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Insulator-to-Metal Transition Induced by Mid-IR Vibrational Excitation in a Magnetoresistive Manganite

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Abstract. Selective vibrational excitation of insulating Pr$_{0.7}$Ca$_{0.3}$MnO$_3$ at 17 µm triggers a transition to a metallic state. A four order of magnitude drop of the sample resistivity and ultrafast, nanosecond-lived reflectivity changes are observed.

1. Introduction

In perovskite manganites the strong correlation between charge, spin, orbital and lattice degrees of freedom produces complex phase diagrams as well as extreme susceptibility to external stimuli, which can alter the balance between competing phases of the system. Pr$_{0.7}$Ca$_{0.3}$MnO$_3$ has been a testing ground for a variety of phase-control concepts, whereby the transition from the charge-orbital ordered states to the ferromagnetic metallic state has been driven by external stimuli such as magnetic or electric field, high pressure and photoexcitation by x-rays or visible light [1]. Here we study the effect of selective excitation of an infrared active phonon mode, which is clearly shown to drive the solid toward the metallic phase. To our knowledge, this is the first example of an ultrafast insulator-metal transition driven in the electronic ground state.

A unique example among various manganites, Pr$_{1-x}$Ca$_x$MnO$_3$ shows insulating behaviour over the whole composition (x) range. This is a consequence of a more pronounced orthorhombic distortion, resulting in a small tolerance factor (see Fig. 1) [2]. The tolerance factor is a geometrical factor depending on the (Pr,Ca)-O and Mn-O distances. It plays a key role in the physics of manganites, being related to the hopping probability of carriers between neighbouring Mn sites [2]. The infrared spectrum of Pr$_{0.7}$Ca$_{0.3}$MnO$_3$ exhibits a typical pattern for manganites [3], with the highest frequency phonon assigned to a Mn-O stretching vibration (Fig. 1). This mode modulates the geometrical parameters determining the tolerance factor and it is thus expected to have a strong coupling to the electronic system.
2. Electrical Measurements

Changes in the sample conductivity were monitored by directly measuring the sample resistance following mid-infrared excitation at 17 µm (200 fs, 1 µJ). The laser spot fully covered the space between two gold electrodes deposited on the sample surface. The sample was biased at a DC voltage of 27 V and cooled to 40 K (charge-ordered, anti-ferromagnetic insulator phase). Mid-infrared excitation induces a 50-fold current increase, corresponding to a resistance drop from 700 MΩ to 15 MΩ. The sample resistivity drops by more than four orders of magnitude, from the 40 K value of 3×10⁻⁷ Ω⋅cm to about 10⁻³ Ω⋅cm (see Fig. 2). The effect vanishes when the infrared pump wavelength is tuned outside the phonon resonance at 17 µm, including radiation at 8.5 µm with twice as much energy/photon. The temperature jump due to laser excitation (< 2 K) can be ruled out as origin of such a dramatic resistivity drop.

3. Pump-probe Experiments

The ultrafast dynamics of the photoinduced phase transition in Pr₀.7Ca₀.3MnO₃ have been previously studied by means of femtosecond reflection spectroscopy [4]. The formation of the metallic phase following electronic excitation results in ultrafast (< 230 fs), long lived reflectivity changes in the near IR. Here, we show that vibrational excitation has comparable effects on the dynamics of the
reflectivity changes. The sample was excited by 17 µm pulses (fluence: 1mJ/cm²) and the reflectivity was probed by delayed 800 nm pulses. Fig. 3(a) shows the relative change of reflectivity (∆R/R) as a function of pulse delay, evidencing an ultrafast reflectivity drop which persists up to the nanosecond timescale. Identical dynamics are observed following electronic excitation. Fig. 3(b) compares the spectrum of the reflectivity changes measured 1 ps after vibrational excitation with the absorption spectrum around the phonon peak at 17 µm, showing that reflectivity changes depend on resonant infrared excitation of Mn-O modes.

![Fig.3.](image-url)

**Fig.3.** (a) Relative change of reflectivity at 800 nm (∆R/R) as a function of pulse delay following electronic excitation (dotted line, normalized) and vibrational excitation at 17 µm (solid line). (b) ∆R/R measured 1 ps after excitation (circles) and absorption spectrum around the 17 µm phonon resonance (solid line). In order to compare the two curves, the phonon spectrum was convoluted with the spectrum of the broad bandwidth pump pulses.

4. Conclusions

The excitation of a specific phonon mode has been demonstrated as a tool to drive the solid in the electronic ground state toward a competing phase of the system. Ultrafast vibrational control of the collective phase of correlated electron systems opens a new window on the interaction between lattice structure and electronic or magnetic properties. This approach may extend well beyond the case of CMR manganites, providing new insight into the behavior of complex matter, including the controversial nature of High-T_{c} superconductivity.

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References