Monitoring ultrafast metallization in LaCoO₃ with femtosecond soft x-ray spectroscopy

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The study of ultrafast dynamics is a new tool to understand and control the properties of correlated oxides. By enhancing some properties and realizing new dynamically excited phrases, this tool has opened new routes for technological applications. LaCoO₃ is one paradigmatic example where the strong electron, spin, and lattice coupling induced by electronic correlations results in a low-temperature spin transition and a high-temperature semiconductor-to-metal transition that is still not completely understood. Here, we monitor ultrafast metallization in LaCoO₃ using time-resolved soft x-ray reflectivity experiments. While the process is entangled at the Co L₃ edge, the time information of the different channels is decrypted at different resonant energies of the O K edge. Metallization is shown to occur via transient electronic, spin, and lattice separation. Our results agree with the thermodynamical model and demonstrate the potential of femtosecond soft x-ray experiments at the O K edge to understand correlated oxides.

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The phase diagram of LaCoO$_3$ (LCO) as a function of temperature with its low-temperature (~100 K) spin transition (ST) and a high-temperature (300–600 K) broad semiconductor-to-metal transition (SMT) has been derived over the years through detailed analysis of magnetic susceptibility, resistivity, atomic structure, phonon spectrum, and heat capacity experiments$^{1-12}$. The complex behavior of the system with temperature is thought to be the consequence of the strong interplay between charge, spin, and lattice degrees of freedom. The change in spin required to explain the magnetic susceptibility anomalies$^{13}$ is accompanied by an increase of the ionic radius$^8$, which, in turn, induces an anomalous lattice expansion responsible for the SMT$^{12}$. Although the spin state has been considered the driving mechanism responsible for all changes in the system with temperature$^{6,12-15}$, the ultimate understanding of the system has remained elusive due to the lack of consensus regarding the strength of electronic correlations. As a result, a longstanding debate has split the community around two models. The first, derived empirically by Goodenough$^{13}$ from structural and transport measurements, proposed that the large $t_{2g}$ of the Co $d^6$ atom in an octahedral oxygen environment, where the $t_{2g}$ and $e_g$ orbitals are split by the crystal field $\Delta$. The ST and the SMT were interpreted as a gradual temperature transition from a low spin (LS) $[t_{2g}^3, e_g^\ast, S = 0]$ to a high spin (HS) $[t_{2g}^3, e_g, S = 2]$ and from an HS to an intermediate spin state (IS) $[t_{2g}^6, e_g^\ast, S = 1]$ state, respectively. The increase of the ionic radius from $r_{LS} = 0.545$ Å to $r_{HS} = 0.610$ Å$^8$ would explain the anomalous lattice expansion below room temperature (ST) and the increase of the $e_g$–O$_{2p}$ overlap involving metalization above room temperature (SMT). Theory refinements to the ionic picture included the trigonal distortion of the R3c structure and the spin–orbit coupling. The resulting energy diagram has the first two states around the Fermi energy ($E_F$) separated from each other by 11 meV (corresponds to 100 K), the ground state being non-magnetic, while the first excited state has a moment of $m = 2.32$ $\mu_B$.$^{12}$

The second model emerged theoretically in an attempt to include electronic correlations and Co–O hybridization within the density function theory (DFT) with large on-site Coulomb interaction, $U$, DFT+$U$ formalism. Using $U = 7.8$ eV, the IS state appears at lower energy than the HS state, thus explaining the ST as a LS–IS one$^{14}$. According to this model, the system would also exhibit orbital ordering to prevent the metallic ground state as a consequence of the large $U$ value, as shown by local spin density approximation (LSDA)+$U$ calculations$^{16}$. A semiconducting ground state with a gap close to the experimental one requires lower $U$ values, $U \sim 3$ eV or $U \sim 6$ eV$^{16}$. Following this route, combined cluster calculations with $U = 5.5$ eV and experiments seem to have balanced the discussion about the ST in favor of the LS–HS crossover with a gradual increase of the HS population with temperature$^{15}$. This result has been supported by advanced DFT including correlations within the dynamical mean-field theory (DMFT), DFT+DMFT$^{17-19}$. Furthermore, dynamical charge fluctuations favor short-range order at room temperature in a configuration where the two cobalt (Co) atoms in the unit cell connected by an oxygen (O) atom are in a different spin state, Co(LS)–O–Co(HS)$^{19}$. This configuration was shown to be favored by the charge imbalance that spontaneously appears in the system when the constraint of equal charge between the two Co atoms in a unit cell is relaxed in calculations with moderate $U \sim 3$ eV$^{20}$. The SMT is explained naturally as a transition from the room temperature Co(LS)–O–Co(HS) state to the HS metallic state at high temperature. (Recent resonant inelastic x-ray experiments and theoretical calculations have proposed a new scenario in which mobile IS excitons propagating in a LS background contribute to the low-temperature physics of the system, see refs. $^{21,22}$.) As an attempt to verify the role of charge and spin fluctuations in the properties of LCO, soft x-ray time-resolved spectroscopic studies with picosecond resolution across the SMT were performed$^{23}$. The high-temperature phase transition was chosen due to its technological relevance in numerous environmentally friendly energy production domains$^{24-31}$. Laser excitation with 800 nm photons allowed the realization of a transient metallic state with Co charge disproportionation and a higher spin state than the thermally stabilized one. This observation was deduced by comparing the evolution with temperature and time delay of soft x-ray reflectivity measurements at the O K and Co L edges$^{23}$.

In this contribution, the SMT in LCO is addressed by means of time-resolved optical pump—soft x-ray probe experiments with femtosecond resolution and DFT+DMFT calculations of the optical reflectivity. Calculations of the measured time-resolved soft x-ray spectra including electronic correlations and lattice dynamics are still not theoretically available. They would require a multi-orbital Schwinger–Keldysh–formalism for interacting electron systems, while only model single-band attempts exist$^{32}$. There are attempts to calculate realistic dynamical core–hole screening in the x-ray absorption spectra, but only in non-correlated DFT approach$^{33}$. Since LCO is definitely correlated system, we did a best possible theoretical DFT+DMFT analysis trying to understand the gap collapse, the charge disproportionation, enhancement of the spin and lattice deformations upon laser excitation. The combined study has allowed the understanding of the properties of the laser-induced transient metallic state disentangling the interplay between the electronic, spin, and lattice degrees of freedom upon metallization. The 800 nm optical laser excitation depopulates electrons from the pseudo-$t_{2g}$ orbitals into the pseudo-$e_g$ ones (see Methods). As a consequence, charge transfer between the pseudo-$e_g$ and the O–$2p$ orbitals is enhanced and the HS state population increases. The larger ionic radius of the HS state induces a lattice distortion with a deformation of the bond angles and a change of interatomic distances, explaining the metallic character of LCO at high temperature. The emerging picture presents the transient metallization as a four-step process in a way similar to the ionic model proposed by Goodenough$^{8}$ to explain the evolution of LCO with temperature. Thus, our studies become the first experimental demonstration of Goodenough metallization model$^{6,13}$. Insight into the electronic properties of the laser-excited transient metallic state and the optical excitation mechanism have been derived from DFT+DMFT calculations of the optical conductivity.

**Results**

**Femtosecond time-resolved reflectivity.** Optical pump—soft x-ray probe reflectivity experiments with femtosecond time resolution were performed at room temperature using the FemtoSpeX-scattering experimental station at the storage ring-based fs-slicing source FemtoSpeX of the BESSY II synchrotron at Helmholtz Zentrum Berlin (HZB). The two beamlines, the PGM beamline ($E/\Delta E > 5000$) and the ZPM beamline ($E/\Delta E \sim 500$) of the FemtoSpeX facility, have been used to record laser-induced changes at the O K and the Co L$_2$ edges with 50 ps and 120 fs time resolution, respectively. Despite the reduced energy resolution of the ZPM beamline, all spectral features seen at the PGM beamline can clearly be assigned to the ZPM spectra (see Supplementary Fig. 1). The dynamics of LCO was characterized by measuring time-delay scans at the photon energies resonant with the relevant spectroscopic signatures of the SMT transition$^{12,15,21}$. At the
O K edge, we have focused on the resonant energies probing oxygen hybridized with different Co orbitals, namely the pseudo-$t_{2g}$ orbitals, the pseudo-$e_g$ orbitals, and the Co-4sp. From the measured spectra, the sum O–Co-pseudo-$e_g$ + O–Co-pseudo-$t_{2g}$ and difference O–Co-pseudo-$e_g$–O–Co-pseudo-$t_{2g}$ were calculated to understand the underlying physics in the system. The spectra are depicted in Fig. 1: O–Co-pseudo-$t_{2g}$ (Fig. 1a), O–Co-pseudo-$e_g$ (Fig. 1b), sum spectra (Fig. 1c), O–Co-4sp (Fig. 1d), and difference spectra (Fig. 1e). At the Co L3 edge, both the white line (WL) and its high-energy shoulder (HSS) ascribed to the HS state population change (Co-HS) have been investigated (Fig. 1f). The data show intensity variations taking place at different time delays, depending on the different resonant energies. The effect can be clearly seen when plotting all the measured data together (see Supplementary Fig. 2). This observation provides clear evidence that the relaxation of the excited states occurs at different time scales for different resonant energies.

**Time constants.** To determine the different time scales observed at different resonant energies, a fitting to all the delay scans in Fig. 1 was done. In the first step, a modified three-temperature model (3TM), previously applied to isostructural manganites, was used. However, the $\chi^2$ value of the fits could be improved by a factor of three by allowing the exponential relaxation of a given component to be delayed by the time constant of the previous one. This cascade decay implies that each relaxation channel becomes the source of the next (see Methods). A fit to all the delay scans is possible with five time constants. For some delay scans, only two components were necessary. The time constants were obtained from a self-consistent fit of all delay scans. The fifth time constant $\tau_5$ was used to account for the slow relaxation of the system observed in the picosecond experiments. The time constant values obtained from the self-consistent fit are $\tau_1 = 0.17 \text{ ps, } \tau_2 = 0.52 \text{ ps, } \tau_3 = 0.91 \text{ ps, } \tau_4 = 1.5 \text{ ps, and } \tau_5 = 45 \text{ ps.}$ The fit laser pulse duration was 0.125 ps, which matches the 0.13 ps laser pulse duration measured experimentally. The final fit curves are displayed in Fig. 1a–f as solid lines, whereas the individual components are displayed as filled curves. The characteristic time scales contributing to each spectra are indicated in the inset labels.

The delay scan at the O K edge in the O–Co-pseudo-$t_{2g}$ region (Fig. 1a) can be fitted with the time constants $\tau_1 = 0.17 \text{ ps, } \tau_2 = 0.52 \text{ ps, and } \tau_4 = 1.5 \text{ ps.}$ The first two components have the dominating intensity corresponding to the sub-picosecond intensity peak. This first increase of the reflectivity corresponds to the initial depopulation of the O–Co-pseudo-$t_{2g}$ orbitals upon laser excitation. It is followed by a decay, which stabilizes around $\tau_4 = 1.5 \text{ ps.}$ The best fit of the time delay scan at the O–Co-pseudo-$e_g$ resonant energy (Fig. 1b) was obtained with the ultrashort components $\tau_1 = 0.17 \text{ ps, } \tau_3 = 0.91 \text{ ps, and } \tau_4 = 1.50 \text{ ps}$ and the slow relaxation channel, $\tau_5 = 45 \text{ ps.}$ The shortest time delay fits the first drop of the reflectivity and corresponds to the population of the O–Co-pseudo-$e_g$ orbitals. It is the counterpart of the depopulation of the O–Co-pseudo-$t_{2g}$ orbitals observed in Fig. 1a. After a very weak recovery, a second broad minimum at around $\tau_4 = 1.50 \text{ ps can be observed.}$ It is much more pronounced than in the delay scan measured in the O–Co-pseudo-$t_{2g}$ region. The fact that the initial reflectivity value does not recover after 3 ps is due to the contribution of the slow relaxation channel, $\tau_5 = 45 \text{ ps,}$ obtained from picosecond studies. The reflectivity changes at the shortest time scale, $\tau_1$, indicate a perfect redistribution of the population between the O–Co-pseudo-$t_{2g}$ and O–Co-pseudo-$e_g$ orbitals, as required to increase the HS of the system. At larger time scales, the reflectivity changes at the two delay scans no longer compensate. The need for different time constants and intensities to fit each scan indicates that other channels besides electron redistribution...
contribute to the changes in reflectivity at these resonant energies. Indeed, the [O–Co–pseudo-ε_g+O–Co–pseudo-t_{2g}] sum spectra (Fig. 1c) shows that the intensity cancels out for time delays below ~0.5 ps. From larger time delays, the intensity continues decreasing until it reaches a minimum around the time delay constant, \( t_4 = 1.50 \) ps, followed by a slow recovery. A good fit of the sum spectra can be done with only \( t_4 = 1.50 \) ps and \( t_5 = 45 \) ps components. The same time constants are required to fit the delay scan measured at the O–Co–4sp resonant energy (Fig. 1d). As in the sum delay spectrum, the ultrafast excitation below 0.20 ps is hardly visible. The fact that changes at the O–Co–4sp resonant energy is provided to indicate the expected bond angle in the excited state. The lattice structures, rhomboedral R3̅m, a = 1.55 eV, \( \epsilon_g \) and \( \epsilon_{2g} \), excites the sample, producing carriers and instantaneous gap collapse. (II) Charge transfer enhancement: After carrier relaxation, the charge drop at short time delays \( \tau_4 \) at the O–Co–3d pseudopotential energy related to modifications in bond length and angles \( 12,23 \) indicates that the time delay \( t_4 \) is related to structural lattice modifications.

In Fig. 1e, the [O–Co–pseudo-ε_g–O–Co–pseudo-t_2g] difference delay scan is displayed. The intensity changes as a function of time delay reproduce quite well those measured at the Co L_3 WL and HSS resonant energies (Fig. 1f). Indeed, the same time constants \( t_1 = 0.17 \) ps, \( t_5 = 0.91 \) ps, \( t_4 = 1.50 \) ps, and \( t_5 = 45 \) ps with the same intensities are able to fit all delay scans. The first drop at short time delays \( \tau_5 = 0.17 \) ps can be related to bleaching resulting from ultrafast carrier excitation.\(^{35} \) Afterwards, a second broad minimum with almost the same intensity reduction is observed around the time delay \( \tau_5 = 0.91 \) ps. The intensity of this time constant is much larger than that observed at the O–Co–pseudo-\( \epsilon_g \), indicating a relation with the increase of the HS population in the system. From there on, the reflectivity recovers slowly to \( \lambda \) of the initial drop after 2.5 ps. To fit this part of the reflectivity, components with time delays \( t_2 = 1.50 \) ps and \( t_5 = 45 \) ps are required. The evolution at short time delays is in agreement with recent results obtained for ferromagnetic LCO thin films.\(^{36} \) Moreover, the saturation-like behavior after three picoseconds reproduces the results measured with picosecond resolution.\(^{23} \) The decrease of intensity with time delay can be explained by the increase of the HS population upon laser excitation that is enhanced by charge disproportionation as deduced previously.\(^{23} \)

The similarity of the [O–Co–pseudo-\( \epsilon_g \)–O–Co–pseudo-t_2g] difference delay scan with the delay scans measured at the Co L_3 edge and that of the [O–Co–pseudo-\( \epsilon_g \) + O–Co–pseudo-t_2g] difference delay scan with the delays scan at the O–Co–4sp resonant energy can be understood by assuming that the scans at the O–Co–pseudo-t_2g and at the O–Co–pseudo-\( \epsilon_g \) can be decomposed in five contributions: orbital electron transfer [OT], charge transfer enhancement [CE], spin stabilization [SS], lattice modification [LM], and slow relaxation [SR]. Assuming O–Co–pseudo-t_2g \( \propto \) [OT + CE – SS – LM] and O–Co–pseudo-\( \epsilon_g \) \( \propto \) [–OT + CE – SS + LM + SR], we obtain O–Co–pseudo-\( \epsilon_g \) + O–Co–pseudo-t_2g \( \propto \) [CE + LM + SR] and O–Co–pseudo-\( \epsilon_g \) – O–Co–pseudo-t_2g \( \propto \) [–OT – SS + SR]. The proposed decomposition agrees with the five time constants required to fit the different delay scans and allows the following assignment: \( \tau_4 \) corresponds to the orbital electron transfer contribution, OT; \( \tau_2 \) to the charge transfer enhancement, CE; \( \tau_5 \) is related to the spin stabilization, SS; and \( \tau_4 \) to the lattice modification, LM. The last constant indicates the slow relaxation, SR. All together, the presence of different time scales points toward an ultrafast metallization mechanism in which the charge, spin, and lattice degrees of freedom evolve with distinct time scales. For the charge channel, the analysis is able to distinguish between the redistribution of electrons between different orbitals and the change in hybridization.

**Metallization model.** The understanding of the mechanism leading to an ultrafast time separation of the charge, spin, and lattice is sketched in Fig. 2. It can be formulated in four steps, indicated by panels I–IV. For each panel the top figures, Fig. 2a, c,
the changes in the DOS obtained from the DFT and their evolution upon laser excitation. Figure 2b, d, f, h display carrier excitation as it has been recently observed in other oxides, like V2O3. Since the semiconducting state, is excited by the femtosecond pulse, the intensity indicative of bleaching. The hot carriers have much energy resulting in an ill-defined out-of-equilibrium spin state. This process, panel II—Fig. 2c, d, takes places after carrier thermalization. It corresponds to the increased change transfer in the system. Its characteristic time scale is \( \tau_2 = 0.52 \) ps and can be deduced from the fact that the reflectivity in the [O–Co-pseudo-\( e_g \) + O–Co-pseudo-\( t_{2g} \)] sum spectrum (Fig. 1c) does not cancel out for time delays larger than \( \tau_2 \). This effect can be understood by the larger hybridization of the O–Co-pseudo-\( e_g \) orbitals pointing towards the O-2p orbitals.

Spin stabilization. The third step, panel III—Fig. 2e, f, corresponds to the stabilization of the spin system and is represented by the time scale \( \tau_s = 0.91 \) ps. The process starts after the breaking of the population balance around a time delay of 0.52 ps and stabilizes at around a time delay of 0.91 ps. In the new spin configuration, both Co-excited atoms will be in a charge-disproportionated HS state, as determined from the comparison to the thermally excited state. To get insight into the nature of the disproportionated HS state, DFT+DMFT calculations of the optical conductivity have been performed following the same approach as in Karolak et al. and benchmarked with the experimental results referenced therein. The equilibrium state of the system at room temperature is a 50% incoherent mixture of LS and HS states located on both sides of the oxygen atom—Co(LS)–O–Co(HS)—with no long-range order. This configuration is favored by the charge imbalance that appears in the system when the constraint of equal charge between the two Co atoms in a unit cell is relaxed. It also explains the semi-conducting paramagnetic state of the system and the absence of orbital ordering at room temperature. The charge imbalance induces strong modifications of the optical conductivity in the region below 2 eV (Fig. 3a). At the laser excitation energy used in the experiments, 1.55 eV (800 nm), the optical conductivities of the LS and HS states are very similar to each other. This contrast with the calculations obtained for pure LS and HS states that result in a weaker excitation of the LS state compared to the excitation for the HS state. The different conductivities can be understood by taking into account the contribution of \( d^6 \), \( S = 1/2 \) states to the LS state, and of \( d^7 \), \( S = 3/2 \) states to the HS configuration. Both contributions result in a large modification of the density of states (DOS) and a reduction of the gap compared to the scenario of LS and HS in a pure \( d^6 \) electronic configuration (Fig. 3b, d). In the pure LS state, the large gap, slightly larger than the laser photon energy, explains the very low intraband transition probability. In the case of the HS state, the gap is much smaller and therefore the transition is more efficient and the conductivity much higher. In the case of the Co(LS)–O–Co(HS) configuration, the small equilibrium charge disproportionation makes possible both LS and HS intraband transitions. Furthermore, interband transitions between the two atoms will also be allowed. The ones from occupied HS states to unoccupied LS states will have higher probably for two reasons: the smaller energy gap between these states, on the one hand, and the Pauli exclusion principle, on the other. The latter will make all transitions from occupied HS O–Co-pseudo-\( t_{2g} \) to empty LS O–Co-pseudo-\( e_g \) possible, whereas only Pauli-satisfying transitions will be available from the occupied LS O–Co-pseudo-\( e_g \) states to the partially empty HS O–Co-pseudo-\( t_{2g} \) orbitals.

As a consequence, the laser-excited state is expected to have a transient metallic state in which the change disproportionation is enhanced by the laser excitation. Furthermore, the spin state of the excited fraction of the system will have the two atoms in a HS state. This fact, together with the increase of charge disproportionation, explains the differences in the thermally induced metallization of the system. This conclusion is expected to hold...
even assuming any of the other initial state models considered for the system\textsuperscript{12}. An alternative explanation to the spin change could be done in terms of laser-induced correlation enhancement\textsuperscript{35}. In this case, the increase of $U$ could result in an excited metallic IS state similar to that proposed initially by LSDA+$U$ calculations\textsuperscript{14}.

**Lattice modification.** The last step of the process, panel IV—Fig. 2g, h, takes places at around $t_1 = 1.50 \text{ ps}$. It corresponds to the stabilization of the metallic state through the structural lattice relaxation. This occurs due to the change in hybridization and the increase of the ionic radius of Co, a consequence of the higher HS content and disproportionation of the system. In Fig. 2—IV, atoms 1 and 2 have $d$ occupancies of 5.83 and 6.17, respectively, and are both in a HS state. This state will be long living, therefore explaining the slow relaxation constant of 45 ps. The ionic radius of the IS and HS states in a pure $d^6$ configuration are $r_{1,5} = 0.545 \text{ Å}$ and $r_{\text{HS}} = 0.610 \text{ Å}$. Increasing the HS percentage will result in a lattice change consequence of the CoO$_x$ distortion related to the higher ionic radius. This effect will be enhanced, taking into account that the increase of the charge disproportionation implies an increase of the $d^5 (\text{Co}^{2+})$ and $d^7 (\text{Co}^{2+})$ population. The ionic radius of the HS Co$^{4+}$ and Co$^{2+}$ states are $r_{\text{HS},\text{4+}} = 0.530 \text{ Å}$ and $r_{\text{HS},\text{2+}} = 0.745 \text{ Å}$, respectively\textsuperscript{40}. As a consequence, an increase of the ionic radius will result in a lattice deformation, both bond angles and lengths, therefore explaining the dynamics observed at the Co-4p region. In Fig. 2—IV, the bond angle of the thermally activated phase at 850 K is provided to indicate the expected increase in bond angle required to stabilize metallization. The change in both length and angle is expected to improve the hybridization with the O 2$p$ orbitals. This will further enhance the charge transfer and will contribute to the stabilization of the metallic excited state of the system. A similar interpretation can be done assuming that enhanced correlations stabilize the IS state spin configuration since the radius is larger and some other lattice deformations, like Jahn–Teller distortions, can be stabilized\textsuperscript{14}.

The slow relaxation constant required to fit the data, $t_2 = 45 \text{ ps}$, is related to laser-induced excitation and how the laser-induced transient metallic system goes back to the equilibrium state. Therefore, it does not need to be considered as part of the ultrafast metallization process.

**Relation to static phonon experiments.** The time scales extracted from the reflectivity data are coherent with recent modifications observed in some phonon modes with inelastic X-ray scattering (IXS)\textsuperscript{41}. More concretely, the $E_g$ $A_{2g}$ phonon modes at 24 meV corresponding to La vibration have shown to soften upon increasing temperature. This phonon energy corresponds to the fastest time scale of 0.17 ps observed in the experiments reported here. The softening of the phonon mode is coherent with the depopulation of the O–Co–pseudo-$t_{2g}$ orbitals pointing towards the La atoms and also with the increase of the ionic radius of the HS state. The second phonon mode that has shown modifications upon increasing temperature across the semiconductor to metal transition is the $E_p$ mode. This mode, corresponding to a rotational mode of O atoms (or a vibration of the CoO$_x$ octahedra), hardens upon increasing the temperature. This hardening is coherent with the increase of the oscillation amplitude of the oxygen atoms perpendicular to the Co–O recently reported and also with the increase on the ionic radius of the Co atoms as the HS spin population of the system increases\textsuperscript{41}. The energy of this mode, 5 meV, corresponds to a time scale of 0.827 ps. This time scale matches the 0.91 ps time scale ascribed to the increase of the HS state and to the change of ~50% in the population unbalance, a consequence of the enhanced hybridization.

The proposed ultrafast metallization model of LCO can be understood by taking into account that the structure is naturally strained by the large size of the La$^{3+}$ ions, which forces a rhombohedral lattice, and by the large variations of the Co ion size with spin configuration. In this respect, the results prove that both the thermodynamical and the experimental DFT+DMFT calculations have been used to address the ultrafast metallization of LCO upon laser excitation. The latter shows intensity variations at different time delays for different resonant energies that allow extracting the relevant time constants with less uncertainty on the fitting procedure. Metallization stabilizes within less than 3 ps in a four-step process in which the increase of ionic radius accompanying the HS state stabilization after carrier thermalization modifies bond lengths in the CoO$_x$ octahedra and increases the Co–O–Co angle, as happens upon thermal excitation of the system. The experiments support the theoretical model within DFT+DMFT with moderate $U$ that favors correlation-induced charge and spin fluctuations. Our findings are the first experimental demonstration of the adiabatic thermal transition model in LCO widely accepted in the literature. Furthermore, the results indicate that a comprehensive analysis of the ultrafast dynamics of perovskites can be obtained by performing time-resolved experiments with soft X-rays at the O K edge. Most of the limits imposed by femtolasers experiments at synchrotrons—reduced time delay range and energy resolution—are overcome at x-ray free-electron lasers.

**Methods**

**d orbitals in a rhombohedral environment.** The atomic structure of LCO is rhombohedral (R3$\bar{m}$). This means that the description of the $d$ orbitals in terms of the octahedral $t_{2g}$, $e_g$ basis is not rigorous. Instead linear combinations of them are required. They will be referred to in the following as pseudo-$t_{2g}$ and pseudo-$e_g$.

**Soft x-ray time-resolved experiments.** Time-resolved soft x-ray reflectivity studies at room temperature were performed at the femtolasers beamline of the BESSY II synchrotron radiation source at the Humboldt-Zentrum Berlin (HZB)\textsuperscript{42–44}. Experiments were carried out in (001) terminated, 4 × 4 mm$^2$ single crystal cuts from a well-characterized large crystal\textsuperscript{45}. A ($\theta$, $2\theta$) = (5°, 10°) geometrical configuration was used for the experiments. The pump pulses were produced by a Ti:Sapphire laser at $\lambda = 800 \text{ nm}$ (1.55 eV), 6 kHz maximum repetition rate, 50 fs pulse duration, and 2 mJ maximum pulse energy. The x-ray pulse duration of about 10 fs in the femtolasers operation mode ensures a global time resolution of 130 fs\textsuperscript{42–44}. The probe soft x-ray femtolasers photons were produced using Bragg Fresnel reflection zone plate monochromators (ZPMs). One of the ZPMs was optimized to operate in the energy range of the O K edge, 2.21–2.36 nm (560.9–525.3 eV). The second one covered the Co L edge energy range, 1.54–1.60 nm (774.87–805.06 eV).

The penetration length of the excitation laser with $\lambda = 800 \text{ nm}$ is ~359 nm, as obtained from Nomerovannaya et al.\textsuperscript{46}. This value is larger than the penetration depth of the soft x-rays at the O K edge (<340 nm) and the Co L edge (<298 nm) obtained from the Center for X-ray optics (http://www.cxro.lbl.gov). The values indicate that the sampled area was fully excited by the pump pulse and therefore that an interpretation of the data based in the ultrasonic propagation of the metallic phase in a system where both semiconducting and metallic phases coexist seems unlikely\textsuperscript{47}.
Data fitting. The fit was done using a Levenberg–Marquardt algorithm to search for the minimum value of $\chi^2$ defined as: $\sum (y - y_\text{fit})^2/\sigma^2$. We initially model the double counting, optical retransitions. Since ED allows direct evaluation of the self-energy, we employ the parametrization via Slater integrals connected to the quantum Monte Carlo calculations shows that the spectra of both methods agree together was obtained. The laser pulse duration was estimated to be 0.125 ps, which methodology was shown in P. Wissgott et al. to work well for the low-frequency part of the optical conductivity, i.e. for the case of transition metal oxides elements of the momentum operator, and the corresponding hybridization $\Delta(\omega)$ with a finite cluster containing $t_k$ sites and its corresponding hybridization $\Delta(\omega_k)$ is:

$$\Delta(\omega_k) \approx \Delta(\omega_k) = \sum_k \frac{|V_{\text{mell}}|^2}{\omega - \omega_k - \xi_k}$$

The optimal set of bath parameters $\xi_k$ and $V_{\text{mell}}$ is found by fitting the full lattice hybridization function. We have employed a five-site cluster, which may appear to be a very crude approximation. However, comparison with continuous time quantum Monte Carlo calculations shows that the spectra of both methods agree very well. The reason is that the states of the Co ions are dominated very strongly by a few local atomic states that we include exactly in the ED and not so much by itinerant states that we approximate. Since ED allows direct evaluation of the self-energy matrix $\Sigma(\omega)$ on the real energy axis, we can quickly calculate the spectral matrices $A(\vec{k}, \omega)$, as needed for the evaluation of optical properties (see below). The calculation of the optical conductivity in DFT+DMFT requires the evaluation of the following expression:

$$\text{Re} \left( \sigma_{\alpha\beta}(\omega) \right) = \frac{1}{\pi} \sum_{\vec{k}} \text{Im} \left( \frac{1}{\omega - \varepsilon(\vec{k}, \omega)} \right) \times \text{Tr} \left[ A(\vec{k}, \omega) \sigma_{\alpha\beta}(\omega) A(\vec{k}, \omega) \right]$$

where $V$ is the unit cell volume, $f$ denotes the Fermi function, $\sigma_{\alpha\beta}(\vec{k}, \omega)$ are the matrix elements of the momentum operator, and $A(\vec{k}, \omega)$ is the momentum-resolved spectral matrix. For the quick calculation of the momentum-resolved spectra on any grid, we have used ED, as explained above. The remaining problem of the calculation of the momentum matrix elements is tackled using the improved Peierls approximation as implemented in the “woptic” package. This methodology was shown in P. Wissgott et al. to work well for the low-frequency part of the optical conductivity, i.e. for the case of transition metal oxides dominated by $d-d$ transitions. The package uses an adaptive $k$ sampling procedure, which calculates the necessary quantities on iteratively improved meshes in the Brillouin zone.

Data availability. All data generated or analyzed during this study are included in this published article (and its Supplementary Information files).

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