Ultrafast Melting of Spin Density Wave Order in BaFe$_2$As$_2$ Observed by Time- and Angle-Resolved Photoemission Spectroscopy with Extreme-Ultraviolet Higher Harmonic Generation

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Transient single-particle spectral function of BaFe$_2$As$_2$, a parent compound of iron-based superconductors, has been studied by time- and angle-resolved photoemission spectroscopy with an extreme-ultraviolet laser generated by higher harmonics from Ar gas, which enables us to investigate the dynamics in the entire Brillouin zone. We observed electronic modifications from the spin-density-wave (SDW) ordered state within $\sim 1$ ps after the arrival of a 1.5 eV pump pulse. We observed optically excited electrons at the zone center above $E_F$ at 0.12 ps, and their rapid decay. After the fast decay of the optically excited electrons, a thermalized state appears and survives for a relatively long time. From the comparison with the density-functional theory band structure for the paramagnetic and SDW states, we interpret the experimental observations as the melting of the SDW. Exponential decay constants for the thermalized state to recover back to the SDW ground state are $\sim 0.60$ ps both around the zone center and the zone corner.

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were measured using a Scienta R4000 hemispherical electron energy analyzer. The total energy resolution was 250 meV. All the measurements were done at 10 K, which was well below the $T_e$ of the equilibrium SDW transition. The first BZs for the PM and antiferromagnetic (AFM) states are shown in Fig. 1 (c). For simplicity, we shall use the notation of the PM BZ to specify positions in the momentum space. In-plane ($k_x, k_y$) and out-of-plane ($k_z$) momenta are expressed in units of $\pi/a$ and $2\pi/c$, respectively, where $a = 3.97$ Å and $c = 13.00$ Å are the in-plane and out-of-plane lattice constants, respectively.

![Image 1](image.png)

**FIG. 1.** (Color online) (a) Temporal evolution of angle-integrated photoemission intensity around the $(0,0,6.5)$ point of the Brillouin zone (BZ). (b) Difference spectra of (a) from the average intensity before the arrival of pump pulse. (c) First BZs of the paramagnetic (black) and antiferromagnetic (red) states. We shall use the paramagnetic notation below. Solid and dotted arrows indicate the momentum cuts for the $(0,0,6.5)$ and $(1,1,6.1)$ points, respectively. (d) Temporal evolution of the electronic temperature ($T_e$). $T_e$ is evaluated by fitting the photoemission intensity to the Fermi-Dirac distribution function convoluted with the instrumental gaussian function. Black dotted line shows the fit to a decay function of the form $T_e(t) = A + B \Theta(t) \exp(-t/\tau_{\text{decay}})$, where $\Theta(t)$ is the Heaviside step function.

Figure 1 (a) shows the temporal evolution of angle-integrated photoemission (PES) spectra around the zone center. With $h\nu = 28$ eV probe photons the momentum cut is on the $k_z = 6.5(2\pi/c)$ plane as shown by a solid arrow in Fig. 1 (c). One observes that upon the arrival of a pump pulse at $t = 0$, the electron population above the Fermi level ($E_F$) suddenly increases and it decays afterwards. To improve the visibility of the data, we show the difference spectra in Fig. 1 (b). Here, the spectrum average before $t = 0$ has been subtracted. It is clear that just after $t = 0$ the population above $E_F$ increases while that below $E_F$ decreases, and that this population modification decreases with delay time. In order to quantify the transient state, we show in Fig. 1 (d) the electronic temperature $T_e$ as a function of delay time. $T_e$ was determined by fitting the PES intensity at each delay time to a Fermi-Dirac distribution function convoluted with a Gaussian with the instrumental resolution of 250 meV. After the pump pulse, $T_e$ reaches $\sim 1400$ K. $T_e$ was fitted to a decay function of the form $T_e(t) = A + B \Theta(t) \exp(-t/\tau_{\text{decay}})$, where $\Theta(t)$ is the Heaviside step function. The best fit is shown by a dotted curve. We obtained a decay constant of $\tau_{\text{decay}} = 0.58$ ps around the $(0,0,6.5)$ point.

In order to study the momentum-resolved spectral function during the relaxation, we show in Fig. 2 (a) the TrARPES spectra taken at different delay times. The momentum cut is the same as that of Fig. 1. To highlight the change of the spectra, we also show the difference TrARPES spectra in Fig. 2 (b). Here, the spectrum at -0.47 ps has been subtracted from each spectrum. Before the pump arrival (-0.47 ps), the spectrum should represent that of the SDW state, although the identification is not obvious under the present experimental resolution. We later argue that the spectrum represents that of the SDW state. After the arrival of pump pulse ($t \geq -0.02$ ps), electrons start to occupy states above $E_F$. At 0.12 ps, high-energy states are populated above the gap ($E - E_F > 0.6$ eV) and the states have parabolic, electron-band-like intensity distribution. At 0.41 ps, this population disappears, indicating the fast decay of electrons in the high energy state. As for the intensity between $0 < E - E_F < 0.4$ eV, it already exists at -0.02 ps, reaches maximum around 0.12 ps, and slowly decays as a function of time. At 1.42 ps, there is almost no separation of positive and negative area in the difference spectrum, indicating that the electronic system has relaxed back to the equilibrium state.

Here we discuss the implications of the present data for an ultrafast phase transition. In the following, we argue that the observed hot electron states are those of the PM states caused by the melting of the SDW state. We compare in Figs. 3 (a) and (b) the difference spectrum at 0.12 ps with density-functional theory (DFT) band structure at $k_z = 6.5 (2\pi/c)$ in the PM state calculated within the generalized gradient approximation (GGA) [17]. The intensity increase is shown by red in the difference spectrum in Fig. 3 (a). We can confirm high-energy population above 0.6 eV and a gap between $0.4 < E - E_F < 0.6$ eV. This is qualitatively consistent with the calculated PM band structure shown in Fig. 3 (b), and quantitatively the smaller gap in experiment can be ascribed either to the renormalization of the band structure or to the experimental resolution. We also show the DFT band structures for the SDW state in Fig. 3 (c). Since the Ba122 samples were not detwinned, we show the results for two inequivalent domains. One observes that the band structure near $E_F$ in the SDW state is more entangled than that in the PM state, due to the band folding caused by the AFM ordering. In particular, the band bottoms of domain 1 at 0.3 eV and the flat band at 0.6 eV of domain 2 are not clearly resolved in the present measurement. From these comparisons, the electronic states at 0.12 ps can be identified as that of the PM state, as expected from the high $T_e > 1000$ K after the optical pump.
FIG. 2: (Color online) (a) Time-resolved angle-resolved photoemission (TrARPES) spectra around the (0, 0, 6.5) point. (b) Difference TrARPES spectra from the spectrum before the pump arrival ($t = -0.57$ ps).

To gain further insight from the intensity decrease shown in blue, equilibrium spectra taken with $h\nu = 63$ eV (Z point) for the SDW (20K) and PM (150 K) states are shown in Figs. 3(d) and (e), respectively. We note that the blue intensity in panel (a) shows a parabolic, electron-band-like feature. We ascribe this to a parabolic feature observed in the equilibrium the SDW state as indicated by a black dotted line in Fig. 3(e), which does not appear in the PM state [Fig. 3(d)]. This feature originates from the electron band bottom located at the zone corner in the PM state. The disappearance of this feature by raising the temperature from the SDW state to the PM state is also reported in Ref. 13. Thus the data at -0.47 ps can be naturally assigned to those of the SDW state, as expected from the sample temperature of $10 \, K < T_N$.

Next we investigate the dynamics of excited electrons around the zone corner. Figure 4(a) shows the temporal evolution of angle-integrated PES spectra at the (1,1,6.1) point. We employed $h\nu = 28$ eV and the momentum cut in the BZ is indicated by a dotted arrow in Fig. 1(b). The difference spectra are also shown in Fig. 4(b). As in the case of the (0,0,6.5) point, the intensity above $E_F$ increases and that below $E_F$ decreases at $t = 0$, and this change weakens with time. The evolution of $T_e$ is plotted in Fig. 4(c). $T_e$ rises up to $\sim 900 \, K$ just after the pump pulse. We obtain a decay constant of 0.60 ps, close to 0.58 ps around the (0,0,6.5) point.

TrARPES spectra around the (1,1,6.1) point are shown in Fig. 5(a) and the difference spectra are shown in Fig. 5(b). Although the modifications in the spectra are weaker than those in the zone center, we observed an intensity increase after the pump pulse. Note that we did not observe high-energy population well above $E_F$ within our measurement range. To understand the difference between the results for the (0,0,6.5) and (1,1,6,1) points, we show the calculated PM band structure in Fig. 5(c). One of the two bands located around $\sim 1$ eV above $E_F$ is holelike and flat; as a result, the lifetime of excited electrons around the (1,1,6.1) point can be much shorter than around that at (0,0,6.5), because they can easily decay into different momentum states via electron-phonon and/or electron-electron interactions. To support the interpretation, we also show the calculated SDW band structure for two domains in Fig. 5(d). As in the (0,0,6.5) point, there are more bands in the SDW state than in the PM state. In particular, there are three electron band bottoms between $0.2 < E - E_F < 0.6 \, eV$, where long excitation lifetime is expected. The absence of the
more consistent with the band structure calculation of the PM state than that of the SDW state, which is expected from the high $T_c$ above 1000 K. After the fast decay of the optically excited electrons, a thermalized state appears and survives for a relatively long time. Decay constants both around the $\Gamma$ point and the X point are $\sim 0.60$ ps, in agreement with previous TrARPES and time-resolved reflectivity measurements.

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FIG. 4: (Color online) (a) Temporal evolution of angle-integrated PES spectra around the (1, 1, 6.1) point. The momentum cut is indicated by a dotted arrow in Fig. 1 (c). (b) Difference spectra of (a) from the average intensity before the arrival of pump pulse. (c) Temporal evolution of $T_e$. 

observation of transient population in this energy range corroborates our assignment of the transient states between 0.1 and 0.2 ps to the PM state.

While the high-energy electrons above 0.6 eV around (0, 0, 6.5) point decay quickly (< 0.41 ps), the hot electrons close to $E_F$ persist for a longer time, which can be interpreted as the thermalized electronic states. For the time scale of the relaxation of near-$E_F$ states, we compare the present decay constants with those from previous works. The present value $\sim 0.60$ ps for 10 K is close to 0.8 ps for EuFe$_2$As$_2$ [9], 0.38 ps for Ba122 [15]. Also, it is comparable with 0.6-0.8 ps for superconducting Ba$_{1-x}$K$_x$Fe$_2$As$_2$ [18] estimated from time-resolved reflectivity. As for the A$_{1g}$ phonon mode, we did not observe any oscillatory intensity modulation near $E_F$. Mansart et al. [8] found that, while the phonon oscillation is barely visible at $\sim 1.3$ mJ/cm$^2$ in the course of the exponential decay of the reflectivity, it becomes more prominent at higher flux ($\sim 2$ mJ/cm$^2$) and makes the relaxation slower. The absence of oscillations may suggest that our data represents the electronic state in weakly pumped regime. Other possibilities include the polarization and the width of the pump pulse. We need further investigation into the dependence of the transient states on fluence, polarization and pump width in the future.

In conclusion, we have studied the ultrafast dynamics of the BaFe$_2$As$_2$ in the SDW state by TrARPES using a HH from rare gas. We observed electronic modifications from the SDW band structure within $\sim 1$ ps after the 1.5 eV pump pulse. High-energy states above $E_F$ were observed at 0.12 ps at the zone center and they decay rapidly. The 0.12 ps spectrum is
FIG. 5: (Color online) (a) TrARPES spectra around the (1, 1, 6.1) point. (b) Difference TrARPES spectra from the spectrum before the pump arrival ($t = -1.02$ ps). (c), (d) DFT band structures around the (1, 1, 6.1) point for the PM and SDW states, respectively.

Kemper, T. Wolf, Y. D. Chuang, Z. Hussain, C. C. Kao, B. Moritz, Z. X. Shen, T. P. Devereaux, and W. S. Lee, Nat. Commun. 6, 7377 (2015).

[13] M. Yi, D. H. Lu, J. G. Analytis, J.-H. Chu, S.-K. Mo, R.-H. He, M. Hashimoto, R. G. Moore, I. I. Mazin, D. J. Singh, Z. Hussain, I. R. Fisher, and Z.-X. Shen, Phys. Rev. B 80, 174510 (2009).

[14] Y. R. Shen, The principles of nonlinear optics (Wiley-Interscience, Hoboken, 2003).

[15] X. Yang, L. G. Rohde, T. Rohwer, A. Stange, K. Hanff, C. Sohrt, L. Rettig, R. Cortés, F. Chen, L. Feng, D. T. Wolf, B. Kamble, I. Eremin, T. Popmintchev, M. Murnane, M. C. Kapteyn, H. L. Kipp, J. Fink, M. Bauer, U. Bovensiepen, and K. Rossnagel, Phys. Rev. Lett. 112, 207001 (2014).

[16] K. Ishizaka, T. Kiss, T. Yamamoto, Y. Ishida, T. Saitoh, M. Matsunami, R. Eguchi, T. Ohtsuki, A. Kosuge, T. Kanai, M. Nohara, H. Takagi, S. Watanabe, and S. Shin, Phys. Rev. B 83, 081104 (2011).

[17] J. P. Perdew, K. Burke, and M. Ernzerhof, Phys. Rev. Lett. 77, 3865 (1996).

[18] E. E. M. Chia, D. Talbayev, J.-X. Zhu, H. Q. Yuan, T. Park, J. D. Thompson, C. Panagopoulos, G. F. Chen, J. L. Luo, N. L. Wang, and A. J. Taylor, Phys. Rev. Lett. 104, 027003 (2010).