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We employ polarization-resolved Raman spectroscopy to study multi-band stoichiometric $T_c$=35 K superconductor CaKFe$_4$As$_4$. We do not detect Pomeranchuk-like electronic nematic fluctuations which were universally observed in XY (B$_2$g) symmetry Raman response for most families of the Fe-based superconductors. In the superconducting state we observe consistent with nodeless order parameter full spectral weight suppression at low energies and emergence of a composite pair-breaking coherence feature at energies between 12 and 20 meV. We analyze the superconductivity induced phonon self-energy effects and give an estimation for the electron-phonon coupling constant $\lambda^f$=0.0015 which is insufficient to provide attraction for high $T_c$ pairing.

Introduction – Understanding the pairing mechanism in the Fe-based superconductors (FeSCs) remains in a focus of research not only due to a high superconducting transition temperature $T_c$, but also because of the interplay of superconductivity with other electronic degrees of freedom, nematicity and magnetism in particular [1–6].

Electronic nematicity has been universally observed for many families of FeSCs [7, 8]. Furthermore, compositions that are in proximity to the nematic quantum critical point (QCP) often show the highest $T_c$ [7–11]. The $T_c$ enhancement near QCP scenario was recently studied by several theoretical proposals [12–14].

Raman spectroscopy has been often used to study the superconducting (SC) order parameter [11, 15–19], the dynamics of electronic nematic fluctuations [10, 11, 20, 21], as well as the electron-phonon interactions [22–25]. The FeSCs are well suited to study the mechanisms of $T_c$ enhancement from an experimental point of view.

Recent discovery of a new class of stoichiometric and strictly tetragonal superconductors Ca$A$Fe$_4$As$_4$ ($A$ = K, Rb, Cs) with rather high $T_c$ (31-36 K) provides an ideal platform for spectroscopic investigation of FeSCs in a clean limit, and to decide if the nematicity is a necessary condition for high $T_c$ superconductivity in FeSCs [26–30]. Although the pristine Ca$A$Fe$_4$As$_4$ compounds do not show a long range magnetic order, an electron doping via substitution of Co or Ni for Fe, and/or application of pressure, suppresses $T_c$ and induces an exotic spin-vortex crystal order, while the system remains tetragonal even for the magnetically ordered phase [31–33].

Crystallographic structure – The CaKFe$_4$As$_4$ structure may be considered as a modification of the extensively studied tetragonal 122 family of FeSCs with the body-centered structure I4/mmm. For CaKFe$_4$As$_4$, every other plane of Ca atom is replaced by K, reducing the crystallographic space group to primitive $P4/mmm$ (point group $D_{4h}$) and therefore doubling the number of atoms in the primitive cell. In Figs. 1(a-b) we compare the tetragonal CaFe$_2$As$_2$ and CaKFe$_4$As$_4$ lattices.

![Figure 1](attachment://fig1.png)

FIG. 1. (a) The comparison between CaFe$_2$As$_2$ and CaKFe$_4$As$_4$ lattice structures. For CaKFe$_4$As$_4$, the Fe sites are shifted away from the high symmetry $z$=1/4 and 3/4 planes, causing two distinct As-Fe bond distances (shown in blue and orange). (b) The reduction of the Fe site-symmetry from $D_{2d}$ for CaFe$_2$As$_2$ to $C_{2v}$ for CaKFe$_4$As$_4$. (c) For CaKFe$_4$As$_4$, a sketch of the partially occupied $d_{xz}$$d_{yz}$ orbital order (the upper panel for real space), and derived two FS pockets in the vicinity of the $\Gamma$ point (the lower panel for momentum space). Solid and dotted lines denote differences in the orbital occupation which induce a static quadrupole moment on the Fe sites with a checkerboard order.

For the body-centered CaFe$_2$As$_2$ structure the Fe layers are confined to the high symmetry $z$=1/4 and 3/4 planes. All the Fe positions have $D_{2d}$ site-symmetry with $S_4$ axis along z-direction, which imposes the degeneracy condition for Fe $3d_{xz}$ and $3d_{yz}$ orbitals. Such orbital degeneracy is a common feature for most of FeSC structures [34, 35]. It has been conjectured that degeneracy of the pa-
tially filled $d_{xz}/d_{yz}$ orbitals is a necessary condition for anomalously strong nematic effects in FeSCs: such degeneracy enables dynamical charge oscillations in sub-THz frequency range giving rise to fluctuating charge ferroquadrupole moment with an amplitude proportional to the local oscillating charge imbalance $n_{xz} - n_{yz}$ [11]. The soft ferro-quadrupole fluctuations often show critical behavior leading to a $d$-wave Pomeranchuk-like instability [11]. These fluctuations most dramatically manifest themselves in the low-frequency part of XY-symmetry Raman response as an overdamped quasi-elastic feature in the normal state [10, 11, 20] which undergoes a metamorphosis into a coherent in-gap collective mode below $T_c$ [11, 17–19, 37, 38].

In contrast to doped CaFe$_2$As$_2$-structure-like Ba$_{1-x}$K$_x$Fe$_2$As$_2$ where K$^+$ substitute ions randomly occupy the host Ba$^{2+}$ sites, for CaKFe$_4$As$_4$ the Ca$^{2+}$ and K$^+$ ions are ordered in two distinct layers above and below Fe-As layers. This layering of Ca$^{2+}$/K$^+$ ions causes the planes of Fe ions shift vertically away from the high symmetry positions [39, 40]. The shift results in nonequivalent Fe-As bond distances for the As atoms above and below the Fe layer, thus, the Fe site symmetry is reduced from $D_{4d}$ to $C_{2v}$, Fig. 1(a-b).

The removal of the $S_4$ symmetry on the Fe sites takes away the degeneracy of the partially occupied Fe $d_{xz}/d_{yz}$ orbitals [39], which gives rise to a static charge imbalance between these two orbitals, and, hence, creates a static $d_{x^2-y^2}$-symmetry quadrupole moment on each Fe site. Because the orbital character of the lower energy state flips between the two neighboring Fe sites, the structure forms a static checkerboard anti-quadrupole order, Fig. 1(c). The stiffness of the static anti-quadrupole order parameter precludes Pomeranchuk-like fluctuations for the CaKFe$_4$As$_4$ compound.

In this work we study the charge dynamics in CaKFe$_4$As$_4$ single-crystals by polarization-resolved Raman spectroscopy. We demonstrate that in contrast to the data from many FeSCs for which strong Pomeranchuk-like electronic nematic fluctuations which give rise to intense electronic Raman continuum in the XY-quadrupole symmetry channel, the electronic Raman signal for CaKFe$_4$As$_4$ is isotropic and generally weak. This data indeed support the notion that the quadrupole fluctuations for CaKFe$_4$As$_4$ structure are quenched. Below $T_c$, we observe complete suppression of the low-frequency spectral intensity indicating that all FS pockets are fully gapped, and transfer of the spectral weight to a composite SC coherence feature between 12 and 20 meV in the XY symmetry electronic Raman response. We also observe superconductivity induced phonon self-energy effect and estimate the electron-phonon ($\epsilon$-$\pi$) coupling to be weak, with coupling constant $\lambda^\prime = 0.0015$.

Experimental – Polarization-resolved Raman scattering measurements were performed in quasi-back scattering geometry from the natural cleaved (001) surface. We use $\mu \nu = XX$, XY and X’Y’ scattering geometries [41]. These geometries allow coupling to $A_{1g} + B_{1g}$ and $B_{2g}$ symmetry excitations correspondingly.

The CaKFe$_4$As$_4$ single crystals ($T_c = 35$ K) used in this study were synthesized by flux method [26, 28]. The crystals were loaded into a continuous helium flow optical cryostat immediately after being cleaved in a connected to the cryostat nitrogen gas glove bag. For excitation we used the 647 nm line of a Kr$^+$ laser, where the laser beam was focused to a $50 \times 50$ $\mu$m spot. The laser power was kept below 10 mW in the normal state and 2.3 mW in the SC state. All referred temperatures are corrected for the laser heating. The Raman signal was collected and analyzed by a triple-grating spectrometer with 1.5 cm$^{-1}$ spectral resolution. Spectra were corrected for spectral response and background determined from the X’Y’ symmetry electronic continuum to obtain the Raman scattering intensity $I_{\mu \nu}(\omega, T)$ [42]. The Raman response function was derived as $\chi''_{\mu \nu}(\omega, T) = I_{\mu \nu}(\omega, T)/(1 + n(\omega, T))$, here $n(\omega, T)$ is the Bose distribution factor.

FIG. 2. Raman spectra for single crystal CaKFe$_4$As$_4$ at 40 and 7 K. The XY and XX polarized spectra are offset by 1 and 2 units, respectively. Inset: top view of the Fe-As layer and axes notation. Raman continuum and the superconducting gap – In Fig. 2 we compare Raman response above and below $T_c$ for three polarizations. Above $T_c$, besides four sharp phonon modes, spectra show a weak featureless electronic continuum for all polarizations. For the XX and X’Y’ scattering geometries, the electronic continuum shows surprisingly little change across $T_c$, while for XY po-
ARPES data [45], is comprised of three pair-breaking coherence peaks due to isotropic gaps on three FS pockets $\chi''(\omega,T) = 4\Delta_i^2/\omega \sqrt{\omega^2 - 4\Delta_i^2}$. $L(\omega, T)$ denotes BCS coherence peaks with SC gap energies $2\Delta_i$ convoluted with $L(\omega, T)$ that accounts for temperature and instrumental broadening, and $\alpha_i$ denotes the weight of Raman coupling [15, 43].

- The second model (in blue) is comprised of a collective mode (CM) due to electron-hole or particle-particle attraction in $d_{xy}$-symmetry channel and a single pair-breaking coherence peak.

The energies resulted from the models are collected in Table I. The models are difficult to tell apart solely based on the Raman data. However, the second model is hard to reconcile with one-electron spectroscopy data [45], and even if the second model containing a CM would apply, the interactions in the sub-dominant $d_{xy}$-wave channel cannot be strong because the mode’s small binding energy. This is in contrast to Ba$_{1-x}$K$_x$Fe$_2$As$_2$ at optimal doping [18, 19], or to Na-111 [11] for which sharp in-gap CMS with significant binding energy have been reported.

**Phonon self-energy effects** – CaKFe$_4$As$_4$ crystal (point group $D_{4h}$) contains 10 atoms in a primitive cell. For the $\Gamma$ point phonons, group-theoretical symmetry decomposition yields $A_u + E_u$ acoustic modes, $4A_{2u} + 5E_u$ infrared active modes, $3A_{1g} + B_{1g} + 4E_g$ Raman active modes, and a $B_{2u}$ silent mode. For the scattering experiments from the (001) surface, we observe all the three $A_{1g}$ and a $B_{1g}$ phonons (Fig. 2). The phonon energies and atomic displacements are summarized in Table II.

Above $T_c$ all modes exhibit a conventional temperature dependence: hardening and narrowing upon cooling due to anharmonic decay [49–51]. However, we note that the behavior for the $B_{1g}$ phonon mode in the SC state is anomalous: the mode’s energy and line width increase upon cooling, as shown in Fig. 4(a).

Similar phonon anomalies upon entering into SC state were reported for MgB$_2$ [25] and for cuprate superconductors [22, 23, 52–55]. The behavior was explained by Zeyher-Zwicknagl’s model [24] which implies that in the presents of electron-phonon coupling, phonon self-energy is upward renormalized when the SC gap opens and the electronic density-of-states is pushed to the proximity of the phonon mode above the gaps energies.

It is interesting to note that in contrast to the 67 cm$^{-1}$ $A_{1g}$ phonon (Fig. 2) that exhibits an asymmetric Fano line shape, the $B_{1g}$ phonon shows nearly perfect Lorentzian shape. We attribute this to weak Raman coupling to the $B_{1g}$ symmetry electronic continuum: the

**TABLE I. Summary of fitting result at 7 K, units in meV**

<table>
<thead>
<tr>
<th>Model 1: three SC gaps</th>
<th>Model 2: CM+SC gap</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta_1 = 13.6 \pm 0.6$</td>
<td>$\omega_{CM} = 16.4 \pm 0.21$</td>
</tr>
<tr>
<td>$\Delta_2 = 16.8 \pm 0.1$</td>
<td>$\Delta_1 = 18.9 \pm 0.4$</td>
</tr>
<tr>
<td>$\Delta_3 = 20.2 \pm 0.2$</td>
<td></td>
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</tbody>
</table>
e-p interaction only renormalizes the phonon self-energy without showing a Fano interference in the spectra.

To quantify the superconductivity induced self-energy effects and the e-p interaction strength, we fit the data with $\chi''(\omega) \propto 4\omega_0\Sigma''(\omega^2 - \omega_0^2 - 2\omega_0\Sigma' + 4(\omega_0\Sigma'')^2)^{-1}$, where $\omega_0$ is the bare phonon frequency, and $\Sigma = \Sigma' + i\Sigma''$ is complex phonon self-energy [24]. Thus, if $\Sigma$ is small, the mode appears at $\omega_{ph} = \sqrt{\omega_0^2 + 2\omega_0\Sigma'}$ frequency. The fitting results are displayed in Figs. 4 (b-c) [51].

We calculate the coupling constant $\lambda_{B1g}^2$, around $\Gamma$ point [56]: $\lambda = -\kappa \sin u/u$, where $\kappa = [(\Sigma'(7K) - \Sigma'(40K)) - i(\Sigma''(7K) - \Sigma''(40K))] / \omega_{ph}(40K)$ and $u \equiv \pi + 2i\cosh^{-1}[\omega_{ph}(40K)/2\Delta]$. Using the energy of the strongest pair breaking peak $2\Delta = 135$ cm$^{-1}$, we acquire weak e-p coupling constant $\lambda_{B1g}^2 \approx 0.0015$. The $A_{1g}$ phonons do not show measurable renormalization, Figs. 4(d-e). Therefore, the $B_{1g}$ mode is the only phonon that exhibits the SC induced self-energy effects. Thus, we use $\lambda_{B1g}^2$ to approximate the total e-p coupling constant $\lambda$ [57].

In comparison, for a conventional phonon-mediated superconductor MgB$_2$ with similar $T_c = 39$ K, a much larger coupling constant $\lambda = 0.2$ was derived from SC induced phonon renormalization [25]. Thus, for CaKFe$_4$As$_4$, the value $\lambda$ is far from being sufficient to result in superconductivity with $T_c$ at 35 K.

Conclusions - In summary, we report polarization-resolved Raman spectroscopic study of the single-crystal CaKFe$_4$As$_4$ superconductor with $T_c$ at 35 K. Above $T_c$ we do not detect Pomeranchuk-like electronic nematic (quadrupole) fluctuations, which implies that the electronic nematicity may not be essential for high-$T_c$ superconductivity in FeSCs.

Below $T_c$ we observe development of a composite coherence feature between 12 and 20 meV in the $B_{2g}$ symmetry channel and a complete suppression of low-frequency spectral weight, which implies that all the FS pockets remain nodeless. We do not detect any sharp in-gap collective modes which were commonly observed for FeSCs which exhibit strong electronic nematic fluctuations.

We also study the SC induced self-energy effects for Raman-active phonons and provide an estimate of the electron-phonon coupling constant $\lambda^2 = 0.0015$, which is very small for a superconductor with $T_c$ at 35 K.

Because both the electronic nematic fluctuations and the superconductivity induced self-energy effects for even-symmetry phononic modes are negligibly small, we conclude that the spin-fluctuations are responsible for primarily pairing interaction in CaKFe$_4$As$_4$. In this case, the expected pairing symmetry is $s_{\pm}$, which is consistent with the observation of the spin-resonance mode at nesting vector $(\pi, \pi)$ by inelastic neutron scattering [58].

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Note added in revision: While this paper was being reviewed, D. Jost et al. published related Raman study [47]. The authors of Ref. [47] interpret features at 134 cm$^{-1}$ (16.6 meV) and at 50 cm$^{-1}$ (6.2 meV) as Bardasis-Schrieffer (BS) collective modes and argue that appearance of the modes imply subdominant $d$-wave pair-

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**TABLE II. Summary of Raman active phonons mode energies and atomic displacements for CaKFe$_4$As$_4$.**

<table>
<thead>
<tr>
<th>Symmetry</th>
<th>Energy at 40 K</th>
<th>Atomic displacements</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A_{1g}(1)$</td>
<td>67 cm$^{-1}$</td>
<td>Fe(z)+As(1)(z)+As(2)(z)</td>
</tr>
<tr>
<td>$A_{1g}(2)$</td>
<td>185 cm$^{-1}$</td>
<td>Fe(z)+As(1)(z)+As(2)(z)</td>
</tr>
<tr>
<td>$A_{1g}(3)$</td>
<td>286 cm$^{-1}$</td>
<td>Fe(z)+As(1)(z)+As(2)(z)</td>
</tr>
<tr>
<td>$B_{1g}(1)$</td>
<td>213 cm$^{-1}$</td>
<td>Fe(z)</td>
</tr>
</tbody>
</table>
ing interactions. We note here that none of the referred features in Raman spectra have a structure of a defined long-lived resonance. As such, assignments to any collective mode is difficult to support. Furthermore, the statement that there exist two independent attractive subdominant XY-symmetry pairing channels is unjustified.

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[40] µν is short for $Z(\mu\nu)Z$ in Porto’s notation.
[41] The background is determined from the XY′ symmetry scattering intensity after subtracting the B1g phonon, see supplement I in [59].
\[ \omega_{ph}(T) = \omega_{ph}(0) - \omega_1 (1 + 2/(\exp(\omega_{ph}(0)/k_B T) - 1)), \Sigma''(T) = \Sigma''(0) + \Gamma_1 (1 + 2/(\exp(\omega_{ph}(0)/k_B T) - 1)) \] are used to fit \( \omega_{ph}(T) \) and \( \Sigma''(T) \) for the normal state [49]. The dashed curves in Figs. 4b-c are plotted with the fitting result \( \omega_{ph}(0) = 216.0 \pm 0.1 \text{ cm}^{-1}, \omega_1 = 3.1 \pm 0.1 \text{ cm}^{-1}, \Sigma''(0) = 2.2 \pm 0.3 \text{ cm}^{-1}, \) and \( \Gamma_1 = 0.6 \pm 0.3 \text{ cm}^{-1}. \)