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## Role of equilibrium fluctuations in light-induced order

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Engineering novel states of matter with light is at the forefront of materials research. An intensely studied direction is to realize broken-symmetry phases that are "hidden" under equilibrium conditions but can be unleashed by an ultrashort laser pulse. Despite a plethora of experimental discoveries, the nature of these orders and how they transiently appear remain unclear. To this end, we investigate a nonequilibrium charge density wave (CDW) in rare-earth tritellurides, which is suppressed in equilibrium but emerges after photoexcitation. Using a pump-pump-probe protocol implemented in ultrafast electron diffraction, we demonstrate that the light-induced CDW consists solely of order parameter fluctuations, which bear striking similarities to critical fluctuations in equilibrium despite differences in the length scale. By calculating the dynamics of CDW fluctuations in a nonperturbative model, we further show that the strength of the light-induced order is governed by the amplitude of equilibrium fluctuations. These findings highlight photoinduced fluctuations as an important ingredient for the emergence of transient orders out of equilibrium. Our results further suggest that materials with strong fluctuations in equilibrium are promising platforms to host "hidden" orders after laser excitation.

In a symmetry-breaking phase transition, fluctuations of the order parameter provide important information about the way an ordered state develops. Near the transition temperature  $T_c$ , fluctuations exhibit a diverging correlation length and correlation time, whose critical exponents define the underlying universality class. In contrast to the equilibrium situation, the role of order parameter fluctuations remains unclear if a phase transition proceeds under nonequilibrium conditions. Of particular interest are transitions instigated by an intense laser pulse, which has led to discoveries of many "hidden" orders that are not accessible in thermal equilibrium, such as light-induced superconductivity [1-3], charge or spin density waves [4–7], and ferroelectricity [8, 9]. These outof-equilibrium orders are often short-lived, raising the question of whether they exist in the form of fluctuations and if so, how they are related to fluctuations in equilibrium.

Empirically, several material classes that host transient states also display strong equilibrium fluctuations of the associated order [3, 8-14]. In underdoped cuprates where light-induced superconductivity was discovered [10–13], pronounced superconducting fluctuations are expected due to the small phase stiffness and poor screening [15]. In  $\kappa$ -type organic salts where light-induced superconductivity was observed above  $T_c$ , Nernst effect measurements also pointed towards large fluctuations due to a nearby Mott criticality [3, 16, 17]. In cases where equilibrium fluctuations do not yield an ordered state at finite temperature, such as in the quantum paraelectric phase of SrTiO<sub>3</sub>, a terahertz pulse can induce a ferroelectric state in a metastable fashion [8, 9, 18]. These observations suggest that photoinduced orders may be a special manifestation of equilibrium fluctuations, but experimental evidence is lacking to formally establish a link between the two entities.

Here, through a side-by-side comparison, we show that a newly-discovered photoinduced charge density wave (CDW) [4, 5] shares the key characteristics of the CDW fluctuations at  $T_c$  even though the former does not have a diverging correlation length. The comparison was enabled by a pump-pump-probe scheme with ultrafast electron diffraction, which gives a direct measurement of fluctuations through diffuse scatterings. Using a nonperturbative calculation, we further demonstrate that the intensity of the photoinduced CDW peak increases with the strength of the CDW fluctuations in equilibrium. The positive correlation suggests that photoinduced "hidden" state is more likely found in systems with significant equilibrium fluctuations, paving the way forward as we search



FIG. 1. Competing charge density waves in rare-earth tritellurides. (a) Left: schematic of the layered structure of  $RTe_3$ , where dashed lines indicate the primary unit cell. Right: Enlarged view of the nearly square-shaped Te sheets that host the CDW instabilities. (b) Schematic phonon dispersion right above  $T_c$  along  $\Gamma$ -X and  $\Gamma$ -Z, featuring two Kohn anomalies at  $q_{\text{soft}}$ . (c) Schematic of fluctuating CDWs right above  $T_c$ . (d)(e) Equilibrium electron diffractions of DyTe<sub>3</sub>  $(T_c = 306(3) \text{ K [19]})$  taken at 100 K (d) and 307 K (e). (f)(g) Time-resolved diffractions of LaTe<sub>3</sub> before (f) and 1.6 ps after (g) photoexcitation by an 80-fs, 800-nm laser pulse with an incident fluence of  $2.1 \text{ mJ/cm}^2$ , measured at 307 K. Blue and red arrows indicate the CDW peaks along the c- and aaxis, respectively. Difference in intensities of lattice Bragg peaks in (d) and (e) results from slight sample drift and tilt during the warm-up process.

for novel nonequilibrium orders.

The charge density wave is hosted by the rare-earth tritelluride  $(RTe_3)$  family. All members possess a layered structure and the CDW instability is found in the nearly square-shaped Te sheets [Fig. 1(a)]. The quasi-two-dimensional nature of the crystals leads to a much reduced  $T_c$  compared to the mean-field transition tem-

perature. This gives rise to significant CDW fluctuations above  $T_c$ , as evidenced by Raman spectroscopy [20] and inelastic X-ray scattering [21]. The near- $C_4$  symmetry of the Te sheets leads to two competing CDWs: The dominant one has a modulation along the *c*-axis while the subdominant one along the orthogonal *a*-axis [22]. Here, we focus on LaTe<sub>3</sub> ( $T_c \approx 670$  K) and DyTe<sub>3</sub> ( $T_c = 306(3)$  K) [22, 23]. They share nearly identical properties except for the different transition temperatures [24]. Hence, under similar experimental conditions, we have access to CDW fluctuations in the critical regime near  $T_c$  (DyTe<sub>3</sub>) as well as a state with only the dominant *c*-axis CDW (LaTe<sub>3</sub>).

Figure 1(d)(e) shows the equilibrium electron diffraction patterns of DyTe<sub>3</sub> in the (H, 0, L) plane, taken below and near  $T_c$  (see [25] for experimental details). At 100 K, pairs of CDW satellite peaks are found along the *c*-axis at a wavevector  $q_c = 0.294(1)c^*$  (blue arrows), but no satellite peaks are observed along the orthogonal a-axis [26]. When the sample is heated to  $T_c$ , the *c*-axis peaks significantly weaken but remain visible [Fig. 1(e)]; in the meantime, diffuse spots arise along the *a*-axis (red arrows). Notably, the diffraction pattern appears symmetric between the *c*- and *a*-axis, as highlighted by three observations: (i) brighter  $(H \pm q_a, 0, L)$  satellites are found along the c-axis than along the a-axis; vice versa for the  $(H, 0, L \pm q_c)$  peaks; (ii) the CDW wavevectors are similar,  $q_a \approx q_c$ ; (iii) the satellite intensities are comparable for the two CDWs. Transverse atomic displacements associated with both CDWs account for the intensity pattern in (i) [25]. Observations (ii) and (iii) preclude the possibility of a long-range CDW along the *a*-axis that is known to occur in DyTe<sub>3</sub> at 68 K  $\ll T_c$  [21] because this low-temperature *a*-axis peak has a markedly different wavevector and a much weaker diffraction intensity compared to its c-axis counterpart [4, 27]. The symmetric appearance of the diffuse spots in Fig. 1(e) is a signature unique to the critical regime near  $T_c$ . Below  $T_c$ , such symmetry is broken by the long-range *c*-axis CDW. At temperatures significantly exceeding  $T_c$ , fluctuations are weak, rendering any diffuse scattering invisible under the background intensity.

We now turn to LaTe<sub>3</sub> and study the behavior of the CDWs out of equilibrium. Figure 1(f)(g) show the electron diffraction patterns taken 0.8 ps before and 1.6 ps after the incidence of an 80-fs, 800-nm laser pulse. After photoexcitation, the long-range CDW order along the *c*-axis is suppressed (blue arrows) while new peaks appear along the *a*-axis (red arrows), whose intensity increases monotonically with pump laser fluence [4, 25]. Remarkably, the CDW superlattice spots in this transient snapshot of the photoexcited state are visually indistinguishable from those in the equilibrium diffraction pattern recorded at  $T_c$  in DyTe<sub>3</sub> [Fig. 1(e)(g)]. In particular, the transient CDW satellites along both axes share a similar intensity and wavevector, hinting at a restored symmetry between the two CDWs.

The similarity between Fig. 1(e) and 1(g) allows us to interpret the light-induced CDW state using an equilibrium picture close to  $T_c$ . In momentum space, the diffuse satellite peaks are indicative of the population of transient soft phonons along the  $a^*$ - and  $c^*$ -axis [Fig. 1(b)]. In real space, this critical regime is characterized by shortrange CDW patches in both directions [Fig. 1(c)], with the correlation length inversely proportional to the momentum width of the Kohn anomaly [25]. From inelastic X-ray measurements [21], the phonon energies at  $q_a$  and  $q_c$  are approximately 1 to 2 meV, corresponding to a fluctuating timescale of 2 to 4 ps for these CDW patches. A similar timescale is observed as the lifetime of the lightinduced a-axis CDW [Fig. S4(a)]. This energy-time correspondence suggests that the light-induced a-axis CDW is indistinguishable from a soft phonon at the corresponding wavevector, confirming the intimate link between the photoexcited state and the critical regime near  $T_c$ .

The comparison between the photoexcited and the critical state suggests that the photoinduced *a*-axis CDW in LaTe<sub>3</sub> does not have long-range order and remains fluctuating. While the statement can be rigorously proven by simple theoretical arguments [25], here we give an estimate of the *finite* correlation length of the *a*-axis CDW. Based on the diffraction peak width w [Fig. 1(g)], which is limited by instrumental resolution, the correlation length has a lower bound of  $1/w \sim 3.5$  nm, or 8 crystallographic unit cells (u.c.). Given the approximate CDW lifetime  $\tau$  of 4 ps [Fig. S4(a)], the correlation length is at most  $v\tau \sim 10 \text{ nm}$  (23 u.c.), where v = 2500 m/s is the speed of sound along the a-axis [28]. This upper bound is a testament that each fluctuating patch cannot establish phase coherence with its neighbors at a speed faster than phonon propagation. Compared to the correlation length of the dominant *c*-axis CDW in equilibrium, which is estimated to be at least 1.8  $\mu$ m within Te planes [22], the particularly small value of  $v\tau$  hence confirms the absence of long-range order along the *a*-axis and suggests that the light-induced CDW consists entirely of short-range fluctuations.

An almost square-symmetric diffraction pattern after photoexcitation and at equilibrium  $T_c$  is suggestive of a close connection between the two states. To further elucidate their relationship, we investigate their response to an external perturbation. By comparing the respective dynamics of the order parameter fluctuations, we can gain some crucial insights into the similarities and differences between the two regimes. To this end, we apply a second laser pulse to LaTe<sub>3</sub> right after the emergence of the *a*-axis satellite peak and record the intensity evolution of the CDW fluctuations along both axes. As a reference, we also photoexcite DyTe<sub>3</sub> at its CDW transition temperature, where fluctuations of both density waves abound.

We first examine the laser-induced response in DyTe<sub>3</sub> at its  $T_c$  [Fig. 2(a)]. After photoexcitation, the diffuse



FIG. 2. Response of CDW fluctuations to photoexcitation. (a)(b) Schematic setups for DyTe<sub>3</sub> and LaTe<sub>3</sub>. Both samples were kept at T = 307 K. The incident fluence was  $3.3~{\rm mJ/cm^2}$  in (a) and  $1.0~{\rm mJ/cm^2}$  for each pump in (b). (c)-(h) Changes in the integrated intensities for thermal diffuse scattering  $[I_{\text{TDS}}, (c)(d)]$ , *a*-axis diffuse CDW peak  $[I_a,$ (e)(f), and c-axis diffuse CDW peak  $[I_c, (g)(h)]$ . Integration areas are marked by solid circles in the insets. Traces are normalized by the average value of  $I_c$  before photoexcitation. In (d)(f)(h), vertical lines indicate the arrival time of pump<sub>2</sub> at  $\Delta t = 1.3$  ps. For reference, dynamics in the absence of pump<sub>2</sub> is shown in gray. (i)(i) Enlarged view of dashed rectangles in (e)–(h) after subtracting the respective thermal diffuse background  $[I_{\text{TDS}} \text{ in } (c)(d)]$ . In (i), traces are normalized by the average values at t < 0. In (j), traces are plotted as a function of the relative delay between the probe pulse and  $pump_2(t')$ , where intensities are normalized by the average value in the interval  $t' \in [-0.7 \text{ ps}, -0.2 \text{ ps}]$ . A slightly larger reduction along the *c*-axis is attributed to a mismatch between pumped and probed volumes, leading to additional melting of residual long-range CDW by the second pulse. Solid curves in (c)-(j)are fits to a phenomenological model in Eq. (S1). The black fitted curve in (i) uses the averaged data along the a- and c-axes.

satellite spots display an initial dip in intensity followed by a fast recovery, a trend perfectly mirrored in both axes [Fig. 2(e) and (g)]. These dynamics are in stark contrast to diffuse scattering intensities at other momenta away from Bragg or CDW peaks, where only a singleexponential rise is observed [Fig. 2(c)]. The dip can be understood in two equivalent ways. From the phonon perspective, it represents a transient stiffening of the soft mode [25, 29]. As electrons are excited to high energy, there is a transient reduction in the electronic band occupation near the Fermi energy that interacts with the lattice ions. This reduction leads to an increase in the renormalized phonon frequency and hence a decrease in the phonon population, as suggested by the equipartition theorem. An alternative viewpoint is based on the classical description of phonons as atomic displacements in real space. In each frame diffracted from a single electron pulse, we capture a snapshot of the system, such as the one depicted in Fig. 1(c). The dip hence indicates a smaller lattice distortion amplitude in the fluctuating CDW patches, averaged over space and over all snapshots at the same pump-probe delay. The second perspective naturally connects the photoinduced melting of fluctuating CDWs to the melting of a long-range CDW. Locally, there is minimal distinction between the two processes and both occur over  $\sim 0.4$  ps, a timescale dictated by the phonon period associated with the CDW distortion [30, 31]. In Fig. 2(e)(g), we observe that the intensities quickly rise after the dip, indicating an increased phonon population from laser-induced heating. After subtracting the thermal diffuse contribution, the dip only partially recovers [Fig. 2(i)], suggesting an elevated lattice temperature above  $T_c$ , where the Kohn anomaly becomes less pronounced.

Next, we study the dynamics in the photoexcited state of LaTe<sub>3</sub>. As illustrated in Fig. 2(b), we use the first laser pulse to bring the material into a nonequilibrium state, where we have observed a symmetric appearance of diffuse satellite spots along both a- and c-axes. We then apply a second pulse to perturb this transient state and look at the response of the two competing CDW fluctuations. In the experiment, the two pump pulses share the same *incident* fluence. To assess the *absorbed* fluence, we note that the maximum value attained in thermal diffuse scattering doubles after the second pulse [Fig. 2(d)]. This observation affirms that energy absorption is minimally affected by the presence of excited carriers after the first pulse. We now move on to analyze the CDW peaks, shown in Fig. 2(f) and (h). Unlike their distinct behavior upon the initial photoexcitation, the intensity evolution of the peaks along both axes share almost identical trends after the second pulse. For a direct comparison between the two orders, we zoom in to their dynamics right after the second pulse and plot them together in Fig. 2(j), where intensities from thermal diffuse scattering have been subtracted using the same procedure applied to DyTe<sub>3</sub>. Similar to the fluctuating CDWs in  $DyTe_3$  near  $T_c$ , the two diffuse peaks in LaTe<sub>3</sub> feature a transient reduction in the fluctuation amplitude, fol-

lowed by a recovery that lasts for more than 2 ps. Unlike  $DyTe_3$ , the satellite intensities in LaTe<sub>3</sub> are fully recovered compared to their values just before the second pulse, suggesting the nonthermal nature of these density wave fluctuations.



FIG. 3. Simulated dynamics of photoinduced CDW and its relation to equilibrium fluctuations. (a) Evolution of integrated intensities of the *c*-axis (blue) and *a*-axis (red) CDW peak upon photoexciting the unidirectional CDW state. Triangle marks the maximum intensity of the light-induced CDW. The nonzero value of the *a*-axis peak at t = 0 originates from thermal fluctuations. (b) Maximum intensity change of the photoinduced *a*-axis CDW peak  $[\Delta I_a(t)]$  as a function of equilibrium diffuse intensity at a fixed temperature above  $T_c$ , the latter of which quantifies thermal fluctuations and is indistinguishable between the two axes.  $\Delta I_a(t)$  is normalized by  $I_a(t=0)$  (see Fig. S8).

The similarities between the excited state in LaTe<sub>3</sub> and the critical state in DyTe<sub>3</sub> – both in their diffraction snapshots (Fig. 1) and in their photoinduced dynamics (Fig. 2) – suggest that the light-induced CDW is a special manifestation of critical fluctuations. While the equilibrium fluctuations near  $T_c$  are thermal and follow the scaling relations prescribed by the theory of renormalization group [32], the light-induced fluctuations may not conform to a thermodynamic distribution [33]. To understand how the strength of equilibrium fluctuations affect the appearance of the light-induced CDW, we developed a time-dependent Ginzburg-Landau formalism within the Gaussian approximation (see [25] for derivation). This approach gives a nonperturbative solution to the lightinduced dynamics, yielding quantities that have a oneto-one correspondence to the observables in our timeresolved diffraction experiments. Unlike N-temperature models [34], here we do not need to artificially assign a temperature to each degree of freedom in the system.

To assess the validity of the model, we first calculate intensity evolution of a- and c-axis CDW peaks after photoexcitation [Fig. 3(a)]. The simulated trends successfully reproduce the experimental observations (Fig. S4). The transient enhancement of intensity along the a-axis is solely the result of CDW fluctuations without longrange order [25]. In Fig. 3(b), as we reduce the order parameter stiffness to increase the amplitude of equilibrium fluctuations above  $T_c$ , the strength of the transient CDW order also increases under identical photoexcitation conditions. This positive correlation suggests that strong fluctuations in equilibrium constitute an important factor for observing light-induced ordering phenomena out of equilibrium.

Despite the similarities between the light-induced CDW and the critical fluctuations, there exist important differences [25]. For example, the transient lattice temperature of LaTe<sub>3</sub> stays far below its equilibrium  $T_c$ , and there is no change in the in-plane lattice anisotropy after photoexcitation, distinct from the evolution of a and c lattice parameters across  $T_c$  [4, 22]. Importantly, the light-induced CDW has a finite correlation length for all time delays but at the critical point in equilibrium, correlation length diverges with fluctuations occurring at all length scales. Hence, strictly speaking, the photoexcited state is not truly critical as described in a thermodynamic transition.

By leveraging the symmetry between two competing CDWs in  $RTe_3$ , we have elicited the correspondence between a photoinduced order and critical fluctuations in equilibrium. The parallels provide a nonthermal pathway to access hidden symmetries of a system even if  $T_c$ is unattainable under equilibrium condition. The similarities also hint at the existence of universal scaling laws that govern the dynamics of a highly nonequilibrium system [33], which have been detected in scattering experiments with high momentum resolution and an extended time delay [35–37]. Furthermore, our results offer a generic mechanism for the creation of photoinduced states, which can emerge as order parameter fluctuations in the absence of long-range order. This insight suggests that one should look for material classes that exhibit strong order parameter fluctuations in equilibrium in order to look for "hidden" states out of equilibrium. Experimental signatures for such strong fluctuations depend on the order parameter, ranging from diffuse peaks in a charge or spin density wave system to Nernst effect in a superconductor [25]. We expect the connection between equilibrium fluctuations and out-of-equilibrium ordering to hold regardless of microscopic details, providing a guiding principle in our search for other light-induced states.

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