum_{j} (c_{\alpha j}^\dagger c_{\alpha j} - H.c.)\) is the current operator, \(|0\rangle\) is the wave function before pumping, \(E_0 = |0\rangle \langle \mathcal{H}_\text{DE} |0\rangle\), and \(\mathcal{N}\) is a normalization factor. Transient excitation spectra are calculated by the linear-response theory. For example, the one-particle excitation spectra are given by a sum of the electron and hole parts, \(A(q, \omega) = A_{\text{ele}}(q, \omega) + A_{\text{hole}}(q, \omega)\), where the first term is obtained as [14]

\[
A_{\text{ele}}(q, \omega) = -\frac{1}{\pi} \sum_a \text{Im}(c_{qa}^\dagger c_{qa} - \omega - \mathcal{H}_\text{DE} + E) \frac{1}{\gamma_{qa}} \tag{3}
\]

We introduce the Fourier transform (FT) of the operator, \(c_{qa}\), the energy \(E = \mathcal{H}_\text{DE}\), and a damping factor \(\gamma\). A bracket \(\langle \cdots \rangle\) implies the expectation value with respect to \(|0\rangle\) in the case before pumping, and that with \(|\Psi(\tau)\rangle\) after pumping. The pseudo-momentum in the open boundary condition is defined as \(O_q = \sqrt{2/(L+2)} \sum \sin(qr_i) O_i\) where \(O_i (O_q)\) is a physical quantity in the real (momentum) space.

In the HF method, the mean-field (MF) decoupling is introduced in the many-body terms in \(\mathcal{H}_\text{DE}\), such as \(n_{i\uparrow} n_{i\downarrow} \rightarrow \langle n_{i\uparrow} \rangle n_{i\downarrow} + n_{i\uparrow} \langle n_{i\downarrow} \rangle - c_{i\uparrow}^\dagger c_{i\downarrow}^\dagger c_{i\uparrow} c_{i\downarrow} - c_{i\uparrow}^\dagger c_{i\downarrow}^\dagger c_{i\downarrow} c_{i\uparrow} + \text{const}\). where \(\langle n_{i\uparrow} \rangle\) is a site-dependent MF. The two-dimensional \(N_L = L \times L (L = 6 - 10)\) site cluster with the periodic-boundary condition is adopted. The electron number is \(N_{\text{ele}} = L^2/2\). The localized spins are treated as classical vectors with amplitude of 3/2. The pump-photon irradiation is simulated as an electronic excitation from the highest occupied HF level to the lowest unoccupied one without changing spin. The time evolution of the electronic wave function is obtained by the time-dependent HF method [15, 16]:

\[
\phi^{(l)}(\tau + \Delta \tau) = T \exp \left[ -i \int_\tau^{\tau + \Delta \tau} \mathcal{H}_\text{HF}^{\text{relax}}(\tau) d\tau \right] \phi^{(l)}(\tau), \tag{4}
\]

where \(\phi^{(l)}(\tau)\) is the \(l\)-th HF wave function, and \(\mathcal{H}_\text{HF}^{\text{relax}}(\tau)\) is the HF Hamiltonian for Eq. (1). Dynamics of the localized spins is described by the Bloch-type equation:

\[
\partial_\tau S_i = -H_{i} \times S_i + \partial_\tau S_i |_{\text{relax}}, \tag{5}
\]

where \(H_{i}\) is the MF at site \(i\) defined by \(H_{i} = J_H \langle s_i \rangle + J_S \sum_{\delta} \langle s_{i+\delta} \rangle\). The last term in Eq. (5) represents the spin relaxation and dephasing which will be explained later.

First, we show the results obtained by the ED method. The parameter values are chosen to be \(U = 10, J_H = 8, V = 5, J_S = 0.4, \omega_0 = 3.72\) and \(\gamma_0 = 0.4\). We take \(\alpha = 1\) except for the results in the inset of Fig. 5. Somewhat larger energy parameter values than the realistic ones for manganites are used to reproduce the AFM/CO insulating ground-state in one-dimension. Thus, our analyses in the ED method are restricted to qualitative properties in the photo-induced dynamics. First we focus on the charge dynamics. An insulating gap and a tendency of the Brillouin-zone doubling shown in \(A(q, \omega)\) [see Fig. 1(a)] imply an alternating charge alignment. Just after photon pumping [see Fig. 1(b)], a photo-carrier band appears inside of the CO gap in \(A(q, \omega)\). Large amounts of the spectral intensity in the upper and lower bands still remain. These results imply that the long-range charge alignment is collapsed, but a short-range correlation survives. It is worth to note that the band width of the in-gap band is broaden with time evolution [see Figs. 1(b) and 1(c)]. The optical absorption spectra \(\alpha(\omega) = \sum (\pi N_L)^{-1} \text{Im}(j(\omega - \mathcal{H}_\text{DE} + E + i\gamma)^{-1} j)\) are presented in Fig. 1(d). After photon pumping, spectral weight appears inside of the optical gap, and grows up with increasing in time. We confirm, through the analyses in a small cluster system, that the lowest component of the in-gap spectral weight corresponds to the Drude component in the thermodynamic limit. These time evolutions in \(A(q, \omega)\) and \(\alpha(\omega)\) are not observed in both the spin-less V-t and Hubbard models. That is, the localized spins and its coupling with conduction electrons play a central role on these phenomena.

Change in the spin sector is monitored by the static spin correlation function, \(S(q) = \langle S_{-q} S_q \rangle\), where \(S_q\) is FT of \(S_i\) [see Fig. 2]. Before pumping, the spin structure is ferrimagnetic one where the localized spins are aligned antiferromagnetically and the total spin is \(S^{\text{tot}} = 1 + L\). This is seen in large intensity of \(S(q \sim \pi)\). After pumping, the large AF correla-
tion is rapidly suppressed, the momentum dependence in $S(q)$ is almost smeared out, and the system is nearly paramagnetic.

Time evolutions of both the conduction electron and localized spins are summarized in Fig. 3. We plot the second moment $W$ of $A(q, \omega)$ reflecting the band width of the in-gap state, the integrated spectral weight $D$ inside of the optical gap in $\alpha(\omega)$, and the NN spin correlation $K_S$ for the localized spins. These are defined by $W = \int_0^{\omega_f} \sum_q A(q, \omega)(\omega - \omega_i)^2 d\omega$ with the center of the in-gap band $\omega_i$ and the upper (lower) band edge $\omega_l (\omega_u)$. $D = \int_0^{\omega_f} \alpha(\omega) d\omega$ with the upper (lower) edge of the in-gap component $\alpha_l (\alpha_u)$, and $K_S = N_B^{-1} \Sigma_{ij} \langle S_i \cdot S_j \rangle$ with the number of NN bonds $N_B$. An almost identical time dependence is shown in the three curves; they increase linearly after pumping and are saturated around $t = 10/\tau$. The characteristic time scale where $K_S$ shows a shoulder is denoted as $\tau_S$. In the inset of Fig. 3 $K_S$’s for different values of the electron transfer are presented. With increasing $\alpha$, a slope of the curve in the region of $\tau < \tau_S$ increases, and the AFM correlation collapses rapidly. When $K_S$’s are plotted as functions of $\tau \alpha t$, the slopes for several values of $\alpha$ are almost identical. We conclude that photo dynamics in the charge and short-range spin sectors are strongly correlated with each other, and are mainly governed by the electron transfer. These results are interpreted as follows; motion of the conduction electrons, excited directly by photon, destroys the charge and AFM orders. The collapse of the AFM spin alignment further promotes motion of the conduction electron, and finally the two systems steady down cooperatively.

In the calculations by the ED method explained so far, the spin-angular momentum is conserved. However, it is expected that, in the simulation for longer time scale, the spin relaxation leads the system to the lower excited state on the energy surface. Microscopic mechanisms for breaking of the spin conservation in manganites are not clear yet. In photo-excited states where the Mn 3$d e_g$ orbitals are mainly concerned, the orbital angular momentum, $L_z$, is basically quenched. This is in contrast to the magnetic semiconductors, such as (Ga,Mn)As where a large spin-orbit coupling provides a spin relaxation [17]. Possible relaxation mechanisms are mixing of the on-site $e_g$ and $t_{2g}$ orbitals by the off-diagonal components of $L_z$, and that induced by lattice distortions with the $T_{1u}$ symmetry. A mixing of the inter-site $e_g$ and $t_{2g}$ orbitals by the GdFeO$_3$-type distortion is another candidate. We predict that the photo-induced magnetization is promoted by excitation of the phonons concerning these lattice distortions.

We introduce the spin relaxation phenomenologically in the HF formulation, and examine roles of the spin conservation on time evolution. In Eq. 5, we introduce the parallel and perpendicular spin components to the total magnetization $\mathbf{S}$, and that induced by lattice distortions with the $T_{1u}$ symmetry. A mixing of the inter-site $e_g$ and $t_{2g}$ orbitals by the GdFeO$_3$-type distortion is another candidate. We predict that the photo-induced magnetization is promoted by excitation of the phonons concerning these lattice distortions.

The results by the HF method are presented in Fig. 4. The parameter values are $U = 8, J_H = 6, V = 1$ and $J_S = 0.03$. We take $\alpha = 1$ except for the inset of Fig. 4(a). Before pumping, CO associated with the canted-AFM alignment in the localized spins is realized. Two time scales are recognized in the NN spin correlations for the localized spins $K_S$ shown in Fig. 4(a); a sharp increase in a short-time scale ($\sim 200/\tau$) depending on the transfer integral [see the inset of Fig. 4(a)], and a slow increase in a long-time scale depending on $\Gamma_L$. The short-time scale corresponds to $\tau_S$ introduced previously. The long-time scale termed $\tau_L$ is defined as a time when an amount of change in the FM spin correlation $S(0)$ becomes $1/e$ of its total value; $\tau_L = 645/\tau$ and $250/\tau$ for $\Gamma_L = 0.002$ and $0.008$, respectively. In the case without relaxation, a $K_S$ v.s. $\tau$ curve is qualitatively similar with that in the ED method [see Fig. 3]. This implies that with regard to the short-time dynamics with $\Gamma_L = 0$, a qualitative feature commonly ob-
FIG. 4: (color online) (a) NN spin correlation function $K_S$ for several values of the relaxation constant $\Gamma_L$. The inset shows $K_S$ for the several transfer integrals $\alpha t$ and $\Gamma_L = 0.002$. (b) Spin correlation functions $S(0)$ for several $\Gamma_L$ (bold lines), and $S(\pi, \pi)$ for $\Gamma_L = 0.002$ (dotted line). The dotted arrows represent $\tau_S$. (c) Charge correlation function $N(\pi, \pi)$ and NN charge correlation $K_N$.

erved by the two methods is reliable. A quantitative difference between the $\tau_S$ values in the ED and HF calculations is mainly attributed to the different photon densities $x_{ph}$: the present $x_{ph} = L^{-2} = 0.01$ is smaller than $x_{ph} \sim 0.1$ in the ED calculation. For small $x_{ph}$, a system takes long time to be settled down to a stationary state. We checked the above points by the HF calculations in one-dimension; a linear increase and a saturation are observed in $K_S$, and $\tau_S$ increases by increasing $L$ from 10 to 30.

We focus on the results for $\Gamma_L = 0.002$. As shown in Fig. 4(b), around $\tau_S$, $S(\pi, \pi)$ disappears, but only a subtle change is seen in $S(0)$; the system becomes almost paramagnetic. In the long-time scale beyond $\tau_S$, a remarkable change is seen in $S(0)$ which grows up monotonically toward the full-spin polarization, in contrast to $K_S$ and $S(\pi, \pi)$. Charge dynamics, i.e. the long-range charge correlation $N(q) = \langle \Delta n_q \Delta n_{-q} \rangle$ and the NN correlation $K_N = (2/Na) \sum_{\{ij\}} \langle \Delta n_i \Delta n_j \rangle$, are shown in Fig. 4(c). We define $\Delta n_j = n_i - 1/2$ and its FT, $\Delta n_q$. Time dependences of the long- and short-range charge correlations are similar to that in $K_S$, not in $S(0)$. These are nearly saturated around $\tau_S$ and do not show a remarkable change beyond $\tau_S$. Changes in $S(0)$ by spin relaxation are almost irrelevant to the charge sector. This weak correlation between the charge correlations and $S(0)$, corresponding to magnetization, is not expected from the equilibrium state where these are combined strongly. The key ingredient in this dynamical separation between the spin and charge sectors is spin conservation; magnetization dynamics is governed by the spin relaxation rather than the charge motion. The present case is in contrast to the photo-induced dynamics in electron-phonon systems where conserved quantities in a phonon sector does not exist.

From the calculated results, we provide a whole picture of the photo-induced spin and charge dynamics. In the short-time scale, charge motions governed by the electron transfer destroy the CO and short-range AFM correlations cooperatively. Excess energy given by the pump photon flows from the conduction electrons to the localized spins. This state is further relaxed through the spin relaxation in long-space and long-time scales. Changes in the charge sector is almost saturated in the short-time scale and do not follow the spin relaxation. This scenario provides a microscopic interpretation for the experimentally observed multiple time scales in $\Delta R$ and $\Delta \theta$. The present dynamical coupling/separation in the multiple degrees of freedom is not expected from a conventional picture in equilibrium state and are key issues to reveal the photo-induced dynamics in correlated system.

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and without this approximation is less than 10 percent.

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