Origin of the Spin Density Wave Instability in $AFe_2As_2$ ($A = Ba, Sr$) as Revealed by Optical Spectroscopy


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We performed optical spectroscopy measurement on single crystals of BaFe$_2$As$_2$ and SrFe$_2$As$_2$, the parent compounds of FeAs-based superconductors. Both are found to be quite metallic with fairly large plasma frequencies at high temperature. Upon entering the spin-density-wave state, the formation of partial energy gaps was clearly observed with the surprising presence of two different energy scales. A large part of the Drude component was removed by the gapping of Fermi surfaces. Meanwhile, the carrier scattering rate was even more dramatically reduced. We elaborate that the spin-density-wave instability is more likely to be driven by the Fermi surface nesting of itinerant electrons than a local-exchange mechanism.

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The recent discovery of superconductivity with a transition temperature $T_c$ above 50 K in $R$FeAs$_{1-x}$F$_x$ ($R = La, Ce, Sm, Pr, Nd, etc.$) has created tremendous interest in the scientific community [1–4]. Those compounds crystallize in a tetragonal ZrCuSiAs-type structure, which consists of alternate stacking of edge-sharing $Fe_2As_2$ tetrahedral layers and $R_2O_2$ tetrahedral layers along the c axis. The parent compound LaFeAsO itself is not superconducting but shows strong anomalies near 150 K in resistivity, magnetic susceptibility, specific heat, etc. Based on experimental observations and first-principles calculations, it is suggested that the ground state is a spin-density-wave (SDW) ordered state with a stripe-type (or collinear) spin configuration [5]. The predicted magnetic structure was confirmed by a subsequent neutron diffraction experiment, although the neutron data indicated that a subtle structural distortion occurs first near 150 K, and the SDW long range order establishes at a slightly lower temperature [6]. With fluorine doping, the SDW order is suppressed and superconductivity emerges [2,5]. The very closeness of the superconducting phase to the SDW instability suggests that the magnetic fluctuations play a key role in the superconducting pairing mechanism.

Investigating the origin of the antiferromagnetic (AFM) SDW instability in the parent compound is an essential step towards understanding the mechanism of superconductivity in doped systems. The stripe-type AFM order was first suggested to result from the nesting between the hole and electron Fermi surfaces (FSs) of itinerant electrons [5]. Alternatively, it was proposed that the superexchange interaction mediated through the off-plane As atom plays a key role in the spin configuration formation [7–12]. A stripe-type AFM would arise when the next nearest neighbor exchange becomes larger than half of the nearest neighbor exchange interaction. Whether an itinerant picture or a local superexchange mechanism should be taken as a starting approach becomes one of the most important issues for those systems [13].

Very recently, it was found that the ThCr$_2$Si$_2$-type ternary iron arsenide BaFe$_2$As$_2$, which contains identical edge-sharing $Fe_2As_2$ tetrahedral layers as in LaFeAsO, exhibits a similar SDW instability at 140 K [14]. It is therefore suggested that BaFe$_2$As$_2$ could serve as a new parent compound for ternary iron arsenide superconductors. Shortly after that, the superconductivity with $T_c = 38$ K was found in $K$-doped BaFe$_2$As$_2$, which was suggested to be a hole-doped iron arsenide superconductor [15]. The SDW instability was also found in SrFe$_2$As$_2$ with a higher transition temperature near 200 K [16,17], while $K$ doping again introduces the superconductivity with $T_c \sim 38$ K [17]. For both parent compounds, the same stripe-type AFM order was confirmed by neutron experiments [18,19]. A great advantage of those ternary iron arsenide compounds is that it is much easier to grow large size single crystals [20].

In this Letter, we present an optical study on both BaFe$_2$As$_2$ and SrFe$_2$As$_2$ single crystals. We find that the undoped compounds of Fe-pnictides are quite metallic with rather high plasma frequencies $\omega_p \approx 1.5$ eV. Upon entering into the SDW state, formation of energy gaps was clearly observed. Surprisingly, the optical measurement revealed two distinct energy scales in the gapped state. Associated with the gapping of the FS, a large part of the Drude component is removed; meanwhile, the carrier scattering rate shows even steeper reduction. Beyond the characteristic energy for SDW gap, another spectral suppression in $R(\omega)$ which covers a much higher energy scale is present even above SDW transition temperature. The physical implications of those results were discussed. Our study favors an itinerant electron approach for the driving mechanism of SDW instability.

Single crystals of BaFe$_2$As$_2$ and SrFe$_2$As$_2$ (space group $I4/mmm$) were grown from the FeAs flux method [21]. The platelike crystals could be easily cleaved, resulting in very shiny surface. The $T$-dependent dc resistivity was measured by the four-contact technique in a Quantum
Design physical property measurement system. As shown in Fig. 1, both samples show metallic $T$ dependence in the whole measurement temperature range. The resistivity drops sharply at 138 and 200 K for BaFe$_2$As$_2$ and SrFe$_2$As$_2$, respectively, which is ascribed to the formation of SDW order.

The optical reflectance measurements were performed on a combination of Bruker IFS 66v/s, 113v, and grating-type spectrometers on newly cleaved surfaces (ab plane) for AFe$_2$As$_2$ ($A = Ba$, Sr) single crystals in the frequency range from 40 to 50,000 cm$^{-1}$. An in situ gold and aluminum overcoating technique was used to get the reflectivity $R(\omega)$. The real part of conductivity $\sigma_1(\omega)$ is obtained by the Kramers-Kronig transformation of $R(\omega)$.

The main panels of Figs. 1(a) and 1(b) focus on the low frequency $R(\omega)$ up to 1800 cm$^{-1}$. For both compounds, $R(\omega)$ exhibits a metallic response and approaches to unity at zero frequency. The most prominent feature is a substantial suppression in $R(\omega)$ for $T < T_{SDW}$, which is strong optical evidence for the formation of energy gaps. The low-$\omega$ reflectance increases faster towards unity at zero frequency. The most prominent feature is a sub-gap structure at $\omega = 0$. For both compounds, $R(\omega)$ can be seen for $T < T_{SDW}$ and the compounds are still metallic below $T_{SDW}$. The change of $R(\omega)$ from an overdamped linear-$\omega$ dependent behavior to a well-defined reflectance edge upon cooling the sample into a SDW ordered state immediately suggests a dramatic reduction of the carrier scattering rate, while its low-energy location implies a considerable reduction of carrier density. A quantitative analysis will be given below.

It is noted that the low-$T$ $R(\omega)$ displays an almost linear-$\omega$ dependence over a certain frequency range below the suppression. This special shape of the suppression leads to the two-peak structure in optical conductivity.

The middle panels of Fig. 1 show the conductivity spectra $\sigma_1(\omega)$ below 2500 cm$^{-1}$. The Drude-like conductivity can be observed for all spectra at low frequencies. For BaFe$_2$As$_2$ ($T_{SDW} = 138$ K), a weak feature around 890 cm$^{-1}$ develops for $T = 130$ K in $\sigma_1(\omega)$, and then the spectra are severely suppressed at low frequencies for 60 and 10 K, resulting in a pronounced double-peak character at 360 and 890 cm$^{-1}$. Associated with the low-$\omega$ reflectance edge, a very sharp and narrow Drude component emerges below the double peaks. Very similar features can be seen for SrFe$_2$As$_2$ crystal, but the double-peak features appear at higher energies, i.e., 500 and 1360 cm$^{-1}$, consistent with the higher $T_{SDW}$ for SrFe$_2$As$_2$. The electrodynamics of broken symmetry ground states, such as the superconducting and density wave states, have been well explored and understood. Because of different coherence factors, a density wave state behaves different from an s-wave superconductor at the gap frequencies in optical conductivity. In an s-wave superconducting state at $T = 0$, the absorption smoothly rises at the gap frequency, while for a density wave state, a sharp maximum appears in conductivity at the gap frequency [22,23]. Based on those studies, we can identify the double-peak energies as the two SDW gaps. The observation of two distinct SDW gaps should be associated with different Fermi surfaces and reflect the multiband property in a FeAs-based compound. From the gap values and SDW

![Diagram](257005-2)
transition temperatures, we obtained the ratio of $2\Delta/k_BT_{SDW} = 3.5–3.6$ for the smaller gap and 9–9.6 for the larger gap for the two compounds. The smaller gap coincides roughly with the gap value expected by the conventional BCS relation, while the large one is very different. Similar two-gap behaviors were also found in optical measurement for the itinerant SDW metal Cr [24].

Because of the presence of two different gap values in the SDW ordered state, direct information on where the FS sheets are gapped is highly desired. Very recently, two angle resolved photoemission spectroscopy (ARPES) investigations on BaFe$_2$As$_2$ single crystals were reported; however, they yielded completely different results from optics. Yang et al. [25] reported a complete absence of gap opening for all bands at the Fermi level. Liu et al. [26] observed a small circular-shaped hole pocket centered at $\Gamma$ and a large electron pocket at the $X$ point below $T_{SDW}$ (at 100 K) on BaFe$_2$As$_2$ which matches well with full-potential linearized plane wave calculations. However, the gap opening is also absent in this work. Apparently, the available ARPES data are in sharp contrast to our results. We emphasize that optical measurement is a bulk probe, while ARPES measurement strongly depends on the surface quality. We also noted that the large area of FS seen by Liu et al. below $T_{SDW}$ is incompatible with the quantum oscillation experiment which reveals a rather small residual FS (occupying only 2% of the Brillouin zone) [27]. More experiments are needed to resolve the inconsistency.

It is well known that the low-$\omega$ Drude component comes from the itinerant carrier contribution. The Drude spectral weight determines the $\omega_p^2$ ($\omega_p$ is the plasma frequency), which is proportional to $n/m_{\text{eff}}$ (where $n$ is the carrier density and $m_{\text{eff}}$ is the effective mass), while its width reflects the carrier scattering rate $1/\tau$. To qualitatively analyze the $T$ evolution of the free-carrier component, we fit the optical response by the Drude-Lorentz model for the whole $T$ and frequency range:

$$\epsilon(\omega) = \epsilon_\infty - \frac{\omega_p^2}{\omega^2 + i\omega/\tau} + \sum_{i=1}^N \frac{S_i^2}{\omega^2 - \omega_i^2 - i\omega/\tau_i}.$$ (1)

Here $\epsilon_\infty$ is the dielectric constant at high energy, and the middle and last terms are the Drude and Lorentz components, respectively. We use one Drude component for the free-carrier response and several Lorentz terms to fit the high frequency $\sigma_1(\omega)$, including the double-peak SDW gap below $T_{SDW}$, a pronounced midinfrared feature near 5000 cm$^{-1}$ [Fig. 2(b)], and some Lorentz components above 16 000 cm$^{-1}$ for interband transitions. The experimental $\sigma_1(\omega)$ data could be well reproduced by the fit [see the insets in Figs. 1(c) and 1(d)].

We are now mainly concerned with the evolution of the itinerant carriers. For BaFe$_2$As$_2$, the plasma frequency $\omega_p = 12 900$ cm$^{-1}$ and scattering rate $1/\tau = 700$ cm$^{-1}$ at 300 K reduce to 4660 and 55 cm$^{-1}$ at 10 K, respectively. For SrFe$_2$As$_2$, $\omega_p = 13840$ cm$^{-1}$ and $1/\tau = 950$ cm$^{-1}$ at 300 K reduce to 4750 and 40 cm$^{-1}$ at 10 K, respectively. Figures 1(e) and 1(f) show the variations of $1/\tau$ and $\omega_p^2$ with temperature for the Drude term. Both parameters are normalized to their 300 K value. Provided the effective mass of itinerant carriers does not change with temperature, then the residual carrier density is only 12% of that at high temperature for both compounds. This means that roughly 88% of the FS is removed by the gapping associated with SDW transitions. On the other hand, the scattering rate was reduced by about 92%–96%. Therefore, the opening of the SDW partial gap strongly reduces the scattering channel, leading to a metallic behavior with enhanced dc conductivity in the gapped state.

The above observations have important implications for the driving mechanism of SDW instability. As mentioned above, the key issue here is whether an itinerant picture based on FS nesting or a local superexchange mechanism is a proper approach. Our optical studies clearly demonstrate that the parent compound has a high itinerant carrier density with the plasma frequency a bit higher than 1.5 eV before SDW transition and is rather metallic both above and below SDW ordering temperatures. Furthermore, the partial gap openings below SDW ordering temperatures are consistent with the expectation of a nesting scenario where the temperature dependence of the gap should resemble

![Figure 2](color online). (a) $R(\omega)$ for BaFe$_2$As$_2$ below 8000 cm$^{-1}$. Inset: $R(\omega)$ below 50 000 cm$^{-1}$. (b) $\sigma_1(\omega)$ below 10 000 cm$^{-1}$ (together with the low-$\omega$ extrapolation based on the dc conductivity). Inset: $\sigma_1(\omega)$ below 20 000 cm$^{-1}$ ($T = 300$ K), the intra- and interband terms, and the midinfrared peak from the Drude-Lorentz fit. (c) The spectral weight below 10 000 cm$^{-1}$. Inset: The spectral weight up to 50 000 cm$^{-1}$. 

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that of BCS theory. On this basis, we think that the local picture is less favored, and the itinerant scenario provides a more reasonable explanation for the driving mechanism.

Besides the dramatic spectral change at low frequencies, both compounds display a very similar and pronounced spectral feature at the midinfrared region. Take BaFe$_2$As$_2$ as an example; the midinfrared component takes up a large spectral weight as shown in the inset in Fig. 2(b). The peak at such a high energy is usually ascribed to the interband transition. However, a puzzling problem is that the spectra below the peak energy exhibit an apparent $T$ dependence that $R(\omega)$ is obviously suppressed with decreasing $T$ below 5000 cm$^{-1}$ [Fig. 2(a)]. Such a gaplike feature is present at all temperatures. Indeed, an analysis for the spectral weight in $\sigma_1(\omega)$ revealed that the suppressed spectral weight below 5000 cm$^{-1}$ is transferred to higher energies. Figure 2(c) plots the spectral weight for $\sigma_1(\omega)$ at 10 and 300 K. Below 200 cm$^{-1}$, the narrowing Drude peak with decreasing $T$ [see the extrapolation data in Fig. 2(b)] yields a growing dc conductivity and thus a larger low frequency spectral weight for $T = 10$ K. Above 200 cm$^{-1}$, the SDW double gap develops which strongly reduces the low $T$ Drude weight, leading to the first suppression below 1000 cm$^{-1}$ in Fig. 2(c). The lost Drude weight fills into the SDW double peak, and the total spectral weight is almost recovered around 2000 cm$^{-1}$ for 10 K. Then the $T$-dependent suppression before the midinfrared peak results in the second spectral weight suppression at 10 K near 3000 cm$^{-1}$. The lost weight finally recovers at about 8000 cm$^{-1}$. We would like to emphasize that this gaplike feature is not directly related to the SDW order. This is because (i) the midinfrared suppression feature is present above $T_{\text{SDW}}$, (ii) similar features exist for K- or Co-doped superconducting samples where the SDW order and the associated low-energy gap structures are completely absent [28], and (iii) the energy scale is much larger than the SDW gaps.

Usually, a gap formation is associated with a broken symmetry state. As the AFe$_2$As$_2$ compounds are in their paramagnetic phase with a tetragonal crystal structure above $T_{\text{SDW}}$, both magnetic and crystal structures are in a very high symmetry state, and one would hardly expect an even higher magnetic or crystal structural symmetry at higher temperatures. Definitely, further experimental and theoretical works are necessary to understand this high energy gaplike behavior.

In summary, the ab-plane optical measurements of AFe$_2$As$_2$ ($A=$ Ba, Sr) single crystals were performed. For the SDW state, our findings indicate that the Fermi surface is largely affected by the SDW transition. Based on a Drude-Lorentz model, we estimate that about 88% itinerant carriers were removed by the gapping of Fermi surfaces. Meanwhile, the carrier scattering rate was reduced by 92%–96%. More importantly, we find two distinct gaps in the SDW ordered state in AFe$_2$As$_2$. For the high frequency regions, $R(\omega)$ show anomalous $T$ dependence even above $T_{\text{SDW}}$ near the 5000 cm$^{-1}$ midinfrared component, which suggests the role of Fe $3d$ electrons in the AFe$_2$As$_2$ system might be very complex and unusual. Since AFe$_2$As$_2$ ($A=$ Ba, Sr) systems have a rather high conducting carrier density at high $T$, a driving mechanism based on an itinerant picture for the SDW instability is thus favored.

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