Correlations between electronic order and structural distortions and their ultrafast dynamics in the single-layer manganite Pr$_{0.5}$Ca$_{1.5}$MnO$_4$


1Swiss Light Source, Paul Scherrer Institute, 5232 Villigen PSI, Switzerland  
2Institute of Physics II, University of Cologne, 50937 Cologne, Germany  
3SwissFEL, Paul Scherrer Institute, 5232 Villigen PSI, Switzerland

(Received 5 December 2019; accepted 24 January 2020; published 18 February 2020)

Time-resolved x-ray diffraction experiments on the half-doped single-layered manganite Pr$_{0.5}$Ca$_{1.5}$MnO$_4$ are used to monitor the ultrafast photoinduced dynamics of the structural distortion associated with the charge and orbital ordering (CO/OO). As in the nonlayered three-dimensional counterpart, the ordered phase melts in less than 100 fs after 800-nm photoexcitation and subsequently partially recovers due to thermal equilibration of electronic and vibrational systems. Photoexciting Pr$_{0.5}$Ca$_{1.5}$MnO$_4$ below the transition temperature of a second structural phase transition that occurs around 146 K (deep inside the CO/OO phase) releases this structural transition, but progresses on a much slower timescale. This additional reduction of crystal symmetry, which we ascribe to a further tilt of the oxygen octahedra, can thus be considered to be only weakly coupled to CO/OO. Furthermore, static hard-x-ray and resonant soft-x-ray diffraction at the Mn$K_{\alpha}$ edges experiments identify correlations between structural distortions and electronic order in thermal equilibrium.

DOI: 10.1103/PhysRevB.101.075119

I. INTRODUCTION

Doped three-dimensional manganites have attracted considerable interest [1] as they exhibit metal-insulator transitions and colossal magnetoresistance effects. Strong coupling of lattice, spin, charge, and orbital degrees of freedom in mixed valence manganites leads to complex phase diagrams. In particular, the Jahn-Teller distortion (JTD) couples electronic charge and orbital order (CO/OO) with a symmetry breaking of the crystal lattice [1]. Information on the couplings can be obtained by studying the temperature dependence of the individual degrees of freedom. In manganites, structural distortions go hand in hand with electronic ordering phenomena, and which degree of freedom drives the CO/OO phase transition remains unknown. Resonant soft-x-ray diffraction of superlattice reflections combines spectroscopic information on the OO and JTD and is therefore a powerful tool to address these couplings [2]. In particular, a detailed understanding of the electronic ordering has been obtained from the study of superlattice reflections at the Mn$L_{2,3}$ and the O $K$ edges [3–7]. Theoretical calculations predicted a different spectral behavior of reflections sensitive to the OO and those sensitive to the JTD even before the first experiments were performed [8]. Experimentally, different temperature dependencies at various spectral features give information on the coupling, which is connected to the driving force of the transition. In particular, investigations of the orbital ($1/4\ 1/4\ 0$) reflection (c stands for cubic notation) in layered manganite La$_{0.5}$Sr$_{1.5}$MnO$_4$ have been interpreted in terms of OO being the main order parameter of the phase transition [9]. Similar investigations have subsequently been reported in another manganite system [10].

The coupling of these electronic degrees of freedom and the crystal structure has gained renewed interest as ultrafast probes allow us now to disentangle the dynamics of the individual degrees of freedom in the time domain [11]. The ultrafast dynamics of electronic and magnetic phase transitions in manganites have been extensively studied [12–19]. In particular, resolving the structural dynamics in three-dimensional (3D) manganites following the photoinduced melting of the charge and orbital ordered (CO/OO) phase has been addressed in detail [11,20–24]. It has been shown that a single order parameter that depends only on the electronic excitation is sufficient to describe the photoinduced ultrafast phase transition of the entire system, including charge and structural degrees of freedom [11]. The excitation suppresses the electronic order instantly [11,25] and the response of both the underlying antiferromagnetic order [14] and the JTD [11] occurs on the same timescale of about 100 fs. The atomic rearrangement of the heavy ions within the unit cell is completed within 1 ps, and the lattice constants readjust within 10–20 ps [23].

---

1Present address: Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, 14195 Berlin, Germany.  
2Present address: Stanford Institute for Materials and Energy Sciences (SIMES), SLAC National Laboratory, Menlo Park, CA 94025, USA.  
3Present address: Fraunhofer-Institut für Naturwissenschaftlich-Technische Trendanalyse, Euskirchen, Germany.  
4urs.staub@psi.ch
FIG. 1. Temperature dependence of the orthorhombic lattice parameters in PCMO4 determined by conventional powder x-ray diffraction. The splitting of the two in-plane parameters, shown in panel (a), sets in at the HTT-LTO transition at 466 K and becomes enhanced at the charge and orbital ordering transition at 320 K; the orthorhombic splitting is reduced at the LTO-LTLO transition at \( T^* \sim 146 \) K (see the text for more information). The black line denotes the average of the in-plane lattice parameters. The inset shows the \( B_{meb} \) unit cell for \( T > 330 \) K (green spheres: Pr/Ca). The HTT-LTO transition is also observed as a kink in the \( c \) parameter, shown in panel (b), and the charge and orbital ordering causes a change in the temperature slope. In contrast, there is no anomaly at \( T^* \). Panel (c) shows the orthorhombic splitting and the \( c/a \) ratio, which also exhibits anomalies at the respective transition temperatures.

Most investigations on electronic phase transitions in manganites concentrated on the three-dimensional distorted perovskites \( R_{1-x}T_xMnO_3 \), with \( R \) being a trivalent light lanthanide (La, or Pr ion) and \( T \) a divalent metal ion such as Ca or Sr. However, much less is known on the structural dynamics of the layered counterparts, which have basically the same CO/OO within the MnO layer [26]. However, the ordering perpendicular to the planes, which have a much larger separation due to the horizontal displacement of the oxygen octahedra (see, e.g., the inset of Fig. 1), exhibits significantly weaker correlations. Though the photoinduced CO/OO dynamics and the dynamics of the antiferromagnetic ordering have been investigated for the single-layer \( La_{0.5}Sr_{1.5}MnO_4 \) (LSMO4) system [15,24], nothing is known on the ultrafast behavior of single-layer manganite \( Pr_{0.5}Ca_{1.5}MnO_4 \) (PCMO4). The CO/OO transition for PCMO4 occurs at \( T \sim 320 \) K. Associated with the metal-insulator-type transition, the high-temperature crystal structure (space group \( B_{meb} \)) transforms to a structure that has an in-plane doubled unit cell with likely \( Pnma \) symmetry below \( T_{CO/OO} \) [27,28]. At 127 K, there is an additional magnetic phase transition where the Mn spins order antiferromagnetically [29]. The lattice constants remain approximately constant between 130 and 200 K with \( a = 5.37 \) Å, \( b = 5.42 \) Å, and \( c = 11.8 \) Å (\( B_{meb} \) symmetry) [28], which will be used further as a reference. An additional phase transition that is presumably structural in origin has been observed at \( T^* = 146 \) K, below which additional reflections forbidden in \( B_{meb} \) symmetry appear in x-ray and neutron diffraction experiments, such as, e.g., the (1 1 0) reflection, indicating a further lowering in crystal symmetry [27].

In our present study, we clarify how the reduced dimensionality of the layered \( Pr_{0.5}Ca_{1.5}MnO_4 \) (PCMO4) [28,29] influences the melting and recovery dynamics of the structural distortion associated with the CO/OO phase. In addition, we investigate the correlation between the \( T^* \) transition and the CO/OO order. To this end, we compare the ultrafast decay dynamics of a reflection sensitive to the \( T^* \) transition with the structural dynamics associated with the suppression of the CO/OO phase. For a qualitative understanding of the equilibrium coupling between electronic order and structural distortions, the temperature dependence of the \( (0 1/2 0) \) superlattice reflection intensity was collected through static resonant soft-x-ray diffraction experiments. These results are compared to nonresonant hard-x-ray diffraction (HXRD) experiments on various superlattice reflections.

II. EXPERIMENTS

Single crystals of PCMO4 were grown using the floating zone technique as described in Refs. [27,30]. A half-moon-shaped disk was cut out of the rod. The sample quality was verified with standard x-ray diffraction characterization techniques and SQUID magnetometry. Temperature-dependent measurements of the lattice constants were performed on a D5000 x-ray powder diffractometer equipped with a liquid-helium cryostat and an Anton Paar furnace using a sample
obtained by crushing a part of the single crystal. For the
time-resolved experiments, a (2 7 0) surface was polished.
The time-resolved x-ray experiments were carried out at the
FEMTO slicing source at the Swiss Light Source [31]. This
source produced hard-x-ray pulses of ∼100 fs duration. Using
a double Mo/B4C multilayer monochromator, an x-ray energy
of 7.1 keV was selected, yielding a good compromise between
the optimal source flux and minimum air-transmission losses.
The penetration depth of the pump and the probe was ap-
proximately matched by using a small 0.5° grazing incident
angle of the x-ray beam. The estimated pump and probe
depths are approximately 40 and 50 nm [32], respectively.
The footprint of the x-ray probe beam on the sample surface
was minimized by reducing the vertical beam size to 10 μm
[33]. The beam remained unfocused horizontally with a width
of approximately 300 μm. The sample was excited with
weakly focused (520 × 620 μm²) p-polarized 100-fs pulses
at 800-nm wavelength entering the sample at 10° grazing
incidence. The angle mismatch between the pump and the
probe increases the time resolution to approximately 200 fs.
The repetition rates of the optical pump and the x-ray probe
were 1 and 2 kHz, respectively. The probed diffracted inten-
sity thus alternates between pumped and unpumped states of
the sample, allowing us to compensate for slow drifts in the
x-ray source, pump laser, detector, and electronics [34]. The
diffracted intensity of superlattice (SL) peaks was measured
with an avalanche photodiode (APD), providing enhanced
sensitivity for weak diffraction signals. The x-ray energy is
above the Mn K-edge and therefore x-ray fluorescence con-
tributes significantly to a homogeneous background, which
has been estimated from the constant baseline of rocking
curve scans and was corrected for in the analysis. The crystal
was cooled with a N2 cryoblower and the temperature has
been set to T ∼ 100 K. Static x-ray diffraction experiments
were performed at the surface diffractometer of the X04SA
Material Science beamline of the Swiss Light Source with an
x-ray energy of 10.8 keV [35]. The sample was either cooled
with a N2 cryoblower or glued on the cold finger of a He
flow cryostat reaching 100 and 15 K, respectively. Data were
collected with a Pilatus pixel detector [36]. The resonant soft-
x-ray diffraction experiments were performed at the RESOXS
endstation [37] of the SIM beamline [38] in the vicinity of
the Mn L2,3 edges. The sample was mounted with the [110]
and [001] directions in the scattering plane. A continuous-flow
helium cryostat was used to achieve sample temperatures
between 50 and 350 K. Experiments were performed with σ
and π incident polarizations, corresponding to an incoming
polarization perpendicular and parallel to the scattering plane.
The outgoing polarization was not analyzed.

III. RESULTS AND DISCUSSION

A. Structural transitions of PCMO4

The temperature dependence of the lattice parameters that
has been extracted from x-ray powder diffraction over a
large temperature interval illustrates the various structural
phase transitions occurring in PCMO4; see Fig. 1. The high-
temperature tetragonal (HTT, space group I4/mmm) phase transforms into the low-temperature orthorhombic phase
(LTO, space group Bmeb) at 466 K due to the tilting of the
MnO₆ octahedra, which commonly occurs in layered
perovskites. In addition to the onset of orthorhombic splitting,
there is a kink in the c lattice parameter and an even more
pronounced one in the c/a ratio, because the average in-plane
lattice constant expands in the LTO phase. The onset of charge
and orbital ordering at T_{CO/OO} enhances the orthorhombic
splitting, but the impact on the average in-plane constant and
on c is opposite to that at the onset of the tilting. At the
lower structural transition T⁺ the orthorhombic splitting is
reduced, but this effect is not associated with the onset of
AFM ordering that occurs at even lower temperature. Below
T⁺ in neutron diffraction, an additional reflection emerges,
(110) in orthorhombic notation [26,27,39], that signals further
symmetry reduction. In accordance with the observation in
various cuprates [40] as well as in nickelate and cobaltate
compounds [41] of the same layered perovskite structure, we
attribute this low-temperature structural phase transition to the
change of the tilting scheme from LTO to the low-temperature
less orthorhombic (LTLIO, space group Pccn) one. In space
group Bmeb, the (h k 0) with h, k = odd reflections are forbid-
den by the b glide mirror symmetry, but these reflections are
allowed in Pccn [and also in the low-temperature tetragonal
(LTT, space group P4‡/nmc) phase]. The change in the tilting
scheme from LTO to LTLIO is a rather common phenomenon,
and the fact that it occurs clearly above the onset of AFM
order disagrees with the previously given interpretation of
strong magnetoelastic coupling [29].

Doubling of the Bmeb unit cell along the b axis induced by
the CO/OO yields a nonzero structure factor of reflections of
type (h k/2 l) in orthorhombic notation [26,39]. Figure 2
shows the temperature dependence of the (2 3/2 0) reflection
(blue curve), which samples the in-plane structural distortion

FIG. 2. Integrated intensity of the (2 3/2 0) and (−3 3 0) superlattice reflections (Bmeb notation) as a function of temperature normalized to the intensity of the (0 8 0) main structural reflection taken at 100 K.
and is insensitive to any structural distortions perpendicular to the plane as its Miller index $l$ is zero. Its intensity $I$ with changing temperature reflects the square of the order parameter of the CO/OO phase, and it is not visibly affected by the LTO-LTLO transition at $T^*$. This reflection has been chosen for a better comparison to a previous pump-probe study on the 3D manganite LPCMO \cite{42}, which has a similar in-plane CO/OO transition and correspondingly similar in-plane distortions. The temperature dependence of the nonresonant x-ray intensity of the $(\mathbf{1} \mathbf{5}/\mathbf{2} \mathbf{0})$ reflection (forbidden in Bmmb symmetry) is shown in Fig. 2 as red data points. The intensity of the $(\mathbf{3} \mathbf{3} \mathbf{0})$ reflection is more than an order of magnitude smaller than that of the $(\mathbf{2} \mathbf{3}/\mathbf{2} \mathbf{0})$ reflection, in agreement with the weakness of these reflections observed in other layered perovskites exhibiting the same LTO-LTLO transition \cite{40,41}.

Further in-plane reflections are shown in Fig. 3, and they exhibit a different temperature dependence than the $(\mathbf{2} \mathbf{3}/\mathbf{2} \mathbf{0})$ reflection. The in-plane reflection $(\mathbf{0} \mathbf{3}/\mathbf{2} \mathbf{0})$, which is only sensitive to the doubling of the unit cell, saturates around 250 K and the $(\mathbf{1} \mathbf{5}/\mathbf{2} \mathbf{0})$ reflection shows a maximum at approximately 250 K. This reflection is also significantly weaker than the $(\mathbf{2} \mathbf{3}/\mathbf{2} \mathbf{0})$ reflection (see Fig. 2). However, both reflections probe structural distortions induced by the in-plane CO/OO. These strongly different temperature dependences show that there are other crystallographic modes than the JTD mode for properly describing the crystal structure below the CO/OO phase transition as well as for the structural transition $T^*$.

### B. Electronic order

To assess the relation of the unexplored structural transition at $T^* = 146$ K \cite{27} to the CO/OO phase, we will first look directly at the temperature dependence of the orbital order by resonant soft-x-ray diffraction. The OO can be directly accessed at the Mn $L_{2,3}$ edges by studying the energy and temperature dependence of the $(0 \mathbf{1}/\mathbf{2} \mathbf{0})$ reflection, which has been done in detail in 3D crystals \cite{5,43} and thin films of PMCO \cite{44,45} as well as in the single-layer LSMO4 \cite{3,4,6,9}. The spectral shape of the $(0 \mathbf{1}/\mathbf{2} \mathbf{0})$ reflection in PMCO4 is shown in Fig. 4 and is very similar compared to that found for single-layer LSMO4. Therefore, we label the spectral features according to Ref. \cite{9}.

The temperature dependence of this reflection taken at different spectral features (energies) is shown in Fig. 5. As for LSMO4 \cite{9}, the shape of the temperature dependence depends strongly on the x-ray energy with a similar behavior for equivalent spectral features \cite{9}. The intensity of the highest energy feature $E$ does not have a regular order-parameter behavior; it peaks around 170 K for PMCO4. In LSMO4 this feature has a smooth temperature dependence and is only weakly temperature-dependent below 150 K with a broad maximum around 125 K. The feature with the largest intensity at the $L_3$ edge ($B$) shows in both cases a more regular order-parameter
behavior, i.e., it increases continuously for decreasing temperatures. The deviation from a regular order-parameter behavior is more pronounced for PCMO4 compared to LSMO4. These deviations are expected to be related to the differences in crystal structure of the two single-layer compounds. Whereas the crystal symmetry remains “quasi” tetragonal for LSMO4, the ground state of PCMO4 is clearly orthorhombic. In addition, PCMO4 shows an additional structural phase transition at $T^\ast$. The transition temperature $T^\ast$ lies lower in temperature than the maximum in the temperature dependence of the electronic spectral feature $F$, which makes a relation to $T^\ast$ questionable. However, the turnover in the temperature dependence of the spectral feature $F$ is likely related to structural modifications that are sensitive to the in-plane $b$ axis such as the JTD predicted in Ref. [9].

C. Ultrafast dynamics of the CO/OO induced structural distortion

Figure 6 shows the normalized intensity of the $(2\,3/\,2\,0)$ superlattice reflection as a function of the pump-probe delay $t$ for a series of excitation fluences $\Phi$ measured at $T = 100 \, K$. Independent of $\Phi$, the optical excitation leads to a fast drop of the diffraction intensity on a timescale comparable to the experimental time resolution. Subsequently, the diffracted intensity partially recovers. Furthermore, we observe weak oscillatory dynamics superimposed on the recovery, which we resolve with better statistics for a selected excitation fluence ($\Phi$) of 3.5 mJ/cm$^2$ at $T = 100 \, K$. The line represents the fit as explained in the text.

If $n > n_c$, the phase transition to the CO/OO disordered state occurs and the order parameter vanishes. The time-dependent model [11] accounts for a partial recovery of the order parameter, which is enabled by the energy dissipation from the electronic to the phonon system for deposited energy densities $n$ below $n_c$. To account for the process of energy deposition by the laser pulse and the subsequent dissipation of energy from the electronic to the lattice system, the energy density in the electronic system is described as being time-dependent [11,46]:

$$n(t) = n_0(t) - \alpha n_c e^{-\frac{t}{\tau_L}} + \alpha n_c$$

with

$$\alpha = 1 - \left(1 - \frac{n_0(t)}{n_c}\right)^{\frac{\gamma}{\gamma_0}}$$

and

$$n_0(t) = \frac{1}{2} n_0 \left[1 + \text{erf}\left(\frac{t}{\tau_L}\right)\right].$$
Here, $\tau$ is an effective relaxation time describing the equilibration of the electron and phonon system, and $n_0$ is the total energy deposited by the pump laser pulse of duration $t_1$. Furthermore, $\gamma_0$ and $\gamma$ are the critical exponents with respect to the initial excitation ($t \sim 0$) and after equilibration ($t \gg \tau$).

Here we do not imply a linear dependence between the reduction of the scattering intensity $I$ and $n_0$ for $t \sim 0$ (note that $I \propto n^2$). This is achieved by leaving $\gamma_0$ as a free parameter of the model ($\gamma_0 = 1/2$ in Refs. [11,46]).

We follow the approach of Ref. [11] and model the transient reflection intensity considering a depth-dependent excitation density using an estimated penetration depth of 51 and 40 nm for the x-ray probe and the laser excitation, respectively. The total depth considered by the present model is 400 nm and consists of 400 layers. This allows us to fit the dynamics for all excitation fluences shown in Fig. 6 simultaneously (Fig. 6, solid lines). We account for the experimental time resolution of $\sim$200 fs by convolution of $\eta^2(t)$ with a Gaussian function. We do not include the coherent oscillations of the Jahn-Teller soft mode by assuming that the JTD adiabatically follows the transient minimum position of the atomic potential (i.e., the JT model is considered massless).

Fitting of the model yields $n_c = 9 \times 10^7 \text{J/cm}^3$, which is about twice as high as in the 3D system [11]. This is in line with the higher ordering temperature of PCMO4, which suggests that more energy is needed to melt the ordered phase. The subsequent recovery occurs on a timescale of $\tau = 1.85 \pm 0.5$ ps, which is slightly slower than in the 3D system. This could indicate a smaller $e$–ph coupling of the layered system. Furthermore, we extract a critical exponent of the thermalized system of $\gamma = 0.29 \pm 0.03$ and for the initial excitation of $\gamma_0 = 0.69 \pm 0.05$. While $\gamma$ is comparable to the 3D system, a higher value of $\gamma_0$ is required to obtain a good fit to the experimental data for PCMO4.

The fast-initial drop after photoexcitation occurs within the time resolution of the experiment. As in the 3D system, the fast dynamics indicates that the structural distortion is promptly released by the suppression of CO/OO. The critical behavior of the squared order parameter against the excitation fluence is shown in Fig. 8, which compares the experimental scattering intensity at early and late times. We find the scaling to be qualitatively similar to that for 800 nm excitation in 3D manganites [11] and nickelates [46]. The deviation of the model to the highest-fluence data is likely due to the incomplete recovery between successive pump pulses, as was recently observed for 3D PCMO [23]: For fluences higher than required to fully melt the CO/OO order at the surface, the diffraction intensity no longer completely recovers between successive pump pulses due to the first-order character of the CO/OO transition, which results in slow dynamics of domain growth. The highest fluence time trace where this effect is largest was thus not included in the fitting procedure.

Figure 7 shows the trace for a pump fluence $\Phi$ of 3.5 mJ/cm$^2$ with improved statistics, giving clear evidence of a displacive lattice excitation. We employ a phenomenological fit function to extract the oscillation frequency:

$$I(t) = 1 - \Theta(t - t_0)(Ae^{-(t - t_0)/\tau} + Be^{-(t - t_0)/\tau_0}\cos(2\pi\nu(t - t_0))),$$

where $\Theta$ is the Heaviside function, and $A$ and $B$ represent amplitudes of the recovery and the phonon mode, respectively. We find a frequency of the displacive mode $\nu = 3.15$ THz that is approximately 30% higher than in 3D Pr$_5$Ca$_{0.5}$MnO$_3$ [11], where it has been interpreted to contain motions of the heavy Pr ions [20], which seems similar to PCMO4 [47]. The slightly higher frequency as compared to the 3D system might be explained by a larger Coulomb repulsion caused by shorter in-plane bond distances. Note that the average in-plane lattice constants are reduced and more anisotropic [48] compared to that of the 3D analogon. Similar to the 3D compound, we assign this coherent modulation to atomic motions along the transition coordinates, triggered by the partial melting of the CO/OO and subsequent relaxation of the JTD.

D. Ultrafast dynamics of structural distortion below $T^*$

To further address a possible correlation between the CO/OO and the structural second-order-like phase transition around $T^*$, we investigated the photinduced ultrafast dynamics of the (−3 3 0) reflection, which appears below $T^*$. The time trace of the reflection intensity after photoexcitation is shown in Fig. 9. As this reflection is much weaker, it required longer integration times to achieve reasonable statistics in the data set. This reflection shows a transient suppression that is significantly slower compared to the electronically driven relaxation of the distortion associated with the CO/OO, where the observed intensity drop was limited by the time resolution of the experiment. A fit using single exponential decay yields a time constant of 0.5 ps. This much slower structural relaxation dynamics is consistent with the absence

**FIG. 8.** Normalized x-ray scattering intensity of the (2 3/2 0) reflection averaged in the vicinity of 0.2 and 5 ps time delay as a function of the pump fluence. Error bars indicate the standard deviation of data points within the averaging windows (0.1–0.3 and 4.5–5.5 ps, respectively). The lines are fits of the model to the data as explained in the text.
of a significant coupling to the CO/OO, which agrees with the interpretation of the resonant soft-x-ray diffraction data. Assuming the transition at $T^*$ is displacive in nature, the slow relaxation indicates a strongly damped, very low energy soft mode associated with this phase transition. Such a behavior is in accordance with the Pccn LTLO symmetry that bridges the LTO and LIT phases and that thus exhibit a soft tilting mode. Interestingly, the observed timescale here is much slower than that observed for a phonon-driven phase transition in simple perovskites. Whereas for SrTiO$_3$ [49] a much faster decrease of the corresponding superlattice reflection has been found, an ultrafast increase of the reflection has been found in EuTiO$_3$, indicative of a transient increase in the structural order parameter [50]. In both cases, however, the dynamics is driven by changes of the structural soft-mode potential due to photodoping, most likely in contrast to the structural soft-mode potential due to changes of the structural soft-mode potential due to changes of the structural soft-mode potential due to the interpretation of the resonant soft-x-ray diffraction data.

ACKNOWLEDGMENTS

We thank D. Meister and A. Oggenfuss for technical support at the X04SA and FEMTO beamlines, and D. Grollmund for supporting the time-resolved “femtoslicing” experiments at the X05SA beamline. We further acknowledge help by E. Pomjakushina and S. Stutz with preparation of the sample surface. The research leading to these results has received funding from the Swiss National Science Foundation and its National Center of Competence in Research, Molecular Ultrafast Science and Technology (NCCR MUST) and Materials’ Revolution: Computational Design and Discovery of Novel Materials (NCCR MARVEL). L.R. acknowledges funding from the DFG in the Emmy Noether program under Grant No. RE 3977/1. E.M.B. acknowledges funding from the European Community’s Seventh Framework Programme (FP7/2007-2013) under Grant Agreement No. 290605 (PSI-FELLOW/COFUND). We acknowledge financial support of the Deutsche Forschungsgemeinschaft (DFG) through CRC 1238 (Project No. 277146847, subprojects A02, B04, and B05).

IV. CONCLUSION

Ultrafast x-ray diffraction was used to study the time dependence of superlattice reflections sensitive to the CO/OO ground state of layered PCMO$_4$. They show similar dynamics compared to three-dimensional PCMO after an ultrashort optical pulse excites the system. It indicates that the CO/OO melting and recovery dynamics is tightly coupled to the JTD distortion also in layered manganites. We further argue that the additional structural phase transition, occurring deep inside the charge order phase (at $T^*$), can be attributed to the change of the tilting scheme from LTO to LTLO. This structural transition behaves differently when probing the respective symmetry-breaking reflections. These characteristic LTLO reflection intensities are suppressed on significantly slower timescales, indicative of a decoupling of this structural transition from the CO/OO. The static resonant soft-x-ray diffraction experiments on OO superlattice reflections find a temperature dependence of the reflection intensity, which depends on the energy of the x rays, but they do not show any indication of an anomaly at $T^*$. This supports the interpretation of insensitivity of the LTO-LTLO transition to the CO/OO. The variation in the temperature dependence of the reflection taken at various x-ray energies at the absorption edge shows that other structural modes couple to the OO in addition to the JTD. This view is supported by the differences in the temperature dependence between structural reflections sensitive to different structural modifications. In summary, it shows that the layered manganites based on Pr have a much richer interplay of crystal structure and its electronic properties than three-dimensional half-doped manganites.

REFERENCES


