Formation of partial energy gap below the structural phase transition and the rare-earth element substitution effect on infrared phonons in ReFeAsO (Re=La, Nd, and Sm)

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Single crystals of LaFeAsO, NdFeAsO, and SmFeAsO have been prepared by means of a NaAs flux growth technique and studied by optical spectroscopy measurements. We show that the spectral features corresponding to the partial energy gaps in the spin-density-wave (SDW) state are present below the structural phase transition. This indicates that the electronic state below the structural phase transition is already very close to that in the SDW state. We also show that in-plane infrared phonon modes display systematic shifts towards high frequency upon rare-earth element substitutions for La, suggesting a strong enhancement of the bonding strength. Furthermore, an asymmetric line-shape of the in-plane phonon mode is observed, implying the presence of an electron-phonon coupling effect in Fe-pnictides.

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I. INTRODUCTION

The discovery of superconductivity at 26 K in F-doped LaFeAsO has created tremendous interests in the scientific community. Shortly after this discovery, the superconducting transition temperature \( T_c \) was raised beyond 50 K through the substitution of La by rare-earth elements. \( T_c \) is found to be 41 K for F-doped CeFeAsO, 52 K for F-doped NdFeAsO and 55 K for F-doped SmFeAsO. In the so-called 1111 structural type series, the undoped parent compounds were commonly found to have a spin-density-wave (SDW) ground state with a collinear antiferromagnetic (AFM) spin configuration. A structural phase transition occurs prior to the magnetic transition. Because superconductivity appears in the vicinity of the magnetic ordered phase, it is widely believed that the spin fluctuations play a crucial role in the superconducting pairing of electrons, while the electron-phonon interactions could not explain the superconductivity in those materials. A theoretical calculation even indicates that the electron-phonon coupling could not lead to the superconductivity with \( T_c \) higher than 1 K in those materials.

A critical question here is why the rare-earth element substitutions for La in the 1111 series could significantly enhance the superconducting transition temperature? For La or different rare-earth based parent compounds, the structural and magnetic phase transitions were found to take place roughly at the same temperatures, then the magnetic interactions should not have much difference. On the other hand, the rare-earth element substitutions lead to certain change in the lattice structure. Some correlations between superconducting transition temperatures and the Fe-As bond length or the Fe-As-Fe angle for the series have been found. However, there is still a lack of a complete understanding of the problem. In this work, we performed optical spectroscopy study on single crystal samples of the several different 1111 compounds, including LaFeAsO, NdFeAsO, and SmFeAsO. We focus our attention on two issues. One is the relationship between the structural and magnetic phase transitions. We identify that the charge excitation gaps have already opened at the structural phase transition, which then discloses the intimate relation between the driving mechanisms of the two transitions. The other is the effect of rare earth element substitutions on the lattice modes. We found that the in-plane phonon modes display systematic shifts towards high frequency with the substitution of La by Nd, and Sm. We also observed asymmetric line-shape of the in-plane phonon mode, providing direct evidence for the presence of electron-phonon coupling. The results may imply that the lattice vibrations play some role for the high temperature superconductivity.

II. EXPERIMENTS AND RESULTS

A. Experiments

The crystal growth in 1111 systems has been proven to be difficult. For a long period, only very small size single crystals could be obtained with typical dimensions less than 300 \( \mu \text{m} \). Millimeter-sized single crystals were obtained only by means of NaAs flux technique until recently. The ReFeAsO (Re=La, Nd, Sm) single crystals used in this study were grown from such a technique, and characterized by the X-ray diffractions (XRD) and dc resistivity measurements. For the Sm-based sample, we put 10% F in the initial composition. However the resultant crystals still have a property of the parent compound. The major change is that the resistivity anomaly corresponding to structural transition is suppressed to about 120 K. Detailed descriptions about crystal growth could be found in Ref. [13].

Figure 1 shows the (00l) XRD patterns for the sin-
single crystal samples of ReFeAsO (Re=La, Nd, Sm) with Cu Kα radiation. The XRD patterns indicate that the samples are of the characteristic of good crystallization along the c axis. The (00l) peaks shift towards higher 2θ angles systematically from LaFeAs to NdFeAsO and SmFeAsO, suggesting a reduction of c-axis lattice parameter. The obtained c-axis lattice parameter is c=8.758 Å for LaOFeAs, 8.595 Å for NdOFeAs, and 8.497 Å. The reduction of the c-axis lattice parameter is apparently due to the smaller ionic radius of the rare earth elements Nd and Sm. Those values are consistent with earlier lattice parameters determined from the polycrystalline samples.

Optical measurement was done on a Bruker IFS 66v/s spectrometer in the frequency range from 40 to 25000 cm\(^{-1}\). An in situ gold and aluminum overcoating technique was used to get the reflectance \(R(\omega)\). The real part of conductivity \(\sigma_1(\omega)\) is obtained by the Kramers-Kronig transformation of \(R(\omega)\).

### B. The electronic spectra

Figure 2 shows the optical reflectance and conductivity spectra up to 6000 cm\(^{-1}\) of LaFeAsO, NdFeAsO and SmFeAsO at two representative temperatures 300 K and 10 K, respectively. The data of LaFeAsO were already presented in ref. 13 where its spectral features and correlation effect were discussed. Here they are used for a purpose of comparison. We can see that all the compounds exhibit similar spectral features in both the paramagnetic and SDW states. At room temperature, the reflectance \(R(\omega)\) drops almost linearly with frequency at low-\(\omega\) region, then merges into the high values of a background contributed mostly from the interband transitions from the mid-infrared to visible regime. The conductivity spectrum \(\sigma_1(\omega)\) displays a Drude-like component at low frequency and a rather pronounced spectral weight in the high frequencies. A broad peak is seen near 4500 cm\(^{-1}\). In the SDW state at 10 K, dramatic spectral change was seen at low frequencies. The reflectance below 1000 cm\(^{-1}\) shows a remarkable suppression, while it is enhanced between 1000 and 2000 cm\(^{-1}\). Then, in the conductivity spectra, the spectral weight is severely suppressed at low frequencies and the missing spectral weight is transferred to absorption peaks at high energies. This gives optical evidence for the gap formation on the Fermi surfaces in the SDW state. Since the reflectance still increases fast toward unity at lower frequencies, residual free carriers or Drude component is still left in the magnetic ordered state. Therefore, the Fermi surfaces are not fully gapped below \(T_N\). Those spectral features are also observed in 122-type AFe\(_2\)As\(_2\) (A=Ba, Sr\(^{15}\)) and 111-type Na\(_{1-x}\)Fe\(_3\)As single crystals\(^{16}\) across the SDW transitions, indicating the generic properties of undoped FeAs-based systems.

As mentioned in the introduction, for 1111 parent compounds, a structural transition from a tetragonal-to-orthorhombic phase occurs prior to the SDW or magnetic transition, whereas the two transitions occur simultaneously for 122-type compounds. The relation between the structural and SDW transitions has been a subject of many discussions\(^{17–20}\). Theoretically, it was suggested that the structural transition is driven by the magnetic transition\(^{17–20}\). Then it is important to detect experimentally whether the charge excitation gaps open in the structurally distorted phase (above the magnetic transition) or not, an issue which has not been addressed by any of previous optical measurements. For this purpose, we present a detailed temperature-dependent \(R(\omega)\) and \(\sigma_1(\omega)\) spectra for one of the parent compounds, Nd-
FeAsO.

Similar to other 1111-type compounds, NdFeAsO exhibits a pronounced anomaly in the resistivity $\rho(T)$ near 150 K, below which $\rho(T)$ decreases rapidly with decreasing temperature. Figure 3 shows the temperature dependent resistivity $\rho(T)$ curve of present single crystal NdFeAsO. Compared with the data of polycrystalline samples, the decrease of the resistivity near 150 K is much faster for single crystal sample. Its temperature derivative, $d\rho(T)/dT$, shows a sharp peak at 140 K. It is well known that the structural phase transition is responsible for the dramatic change in the resistivity curve, while the magnetic transition corresponds to the peak position in $d\rho(T)/dT$. A neutron scattering measurement on polycrystalline NdFeAsO sample indicates $T_{SDW}=141$ K, being close to the peak temperature. The unusual decrease at about 8 K could be attributed to the ordering of the magnetic moment of Nd ions. The overall behavior of the resistivity is very similar to the earlier report on micrometer-sized NdFeAsO crystals.

Figure 4 shows the temperature-dependent $R(\omega)$ and $\sigma_1(\omega)$ spectra of NdFeAsO. The energy gap features are not present above the structural transition (see spectra above 160 K). However, below the structural phase transition, the low frequency spectral weight suppression appears. The spectral features observed at 142 K, a temperature between the structural and magnetic transitions, resemble to that at very low temperatures in the SDW state, although the features are weak at such a high temperature. This clearly indicates that the charge excitation gaps start to open at the structural phase transition, thus demonstrating that the electronic state below near the structural phase transition is already very close to that in the magnetic state. The observation would imply essentially the same driving mechanism for both transitions. Although the long-range magnetic order is not formed at the structural transition, short-range order, either nematic order or very strong spin-fluctuations, would exist between the structural and magnetic transition. We note that, a recent ARPES experiment on a 111-based system NaFeAs, which also has separated structural and magnetic transitions, revealed band-folding effect just below the structural transition; while a NMR study on NaFeAs indicated strong enhancement of magnetic fluctuation. Both are consistent with our optical measurement on 1111 systems.

C. The phonon spectra

Despite the fact that the electronic spectra display similar features for La or different rare-earth substituted 1111 parent compounds, we found that there exists a systematic frequency shift of infrared active in-plane phonon modes, which is the focus in the rest of the present paper. Figure 5 shows the low frequency conductivity spectra of the ReFeAsO (Re=La, Nd, Sm) crystals at 10 K, where two phonon modes could be seen in the frequency range. We shall show later that those phonon peaks do not show clear frequency shift across the structural and magnetic transitions.

It is known that ReFeAsO crystalizes in the tetragonal $P4/nmm$ structure at high temperature. According to symmetry analysis, there are six infrared active phonon modes $A_{2u}$, $E_u$, $E'_u$, $E''_u$, $E''_g$, $E'_g$ along the $c$-axis and in $ab$ planes.
Fe-As ions. The assignments of the two modes should be reversed. The crystal and the polycrystalline samples indicates that the A comparison of the phonon modes between the single only observe two E modes near 94 and 253 cm$^{-1}$ for LaFeAsO in the ab plane response. It is noted that the 248 cm$^{-1}$ mode seen in polycrystalline samples was assigned to the $A_{2u}$ mode along the c-axis in early studies, while another mode with slightly higher frequency at 266 cm$^{-1}$ was assigned to the in-plane $E_u$ mode$^{17,28}$. A comparison of the phonon modes between the single crystal and the polycrystalline samples indicates that the assignments of the two modes should be reversed. The 248 cm$^{-1}$ $E_u$ mode involves the in-plane displacement of Fe-As ions. The 339 cm$^{-1}$ mode, which is particularly strong in intensity, is associated with the in-plane displacement of the O-La(Re) bonds. Its high energy location indicates that this mode is mostly oxygen derived$^{29}$. Several interesting features about the phonon modes are found from Fig. 5: (1) both in-plane $E_u$ modes shift to higher frequencies in Nd- and Sm-based samples. The shift is more pronounced for the oxygen-derived mode; (2) The strength of Fe-As mode increases from LaFeAsO to NdFeAsO and SmFeAsO, while the intensity of the O-Re (Re=La, Nd, Sm) mode decreases; (3) There exist asymmetric line-shapes for the phonon modes (Fano line shape$^{30}$), evidencing sizable electron-phonon coupling effect. An expanded frequency plot for the Fe-As mode of SmFeAsO sample is shown in the inset of Fig. 5.

The increase of the phonon frequency in rare-earth element substituted systems strongly indicates an enhancement of the in-plane Fe-As and O-Re bonding strengths. With the substitution of La by Nd and Sm, the masses of the rare-earth ions increase. If there was no bonding strength change, we would expect a reduction of frequencies of the in-plane O-Re stretching mode. The noticeable increase of the mode frequency suggests a reduction of the bond-length, and it overcomes the effect of the mass change of the rare-earth ions. Not only the O-derived mode shows a substantial increase, but the in-plane Fe-As mode also shows a hardening from La to Nd and Sm substitutions, although less pronounced. It is worth noting that the Raman scattering measurements also indicate a strong hardening of the Raman active O modes.$^{27,28}$ Overall, the observations are consistent with the structural characterization data which show a decreasing of the Fe-As bond length$^{29}$. The above result is not affected by the F-doping, since low level F-doping is found to have little influence on the phonon frequencies of the parent compound$^{31}$.

Figure 6 shows the temperature evolution of the $E_u$ phonon modes across the structural and magnetic transitions for the prototype NdFeAsO crystal. Similar to the case of infrared measurement on the polycrystalline samples$^{26}$, we do not observe any splitting or new phonon lines below the structural transition. According to density function theory calculations on LaFeAsO, the splitting of the infrared active phonon modes would be very small and therefore could not be well resolved by the optical measurement$^{17}$. The intensities of the two modes show significant temperature dependence. They are strongly enhanced below the structural or magnetic transition. This behavior is similar to the observation in BaFe$_2$As$_2$ crystal where two infrared active in-plane modes near 94 and 253 cm$^{-1}$ are observed, and the 253 cm$^{-1}$ mode shows a strong intensity enhancement below the structural transition. A natural explanation is that the screening effect from the conduction electrons is significantly reduced due to the partial gapping of Fermi surfaces. Alternatively, it was proposed that the intensity change could be due to a charge redistributions at each atom leading to a change in bonding between different atoms$^{32}$. This point of view might be useful in interpreting the intensity increase of the Fe-As stretching mode but the decrease of the O-La mode from LaFeAsO.

FIG. 5: (Color online) In-plane infrared phonon modes at 10 K for La-, Nd-, and Sm-based 1111 single crystals. Both the Fe-As and O-derived modes show hardening with the rare-earth element substitutions. Inset shows the expanded frequency region for Fe-As mode of SmFeAsO crystal. An asymmetric behavior is clearly seen, indicating strong electron-phonon coupling.

![Infrared phonon modes at 10 K for La-, Nd-, and Sm-based 1111 single crystals.](image1)

FIG. 6: (Color online) The temperature dependence of the in-plane phonon modes of NdFeAsO single crystal. The noticeable increase of the mode frequency suggests a reduction of the bond-length, and it overcomes the effect of the mass change of the rare-earth ions. Not only the O-derived mode shows a substantial increase, but the in-plane Fe-As mode also shows a hardening from La to Nd and Sm substitutions, although less pronounced. It is worth noting that the Raman scattering measurements also indicate a strong hardening of the Raman active O modes. Overall, the observations are consistent with the structural characterization data which show a decreasing of the Fe-As bond length. The above result is not affected by the F-doping, since low level F-doping is found to have little influence on the phonon frequencies of the parent compound.
to NdFeAsO and SmFeAsO.

The above experimental results indicate a good correlation between the bonding length in the crystal structure and the frequencies of the infrared phonon modes. Since the rare-earth element substitutions for La significantly enhance the superconducting transition temperature in doped compounds, it is of worth to further investigate the role played by the electron-phonon coupling for the superconductivity.

### III. CONCLUSIONS

We have grown single crystals of the parent compounds LaFeAsO, NdFeAsO, and SmFeAsO by means of a NaAs flux growth technique and studied their charge excitations and infrared phonon spectra by optical spectroscopy measurement. We identify that the charge excitation gaps are already present below the structural phase transition between the bonding length in the crystal structure to NdFeAsO and SmFeAsO.

The role played by the electron-phonon coupling for the doped compounds, it is of worth to further investigate the enhancement of the bonding strength. The study also yields evidence for the presence of electron-phonon coupling effect in the compounds.

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