Ultrafast Strain Engineering in Complex Oxide Heterostructures

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We report on ultrafast optical experiments in which femtosecond midinfrared radiation is used to excite the lattice of complex oxide heterostructures. By tuning the excitation energy to a vibrational mode of the substrate, a long-lived five-order-of-magnitude increase of the electrical conductivity of NdNiO3 epitaxial thin films is observed as a structural distortion propagates across the interface. Vibrational excitation, extended here to a wide class of heterostructures and interfaces, may be conducive to new strategies for electronic phase control at THz repetition rates.

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Complex oxide heterostructures have emerged as multifunctional materials of striking flexibility, in which unconventional electronic phases can be realized by engineering the strain field across interfaces [1–5]. This same mechanical coupling is also expected to be effective on the ultrafast timescale, and could be exploited for the dynamic control of materials properties. Here, we demonstrate that a large-amplitude midinfrared field, made resonant with a stretching mode of the substrate, can switch the electronic properties of a thin film across an interface. Exploiting dynamic strain propagation between different components of a heterostructure, insulating antiferromagnetic NdNiO3 is driven through a prompt, five-order-of-magnitude increase of the electrical conductivity, with resonant frequency and susceptibility that is controlled by choice of the substrate material.

Many of the functional properties of ABO3 perovskite oxides are extremely sensitive to rotation and tilting of the BO6 octahedra, which control the hopping amplitudes and the exchange interaction through the B-O-B bond angle and length. It follows that one can engineer the electronic and magnetic properties by designing or actively controlling such distortions [6–8], or by coupling them to other structural instabilities [9,10]. One class of materials in which octahedral rotations and distortions have been linked to changes in the electronic structure are the rare earth nickelates, which display a sharp transition from a high-temperature metallic to a low-temperature insulating state. In the bulk, this electronic phase transition is accompanied by a structural change from an orthorhombic (Pbnm) to a monoclinic (P21/n) symmetry with an increase in the Ni-O-Ni bond bending and the appearance of a charge density wave. Additionally, charge disproportionation between adjacent Ni sites is associated with different Ni-O bond lengths [11,12]. At low temperatures, the nickelates also possess an unusual antiferromagnetic spin arrangement [13].

The metal-insulator transition in nickelates has been investigated so far using static experimental techniques that directly affect the electronic bandwidth through the Ni-O-Ni bond angle, such as chemical [14,15] and hydrostatic pressure [16,17] or epitaxial strain [18]. Recently it was also shown that electrostatic fields can be used to control the metal-insulator transition in these materials [19]. Ultrafast optical experiments have measured carrier relaxation [20] at low irradiation levels, insufficient to drive a photo-induced phase transition.

In this work we use intense coherent femtosecond midinfrared pulses to control the lattice structure and with it the electronic properties along a nonequilibrium path. Indeed strong vibrational excitation [21] is capable of inducing electronic phase transitions, as demonstrated in a recent series of experiments on cuprates [22] and manganites [23,24] where transient superconductivity and metalliclicity were triggered. In all of these cases the B-O stretching mode (∼70–80 meV) of the crystal structure was excited. Here we use epitaxy to mechanically couple NdNiO3 to a substrate, whose B-O stretching mode is selectively driven.

In a first set of experiments, NdNiO3 epitaxial thin films (100 u.c., 33 nm thick) were deposited on (001) LaAlO3 single crystals by off-axis RF magnetron sputtering as detailed in Ref. [19]. Note that NdNiO3 single crystals cannot be grown in large enough size for most experiments. This substrate provides −0.5% compressive strain to the material and reduces the metal-insulator-transition temperature TMi from a bulk value of 200 K to about 130 K, as determined through 4-point dc transport measurements. In our midinfrared pump—optical probe experiments the sample was excited by 150 fs pulses tuned

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to photon energies between 70 and 130 meV. These pulses were generated by optical parametric amplification and difference frequency generation of the output of a Ti:sapphire laser amplifier, as detailed in Ref. [25]. Transient changes in reflectivity probed in the near (800 nm, 1.55 eV) and the far-infrared (1–6 THz, 5–25 meV) were used to characterise the electronic properties of NdNiO₃ after vibrational excitation. Far-infrared probe experiments are performed using normal incidence reflection geometry. Broadband THz pulses were generated in a gas plasma [26] and detected by electro-optic sampling using 800 nm probe pulses in ZnTe and GaP crystal [27].

Figure 1(a) shows transient near-infrared reflectivity changes $\Delta R/R_0$ induced by excitation at 15 $\mu$m wavelength for different base temperatures $T$. At room temperature, where NdNiO₃ is metallic, only a modest reflectivity increase was observed, decaying within 2 ps and likely associated to electronic excitations and relaxation near the Fermi level. For $T < T_{\text{MI}}$ a dramatically different response is observed. Changes in reflectivity as large as $-20\%$ indicate the formation of a metastable electronic phase. No measurable relaxation of the signal could be observed within 100 ps. By extrapolation, we estimate the lifetime to be well in excess of 1 ns. This response is independent of the polarization of either the pump or the probe beams. As evident from the temperature dependence of the photoresponse (discussed below), such large reflectivity changes can only be achieved below the static metal-insulator-transition temperature.

The time-dependent THz response, displayed in Figs. 1(b) and 1(c) for the spectral range between 5 and 25 meV, reveals that this transient phase is metallic. First, the frequency integrated THz reflectivity ($\Delta E/E_0$) displays a long-lived increase, suggestive of a phase with higher conductivity [Fig. 1(b)]. These reflectivity changes have the opposite sign to those observed when probing at 800 nm (see above) pointing to a transfer of spectral weight from high to low frequencies [28]. Second, the frequency dependent THz reflectivity for the unperturbed heterostructure and for the photo-induced state are well traced by the calculated reflectivity of a 33 nm NdNiO₃/LaAlO₃ heterostructure with NdNiO₃ in its insulating and metallic state, respectively, [Fig. 1(c)]. As shown in Fig. 1(d), within 3 ps after vibrational excitation we extrapolate a 5 orders of magnitude increase in quasi-dc conductivity for the NdNiO₃ (for details on the calculations see Supplemental Material [29]). Note that for the fluence of the order 1 mJ/cm² used here, and for the 4% absorption at this wavelength, the temperature increase is estimated to be below 5 K (see Supplemental Material [29]), pointing to a nonthermal mechanism for the observed effect.

As the light induced reflectivity changes scale linearly with the pump fluence (shown below), we can define an effective photosusceptibility as $\chi = |d\rho/df|$, where $f = F(1 - R)$ is the absorbed pump fluence ($F$) and $\rho = \Delta R/R_0$. The dependence of $\chi$ on the pump photon energy measured at $T = 20$ K is shown in Fig. 2(a), and highlights the most striking observation reported here. We find that the susceptibility for the photoinduced insulator-metal transition follows the absorption of the substrate rather than that of NdNiO₃. This phenomenon could be related to two distinct effects. One scenario involves the excitation of a substrate phonon, which may lead to a distortion coupled into the film across the interface. Alternatively, epitaxial strain may shift the resonance frequency of the thin film itself, accidentally matching the frequency of the substrate. In the first case one expects a strong dependence of the photosusceptibility on the
oscillator strength of the substrate phonon. Differently, in the second case, the substrate resonance strength is not expected to influence the amplitude of the response.

To clarify this point, 100 u.c thick NdNiO₃ epitaxial thin films deposited on (110) NdGaO₃ single crystals were analyzed. This substrate provides different mechanical boundary conditions (tensile strain, 1.1%), a different vibrational resonance (74 meV) and a much smaller oscillator strength of the phonon [30,31]. NdNiO₃ thin films deposited on NdGaO₃ can be considered our best representation of the bulk material, since their transition temperature (∼ 180 K) is close to the one observed in ceramics (∼ 200 K [11]). Figure 2 shows the temperature dependence of the 800 nm reflectivity changes ∆R₅₈₃/ROI probed 5 ps after vibrational excitation in NdNiO₃/NdGaO₃ heterostructures. As observed for LaAlO₃ substrates, the transient near-infrared reflectivity changes are negative, long-lived and the temperature at which the reflectivity decreases after excitation corresponds to T_M (Fig. 2). More importantly, the signal amplitude ∆R₅₈₃/ROI is in this case much smaller than the one observed in NdNiO₃/LaAlO₃ heterostructures.

This is clearly highlighted by the dependence of the photosusceptibility on the pump photon energy shown in Fig. 2(c). Also for this heterostructure the photosusceptibility peaks at the phonon resonances of the substrate. Moreover, the data show a clear dependence of the photosusceptibility on the absorption coefficient of the substrate, which is 3 times larger for NdNiO₃/LaAlO₃ [30,31]. This indicates that the modulation of the electronic properties of the thin film occurs via excitation of the crystal lattice of the substrate [Fig. 2(e)].

Insight into the dynamics of the insulator-metal transition can be gained from the pump intensity dependence. Transient reflectivity changes probed using 800 nm pulses after vibrational excitation with pump fluences between 0.2 and 2.1 mJ/cm² and T = 20 K, are reported in Fig. 3(a). For fluences larger than ∼0.8 mJ/cm² the reflectivity change ∆R₅₈₃/ROI increases linearly [Fig. 3(b)] and no saturation of the signal is observed in the experimentally accessible fluence range. As the fluence is reduced below ∼0.8 mJ/cm² the photoinduced state decays exponentially back to equilibrium on the picosecond time scale. In this regime the relaxation time is strongly reduced with lower pump fluences [Fig. 3(c)]. This observation can be interpreted by considering that the photo-induced transition is initiated locally, injecting a number of domains with sizes (or densities) that depend on the pump fluence. For large pump fluences the nucleated domains are large (or dense) enough to reach the percolation threshold, leading to a long-lived phase. Below threshold the metallic domains never coalesce and collapse back to the parent phase. Alternatively, since the vibration of the film is driven
The other possibility is that the excitation will be transferred deep into the film, stabilizing the metallic state throughout the sample. Similar (but static) inhomogeneous distortion profiles induced by mechanical coupling with the substrate have recently been proposed for other perovskite heterostructures [6].

In summary, this work demonstrates how optical excitation of a substrate lattice can control the electronic properties of a thin film through propagation of a dynamic distortion across the interface. This greatly broadens the scope of research of quantum phase control via optical excitation and the field of strain engineering in complex solids. The precise interaction within NdNiO$_3$ responsible for the photo-induced metallic phase remains poorly identified. One possibility is that the electronic transition is caused directly by the excitation of the Ni-O stretching mode that couples to the B-O bond vibration of the substrate, as changes in Ni-O bond lengths are known to occur during the thermally induced metal-insulator transition. The other possibility is that the B-O stretching mode couples to a rotational mode of the oxygen octahedra that would lead to changes in overlap between the Ni 3d and O 2p orbitals. Such a coupling between the stretching and rotational modes has recently been observed in bulk manganites [21]. The most obvious way forward for this area of research is to seek conditions under which bi-directional phase control can be achieved, for example, by incorporating phonon-active layers in superlattices. Understanding and engineering such couplings between structural modes in complex oxide heterostructures opens new possibilities for design and control of their scientifically fascinating and technologically useful properties.

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