Ultrafast transient generation of spin-density-wave order in the normal state of BaFe$_2$As$_2$ driven by coherent lattice vibrations

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The interplay among charge, spin and lattice degrees of freedom in solids gives rise to intriguing microscopic quantum phenomena such as colossal magnetoresistance, multiferroicity and high-temperature superconductivity$^{12-13}$. Strong coupling or competition between various orders in these systems presents the key to manipulate their functional properties by means of external perturbations such as electric and magnetic fields or pressure$^1$. Ultrashort and intense optical pulses have emerged as an interesting tool to investigate elementary dynamics and control material properties by melting an existing order$^{4-6}$. Here, we employ few-cycle multi-terahertz pulses to resonantly probe the evolution of the spin-density-wave (SDW) gap of the pnictide compound BaFe$_2$As$_2$ following excitation with a femtosecond optical pulse. When starting in the low-temperature ground state, optical excitation results in a melting of the SDW order, followed by ultrafast recovery. In contrast, the SDW gap is induced when we excite the normal state above the transition temperature. Very surprisingly, the transient ordering quasi-adiabatically follows a coherent lattice oscillation at a frequency as high as 5.5 THz. Our results attest to a pronounced spin–phonon coupling in pnictides that supports rapid development of a macroscopic order on small vibrational displacement even without breaking the symmetry of the crystal.

In the past few years, the pnictides have been found as a new class of unconventional superconductors$^{5,11}$. Like in cuprates, high-temperature superconductivity is implemented by doping of an antiferromagnetic pnictide parent compound with carriers$^{12,10}$. Such a phase diagram with superconductivity emerging from a magnetic ground state has been observed in many other unconventional superconductors$^{14}$. Therefore, the underlying magnetic order has been at the heart of the discussion concerning the mechanism of high-temperature superconductivity in general$^{10-12}$. BaFe$_{2-x}$Co$_x$As$_2$ is one of the most studied Fe-based superconductors owing to the availability of large and high-quality single crystals. An antiferromagnetic SDW represents the ground state in the undoped compound BaFe$_2$As$_2$. Electron doping with Co gradually suppresses the SDW transition temperature $T_{SDW}$ and superconductivity emerges$^{11}$. Interestingly, in the underdoped region the two phases are found to coexist on a length scale smaller than a few nanometres$^{1,11}$. In addition, BaFe$_{2-x}$Co$_x$As$_2$ undergoes a structural phase transition from tetragonal to orthorhombic at or slightly above $T_{SDW}$. Another structural phase transition back to the tetragonal symmetry was reported in the superconducting state in samples close to optimal doping$^5$. Experiments under external pressure have confirmed the intimate connection between the lattice structure and electronic ordering, with increasing pressure suppressing the SDW and inducing superconductivity in a fashion similar to chemical doping$^1$. Therefore, unveiling the interplay among the lattice structure, magnetism and superconductivity seems to be crucial for an understanding of the nature of high-temperature superconductivity in pnictides.

 Femtosecond pump–probe techniques provide new opportunities to investigate various quantum degrees of freedom and their interactions in complex systems. Direct information about the interplay between single-particle electronic states, collective modes, magnetization and lattice structure may be obtained by monitoring the real-time dynamics following perturbation by ultrashort optical pulses$^{4,6,15}$. In particular, time-resolved multi-terahertz techniques extending over the far- and mid-infrared (MIR) spectral windows enable resonant probing of single-particle and collective low-energy excitations with a resolution of a few tens of femtoseconds$^{13}$.

We present a study of the dynamics of the complex conductivity in BaFe$_2$As$_2$ over the frequency range from 10 THz to 26 THz (corresponding to photon energies between 41 meV and 110 meV) following photo-excitation with a 12 fs pump pulse in the near-infrared (NIR) region centred at 1.55 eV. Single crystals of BaFe$_2$As$_2$ exhibiting a SDW transition at $T_{SDW} = 120$ K were grown from Sn flux$^{16}$. Temperature-dependent equilibrium spectra were measured by spectroscopic ellipsometry in the range from 12 meV to 500 meV and for temperatures between 10 K and 300 K (ref. 17). Photo-induced changes in the complex conductivity were obtained by measuring reflection changes with few-cycle MIR pulses directly in amplitude and phase of the electric field and as a function of the delay time $t_d$ after excitation$^{13}$. The spot size of the pump pulses was set to 110 μm and that of the probe pulses to 50 μm.

The equilibrium optical conductivity above and below $T_{SDW}$ is depicted in Fig. 1a by the red and blue lines, respectively. The opening of the SDW gap gives rise to a transfer of spectral weight from below to above the gap edge at 80 meV (ref. 18). Figure 1b shows the time evolution of the optical conductivity change induced by excitation with the NIR pump pulse at an absorbed fluence of

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\[ \Phi = 63 \mu \text{cm}^{-2} \] at a base temperature of the lattice of \( T_b = 10 \text{ K} \).

After excitation, the optical conductivity rapidly increases below the crossing point and decreases above. The extrema are reached at \( t_D = 180 \text{ fs} \), where the optical conductivity agrees well with the equilibrium optical properties in the normal state (green circles in Fig. 1a), indicating that the SDW order has vanished. Figure 1c presents the averaged responses in three distinct spectral regions below (A), around (B) and above (C) the gap edge, as indicated in Fig. 1b. The dynamics fits well to an exponential recovery with a time constant of 0.63 ps (black solid lines in Fig. 1c).

In addition to the exponential decay, weak temporal oscillations with a frequency of 5.5 THz become discernible in the reflectivity changes at higher excitation densities (Fig. 2 and Supplementary Fig. S3). This frequency matches the one of an \( A_{1g} \) mode, which involves the \( c \) axis displacement of arsenic with respect to the Fe square lattice\(^{19} \) (Fig. 3a). Coherent driving of this phonon mode has been observed in time-resolved optical reflection\(^{20} \) and photoemission studies\(^{21} \). When exciting below \( T_{SDW} \), the coherent oscillations appear as small modulations on top of the strong electronic signal due to the collapse and recovery of the SDW gap (Supplementary Fig. S2). Notably, we find that these oscillations become a prominent feature when pumping above \( T_{SDW} \). Figure 2a shows the temporal changes of the MIR optical conductivity in the normal state recorded at \( T_L = 134 \text{ K} \) and \( \Phi = 530 \mu \text{cm}^{-2} \). Instead of the strong dispersive signal due to the gap closure in the SDW state, the conductivity now exhibits an overall increase and strong modulations of the optical conductivity with the \( A_{1g} \) phonon frequency at 5.5 THz. Most importantly, we find that these oscillations have a very characteristic, dispersive behaviour. As demonstrated in Fig. 2b, the oscillations in spectral regions A and C (red and blue circles, respectively) are out of phase with each other whereas the oscillatory response is strongly suppressed in region B (green circles). A quantitative analysis of the oscillatory component reveals that it closely matches the change in the optical conductivity that is induced in the equilibrium state by the SDW: the difference between the stationary conductivities in the normal state at 140 K and in the SDW state at 100 K is shown as a black line in Fig. 2c. Except for a scaling factor of 2.5, the spectral change in conductivity derived by taking the difference between the transient oscillation maxima and minima averaged over delay times from 280 fs and 1.12 ps closely follows this graph (red circles in Fig. 2c). This congruence between the spectral shapes of the oscillatory response and the gap induced by the SDW represents the most important and surprising observation of our study. It indicates that even in the normal state well above \( T_{SDW} \) the coherently driven \( A_{1g} \) phonon periodically induces the SDW. This finding indicates that the macroscopic order parameter underlying the SDW exhibits a fast response to the collective nuclear motion with a modulation frequency as high as 5.5 THz.

In the following, we will develop a qualitative understanding of the coherent induction of SDWs in the normal state of \( \text{BaFe}_2\text{As}_2 \) that is supported by further experimental evidence. As discussed above, the ground state of pnictides may be tuned by means of external pressure and giant magneto–elastic coupling has been pointed out\(^{1,3,22} \). The distance of As from the Fe square lattice or the Fe–As–Fe bonding angle \( \alpha \) marked in Fig. 3a is suggested to represent the...
most important structural parameters. It was demonstrated that the maximum superconducting transition temperature is reached when the FeAs$_4$ unit forms an ideal tetrahedron with $\alpha = 109.47^\circ$ (refs 3,9). As the $A_{1g}$ vibration of the arsenic ions modulates precisely this angle, it might sensitively affect the ground state of the system. It was argued that owing to the large polarizability and magnetic ordering further enhances this effect (ref 24). This prediction is supported by the strong modulations of the density of states around the Fermi level due to the vibration of the arsenic ions modulates $g$ and off. Figure 4a presents the temperature dependence of the modulation amplitude of the MIR conductivity due to the $A_{1g}$ phonon. It exhibits a broad maximum around $T_{SDW}$ followed by a pronounced decrease with increasing temperature. However, the amplitude of the coherent lattice vibrations depends on the Raman susceptibility for the NIR pump light. Hence, it is expected to stay constant without a marked change in the electronic structure (ref 27). This fact is well reflected by the temperature dependence of the oscillatory component of the reflectivity recorded at a probe photon energy of 1.55 eV, that is far above the SDW gap: following a weak anomaly near $T_{SDW}$ in the normal state the amplitude shows only a slight increase towards higher temperature (Fig. 4b). The opposite temperature dependencies of the modulation amplitudes of MIR conductivity and NIR reflectivity support our assignment of the dispersive oscillatory component in the MIR range to a photo-induced ordering of SDWs: despite the fact that the amplitude of the coherent lattice motion does not decrease with increasing temperature it becomes exceedingly difficult to induce the SDW order at $T \gg T_{SDW}$.

To test our model, it is instructive to compare the structural changes described by the phonon vibrations with those induced by temperature or external pressure: the absolute amplitude of the coherent lattice vibrations may be estimated on the basis of the modulation amplitude of the MIR conductivity and NIR reflectivity. Figure 4a shows a broad maximum around $T_{SDW}$ followed by a pronounced decrease with increasing temperature. However, the amplitude of the coherent lattice vibrations depends on the Raman susceptibility for the NIR pump light. Hence, it is expected to stay constant without a marked change in the electronic structure (ref 27). This fact is well reflected by the temperature dependence of the oscillatory component of the reflectivity recorded at a probe photon energy of 1.55 eV, that is far above the SDW gap: following a weak anomaly near $T_{SDW}$ in the normal state the amplitude shows only a slight increase towards higher temperature (Fig. 4b). The opposite temperature dependencies of the modulation amplitudes of MIR conductivity and NIR reflectivity support our assignment of the dispersive oscillatory component in the MIR range to a photo-induced ordering of SDWs: despite the fact that the amplitude of the coherent lattice motion does not decrease with increasing temperature it becomes exceedingly difficult to induce the SDW order at $T \gg T_{SDW}$.
we induce is equivalent to an alternating internal pressure in the range of gigapascals. Whereas the decrease of π by external pressure suppresses the SDW, the As displacement towards a larger angle might enhance ordering. The angle indeed increases by 0.1–0.5° when BaFe2As2 is cooled down from the normal state to the SDW state28,29. This finding indicates that the amplitudes of the coherent phonon vibrations resulting from photoexcitation provide a deformation potential sufficient for the ultrafast ordering phenomena we observe.

A SDW can be established only when the energy of the participating electrons remains smaller than the SDW gap energy. When the pump energy is delivered to the sample, the electrons are prepared in a non-equilibrium energy distribution within a few tens of femtoseconds. If we assume that all of the absorbed energy is initially deposited in the electronic subsystem and efficient electron–electron scattering leads to a rapid thermalization within approximately 100 fs, then the system can be approximately described by an electronic temperature $T_e$ (as, for example, in a two-temperature model30). The fluence of 530 μJ cm$^{-2}$ used in the measurement of Fig. 2 could increase $T_e$ up to 1,100 K and prevent the formation of the SDW (ref. 20). Ultrafast energy relaxation of normal-state electrons with a time constant shorter than 0.2 ps, however, reduces $T_e$ to approximately 500 K within one cycle of the $A_{bg}$ phonon oscillation30. As a result, the electronic energy $k_b T_e$ rapidly becomes much smaller than the modulated SDW gap energy of $2\Delta_0^{m} \approx 76$ meV ($2\Delta_0^{m}/k_b = 880$ K) observed in Fig. 2. It is noteworthy that whereas the static SDW state has been observed only in the orthorhombic structure, electronic anisotropy as well as magnetic fluctuations survive in the tetragonal structure up to temperatures well above the transition to long-range macroscopic order31,32. The phonon vibrations may drive the fluctuation coherently such that the transient order could develop at a higher temperature. Note that despite the ultrafast formation we find for the periodic build-up of the SDW, it is unlikely that this process will proceed instantaneously and some retardation should be expected. Such an inherent timescale may prevent the full development of SDW order and account for the reduction of the spectral weight in the transient case found in Fig. 2c. Further studies on the ultrafast structural dynamics of the system are desired to reveal the detailed interplay between the lattice and SDW order.

Our observation of a coherent lattice vibration directly modulating a macroscopic order parameter on ultrafast timescales represents a qualitatively new form of coherent control of complex solids. Furthermore, we demonstrate a crucial role of the lattice for the appearance of a SDW state in BaFe$_2$As$_2$. The remarkable fact that the magnetic order quasi-adiabatically follows fast lattice motion at a frequency as high as 5.5 THz is compatible with the scenario of a transient SDW order driven by Fermi surface nesting. Finally, despite the fact that band structure calculations33 as well as some experiments20,21 have indicated a weak electron–phonon coupling due to the coherent phonon at 5.5 THz. Amplitudes are obtained from fits to a sinusoidal function and the error bars indicate 95% confidence intervals for the fitting parameter (see Supplementary Information).

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Author contributions

K.W.K., J.D., R.H. and A.L. planned the project; K.W.K. performed ellipsometry measurements; K.W.K., J.D., R.H. and A.L. wrote the paper. All measurements; K.W.K., A.P. and M.P. performed terahertz measurements; H.S. and M.B. performed NIR/visible region measurements; T.W. grew samples; and K.W.K., A.P., C.B., J.D., R.H. and A.L. wrote the paper. All authors contributed to discussions and gave comments on the manuscript.

Additional information

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