Relaxation timescales and electron-phonon coupling in optically-pumped \( \text{YBa}_2\text{Cu}_3\text{O}_{6+x} \) revealed by time-resolved Raman scattering.

N. Pellatz,\(^1,2\) S. Roy,\(^1,2\) J-W. Lee,\(^3\) J. L. Schad,\(^3\) H. Kandel,\(^4\) N. Arndt,\(^4\) C.B. Eom,\(^3\) A.F. Kemper,\(^5\) and D. Reznik\(^1,2\)

\(^1\)Department of Physics, University of Colorado, Boulder, Colorado 80309, USA
\(^2\)Center for Experiments on Quantum Materials, University of Colorado - Boulder, Boulder, Colorado, 80309, USA
\(^3\)Department of Materials Science and Engineering, University of Wisconsin-Madison, Madison, WI 53706, USA
\(^4\)Department of Physics and Mathematics, University of Wisconsin-Parkside, Kenosha, WI 53140
\(^5\)Department of Physics, North Carolina State University, Raleigh, North Carolina 27695, USA

Time resolved measurements provide a new way to disentangle complex interactions in quantum materials due to their different timescales. We used pump-probe Raman scattering to investigate the apical oxygen vibration in \( \text{YBa}_2\text{Cu}_3\text{O}_{6.9} \) under nonequilibrium conditions. Time-dependence of the phonon population demonstrated strong electron-phonon coupling. Most importantly, the phonon shifts to a higher energy due to transient smearing of the Fermi surface in a remarkable agreement with theory. We also discuss new insights into photoinduced superconductivity reported at lower doping that follow from these results.

Driving quantum materials with electromagnetic fields can generate novel phases and states away from thermal equilibrium \([1,5]\). Recently reported signatures of superconductivity at elevated temperatures in photoexcited copper oxides and intercalated fullerenes are particularly interesting but still enigmatic \([6,12]\). In most of these experiments the ultrafast laser pulse (pump) drives or photoexcites the system and another ultrafast pulse (probe) takes snapshots of a specific property as a function of pump-probe time delay. Such experiments also elucidate energy flows between phonons, electrons, and magnons providing a way to determine the strength of different interactions.

This work focuses on relaxation timescales and interactions between electrons and phonons in a prototypical copper oxide superconductor, \( \text{YBa}_2\text{Cu}_3\text{O}_{6+x} \).

Electron-phonon coupling in the copper oxides is still enigmatic. It allows electron-hole recombination and electron scattering with the creation or annihilation of phonons; phonons in turn can decay into electron-hole pairs. As a result, spectroscopic quasiparticle peaks shift and/or broaden due to decreased lifetime. However, electron-phonon scattering is not the only process that gives rise to these effects. Internally, phonon-phonon coupling (anharmonicity) also broadens the phonon peaks; similarly, electron-electron interactions can broaden the electronic quasiparticles, as does disorder \([13]\). Thus, extracting just the electron-phonon coupling strength from the linewidths of quasiparticle peaks is challenging.

To get around this problem, we used time-resolved Raman scattering (TRR) (Fig. 1b)\([1,4,23]\). In TRR an ultrafast optical pump laser pulse first photoexcites the material and Raman scattering from another time-delayed pulse probes the system. In the copper oxides, previous TRR work highlighted ultrafast destruction of the antiferromagnetic order in the undoped parent compound of \( \text{YBa}_2\text{Cu}_3\text{O}_6 \)\([24]\) and nonequilibrium behavior of the superconducting gap in \( \text{Bi}_2\text{Sr}_2\text{CaCu}_2\text{O}_{8+\delta} \) (BSCCO) \([25]\).

In our experiment near infrared (IR) pump pulses create hot electrons, and the time-delayed Raman probe measures the apical oxygen \( A_{1g} \) phonon. Apical oxygen modes recently came to the center of attention due to their impact on electronic \([20]\) and magnetic \([27]\) degrees of freedom of the copper-oxygen planes, interlayer charge transport \([28]\), and photoinduced superconductivity \([9,11,29]\). The \( A_{1g} \) mode is known for a large spectroscopic linewidth \([29,31]\) and an anomaly at the superconducting transition temperature, \( T_c \).

Upon optical pumping the phonon occupation number increased dramatically due to absorption of energy from photoexcited electrons as expected from strong electron-phonon coupling. \([32,37]\). This phonon also hardened at short time delays as a result of the decrease of its electronic self-energy expected from smearing of the Fermi surface due to very high transient electronic temperature. This effect provides a rigorous test of electron-phonon calculations based on electronic structure. Our results were in quantitative agreement with Green’s functions-based theory of Refs. \([33,35]\) (see Fig. 1b).

Data presented in this letter were collected on a 170 nm-thick (110)-oriented \( \text{YBa}_2\text{Cu}_3\text{O}_{6.9} \) thin film prepared by pulsed laser deposition on a (110) \( \text{LaAlO}_3 \) substrate with \( T_c \) of 81 K \([33,40]\). We also obtained a dataset on a similar film with reduced oxygen concentration (\( T_c=40\)K) with the temperature in the cryostat of 100K. Our TRR setup (Fig. 2b) uses 20 kHz 790 nm (1.57 eV) laser pulses from an amplified mode-locked Ti:sapphire laser, which produces intense 40 fs pulses. Second harmonic generation at 395 nm (3.14 eV) was used as the probe light source for Raman scattering.
FIG. 1. Time-resolved Raman scattering (TRR) setup and key results. (a) Schematic of the TRR experiment and the color map showing representative data on the anti-Stokes (AS) side of the spectrum obtained with the sample at 300 K. Negative time corresponds to the probe pulse arriving before the pump pulse when the system is still at thermal equilibrium. Most intensity in the peak is from the apical oxygen phonon whose atomic displacements are indicated by green arrows in the schematic of the YBa$_2$Cu$_3$O$_y$ unit cell to the right of the color map. (b) Comparison of diagrammatic theory [38] and experiment for the dependence of phonon energy as a function of electronic temperature at delay times up to 600 fs. The temperature in the cryostat was 10 K/300 K in the top/bottom panels respectively. Impurity scattering of 20 meV was added to the theory curve in the upper panel, but not to the lower panel, since scattering by phonons at 300 K is greater. Unrenormalized phonon energy was picked to obtain good agreement of theory with experiment at thermal equilibrium [38].

The time-energy uncertainty principle limits the energy resolution of the ultrafast probe. In order to resolve the phonons from the elastic line, the 790 nm pulses were passed through an in-house built time-compensating band-pass filter (Fig. 1b) making both the pump and the probe pulses narrower in energy and broader in time. A cross-correlation measurement gave a time-resolution of 220 fs FWHM (see Supplementary Note 1 in [41]). The time-averaged probe power was below 1 mW, to eliminate self-pumping nonlinearities (in Supplementary information [41], see reference [17] therein).

The scattered light was collected by a pair of parabolic mirrors with a polarizer in the middle, and analyzed on a single-stage McPherson spectrometer equipped with a LN$_2$-cooled CCD detector. A custom-made notch filter blocked elastically-scattered light. The samples were in air or in a cryostat in a He exchange gas. We show results obtained with the pump photon polarization parallel to the ab-plane. Photon polarization along the c-axis gave similar results. Background measured under identical conditions but without the probe was subtracted from raw data.

We read off phonon temperature from the intensities on the Stokes (S) and anti-Stokes (AS) sides, $I_S$ and $I_{AS}$ [14, 15]. They are related by the fundamental principle of detailed balance:

$$\frac{g I_{AS}}{I_S} = \frac{(\omega_0 + \omega_{ph})^4}{(\omega_0 - \omega_{ph})^4} e^{\frac{-\hbar \omega_{ph}}{k_B T_{ph}}},$$

where $\omega_0$ is the laser frequency, $\omega_{ph}$ is the phonon energy, $k_B$ is the Boltzmann’s constant, $T_{ph}$ is the phonon temperature, and $g$ is equal to 1. Introducing $g$ allows us to correct for the systematic error due to imprecise spectrometer calibration. We made $g = 0.9$ to make the phonon temperature at negative times equal to 300 K at room temperature. Following convention, we define the temperature of each bosonic mode, $T_{boson}$, via its relation to the occupation number $n = (e^{\frac{\hbar \omega_{ph}}{k_B T_{boson}}} - 1)^{-1}$. Away from thermal equilibrium different phonons have different occupation numbers, and therefore different temperatures. According to the fluctuation-dissipation theorem, the S and AS Raman intensities are given by $I_S = (n+1) \chi'' (\hbar \omega)$, and $I_{AS} = n \chi'' (\hbar \omega)$, where $\chi''$ is the imaginary part of the Raman response function (polarizability) of the phonon of interest. The color map in Fig. 1b shows the dramatic increase of the AS intensity, indicating an increase of $T_{ph}$ (Eqs. 1, 2), as well as a peak shift to larger energy right after photoexcitation.

The apical oxygen phonon around 500 cm$^{-1}$ and a weaker plane oxygen mode at 440 cm$^{-1}$ (Fig. 2a inset), which may include substantial apical oxygen character [42, 43], dominate the Raman spectrum in the zz geometry where both incident and scattered photons are polarized perpendicular to the copper-oxygen planes (Fig. 2a) [29, 30, 44]. The intrinsic lineshape of the phonon peaks is known to be best describes by characteristic Fano profiles (See inset in Fig. 2a and [29, 30, 44]), however, due to a very broad energy resolution of the experiment, they were fit with Gaussians as shown in the main panel of Fig. 2a. When fitting the time-resolved data where the peaks are not resolved (Fig. 2a, main panel), the intensity of the 440 cm$^{-1}$ peak was fixed at 20% of the 500

[...]

$\Delta E$, $\Delta g$, $\Delta S$, $\Delta AS$
FIG. 2. Raw data and phonon temperature at two doping levels. (a) Background-subtracted TRR spectra at different delay times with the cryostat at 300 K (black circles). The peaks at ±500 cm\(^{-1}\) are Stokes (+) and anti-Stokes (−) phonon peaks. Red line is a guide to the eye representing smoothed data without the pump. Large peak widths are due to the increased energy width of the pulsed laser. (b) Phonon temperature as a function of time after photoexcitation at optimal doping and reduced doping where photoinduced superconductivity was reported earlier [9]. Temperature in the cryostat was 100K. Pump fluence was 8 mJ/cm\(^2\), the same as in Ref. [9]. Note that pump energy is absorbed first by electron-hole pairs, which then thermalize with the apical oxygen phonons, i.e., electronic temperature never drops below the phonon temperatures.

-1000 -500 0 500 1000
Raman Shift (cm\(^{-1}\))
0 500 1000
Counts (arb. unit)

FIG. 3. Phonon temperature and energy together with fits to the two temperature model. Pump (probe) fluence was 1.4 mJ/cm\(^2\) (15 \(\mu\)J/cm\(^2\)). \(T_{\text{cryo}}\) was 250 K. Black line represents hot phonons, red – hot electrons, blue – cold phonons. The electronic temperature curves in (b) and (c) are the same. Inset to (b) illustrates the behavior at large delay times for the sample in air at 300 K. Dashed blue line in (c) is the measured pump-probe cross correlation centered at t=0.

The decay saturates around 5 ps (Fig. 3 inset) when the photoexcited region reaches the internal thermal equilibrium temperature, \(T_{eq} = 300\) K. Inset to Fig. 3b shows \(T_{eq}\) as a function of \(T_{\text{cryo}}\) highlighting increased transient heating of the sample, \(\Delta T\), at lower temperatures due to its smaller heat capacity. Further equilibration with the cryostat exchange gas is much longer than 100 ps (see inset to Fig. 3a). An earlier study attributed this heating to nonthermal effects, but the heating at these time delays is consistent with the pump fluence as discussed below.

Our 1.5 eV pump photons should create electron-hole pairs [45], which thermalize among themselves much faster than the time resolution. Time resolved angle-resolved photoemission (trARPES) showed that electrons reached a maximum temperature of 800 K with 100 \(\mu\)J/cm\(^2\) pump pulses in BSCCO [46]. In graphite, the electronic temperature reached far above 1000 K when pumping with 150 \(\mu\)J/cm\(^2\) [15, 47, 48]. It is reasonable to expect similarly high electronic temperatures in YBCO.

We interpret our results in terms of the two-temperature model where \(f\) is a fraction of phonons with strong electron-phonon coupling, \(\lambda\). Photoexcited electrons first give off energy to the hot phonons [46, 49, 50], which in turn decay into other, cold phonons with the lifetime \(\tau\) until all phonons and electrons thermalize at \(T_{eq}\). The system then slowly equilibrates with the heat bath via propagating (mostly acoustic) phonons.

Our experiments provide detailed information on each step of this process.

We performed a detailed analysis of the time-dependent phonon data obtained with a lower pump fluence (Fig. 3). The phonon temperature dramatically increases within the time-resolution to about 450 K independent of the temperature in the cryostat. This increase is followed by the exponential decay starting from 0.5 ps.
where $n$.

FIG. 4. Dependence on pump power, $P$, and $T_{cryo}$. (a) Phonon hardening at 200 fs ($E_{ph}(200 \text{ fs}) - E_{ph}(-1000 \text{ fs})$) as a function of $T_{cryo}$ at $P = 27 \text{ mW}$. Inset: Phonon hardening at $T_{cryo} = 10 \text{ K}$ as a function of $P$. (b) Phonon temperature at 200 fs and 1 ps as a function of $T_{cryo}$ at $P = 27 \text{ mW}$ (fluence of 1.4 mJ/cm$^2$). Note that $T_{cryo} = T_{ph}$ at 1 ps. Inset: $T_{cryo}$ at 200 fs at $T_{cryo} = 10 \text{ K}$ as a function of $P$. (b) Electron-phonon coupling constant $\gamma$ together with the anharmonic scattering rate $1/\tau$. See Supplementary Notes 2 and 3 for details on how $\gamma$ was taken from measurements in Ref. [51]. If $\Omega$ is a typical phonon energy, then the electronic temperature, $T_{el}$, the hot phonon temperature, $T_h$, and the cold phonon temperature, $T_c$, obey the rate equations

$$\frac{\partial T_{el}}{\partial t} = -\frac{3\Omega_0^2}{\hbar \pi k_B^2} \frac{n(T_{el}) - n(T_h)}{T_{el}} + \frac{P(t)}{c_{el}(T_{el})},$$

(2)

$$\frac{\partial T_h}{\partial t} = \frac{c_{el}(T_{el})}{c_h(T_h)} \frac{3\Omega_0^2}{\hbar \pi k_B^2} \frac{n(T_{el}) - n(T_h)}{T_{el}} - \frac{T_h - T_c}{\tau},$$

(3)

$$\frac{\partial T_c}{\partial t} = \frac{c_h(T_h)T_h - T_c}{c_c(T_c)} \frac{T_h - T_c}{\tau},$$

(4)

where $n(T) = (\epsilon^{T_{cryo}/k_B T} - 1)^{-1}$ and the specific heats are $c_{el} = \gamma T_{el}$, $c_h = 3N_{atm}f \Omega_0 (\partial n_h/\partial T_h)$, and $c_c = 3N_{atm}(1 - f) \Omega_0 (\partial n_c/\partial T_c)$. Here, $N_{atm}$ is the number of atoms per formula unit (= 12.9 for our sample) and the value of $\gamma$ was taken from measurements in Ref. [51]. Equations 2-4 correspond to equations 1-3 of Ref. [60] written down in terms of $\Delta T$, which we measured with high precision.

We solved these coupled differential equations using the Euler method with 1 fs steps. Fig. 3a presents a fit to this model with $\Omega_0 = 60 \text{ meV}$ and $\tau = 478 \pm 25 \text{ fs}$ (see Supplementary Note 3 for fits with different values of $\Omega_0$ and $\tau$ [11]). $f = 0.14$ and $\lambda = 0.065$ were free parameters. Electrons initially heat up to 1000 K similarly to the previous result on BSCCO [66], and then cool quickly to thermalize with hot phonons at 300 fs. Thermalized hot electrons and hot phonons have a larger heat capacity compared with the heat capacity of electrons alone so they equilibrate with cold phonons with the much slower 478 fs time constant at 250 K. This lifetime increases with reduced temperature reaching 700 fs at 10 K (see Supplementary Note 2 and Fig. 4 in [11]). It is much smaller than in graphite [15,21]. The electron-phonon coupling strength is close to $\lambda = 0.07$ at all temperatures, which is smaller than the the LDA value [52].

Thermalization on subpicosecond timescales is controversial. Deviations from the Fermi-Dirac distributions of the electrons have been reported at pump-probe delay times of a few hundred fs in simple metals [63,64]. However, the ARPES measurements of Ref. 40 showed that in BSCCO, thermalization occurs already at 100fs. It also showed that nonthermal distribution at shorter times is characterized by a small (only a few percent) deviation from the Fermi-Dirac distribution. Assuming that YBCO behaves similarly, for our purposes electrons can be treated as internally thermalized [53].

The assumption of internally thermalized hot phonons is valid only if they have similar electron-phonon coupling strength. For example, the 2T model breaks down in graphite where one hot phonon had a significantly larger electron-phonon coupling than another [15]. According to Ref. [69] as well as prior experimental work, the breathing phonons and the buckling phonons have a significantly stronger electron-phonon coupling than the apical oxygen mode [55,66]. We found that adding phonons that have a much stronger electron-phonon coupling than the apical mode to the model can increase $\lambda$ by nearly an order of magnitude, thus $\lambda$ obtained from fits to the apical oxygen phonon occupation data alone is not accurate. Time-dependent electronic temperature, however, was only weakly model-dependent, which allowed us to make a quantitative comparison of the phonon energy vs electronic temperature to theory as discussed below.

We observe a profound shift of the phonon peak position to a higher energy (hardening) at small time delays (Fig. 5a) with the maximum at 200 fs. It is independent of $T_{cryo}$ but decreases with reduced pump power (Fig. 4b). These changes in the phonon energy do not correlate with either phonon occupation or with the electric field of the pump, which should follow the cross-correlation curve in Fig. 4b. Instead phonon hardening closely follows the electronic temperature. A similar effect in graphite was associated with electronic temperatures of over 1000 K [17,57,58].

The phonon self-energy $\Pi(q, \Omega)$ is proportional to electronic polarizability, which depends on the electronic temperature [59,61]. To make this quantitative, we model the system as a band of electrons interacting with
the three strongly coupled phonons: the $A_g$, $B_{1g}$ and breathing modes, which are kept at the cryostat temperature. Electronic quasiparticle excitations renormalize the apical phonon frequency. The increase in the quasiparticle temperature diminishes the softening of the apical phonon (lines in Fig. 1b), leading to an increase in the phonon frequency as a function of the effective electronic temperature [60] (also see Supplementary note 4 therein for the details). The initial decrease of the theory curve at $T_{\text{cryo}}=10\, \text{K}$ is a result of the broadening of the electronic bands due to their phonon-mediated self-energy. At weak interactions (and low T) broadening the bands increases the bubble (phonon self-energy), and thus softens the phonon as temperature increases. This effect competes with the dominant hardening effect due to smearing of the Fermi surface described above. The calculation picks it up due to relatively sharp bands when the lattice temperature is low in the absence of impurity scattering. It is not present at 300K because of the existing broadening by phonons at this temperature. Increasing the impurity scattering rate weakens the kink at $T_{\text{cryo}}=10\, \text{K}$ as shown in Supplementary note 5 [41].

Note that simple heating of the sample leads to phonon softening, not hardening. This is primarily because of thermal expansion caused by increased population of acoustic phonons. However, acoustic phonons stay cold at short pump-probe time delays and lattice expansion does not occur.

We have demonstrated quantitative agreement with theory of the time-varying apical phonon frequency as the electronic system loses its excess energy to the broader phonon bath. It highlights that the dynamics of energy transfer are responsible for the temporal behavior of the electrons and phonons, as well as the disparity between interactions in- and out of equilibrium. Although in strongly correlated materials such as YBCO there are strong Coulomb processes and impurity scattering that dominate the electronic spectra, when it comes to time domain these processes rapidly come to an internal equilibrium and effectively shut off [38]. Our results provide a new, phonon-centered, perspective on previous experiment in Bi$_2$Sr$_2$CaCu$_2$O$_8$ with both time-resolved ARPES [63] and ultrafast electron diffraction [64] that both observed a similar quantitative agreement.

Our work provides new insights into photoinduced superconductivity [65, 66]. Its signatures were recently reported in the optical spectra of underdoped YBCO up to time-delays of about 1ps when pumping with 790 nm near-IR pulses [9] as well as with pulses that resonated with IR-active apical oxygen phonons. Our experiments reproduced the former pumping condition and showed that optimally-doped and underdoped YBCO behave similarly (Fig. 2b). We found that hot and cold phonons were out of thermal equilibrium, but electrons and hot phonons were at or near thermal equilibrium at time delays below 1ps. At these time-delays electronic temperatures were always well above room temperature. We plan to determine transient heating while pumping the IR-active phonons in future experiments.

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[1] Y. H. Wang, H. Steinberg, P. Jarillo-Herrero, and N. Gedik. Observation of Floquet-Bloch states on the surface of a topological insulator. Science, 342(6157):453–457, 2013.

[2] Andrei Kirilyuk, Alexey V. Kimel, and Theo Rasing. Ultrafast optical manipulation of magnetic order. Rev. Mod. Phys., 82:2731–2784, Sep 2010.

[3] Phoebe Tengdin, Wenjing You, Cong Chen, Xun Shi, Dmitriy Zusin, Yingchao Zhang, Christian Gentry, Adam Blonsky, Mark Keller, Peter M Oppeneer, et al. Critical behavior within 20 fs drives the out-of-equilibrium laser-induced magnetic phase transition in nickel. Science Advances, 4(3):9744, 2018.

[4] H. Okamoto, T. Miyagoe, K. Kobayashi, H. Uemura, H. Nishioka, H. Matsuzaki, A. Sawa, and Y. Tokura. Photoinduced transition from mott insulator to metal in the undoped cuprates Nd$_2$CuO$_4$ and La$_2$CuO$_4$. Phys. Rev. B, 83:125102, Mar 2011.

[5] H. Ehrke, RI Tobey, Simon Wall, SA Cavill, Michael Först, Vikaran Khanna, Th Garl, N Stojanovic, D Prabhakaran, AT Boothroyd, et al. Photoinduced melting of antiferromagnetic order in La$_{0.5}$Sr$_{1.5}$MnO$_4$ measured using ultrafast resonant soft x-ray diffraction. Phys. Rev. Lett., 106(21):217401, 2011.

[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. Optically enhanced coherent transport in YBa$_2$Cu$_3$O$_{6.5}$ by ultrafast redistribution of interlayer coupling. Nature Materials, 13(7):705–711, July 2014.
and graphite by time-resolved incoherent anti-Stokes Raman scattering. *Phys. Rev. B*, 81:165405, Apr 2010.

[21] Shiewei Wu, Wei-Tao Liu, Xiaogian Liang, P James Schuck, Feng Wang, Y Ron Shen, and Miquel Salmeron. Hot phonon dynamics in graphene. *Nano letters*, 12(11):5495–5499, 2012.

[22] J. Zhu, R. B. Versteeg, P. Padmanabhan, and P. H. M. van Loosdrecht. Dynamical resonance quench and fano interference in spontaneous Raman scattering from quasiparticle and collective excitations. *Phys. Rev. B*, 99:094305, Mar 2019.

[23] R. B. Versteeg, J. Zhu, P. Padmanabhan, C. Boguschewski, R. German, M. Goedecke, P. Becker, and P. H. M. van Loosdrecht. A tunable time-resolved spontaneous Raman spectroscopy setup for probing ultrafast collective excitation and quasiparticle dynamics in quantum materials. *Structural Dynamics*, 5(4):044301, 2018.

[24] Jihi-An Yang, Nicholas Pellatz, Thomas Wolf, Rahul Nandkishore, and Dmitry Reznik. Ultrafast magnetic dynamics in insulating YBa$_2$Cu$_3$O$_{6.1}$ revealed by time resolved two-magnon Raman scattering. *Nature Communications*, 11(1):2548, May 2020.

[25] R. P. Saichu, I. Mahns, A. Goos, S. Binder, P. May, S. G. Singer, B. Schulz, A. Rusydi, J. Unterhinninghofen, D. Manske, P. Guptasarma, M. S. Williamsen, and M. Rübbelake. Two-component dynamics of the order parameter of high-temperature Bi$_2$Sr$_2$CaCu$_2$O$_{8+}$ superconductors revealed by time-resolved raman scattering. *Phys. Rev. Lett.*, 102:177004, Apr 2009.

[26] Baruch Rosenstein and B. Ya. Shapiro. Apical oxygen vibrations dominant role in cuprate superconductivity and its interplay with spin fluctuations, 2020.

[27] Y. Y. Peng, G. Dellea, M. Minola, M. Conni, A. Amorese, D. Di Castro, G. M. De Luca, K. Kummer, M. Saluzzo, X. Sun, X. J. Zhou, G. Balestrino, M. Le Tacon, B. Keimer, L. Braicovich, N. B. Brookes, and G. Ghiringhelli. Influence of apical oxygen on the extent of in-plane exchange interaction in cuprate superconductors. *Nature Physics*, 13(12):1201–1206, Dec 2017.

[28] Marios H. Michael, Alex von Hoegen, Michael Förster, Andrea Cavalleri, and Eugene Demler. Parametric resonance of Josephson plasma waves: A theory for optically amplified interlayer superconductivity in YBa$_2$Cu$_3$O$_{6.5}$, 2020.

[29] M. F. Limonov, A. I. Rykov, S. Tajima, and A. Yamanaka. Raman scattering in YBa$_2$Cu$_3$O$_7$ single crystals: Anisotropy in normal and superconductivity states. *Physics of the Solid State*, 40(3):367–376, March 1998.

[30] E. Altendorf, J. Chrzanowski, J. C. Irwin, A. O’Reilly, and W. N. Hardy. Electron-phonon interactions of Raman active phonons in YBa$_2$Cu$_3$O$_{6+y}$. *Physica C: Superconductivity*, 175(1):47 – 57, 1991.

[31] K. F. McCarty, H. B. Radousky, J. Z. Liu, and R. N. Shelton. Temperature dependence of the linewidths of the Raman-active phonons of YBa$_2$Cu$_3$O$_7$: Evidence for a superconducting gap between 440 and 500 cm$^{-1}$. *Phys. Rev. B*, 43:13751–13754, Jun 1991.

[32] O Gunnarsson and O Rösch. Interplay between electron-phonon and coulomb interactions in cuprates. *Journal of Physics: Condensed Matter*, 20(4):043201, Jan 2008.

[33] S. Johnston, F. Vernay, B. Moritz, Z.-X. Shen, N. Nagosa, J. Zaanen, and T. P. Devereaux. Systematic study of electron-phonon coupling to oxygen modes across the cuprates. *Phys. Rev. B*, 82:064513, Aug 2010.
[34] A.S. Alexandrov and P.E. Kornilovitch. High-temperature superconductivity and charge segregation in a model with strong long-range electron-phonon and Coulomb interactions. *Physics Letters A*, 299(5):650 – 655, 2002.

[35] Claus Falter, Michael Klenner, and Georg A. Hoffmann. Phonon renormalization and c-axis phonon-plasmon mixing in \(\text{La}_2\text{CuO}_4\). *Phys. Rev. B*, 52:3702–3710, Aug 1995.

[36] C. Falter, M. Klenner, and W. Ludwig. Effect of charge fluctuations on the phonon dispersion and electron-phonon interaction in \(\text{La}_2\text{CuO}_4\). *Phys. Rev. B*, 47:5390–5404, Mar 1993.

[37] O. Rösch, O. Gunnarsson, X. J. Zhou, T. Yoshida, T. Sasagawa, A. Fujimori, Z. Hussain, Z-X. Shen, and S. Uchida. Polaronic behavior of undoped high-\(T_c\) cuprate superconductors from angle-resolved photoemission spectra. *Phys. Rev. Lett.*, 95:227002, Nov 2005.

[38] AF Kemper, O Abdurazakov, and JK Freericks. General principles for the nonequilibrium relaxation of populations in quantum materials. *Physical Review X*, 8(4):041009, 2018.

[39] J. Z. Wu, F. Y. Hsieh, A. V. McGuire, D. L. Schmidt, L. T. Wood, Y. Shen, and W. K. Chu. Anisotropic properties of the high-quality epitaxial \(\text{YBa}_2\text{Cu}_3\text{O}_{7-\delta}\) thin film. *Phys. Rev. B*, 44:12463–12646, Dec 1991.

[40] M. S. Raven, E. E. Inameti, S. Iwama, Y. M. Wan, and B. G. Murray. Epitaxial growth and critical currents in \(\text{YBa}_2\text{Cu}_3\text{O}_{7-\delta}\) films. *Phys. Rev. B*, 52:6845–6853, Sep 1995.

[41] Supplementary Information contains experimental, calculation and fitting details.

[42] C. Ambrosch-Draxl, H. Auer, R. Kourba, E. Ya. Sherman, P. Knoll, and M. Mayer. Raman scattering in \(\text{YBa}_2\text{Cu}_3\text{O}_7\): A comprehensive theoretical study in comparison with experiments. *Phys. Rev. B*, 65:064501, Jan 2002.

[43] O. V. Misochko, E. I. Rashba, E. Ya. Sherman, and V. B Timofeev. On the mixing of vibrational modes in high-\(T_c\) superconductors. *Physics Reports*, 194(5):387–395, 1990.

[44] D. Reznik, M. V. Klein, W. C. Lee, D. M. Ginsberg, and S-W. Cheong. Effect of conduction electrons on the polarized Raman spectra of copper oxide superconductors. *Phys. Rev. B*, 46:11725–11729, Nov 1992.

[45] S. L. Cooper, D. Reznik, A. Kotz, M. A. Karlow, R. Liu, M. V. Klein, W. C. Lee, J. Giapintzakis, D. M. Ginsberg, B. W. Veal, and A. P. Paulikas. Optical studies of the a-, b-, and c-axis charge dynamics in \(\text{YBa}_2\text{Cu}_3\text{O}_{6+\delta}\). *Phys. Rev. B*, 52:8233–8248, Apr 1993.

[46] L. Perfetti, P. A. Loukakos, M. Lisowski, U. Bovensiepen, H. Eisaki, and M. Wolf. Ultrafast electron relaxation in superconducting \(\text{Bi}_2\text{Sr}_2\text{Ca}_x\text{Cu}_{2}\text{O}_{8+\delta}\) by time-resolved photoelectron spectroscopy. *Phys. Rev. Lett.*, 99:197001, Nov 2007.

[47] Y Ishida, T Togashi, K Yamamoto, M Tanaka, T Taniguchi, T Kiss, M Nakajima, T Suzuki, and S Shin. Nonthermal hot electrons ultrafastly generating hot optical phonons in graphite. *Scientific reports*, 1:64, 2011.

[48] Markus Breusing, Claus Ropers, and Thomas Elsaesser. Ultrafast carrier dynamics in graphite. *Phys. Rev. Lett.*, 102:086809, Feb 2009.

[49] J. C. Johanssen, S. Ulstrup, F. Cilento, A. Crepaldi, M. Zacchigna, C. Cacho, I. C. Edmond Turcu, E Springate, F. Fromm, C. Raidel, T. Seyller, F. Parmigiani, M. Grioni, and P. Hofmann. Direct view of hot carrier dynamics in graphene. *Phys. Rev. Lett.*, 111:027403, Jul 2013.

[50] Tobias Kampfrath, Luca Perfetti, Florian Schapper, Christian Frischkorn, and Martin Wolf. Strongly coupled optical phonons in the ultrafast dynamics of the electronic energy and current relaxation in graphite. *Phys. Rev. Lett.*, 95:187403, Oct 2005.

[51] J. W. Loram, K. A. Mirza, J. R. Cooper, and W. Y. Liang. Electronic specific heat of \(\text{YBa}_2\text{Cu}_3\text{O}_{6+\delta}\) from 1.8 to 300 k. *Phys. Rev. Lett.*, 71:1740–1743, Sep 1993.

[52] -P Bohnen, R Heid, and M Krauss. Phonon dispersion and electron-phonon interaction for \(\text{YBa}_2\text{Cu}_3\text{O}_y\) from first-principles calculations. *Europhysics Letters (EPL)*, 64(1):104–110, oct 2003.

[53] W. S. Fann, R. Storz, H. W. K. Tom, and J. Bokor. Direct measurement of nonequilibrium electron-energy distributions in subpicosecond laser-heated gold films. *Phys. Rev. Lett.*, 68:2834–2837, May 1992.

[54] Ulrike Ritzmann, Peter M. Oppeneer, and Pablo Maldonado. Theory of out-of-equilibrium electron and phonon dynamics in metals after femtosecond laser excitation. *Phys. Rev. B*, 102:214305, Dec 2020.

[55] D. Reznik. Phonon anomalies and dynamic stripes. *Physica C: Superconductivity*, 481:75 – 92, 2012.

[56] M. Raichle, D. Reznik, D. Lamago, R. Heid, Y. Li, M. Bakr, C. Ulrich, V. Hinkov, K. Hradil, C. T. Lin, and B. Keimer. Highly anisotropic anomaly in the dispersion of the copper-oxygen bond-bending phonon in superconducting \(\text{YBa}_2\text{Cu}_3\text{O}_7\) from inelastic neutron scattering. *Phys. Rev. Lett.*, 107:177004, Oct 2011.

[57] Eva A. A. Pogna, Stefano Dal Conte, Giancarlo Soavi, Vasyl G. Kravets, Yong-Jin Kim, Stefano Longhi, Alexander N. Grigorenko, Giulio Cerullo, and Giuseppe Della Valle. Ultrafast spectroscopy of graphene-protected thin copper films. *ACS Photonics*, 3(8):1508–1516, 2016.

[58] Manuel Obergfell and Jure Demsar. Tracking the time evolution of the electron distribution function in copper by femtosecond broadband optical spectroscopy. *Phys. Rev. Lett.*, 124:037401, Jan 2020.

[59] D. Lamago, M. Hoesch, M. Krisch, R. Heid, K-P. Bohnen, P. Böni, and D. Reznik. Measurement of strong phonon softening in Cr with and without fermi-surface nesting by inelastic x-ray scattering. *Phys. Rev. B*, 82:195121, Nov 2010.

[60] O. Abdurazakov, D. Nevola, A. Rustagi, J. K. Freericks, D. B. Dougherty, and A. F. Kemper. Nonequilibrium electron dynamics in pump-probe spectroscopy: Role of excited phonon populations. *Phys. Rev. B*, 98:245110, Dec 2018.

[61] G. Coslovich, B. Huber, W.-S. Lee, Y.-D. Chuang, Y. Zhu, T. Sasagawa, Z. Hussain, H. A. Bechtel, M. C. Martin, Z.-X. Shen, R. W. Schoenlein, and R. A. Kaindl. Ultrafast charge localization in a stripe-phase nickelate. *Nature Communications*, 4(1):2643, 2013.

[62] MR Norman. Linear response theory and the universal nature of the magnetic excitation spectrum of the cuprates. *Physical Review B*, 75(18):184514, 2007.

[63] JD Rameau, S Freutel, AF Kemper, Michael A Sentef, JK Freericks, I Avigo, M Ligges, Laurenz Rettig, Y Yoshida, H Eisaki, et al. Energy dissipation from a correlated system driven out of equilibrium. *Nature communications*, 7(1):1–9, 2016.

[64] Tatiana Konstantinova, Jonathan D Rameau, Alexander H Reid, Omadillo Abdurazakov, Lijun Wu, Renkai G. Coslovich, B. Huber, W.-S. Lee, Y.-D. Chuang, Y. Zhu, T. Sasagawa, Z. Hussain, H. A. Bechtel, M. C. Martin, Z.-X. Shen, R. W. Schoenlein, and R. A. Kaindl. Ultrafast charge localization in a stripe-phase nickelate. *Nature Communications*, 4(1):2643, 2013.

[65] MR Norman. Linear response theory and the universal nature of the magnetic excitation spectrum of the cuprates. *Physical Review B*, 75(18):184514, 2007.
Li, Xiaozhe Shen, Genda Gu, Yuan Huang, Laurenz Ret-tig, et al. Nonequilibrium electron and lattice dynamics of strongly correlated Bi$_2$Sr$_2$CaCu$_2$O$_{8+\delta}$ single crystals. Science advances, 4(4):eaap7427, 2018.

[65] C. R. Hunt, D. Nicoletti, S. Kaiser, D. Pröpfer, T. Loew, J. Porras, B. Keimer, and A. Cavalleri. Dynamical decoherence of the light induced interlayer coupling in YBa$_2$Cu$_3$O$_{6+\delta}$. Phys. Rev. B, 94:224303, Dec 2016.

[66] S. Kaiser, C. R. Hunt, D. Nicoletti, W. Hu, I. Gierz, H. Y. Liu, M. Le Tacon, T. Loew, D. Haug, B. Keimer, and A. Cavalleri. Optically induced coherent transport far above $T_c$ in underdoped YBa$_2$Cu$_3$O$_{6+\delta}$. Phys. Rev. B, 89:184516, May 2014.