Decoupling spin-orbital correlations in a layered manganite amidst ultrafast hybridized charge-transfer band excitation

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In the mixed-valence manganites, a near-infrared laser typically melts the orbital and spin order simultaneously, corresponding to the photoinduced $d^0d^0 \rightarrow d^0d^1$ excitations in the Mott-Hubbard bands of manganese. Here, we use ultrafast methods—both femtosecond resonant x-ray diffraction and optical reflectivity—to demonstrate that the orbital response in the layered manganite Nd$_{1-x}$Sr$_{1+x}$MnO$_4$ ($x = 2/3$) does not follow this scheme. At the photoexcitation saturation fluence, the orbital order is only diminished by a few percent in the transient state. Instead of the typical $d^1d^0 \rightarrow d^0d^1$ transition, a near-infrared pump in this compound promotes a fundamentally distinct mechanism of charge transfer, the $d^0 \rightarrow d^1L$, where $L$ denotes a hole in the oxygen band. This finding may pave a different avenue for selectively manipulating specific types of order in complex materials of this class.

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The mixed-valence (Mn$^{3+}$/Mn$^{4+}$) manganites promise great potential for the applications in information technology, e.g., spintronics [7] and skyrmion devices for low power consumption [8]. Fundamentally, these applications rely on the coexistence of, and interplay between, the distinct electronic orders therein [9]. On the experimental front, manipulation of the corresponding order parameters has been extensively explored from quasistatic to ultrafast time scales, using external stimuli including static magnetic [10] or electric field [11], x-ray illumination [12], as well as ultrafast optical pulses [13–19]. Under the manipulation schemes reported so far, all electronic orders will respond simultaneously to an external trigger because of the aforementioned strong correlation effect in materials of this class. For example, while a femtosecond near-infrared (NIR) pump can melt the orbital order via the resonant relaxation of the Jahn-Teller (JT) distortion, the charge and magnetic orders also collapse concomitantly [15–17]. What remains to be realized however is the essence of control in these types of quantum systems: decoupling these strong correlation effects to modify one type of order while leaving others undisturbed. This is of great importance for example for the next-generation multistate logic or memory devices [20].
In this Rapid Communication, we demonstrate the decoupling of strong correlations on an ultrafast time scale. We show that in a mixed-valence manganite, the orbital ordering is robust amidst intense femtosecond laser pulse excitation. This was achieved by employing resonant soft x-ray scattering (RSXS) together with optical spectroscopy to study the ultrafast dynamics in the overdoped layered system Nd$_1$−$x$Sr$_x$MnO$_3$ (NSMO, $x = 2/3$) [21]. We optically pumped the magnetic system, demonstrating a strong response within the magnetic state. We then directly studied the orbital modulation in the transient state—at the photoexcitation saturation fluence—and found an unusual resilience. That is, the RSXS intensity from the orbital order is diminished by no more than 5%. This points to the importance of the $d^0 \rightarrow d^1L$ charge-transfer photexcitation process, where $d^i$ ($i = 0, 1$) and $L$ denote the $e_g$ orbital occupancy of the Mn$^{3+}$/Mn$^{4+}$ cation and the hole in the oxygen anion band, respectively [22], and how this could be exploited to selectively control the order parameters in the transient state. This observation will be important for technological applications and draws a fundamental distinction from the $d^1d^0 \rightarrow d^0d^1$-type excitation which has been reported in past ultrafast experiments on the manganites [13–17]. This is shown schematically in Fig. 1 and will be discussed below.

A single crystal of NSMO ($x = 2/3$) was used in our study and was grown by the floating-zone method [21]. It was polished along the (110) direction. The optical conductivity ($E \parallel ab$) data were calculated by the Kramers-Kronig analysis on the reflectivity spectra measured by a combination of Fourier spectroscopy and grating spectroscopy. The NIR (1.55 eV) pump, RSXS probe measurements were performed on the SXR instrument at the Linac Coherent Light Source using the RSXS end station, an avalanche photodiode, and a recently developed THz optical system [23–26]. The pump and probe pulses propagated collinearly to the sample position with $\pi$ polarization ($E_{\text{pump}} \parallel ab$). The x-ray beam energy was tuned to the Mn $L_3$ edge (641.5 eV) with a bandwidth of 1.3 eV [27], giving an attenuation length of about 70 nm for this geometry. The overall time resolution, calibrated by measuring the response of a sample of gallium phosphide, was about 400 fs [28]. In addition, we have performed time-resolved optical reflectivity measurements. A high-power Ti : sapphire-based laser (1.55 eV) with a 50-fs pulse duration and 120-Hz repetition rate was split and cross-polarized into the pump ($E_{\text{pump}} \parallel ab$) and probe ($E_{\text{probe}} \parallel c$) pulses, which gives a temporal resolution of about 75 fs. The optical penetration depth matches that of the x rays [29], in agreement with previous work on related systems [30].

The NIR optical conductivity $\sigma(\omega)$ with $E \parallel ab$ between 10 and 300 K in the equilibrium state is shown in Fig. 2(a). A gapped mode can be clearly resolved at low temperatures. Upon heating, the optical gap $\Delta$ and conductivity $\sigma(\omega)$ at the NIR pump energy (1.55 eV) undergo a mild suppression across the antiferromagnetic (AFM) ordering temperature $T_N \approx 90$ K [31], while a stronger one occurs around the orbital ordering temperature $T_{OO} \approx 257$ K [Figs. 2(b) and 2(c)] [31,32]. The AFM state in this compound is of charge-exchange type and characterized by the population of zigzag ferromagnetic (FM) spin chains along the (1, 1, 0) direction [31]. As a result, the $d^1d^0 \rightarrow d^0d^1$ hopping depicted in Fig. 1(a) is plausible for the temperature dependence because it is governed by Hund’s rule and short-range FM chains can persist up to $T_{OO}$ in manganites with charge-exchange type antiferromagnetism [33]. However, the AFM state in the overdoped NSMO ($x = 2/3$) has a very short spin-correlation length along the zigzag chains, which quickly disappears around $T_N$ [31]. From this evidence combined with the strong optical response near $T_{OO}$, we conclude that the NIR mode observed here is largely absent of the $d^1d^0 \rightarrow d^0d^1$ process. Indeed, the behavior revealed in Fig. 2 is characteristic of a $d^0 \rightarrow d^1L$ charge-transfer process, which gives rise to an optical gap that gets suppressed at $T_{OO}$ due to the abrupt change in the optical spectra instead of the departure of ferromagnetism [34,35].

The ultrafast response of the optical reflectivity $R(t)$ as a function of pump fluence is shown in Fig. 3(a). These curves were fit to a single exponential decay:

$$ R(t) = \frac{1}{2} \times \left[ \text{erf} \left( \frac{t - t_0}{\tau_0} \right) + 1 \right] \left[ A_0 + A_1 e^{-\left(\frac{t}{\tau_0}\right)^2} \right], \tag{1} $$

where $t_0$ and $A_0$ are the initial decay rate and amplitude, $\tau_0$ is the recovery rate of the fast component, and the constant $A_1$ accounts for the long-lived state associated with the nuclear lattice that recovers on a much slower time scale [~400 ps at 21 mJ/cm², inset of Fig. 3(a)]. All the optical data presented...
here can be well described by Eq. (1), allowing us to quantitatively determine the electronic response to the NIR pump.

The initial reflectivity drop is always resolution-limited within the errors, i.e., \( t_0 = 75 \) fs, indicating a nonthermal excitation process. The fast recovery process is on the time scale between 0.5 and 1.0 ps [Fig. 3(b)]. This component is often assigned to the electron-electron thermalization [36] that partially recovers the electronic order [15,16]. The decay amplitude increases linearly with the fluence until saturating at 16(1) mJ/cm\(^2\) [Fig. 3(b)]. While a NIR pump can also excite free carriers in the system, the linear fluence dependence below the saturation threshold supports the notion that the pump induced reflectivity change at 1.55 eV must mainly come from the above-the-band-gap electronic excitations [15,30,32,37]. As a result, our optical data have revealed that the electronic system has been saturated by the femtosecond NIR pump at \( f_{\text{sat}} = 16(1) \) mJ/cm\(^2\), through the ultrafast \( d^0 \rightarrow d^1 L \) charge-transfer process [Fig. 1(b)].

Typically, the ultrafast dynamics in a manganite under an optical pump are governed by the Mott-Hubbard type \( d^1 d^0 \rightarrow d^0 d^1 \) excitations. This strongly suppresses the underlying orbital order, as has been reported by many authors [13–17]. We implemented RSXS to investigate if the ultrafast orbital order response is consistent with a charge-transfer-type excitation.

Next the ultrafast dynamics of the orbital order under femtosecond NIR excitation were measured. For 15.0 mJ/cm\(^2\), which is slightly below or equal to \( f_{\text{sat}} \) [Fig. 3(b)], the RSXS pump-probe curve is shown in Fig. 4(b). This behavior also fits well to Eq. (1). The initial x-ray-scattering intensity decay is resolution limited, but is given here by the jitter of the optical and x-ray pulse arrival time, i.e., \( t_0 = 400 \) fs [38]. This verifies the nonthermal nature of this photoinduced transition.
For recovery times, the time window is not wide enough to reliably fit the $A_t$ term on the longer time scales. Fixing it to zero leads to $\tau_R = 23(7)\,\text{ps}$. This value is smaller than the spin-lattice thermalization, i.e., the energy transfer from the initially excited electron system to the nuclear lattice; this time scale is usually long ($> 100\,\text{ps}$) in relevant systems [30]. But $\tau_R$ is also considerably longer than the fast electronic recovery process extracted from the optical reflectivity measurements [$\lesssim 1.0\,\text{ps}$, Fig. 3(b)]. This intermediate recovery process, which has been observed in other nonoptimally doped manganites [16], is related to the transient photoinduced phase separation that leads to an inhomogeneous recovery of the electronic order. There seems to be evidence of an oscillation immediately after the pumped orbital state, but as this frequency is much longer than the typical phonon modes found in relevant systems [15,19,39], we believe it is unlikely to be real orbital excitations.

Most importantly, we have found that the orbital order is surprisingly robust against this photoexcitation. Although the electronic system is saturated by the NIR pump at 15.0 $\text{ml/cm}^2$ [Fig. 3(b)], the RSXS intensity scattered by the orbital order is only suppressed by about between 4% and 5% in the transient state. Because we have carefully calibrated the energy density for the different geometries and experiments, this character sharply contrasts the observations in other manganites [29]. In previous experiments on similar systems, the same NIR excitation promotes the Mott-Hubbard type $d^1d^0 \rightarrow d^0d^1$ transitions [Fig. 1(a)] and therefore greatly suppresses the underlying orbital order [15].

According to the Zaanen-Sawatzky-Allen classification scheme, the electronic structure near the Fermi surface (FS) of a correlated transition-metal insulator depends on not only the Hubbard $U$, but also on the charge-transfer energy $\Delta$ [22]. Due to the comparable $U$ and $\Delta$ values in mixed-valence manganites, the above-the-band-gap excitation near the FS can be to be dominated by either the $d^1d^0 \rightarrow d^0d^1$ transition, the $d^0 \rightarrow d^1L$, or both [40–43]. Our RSXS observations rule out a meaningful contribution from the former in NSMO ($x = 2/3$), since the relative orbital order intensity loss of no more than 5% in the transient state is measured close to $f_\text{cat}$. On the other hand, the $d^1L$ orbital state created by the charge-transfer process does not contribute to ordered orbital structure because of the twofold $e_g$ degeneracy [44]. Consequently, the bulk orbital order structure would not be expected to change considerably under this photoexcitation scheme. The robustness of the orbital order therefore strongly indicates that the latter charge-transfer type scheme is responsible for the NIR photoexcitation process in this compound [Fig. 1(b)]. This is consistent with the temperature dependence of the optical spectrum at these energies (Fig. 2).

To obtain more insight about the nature of the large optical spectral weight at 1.55 eV in Fig. 2(a), we measured the ultrafast dynamics of the optical reflectivity at 15 $\text{mJ/cm}^2$ as a function of temperature (Fig. 5). The decay amplitude $A_0$ changes smoothly with temperature below about $T_N$. Upon further heating, $A_0$ is suppressed and falls below the noise level around 110 K [Fig. 5(b)]. This suggests that the electronic excitation process triggered by the NIR pump here is magnetically activated: the increased probability amplitude for the hopping matrix element—only in the magnetic state—originates from the hybridized wave function. Intuitively, this character does not fit the $d^0 \rightarrow d^1L$ scenario depicted in Fig. 1(b) because it does not require magnetic order. However, similar behavior is expected if the $e_g$ band formed by the Mn$^{3+}$ cations is involved. For example, the $d^1d^0 \rightarrow d^0d^1$ process in Fig. 1(a) belongs to this category; it is energetically favored if the adjacent Mn$^{3+}$/Mn$^{4+}$ spins are parallel with each other, a configuration that is only fulfilled in the magnetically ordered region of the relevant manganites [15,17,30].

Since the orbital response and optical spectroscopy imply the $d^1d^0 \rightarrow d^0d^1$ contribution is very weak in NSMO ($x = 2/3$), we propose that the $d^0 \rightarrow d^1L$ process is between the hybridized Mn (3$d$) and O (2$p$) bands near the FS [32]. The $dp$ hybridization occurs when $U$ and $\Delta$ are similar in strength [41,42]. In this modified scenario, the $O^2−/2p$ electrons are spin polarized by the Mn$^{3+}$/Mn$^{4+}$ electrons in the hybridized $dp$ band. Accordingly, the photoinduced charge transfer is allowed along the FM zigzag spin chains along the (1, 1, 0) direction, which is restricted in the AFM state of NSMO ($x = 2/3$) [31]. This scenario naturally explains the ultrafast results presented here as well as the mild change in the static NIR optical spectrum around $T_N$ (Fig. 2).

Furthermore, the $d^0 \rightarrow d^1L$ scenario may also be partially responsible for the ultrafast orbital dynamics in the other layered manganite La$_{1−x}$Sr$_{1+x}$MnO$_4$ ($x = 1/2$) [30]. In this compound, a femtosecond NIR pump melts about 25% of the orbital order reflection probed by RSXS at the saturation fluence, whereas the AFM order is completely suppressed. This finite RSXS intensity was explained by the residual JT distortion after the photoexcitation [30]. A recent femtosecond NIR pump/hard x-ray probe work on Pr$_{1−x}$Ca$_x$MnO$_3$ ($x = 1/2$), where the $d^0 \rightarrow d^0d^1$ physics is clearly dominant, directly revealed that the JT distortion disappears with the orbital and charge order when the electronic system is fully excited [15] and therefore does not fit the model proposed in Ref. [30]. Alternatively, the residual orbital order in La$_{1−x}$Sr$_{1+x}$MnO$_4$ ($x = 1/2$) could indicate that about 75% of the electronic excitations involve the $d^0 \rightarrow d^1L$ charge transfer.
In conclusion, we have investigated the effect of NIR excitation (1.55 eV) on the orbital order in the layered manganite NSMO ($x = 2/3$) through ultrafast techniques. Conducting a systematic ultrafast optical reflectivity study, we found the fluorescence and temperature dependence of the optical reflectivity change. By using RSXS at an x-ray free-electron laser source, we combined this finding with the ability to directly monitor the femtosecond orbital response in the transient photoexcited state. We found that the RSXS intensity arising from the orbital order is diminished by no more than 5% percent while the electronic system is fully saturated. These results strongly suggest that the photoexcitation mechanism in this compound is $d^0 \rightarrow d^1 L$, instead of the commonly assumed $d^1 d^0 \rightarrow d^0 d^1$. In addition, we show that the correct description of the photoexcitation process requires the $d^0$ hybridization near the FS in the magnetically ordered state. Taking advantage of the hybridized $d^0 \rightarrow d^1 L$ charge-transfer mechanism holds great potential for selectively manipulating the electronic order. Furthermore, this shows that other excitation mechanisms at play may be more closely related to that seen in the cuprates, such as the high-energy scale physics associated with Mott-like excitations [45]. The results here point to future work based on exploiting the different transfer mechanisms for specific materials to unravel the emergent excitations in systems with multiple types of order, especially other quantum materials that display nearly degenerate energy scales, such as multiferroics [20].

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