Incoherent Topological Defect Recombination Dynamics in TbTe$_3$

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We study the incoherent recombination of topological defects created during a rapid quench of a charge-density-wave system through the electronic ordering transition. Using a specially devised three-pulse femtosecond optical spectroscopy technique we follow the evolution of the order parameter over a wide range of time scales. By careful consideration of thermal processes we can clearly identify intrinsic topological defect annihilation processes on a time scale $\sim 30$ ps and find a possible signature of extrinsic defect-dominated relaxation dynamics occurring on longer time scales.

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Topological defects are nonlinear objects that can be created any time a symmetry-breaking transition occurs [1–3]. They can be described theoretically as solutions to systems of nonlinear differential equations based on Ginzburg-Landau theory. They are of great fundamental importance in fields such as cosmology where they appear as strings and condensed matter physics where they appear in the form of vortices and domain walls. While a good understanding of static properties of topological defects (TDs) has come from systems such as liquid crystals, the dynamics of TDs is much less understood. Electronic phase transitions in charge-density-wave (CDW) systems [4] are particularly interesting model systems for studying the general behavior of the dynamics of topological excitations. The collective excitations are not overdamped, which allows the observation of both collective and quasiparticle (QP) excitations as they evolve through the transition. In particular, they can be used to investigate the dynamic behavior of topological excitations such as domain walls in real time using ultrafast laser techniques.

Recently, time-resolved experiments have shown that, following a quench created by a strong laser pulse, the order parameter (OP) oscillates coherently, revealing coherent TD dynamics [5]. Domain walls are created parallel to the crystal surface that can coherently annihilate on the time scale of a few picoseconds with the accompanying emission of collective modes that have been detected as modulations of reflectivity upon reaching the surface. In addition to the coherent defect dynamics, incoherent topological defects created by the Kibble-Zurek mechanism [2] are also expected, but very little is known about the incoherent TD dynamics in CDW systems, and in condensed matter systems in general.

In this Letter we investigate the incoherent evolution of TDs using a specially devised three-pulse femtosecond spectroscopy [5] technique, which allows the direct background free observation of the evolution of the OP as a function of time through the electronic ordering transition. In a rapid quench experiment order emerges in different regions of the sample independently and so multiple topological defects can be created. Their presence can be detected in the optical response as a spatial inhomogeneity of the order parameter. The determination of incoherent TD dynamics is a challenging task, however. Because of thermal-diffusion processes, which evolve on similar time scales as topological annihilation and also introduce temperature inhomogeneity, careful temperature calibration from independently measured frequencies is needed to accurately account for thermal effects. We deal with the problem by careful calibration of the transient effective temperature inhomogeneity, which enables us to unambiguously distinguish the incoherent dynamics from thermal-diffusion effects.

In our experiments, we use a three-pulse technique described in Refs. [5,6]: A “destruction” (D) laser pulse at 800 nm excites a cold sample [7] into the disordered state, breaking up the CDW order. We then monitor the evolution of the transient reflectivity $\Delta R(t_{pp})/R$ excited with a weaker pump (P) pulse as a function of time delay $t_{DP}$ between the D and P pulse [the pulse sequence nomenclature is illustrated in the insert to Fig. 1(b)]. The D pulse fluence is adjusted to $\sim 650$ $\mu$J/cm$^2$, which is four times the threshold for causing the destruction of the ordered state [5]. After a quench by the laser pulse, order recovers first through the subpicosecond recovery of the quasiparticle gap leading to coherent oscillations of the OP and the coherent creation of TDs, which decay within 5–8 ps in TbTe$_3$ [5]. Because the CDW coherence length ($\sim 2$ nm) is much shorter than the size of the laser excited volume ($\sim 50$ $\mu$m dia), order emerges with different phase in

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different regions, resulting in the formation of topological defects whose spatial distribution is determined partly by the inhomogeneous excitation and partly by the underlying fluctuations that nucleate the emergence of order by the so-called Kibble-Zurek [2,3] mechanism. The resulting inhomogeneity of the OP leads to observable temporally resolvable effects in the frequency, linewidth and amplitude of the collective amplitude mode, all of which are related to the OP, as shown in previous studies [5].

In Fig. 1(a) we show the raw data on the transient reflectivity \( \Delta R(t_{DP})/R \) of TlTe at different time delays \( t_{DP} \) after the \( D \) pulse. After the initial QP relaxation we observe oscillations due to the coherently excited order-parameter amplitude mode (AM) and other phonons [9–11]. The level of noise is very small, due to the excellent intrinsic properties of the material, which helps us make a detailed quantitative analysis. We analyze the transient reflectivity oscillations using the theory for dispersive excitation of coherent phonons [12]:

\[
\frac{\Delta R(t_{DP})}{R} = \int_0^\infty G(t-u)[A_e \exp(-u/\tau) + A_0]du \\
+ \sum A_i \int_0^\infty G(t-u) \exp(-\gamma_i u) [\cos(\Omega_i u) \\
- \beta_i \sin(\Omega_i u)]du,
\]

where \( \beta_i = (1/\tau - \gamma_i)/\Omega_i, G(t) = \exp(-2t^2/\tau^2) \) and \( \tau_P \) is the laser pulse length. The first integral represents the QP relaxation with the relaxation time \( \tau \), while the sum depicts the response of coherent phonons with frequencies \( \Omega_i \), and effective damps \( \gamma_i \). \( A_e \) corresponds to the QP relaxation amplitude while the residual value at long delays is \( A_B \). To limit the number of fitting parameters we keep only two phonon terms corresponding to the AM at 2.2 THz and the 1.7–THz phonon which strongly interacts with the AM at higher temperatures [9]. Figure 1(b) shows the fast Fourier transform (FFT) of the raw data and of the fit to the data (i) without the \( D \) pulse and (ii) for a \( D-P \) delay of \( t_{DP} = 10 \) ps, clearly showing that Eq. (1) fits the response very well below \( \sim 2.4 \) THz irrespective of \( t_{DP} \). The \( t_{DP} \) dependence of the frequency, linewidth, and amplitude of the AM and 1.7 THz phonon modes are shown in Fig. 2. The linewidth is shown for two sets of data obtained from different samples.

In order to obtain a calibration of the effective temperature \( T_{eff} \) of the photoexcited sample volume we measured independently, by means of a standard pump-probe experiment, the \( T \) dependence of the reflectivity transients in the thermal equilibrium and determine the \( T \)-dependent amplitude, frequency \( \omega_{AM}(T) \) and damping \( \gamma_{AM}(T) \) for the AM [13]. Using these calibrations we are in a position to determine \( T_{eff} \) as a function of time from \( \omega_{AM}(t_{DP}) \) and \( \gamma_{AM}(t_{DP}) \) and take it into account to obtain the thermal inhomogeneity dynamics. The time dependence of the \( T_{eff} \) is shown in Fig. 3(e). We observe that the two effective temperatures \( T_w(t_{DP}) \) and \( T_c(t_{DP}) \) obtained from the \( \omega_{AM}(t_{DP}) \) and \( \gamma_{AM}(t_{DP}) \) systematically differ by more than \( \sim 20 \) K indicating an excess AM linewidth with respect to the thermal equilibrium state.

One of the most obvious contributions to the excess AM linewidth is the inhomogeneous broadening caused by the thermal inhomogeneity. In order to be able to identify and analyze any other contributions to the linewidth we therefore need to determine the contribution of the thermal inhomogeneity to the excess AM linewidth. To do this we first fit a thermal-diffusion model [14,15] (TDM) to the effective temperature obtained from the AM frequency \( T_w(t_{DP}) \), and then use the TDM parameters to calculate the transient optical reflectivity, which fully takes into account inhomogeneity of the temperature in the excited volume [14]. As seen in Fig. 3(e), \( T_w(t_{DP}) \) can be fit very well over 5 decades of time from 30 ps to 4 \( \mu \)s using a one-dimensional [16] TDM, where \( T(t_{DP}) = \Delta T/\sqrt{1 + t_{DP}/\tau_D + t_0} \), and the fit parameter \( \tau_D \sim 170 \) ps represents the characteristic heat diffusion time [14].
between the calculated primarily governed by the 1D heat diffusion process.

FIG. 2 (color online). (a) Frequencies of the AM and the 1.7-THz phonon as a function of \( t_{DP} \). (b) Effective dampings of the AM and the phonon as a function of \( t_{DP} \). Full diamonds represent a measurement from another sample. (c) Amplitudes of the AM and the phonon as a function of \( t_{DP} \). The solid lines are the frequencies, linewidths, and amplitudes calculated using an inhomogeneous temperature distribution model [14]. The actual D-pulse fluence on sample #2 was slightly lower leading to different TDM parameters.

Comparing now the simulation [14] with the experiment in Fig. 2, we see that the validity of the TDM beyond \( \sim 30 \) ps is well supported by the good agreement of the simulation for both the AM and the 1.7-THz phonon parameters. We can thus safely conclude that the recovery of the order parameter on time scales longer than \( \sim 30 \) ps is primarily governed by the 1D heat diffusion process.

Below \( \sim 30 \) ps, however, there is a large discrepancy between the calculated \( T_{eff} \), \( \gamma_{AM} \), and other phonon parameters in comparison to the data, even after carefully taking into account the thermal inhomogeneity. The observed magnitude and the evolution of \( \gamma_{AM}(t_{DP}) \) for \( t_{DP} < 30 \) ps clearly cannot be assigned solely to the temperature inhomogeneity. Subtracting the thermal inhomogeneity contribution from the AM linewidth in Fig. 2(b), we can now isolate the topological-defect inhomogeneity contribution as shown in Fig. 4.

As discussed in the introduction, some of the defects created in the quench process annihilate coherently resulting in an aperiodic modulation of the AM intensity and frequency in the first \( \sim 8 \) ps [5]. The very low level of experimental noise in the raw data allows us to attribute the observed data scatter in Fig. 4 to the coherent defect dynamics in the material, rather than experimental noise.

The modulation of the AM due to the defects that annihilate incoherently is unfortunately not detected directly by our stroboscopic technique. However, the incoherent topological defects give rise to a spatial inhomogeneity of the order parameter and a decoherence of the AM oscillations leading to an increased linewidth \( \gamma_{AM} \) for \( t_{DP} < 30 \) ps which we have detected in our experiments. Concurrently the defects give rise to a softening of the collective mode \( \omega_{AM} \), because of the OP suppression that they cause. The increase of the coupled 1.7-THz phonon effective damping at shorter \( t_{DP} \) as shown in Fig. 2 is presumably also caused by the inhomogeneity of the OP.

FIG. 3 (color online). (a) The temperature dependence of the coherent oscillations spectrum measured in a standard \( Pp \) experiment. Frequencies (b), dampings (c), and amplitudes (d) obtained from the time domain fits (1) as functions of temperature. Circles and diamonds represent the parameters for the AM and the 1.7-THz phonon, respectively. (e) The time dependence of the effective temperature \( T_{eff}(t_{DP}) \) from \( \omega_{AM}(t_{DP}) \) and \( \gamma_{AM}(t_{DP}) \). The solid line is a fit to \( \omega_{AM} \) effective temperature using the thermal-diffusion model. The inverted triangles correspond to the calculated \( T_{TD}^{TDM}(t_{DP}) \) taking into account the inhomogeneous temperature distribution [14].
A further manifestation of the incoherent annihilation is the increase of the amplitudes of the AM and the phonon with respect to the TDM for short $t_{DP}$, which is also consistent with the suppression of the OP when one takes into account that the amplitudes of the modes increase with the decreasing OP amplitude deduced from their $T$ dependence shown in Fig. 3(d).

Apart from intrinsic topological defect annihilation processes, which we have identified on a time scale of ~30 ps, we expect to observe annihilation of domain walls pinned to defects and imperfections at longer times. The time scale of their annihilation may extend well beyond 30 ps. Evidence for such slower extrinsic recombination processes comes from the long time behavior shown in Fig. 4. $\gamma_{AM}$ remains systematically larger than the predicted thermally inhomogeneous linewidth $\gamma_{TDM}$ suggesting a slower recombination of the pinned domain walls.

Recently the absence of topological defects on ultrafast time scales in highly excited charge ordered (CO) nickelate was suggested [17]. A slow relaxation of the CO x-ray diffraction peak intensity, on the time scale of ~60 ps, was, due to the absence of any increase of the diffraction peak linewidth, attributed to a depopulation of the phason mode. While, contrary to Lee et al. [17], our excitation density is clearly high enough [18] to excite topological defects in TbTe$_3$ [5], there exists a possible anharmonic contribution of the highly excited phason mode to the AM linewidth. The anharmonic processes, however, contribute to both the linewidth and frequency renormalization of the AM [19], and cannot lead to the difference between $T_{\omega}$ and $T_{\gamma}$ as observed in the experiment.

In conclusion, these experiments demonstrate the possibility of studying both coherent and incoherent topological defects dynamics in complex materials in which the order parameter can be monitored in real time through the dynamics of the collective mode. The dynamics on a time scale of ~30 ps can unambiguously be associated with intrinsic topological defect annihilation in TbTe$_3$, following a laser quench arising from the time-dependent inhomogeneity and suppression of the order parameter. The inhomogeneity causes an increased effective damping of the amplitude mode while the suppression of the order parameter is indicated by an additional softening of the AM-mode frequency. Beyond ~30 ps we find a predominantly thermal-diffusion governed order-parameter dynamics with a possible signature of extrinsic defect annihilation dynamics.

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[7] The growth of the samples by a self-flux technique and subsequent characterization was described elsewhere [8].
[13] To avoid systematic errors due to possible sample differences and inhomogeneity the $T$ dependence was measured for each sample separately on the same spots as the three-pulse measurements.
[16] The diameters of the beams are much larger than the optical penetration depth, and so on the relevant time scale the heat diffusion is 1D.
Although the fluences used by Lee et al. [17] are comparable or higher than the D-pulse fluence (~650 μJ/cm²) used in the present Letter they were not able to completely suppress the CO diffraction peak. In the case of TbTe₃, however, the CDW destruction threshold fluence at 800-nm photon wavelength is 160 μJ/cm², and so the D-pulse fluence used in our experiment is four times the threshold, which is enough to completely destroy CDW in the probed volume.