Phase stability and large in-plane resistivity anisotropy in the 112-type iron-based superconductor Ca$_{1-x}$La$_x$FeAs$_2$

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(Dated: July 2, 2018)

The recently discovered high-T$_c$ superconductor Ca$_{1-x}$La$_x$FeAs$_2$ is a unique compound not only because of its low symmetry crystal structure, but also because of its electronic structure which hosts Dirac-like metallic bands resulting from (spacer) zig-zag As chains. We present a comprehensive first principles theoretical study of the electronic and crystal structures of Ca$_{1-x}$La$_x$FeAs$_2$. After discussing the connection between the crystal structure of the 112 family, which Ca$_{1-x}$La$_x$FeAs$_2$ is a member of, with the other known structures of Fe pnictide superconductors, we check the thermodynamic phase stability of CaFeAs$_2$, and similar hypothetical compounds SrFeAs$_2$ and BaFeAs$_2$ which, we find, are slightly higher in energy. We calculate the optical conductivity of Ca$_{1-x}$La$_x$FeAs$_2$ using the DFT + DMFT method, and predict a large in-plane resistivity anisotropy in the normal phase, which does not originate from electronic nematicity, but is enhanced by the electronic correlations. In particular, we predict a 0.34 eV peak in the $yy$ component of the optical conductivity of the 30% La doped compound, which corresponds to coherent interband transitions within a fast-dispersing band arising from the zig-zag As-chains which are unique to this compound. We also study the Landau free energy for Ca$_{1-x}$La$_x$FeAs$_2$ including the order parameter relevant for the nematic transition and find that the free energy does not have any extra terms that could induce ferro-orbital order. This explains why the presence of As chains does not broaden the nematic transition in Ca$_{1-x}$La$_x$FeAs$_2$.

PACS numbers:

INTRODUCTION

The discovery of high temperature superconductivity in the iron based materials [1] triggered a large number of investigations [2–5]. The basic ingredient of this class of materials is square nets of iron tetrahedrally coordinated by a pnictide or a chalcogenide. By now this ingredient has been realized in multiple prototypical structures, such as the original realization in the 1111 structure, the 112 structure of LiFeAs, the 11 structure such as that of FeSe, and the 122 structure such as in BaFe$_2$As$_2$.

The possibility of superconductivity in iron based compounds with a 112 structure was suggested based on electronic structure calculations in Ref. [6], where it was pointed out that this structure would support metallic spacer layers which could aid in elucidating the mechanism for high temperature superconductivity. Attempts to synthesize iron pnictide compounds in this structure were not originally successful, but new Mn based materials in this structure were found [7, 8] and it was observed theoretically [7] and experimentally [8–11] that the spacer layers exhibit Dirac cones [12]. Recently, Fe superconductors in the 112 structure were synthesized, (Ca,Pr)FeAs$_2$ [13], and Ca$_{1-x}$La$_x$FeAs$_2$ [14].

These materials form in a structure where the As in the CaAs layers are distorted in zig zag chains, i.e. the space group $P2_1$ or $P2_1/m$ rather than the originally assumed tetragonal structure [13, 14]. Second harmonic generation experiments confirmed the space group $P2_1$ for La doped compounds [15]. More recently, theoretical studies focusing on the spacer layers determined that the As $p_x$ and $p_y$ orbitals are responsible for the Dirac cones, and the spin-orbit coupling can open a gap and induce topological phases on these layers, suggesting the 112 compounds as prime candidates for proximity induced topological superconductivity [16, 17].

These works motivate us to revisit the early theoretical predictions in the light of these experimental developments to address some basic questions. (1) The original density functional theory (DFT) calculations focused on the FeAs layers only. A natural question is what would be the result of a full relaxation of the crystal structure? (2) Parent compounds such as CaFeAs$_2$ have so far not been synthesized, and a rare earth is needed to facilitate
the synthesis [18]. Raising the question of relative stability of these compounds, what is the role of the rare earth like La in stabilizing the structure? (3) Photoemission experiments, confirmed the theoretical prediction of the existence of metallic spacer layers (with Fermi pockets of As \( p_z \) and Ca character, in addition to the Dirac cones) [19, 20], however, it is not clear from them what the role of doping is, since in the 112 structure both the CaAs and the FeAs layers can accommodate carriers. (4) It would also be useful to establish consequences of the anisotropy introduced by the formation of the CaAs chains, which should be visible in the optical response, and elucidate how this anisotropy couples to the nematic order parameter whose origin is a subject of intensive discussion. In this paper we answer these questions and determine how this iron pnictide fits with the other families already studied within dynamical mean-field theory (DFT+eDMFT). We conclude that in spite of the strong anisotropy induced by the As zigzag chains, the 112 compounds are very similar to the rest of the iron pnictide superconductors, indicating the superconductivity resides essentially on the FeAs layers unaffected by spacers.

In this work, we (1) present a detailed explanation of the crystal structure of CaFeAs\(_2\), and elucidate the connection of it with other pnictide superconductors, (2) check the phase stability of CaFeAs\(_2\), as well as SrFeAs\(_2\) and BaFeAs\(_2\) at the level of DFT by finding the optimum first principles structure via an evolutionary structure search, and then building the convex hull, (3) systematically study the electronic structure of Ca\(_{1-x}\)La\(_x\)FeAs\(_2\) using state of the art DFT + embedded DMFT (eDMFT) to understand the effect of carrier doping, and (4) for the first time predict the optical conductivity of this compound at the level of DFT + eDMFT. We show that the presence of metallic As chains on the spacer layer gives rise to a strong in-plane anisotropy even at high temperatures, distinct from that driven by the nematic transition. Comparing DFT and eDMFT results, we further show that this anisotropy is enhanced by the electronic correlations.

**METHODS**

DFT calculations are performed using the projector augmented wave method as implemented in the Vienna Ab Initio Simulation Package (VASP) [21–24]. A plane wave cutoff of 500 eV and the generalized gradient approximation of Perdew-Burke-Ernzerhof (PBE) are used [25].

To obtain the lowest energy structures, an \textit{ab initio} evolutionary search algorithm [26] as implemented in the USPEX package [27] is used in conjunction with VASP. In these calculations, initial structures are randomly generated and the next generations are generated using mutations of the previous ones and new random structures. Once the stable structural phases are obtained from the evolutionary search, corresponding formation energies are calculated with a dense Monkhorst-Pack sampling grid with a resolution of \(2\pi \times 0.02\text{Å}^{-1}\) for the \(k\)-space integrations.

For the phonon dispersion calculations, the PHONOPY code [28] is used to build the supercells, and to find the minimum number of ionic displacements required. The direct method is used to calculate the forces in these supercells, as opposed to the Density Functional Perturbation Theory. The force constants and the dynamical matrices are obtained from the Hellmann-Feynman forces in these \((2 \times 2 \times 2)\) supercells (16 formula units). A \(6 \times 6 \times 2\) \textit{k}-point mesh is used in these supercell calculations.

To treat correlation effects in the Ca\(_{1-x}\)La\(_x\)FeAs\(_2\) compounds, the fully charge self-consistent scheme DFT + eDMFT [29, 30] (for a review see Ref. [31]) is used as implemented in Ref. [32, 33] with the hybridization expansion continuous-time quantum Monte Carlo [34] as the impurity solver as implemented in Ref. [35]. We use the Coulomb interaction \(U = 5.0\) eV and the Hund’s coupling \(J = 0.8\) eV, which were shown to describe Ca\(_{1-x}\)La\(_x\)FeAs\(_2\) compounds [19, 36]. The temperature is set to \(T = 116\) K. The experimental crystal structure determined by X-ray diffraction [14] is used. The virtual crystal approximation is adopted to investigate the electronic structure of the La doped compounds.

**CRYSTAL STRUCTURE OF CaFeAs\(_2\) AND ITS CONNECTION TO OTHER PNICTIDES**

In this section, we explain the details of the crystal structure of Ca\(_{1-x}\)La\(_x\)FeAs\(_2\) and its connection to the 1111 family of superconductors with the ZrCuSiAs-type

![FIG. 1: (Color Online) (a) Tetragonal crystal structure with \(P4/mnm\) of CaFeAs\(_2\). (b) Monoclinic crystal structure with \(P2_1/m\) of CaFeAs\(_2\). The black lines represent the unit cell in both (a) and (b). The As square-net is clearly shown in (a), whereas, the zig-zag As chain appears in (b). Both spacers, As square-net and the zig-zag As chain, are metallic.](image-url)
structure (see Fig. 2 and Table I). The ZrCuSiAs-type structure has the simple tetragonal space group $P4/nmm$ (number #129) with two formula units per primitive unit cell. In the superconducting pnictides with this structure, the Fe ion occupies the same Wyckoff position as Cu in ZrCuSiAs, $2b$ ($1/4, 3/4, 1/2$) with no internal parameter, and forms a layer with the pnictogen in the same Wyckoff position as As in ZrCuSiAs, $2c$ ($1/4, 1/4, z$). The internal parameter $z$ determines the pnictogen height, which is shown to be sensitively linked to superconducting $T_c$ in these compounds (Fig. 14). In this layer, the Fe cations form a square plane with pnictogen ions above and below. The pnictogen ions form edge-sharing tetrahedra around the Fe ions. The other two ions are a cation on the $2c$ position and an anion on the $2a$ ($3/4, 1/4, 0$) position. While in certain compounds these two ions can form a tightly bound layer together, depending on the particular atoms there may or may not be any significant bonding between them, and this gives rise to a wide range of internal parameter $z$ for the cation on the $2c$ position.

Having the same anions on both $2c$ and the $2a$ positions in the ZrCuSiAs-type structure gives rise to the HfCuSi$_2$-type structure. Note that in this structure, while both anions (Si) are the same, their environments are significantly different. The $2a$ position has both atoms in one unit cell on the same layer (with coordinates $(3/4, 1/4, 0)$ and $(1/4, 3/4, 0)$), but the $2c$ position corresponds to atoms on two separate layers (with coordinates $(1/4, 1/4, z)$ and $(3/4, 3/4, -z)$) with the Fe (Cu in HfCuSi$_2$) layer in between. The anions on the $2a$ site form a dense square net, and often have strong covalent bonds between themselves, whereas the anions on the $2c$ site hybridize with the transition metals on the $2b$ site. Compounds with the HfCuSi$_2$-type structure has been studied extensively (for example see [37, 38]), and hypothetical BaFeAs$_2$ and BaFeSb$_2$ with this structure has been proposed to be high $T_c$ superconductors if synthesized [6]. The physics that emerges from this square net of covalently bound anions has drawn particular attention in the literature. It has been shown that this layer can support conventional superconductivity [39], as well as Weyl semimetallic phases [40]. The wide bands that are formed from the $p$ orbitals of the main-block elements on the square net also commonly give rise to Peierls type instabilities that lead to the distortions of the crystal structure [41], which have been studied in detail in various compounds with a similar structure, including various arsenides [42–45]. These studies show that the distortions on the As layers can be tuned by both doping and pressure, and studying the similar distortion present in CaFeAs$_2$ and its effect on superconductivity can be important.
There are two very similar structures reported in the literature for lanthanide doped CaFeAs$_2$: A noncentrosymmetric structure with space group $P2_1$ (number #4) and a centrosymmetric structure with space group $P2_1/m$ (number #11). (The former structure is reported, for example, for the La doped compound in Ref. [14], and the latter is reported, for example, for the Pr doped compound in Ref. [13]. While second harmonic generation verifies the lack of inversion symmetry in the La doped compound [15], no similar study exists for the Pr doped compound.) There is a group-subgroup relationship between these two space groups, and the only difference between the structures is a polar distortion, discussed at the end of this subsection.

The main feature that differentiates the structure of Ca$_{1-x}$RE$_x$FeAs$_2$ from the HfCuSi$_2$-type structure is a Peierls type distortion on the As square net as well (Fig. 5). This distortion is at the $\Gamma$ point of the unit cell and transforms as the $\Gamma_5^+$ irreducible representation (irrep) (Fig. 4 (d)). (Throughout this study, we label the irreps corresponding to structural distortions using their labels for the space group #129.) This distortion form zig-zag chains from the As atoms, and decreases the symmetry of the crystal significantly. (Even though this distortion also displaces the Fe atoms to form Fe-Fe chains in a similar fashion, and the Ca ions are also displaced, the amplitude of these displacements, though nonzero by symmetry, are found to be very small in the experimental structure and we hence ignore them.) The $\Gamma_5^+$ distortion by itself breaks the four-fold rotational symmetry, and renders Ca$_{1-x}$RE$_x$FeAs$_2$ unique in the sense that it is the only pnictide superconductor with broken four-fold rotational symmetry even above the nematic transition. (The effect of this symmetry breaking (or lack thereof) on the nematic transition will be discussed later.)

The structure that is obtained from the high symmetry reference #129 by freezing in the $\Gamma_5^+$ has the space group $P2_1/m$ (#11). This symmetry is low enough that a distortion that transforms as another irrep, $\Gamma_5^-$ (Fig. 4(b)), can also have a nonzero amplitude without reducing the symmetry any further. This irrep involves Fe and As (which is in the CaAs layer) layers' rumpling: For example the two Fe ions in the same unit cell are displaced by $\sim$ 0.01Å in opposite directions along the c axis. While in general it might be important that the Fe ions no longer form a perfect plane, this rumpling is so small that it can safely be ignored. The small amplitude of this rumpling also suggests that there is no driving force for the
The structure that is obtained by the $\Gamma_5^+$ and $\Gamma_5^-$ distortions starting from the tetragonal HfCuSi$_2$-like reference structure is the centrosymmetric structure observed for some of the rare earth doped CaFeAs$_2$ compounds. But, as mentioned before, some of these compounds actually have even lower symmetries. The irrep that corresponds to the other structural distortion present in these structures is the in-plane polar irrep $\Gamma_5^-$ (Fig. 4(c)). This distortion is an overall displacement of all the atoms parallel (or antiparallel) to each other and breaks the inversion symmetry. While, by symmetry, every ion is displaced by this irrep, a symmetry mode amplitude analysis of the experimental structure indicates that this displacement is more than 80% on the Ca layer, i.e. it is the Ca (and the La atoms) that are displaced by far the most according to this irrep. This is a very interesting observation given the fact that Ca is by far the most electropositive element in this compound and has a closed shell configuration, and as a result, is not chemically active and should not be the driving force of a polar distortion. It is neither in a particular coordination geometry that is known to give rise to some type of geometric-ferroelectricity, which would be robust even though the compound is conducting [46, 47].

Given the fact that this polar distortion is quite small on the Fe and As layers, we do not analyse its effect on the electronic structure in detail, and consider a complete study of the driving force behind it beyond the scope of this work. However, in passing, we note that the evolutionary structure search we performed for CaFeAs$_2$, discussed in the following sections, does not predict a polar ground state. Given the electropositivity of La, it would be surprising if it was the driving ion for polarization in this compound. The only possibility left seems to be that the ordering of the La ions in the Ca layer gives rise to the polar distortion. This possibility is consistent with the fact that the ionic radius of Pr is closer to Ca than La and therefore it is less likely to cause cation ordering when substituting for Ca, however, these differences are all very small and it is not possible to go beyond the level of speculation at this point.

Lack of tetragonal symmetry and relation to electronic nematicity

Figure 3 shows various irreducible representations that correspond to structural distortions from the tetragonal $P4/nmm$ structure of Ca$_{1-x}$La$_x$FeAs$_2$. Even though there are orthorhombic phases listed in this figure, these phases are different from (i.e. they belong to different space groups) the orthorhombic phases realized below the structural transitions in, for example, LaFeAsO, where the transition is from the tetragonal $P4/nmm$ to the orthorhombic $Cmna$ (# 67) structure [48]. Similarly, in the 122 compounds that have a tetragonal structure with the ThCr$_2$Si$_2$ type (space group $Fmmm$) at high temperature, an electronic nematic phase breaks the four-fold rotational symmetry and hence induces the structural transition to an orthorhombic space group [49–51]. In the case of Ca$_{1-x}$La$_x$FeAs$_2$, the monoclinic structure is stable up to at least 450 K [14], and no transition to the tetragonal phase is reported. Given the one order of magnitude lower critical temperature of the nematic transition in other iron pnictides, the monoclinic phase in CaFeAs$_2$ certainly does not have a nematic origin. As we discussed before, it is most likely due to a Peierls-like mechanism, active on the As layer that forms the zig-zag chains.

Ca$_{1-x}$La$_x$FeAs$_2$ also undergoes a structural transition near its antiferromagnetic ordering temperature [19]. This transition is from monoclinic to triclinic symmetry. However, the argument for electronic nematicity is not straightforward for this transition either because the four-fold symmetry is already broken above this transition, in the monoclinic structure. We address this problem in the final section and show that the free energy expression for CaFeAs$_2$ does not contain any terms that
can give rise to a different character of the nematic transition than the other iron pnictide superconductors. This justifies the studies such as references [52–54] which compare the phase diagram of this compound with other iron pnictide compounds.

FIRST PRINCIPLES RESULTS

Crystal Structure of CaFeAs$_2$

The hypothetical, tetragonal high-symmetry phase of CaFeAs$_2$ has the P4/nmm symmetry (Fig. 1(a)), and it is the same structure as the one proposed by Shim et al. [6] for the Ba compounds with the same stoichiometry. In order to check the stability of this structure with respect to structural transitions, we calculated its phonon frequencies with DFT. We present the resulting phonon dispersion curves in Figure 5(a). There is a single, but two-fold degenerate Γ-point unstable phonon, which is the chain-forming instability that transforms as $\Gamma^-_5$. There is no unstable $\Gamma^-_2$ mode, consistent with our claim in the previous section that this irrep exists in the lower symmetry structure only because of its coupling with the $\Gamma^-_5$ and not because there is a separate driving force for this distortion.

The evolutionary structure prediction performed by USPEX predicts the lowest energy crystal structure to have the $P2_1/m$ symmetry (Fig. 1(b)). The predicted lattice constants are $a = 3.962\,\text{Å}$, $b = 3.896\,\text{Å}$, $c = 10.057\,\text{Å}$, and $\beta = 91.135^\circ$ ($\alpha = \gamma = 90^\circ$). This result is based on the spin non-polarized GGA functional. Spin polarized GGA and GGA + $U$ ($U = 2$, 4 eV) schemes combined with USPEX give the same space group $P2_1/m$ with the similar (but, of course, different) lattice constants. Note that the existence of the zig-zag As chains is also captured by the evolutionary structure search, but the predicted structure is centrosymmetric, both in line with the phonon calculations.

The experimental structural data for the La-doped compound (Ca$_{1-x}$La$_x$FeAs$_2$) measured by X-ray diffraction has the monoclinic $P2_1$ symmetry with the lattice constants are $a = 3.94710\,\text{Å}$, $b = 3.87240\,\text{Å}$, $c = 10.3210\,\text{Å}$, and $\beta = 91.415^\circ$ ($\alpha = \gamma = 90^\circ$) [14]. The other X-ray diffraction experiment on Pr-doped compounds (Ca$_{1-x}$Pr$_x$FeAs$_2$) were reported to have the monoclinic structure with $P2_1/m$ having the inversion symmetry [13]. (The measured lattice constants are $a = 3.9163\,\text{Å}$, $b = 3.8953\,\text{Å}$, $c = 10.311\,\text{Å}$, and $\beta = 90.788^\circ$.)

The strong optical second-harmonic response was recently observed in Ca$_{1-x}$La$_x$FeAs$_2$, clearly implying that the crystal does not have the inversion symmetry [15], but similar data does not exist for Ca$_{1-x}$Pr$_x$FeAs$_2$ to the best of our knowledge. Hence, the first principles predicted structure is quite similar to the experimental structure, except that it does not capture the possible inversion symmetry breaking.

Thermodynamic stability and convex hull construction

Even though the methods we used so far can predict what the crystal structure of CaFeAs$_2$ will be if it forms, they do not address the possibility of the constituent elements phase separating into different compounds. In order to check the phase stability of CaFeAs$_2$, we build the convex-hull for all known binary and ternary compounds formed by these elements. The convex-hull construction evaluates the stability of a given compound against any linear combination of possible compounds effectively [55–59]. Hence, it is possible to determine whether a given compound is stable or prefers to decompose to other compounds (within the accuracy of DFT and the approximations to the exchange correlation functional). We show in Fig. 6(a) the Ca-Fe-As phase diagram constructed from the calculated GGA(PBE) total energy of all relevant phases listed in the materials database [60]. Since the additional symmetry breaking due to magnetic order (for example, antiferromagnetic order) is not considered in

![Diagram](image-url)
the structural prediction performed by USPEX, we take into account stripe magnetic order in the predicted monoclinic $P_{21}/m$ structure of CaFeAs$_2$. Due to the stripe magnetic order, the monoclinic $P_{21}/m$ structure is further relaxed into a triclinic $P\bar{1}$ structure having lower symmetry. The magnetic moment is 1.95 $\mu_B$/Fe and the total energy is lower by 19.50 meV/atom. CaFeAs$_2$ with the stripe magnetic order is computed to be 13 meV/atom above the convex hull, that is, it has a decomposition energy of 13 meV/atom to CaAs and FeAs phases. Reference [61] has systematically studied the error of DFT in similar predictions, and found that the errors can be modeled by a normal distribution with a mean close to zero and a standard deviation of 24 meV/atom. Using this error bar, our calculation indicates that the undoped CaFeAs$_2$ compound is, within the error bar, on the stability boundary. It is possible that doping it with rare earth ions helps its stability.

Apart from a possible energetic gain, an effect of La doping would be an entropic gain if there is no ordering of the La atoms. (A complete ordering of La is unlikely because of its similar ionic radius to Ca.) The corresponding free energy is $-TS = -k_B T \ln N$, where $T$, $S$, $k_B$, and $N$ are temperature, entropy, Boltzmann constant, and configuration number, respectively. For example, in order to simulate the phase stability of 25% La-doped compounds, we used a $2 \times 2 \times 1$ supercell. This super-cell contains 8 Ca atoms, and there are $(2)^8 = 28$ configurations for substituting 2 La atoms for Ca. Hence, we set the configuration number $N$ as 28 and use $T = 1000$ K, which is an estimate of synthesis temperature [18]. The resulting entropy contribution for the La-doped compound is about $-9$ meV/atom. This could render the compound marginally stable according to our calculations, however, a more detailed study is necessary to address the energetics of doping in this compound.

Hypothetical SrFeAs$_2$ and BaFeAs$_2$ compounds

To the best of our knowledge, there exists no report of the synthesis of the Sr or Ba variants of CaFeAs$_2$ even with doping. This is surprising given that Sr usually easily substitutes for Ca, a fact that is supported by data-mining studies of existing crystal structures [62]. In order to see if it is possible, in theory, to synthesize these compounds, and whether they would have a crystal structure that is favorable for pnictide superconductivity, we also investigate the ground state structure for Sr and Ba compounds using the evolutionary structure search method with several schemes for the DFT part, such as spin non-polarized GGA, spin polarized GGA, and GGA + $U$ schemes. For the Sr compound, the lowest energy crystal structure for the spin non-polarized GGA functional is the monoclinic structure with the space group $P_{21}/m$, which is the same as the Ca compound. This monoclinic structure with $P_{21}/m$ is robust with other spin polarized GGA and GGA + $U$ schemes. Considering the stripe magnetic order in the monoclinic $P_{21}/m$ structure of SrFeAs$_2$, the triclinic $P\bar{1}$ structure with the magnetic moment of 2.09 $\mu_B$/Fe is obtained. The total electronic energy is lower by 23.08 meV/atom compared to the monoclinic $P_{21}/m$ structure with no magnetic order (the nonmagnetic state). As shown in Fig. 6(b), SrFeAs$_2$ with the stripe magnetic order is unstable with 24 meV/atom above the convex hull. This energy above the convex hull is higher than that of the CaFeAs$_2$ compound, which might explain why the Sr compound is not synthesized so far.

The situation for the Ba compound is quite different, and different choices for the DFT scheme gives different results for the preferred structure. The evolutionary structure search with the spin non-polarized GGA functional gives the orthorhombic structure with the space group $I4m2$ (# 44) (Fig. 7(a)) as the ground state structure. The obtained lattice constants are $a = 4.059\AA$, $b = 3.984\AA$, $c = 23.161\AA$, and two FeAs layers and As zig-zag chains are contained in the unit cell. This orthorhombic structure does not have inversion symmetry. Evolutionary search using the spin polarized GGA functional gives a structure that is orthorhombic and has the

FIG. 7: (Color Online) Crystal structures of BaFeAs$_2$ having the lowest energy based on (a) spin non-polarized GGA, (b) spin polarized GGA, and (c) GGA + $U$ ($U = 2, 4$ eV) schemes. All of them are orthorhombic and their space groups are $I4m2$ (# 44), $Cmcm$ (# 67), and $Cmcm$ (# 63), respectively.
FIG. 8: (Color Online) (a) $A(k, \omega)$ of $\text{Ca}_{0.8}\text{La}_{0.2}\text{FeAs}_2$ at $T = 116K$ as computed by DFT + eDMFT. The Brillouin zone is shown in inset in the bottom of (c). (b) $A(k, \omega)$ of $\text{Ca}_{0.7}\text{La}_{0.3}\text{FeAs}_2$ at $T = 116K$ as computed by DFT + eDMFT. (The $A(k, \omega)$ of the 30% doped compound is also presented in Ref. [19] with orbital projection onto the in-plane $p$ orbitals of As ions forming the chains.) (c) Optical conductivities within DFT + eDMFT method of $\text{CaFeAs}_2$ (top), $\text{Ca}_{0.8}\text{La}_{0.2}\text{FeAs}_2$ (middle), and $\text{Ca}_{0.7}\text{La}_{0.3}\text{FeAs}_2$ (bottom). Optical conductivities (interband contributions only) within DFT method are also shown for comparison. (Intraband transitions within DFT give a delta function-like contribution at the zero frequency.) The values of in-plane resistivity anisotropy $\rho_a/\rho_b$ correspond to the eDMFT results. The in-plane average conductivity is shown in inset for comparison between eDMFT, DFT, and experiment. We used 0.01 eV for the broadening of intraband contributions in DFT. Experiment data is for $\text{Ca}_{0.7}\text{La}_{0.3}\text{FeAs}_2$ at $T = 125K$, and digitized from Ref. [63]. For $\text{Ca}_{0.7}\text{La}_{0.3}\text{FeAs}_2$, there is a low-energy peak of 0.34 eV in the optical conductivity, which corresponds to the transition marked by the green arrow in (b).

Space group $\text{Cmcm}$ (# 67). Its lattice constants are $a = 6.617\,\text{Å}$, $b = 5.977\,\text{Å}$, $c = 10.316\,\text{Å}$ (Fig. 7(b)). In this orthorhombic structure, Fe atom is surrounded by As anions tetrahedrally, however these tetrahedra form chains instead of a 2D FeAs layer, which is not a structure that would favor high $T_c$ superconductivity even if it actually formed. Finally, the evolutionary structure search with GGA + $U$ ($U = 2, 4$ eV) gives the orthorhombic structure with space group $\text{Cmcm}$ (# 63) (Fig. 7(c)) as the lowest energy structure. The lattice constants of this structure are $a = 6.086\,\text{Å}$, $b = 5.501\,\text{Å}$, $c = 12.621\,\text{Å}$. Each Fe ion is again coordinated by 4 As ions, but the coordination geometry is square rather than a tetrahedron. This is not a commonly observed geometry for Fe, and this structure, even if it is synthesized, would surely not favor high temperature superconductivity.

We consider the various possible magnetic orderings among several predicted structures in $\text{BaFeAs}_2$ and calculate the total electronic energies. Among them, the antiferromagnetic order (with the magnetic moment of $2.17 \, \mu_B/\text{Fe}$) in the orthorhombic $\text{Imm2}$ (#44) has the lowest energy. This antiferromagnetic order has the spins aligned the same way as a single stripe magnetic order within the $ab$ plane, but they are ferromagnetic along the $c$ axis. Therefore, this order is different from the single stripe magnetic order exhibited in most iron pnictide superconductors, where the spins are ordered in an antiferromagnetic fashion along the $c$-axis as well. The inter FeAs layer distance in the orthorhombic $\text{Imm2}$ is 11.55 Å, so that the quite large inter-layer distance might affect the inter-layer magnetic ordering.

We construct the convex hull for $\text{BaFeAs}_2$ compound with the stripe magnetic order. As shown in Fig. 6(c), $\text{BaFeAs}_2$ compound is unstable with 17 meV/atom above the convex hull. The energy above the hull is somewhat higher than that of $\text{CaFeAs}_2$, however, it is quite re-
duced compared to that of SrFeAs$_2$. We predict that the rare-earth doped BaFeAs$_2$ might be possible to synthesize considering that the rare-earth doping makes Fe 112 phase be more stable (for example, the energetic gain from entropy as discussed before). From Fig. 6, we can get some information about mixtures of the essential compounds to grow the target material. Since the reaction CaAs + FeAs → CaFeAs$_2$ has the minimum enthalpy of formation of 13 meV/atom (Fig. 6(a)) and the rare-earth doping makes this enthalpy of formation be lower, heating (supplying energy) a mixture of CaAs, FeAs, and rare-earth compounds is essential to grow the rare-earth doped CaFeAs$_2$ compound [13, 18]. On the other hand, BaFeAs$_2$ compound would decompose into BaAs, FeAs$_2$, and BaFe$_2$As$_2$.

Electronic structure of Ca$_{1-x}$La$_x$FeAs$_2$

Figure 8(a) shows the momentum-resolved electronic spectral function $A(k, \omega)$ of Ca$_{0.8}$La$_{0.2}$FeAs$_2$ at $T = 116$ K as computed by DFT + eDMFT. A fast-dispersing band near the X point has the dominant character of the zig-zag As chain [19, 20], and this band goes down below the Fermi level upon doping as shown in Fig. 8(b), indicating that the spacer of the zig-zag As chain has a crucial role in doping [19]. The electronic anisotropy is evident in the band structures, especially along \( \Gamma - X \) and along \( \Gamma - Y \) directions, as shown in Figs. 8(a) and (b). In order to clarify the origin of the anisotropy, we calculated the orbital polarization between the Fe \( d_{xz} \) and \( d_{yz} \) orbitals, \( \phi = (n_{xz} - n_{yz}) \), where \( n_{xz} \) (\( n_{yz} \)) denotes the occupation of the Fe \( d_{xz} \) (\( d_{yz} \)) orbital. \( \phi \) is zero for all compositions at given temperature $T = 116$ K (Fig. 9). Therefore, the anisotropy does not originate from the electronic nematic phase and it is due to the structural anisotropy arising from the zig-zag As chain.

This anisotropy is also reflected in the optical conductivity. Figure 8(c) illustrates the doping dependence in the optical conductivity. Comparing the optical conductivity of CaFeAs$_2$ to that of Ca$_{0.8}$La$_{0.2}$FeAs$_2$, the \( xx \) and \( zz \) components does not change much upon doping. However, there is a significant change in the \( yy \) component upon doping, and the dc conductivity is enhanced significantly. This results in a large in-plane resistivity anisotropy as much as $\rho_a/\rho_b \approx 4.5$, where $\rho_a$ and $\rho_b$ are dc resistivities along \( a \)-axis (\( x \)-axis) and \( b \)-axis (\( y \)-axis), respectively. The in-plane resistivity anisotropy has only a structural origin, and it is different from a electronic nematicity driven anisotropy which was found in Ba(Fe$_{1-x}$Co$_x$)$_2$As$_2$ [49]. The metallic zig-zag As chain is formed along the \( b \)-axis, so that it gives the higher conductivity along the \( b \)-axis. Note that the average in-plane optical conductivity for Ca$_{0.77}$Nd$_{0.23}$FeAs$_2$ was recently reported experimentally [63], and it is consistent with our calculated optical conductivity for Ca$_{0.8}$La$_{0.2}$FeAs$_2$ as shown in inset of Fig. 8(c).

For further La-doping (Ca$_{0.7}$La$_{0.3}$FeAs$_2$), only the in-plane \( xx \) component of the dc conductivity changes sig-
nificantly. Besides, a low-energy peak of 0.34 eV appears in the $yy$ component of the optical conductivity. The low-energy peak of 0.34 eV corresponds to the coherent interband transition within the zig-zag As chain, which is marked by the green arrow in Fig. 8(b).

Since the parent compound of $\text{Ca}_1-x\text{La}_x\text{FeAs}_2$ is regarded as $\text{Ca}_0.7\text{La}_0.3\text{FeAs}_2$ [19], the superconductivity in $\text{Ca}_0.8\text{La}_0.2\text{FeAs}_2$ arises from the hole doping through Ca substitution on the La sites. The resistivity anisotropy is a nonmonotonic function of doping as shown in Fig. 8(c), and it reaches a maximum near the superconducting dome [49].

The plasma frequencies [64] $\omega_{p,xx}$, $\omega_{p,yy}$, and $\omega_{p,zz}$ in the $x$, $y$, and $z$ directions obtained by DFT calculations are 2.47 (2.56), 3.36 (3.56), and 0.57 (0.52) eV, respectively, for $\text{Ca}_0.8\text{La}_0.2\text{FeAs}_2$ ($\text{Ca}_0.7\text{La}_0.3\text{FeAs}_2$). The estimated in-plane resistivity anisotropy in DFT calculations $\rho_a/\rho_b \simeq \omega_{p,yy}^2/\omega_{p,xx}^2$ is about 1.85 (1.94) for $\text{Ca}_0.8\text{La}_0.2\text{FeAs}_2$ ($\text{Ca}_0.7\text{La}_0.3\text{FeAs}_2$). The in-plane resistivity anisotropy is diminished compared to the eDMFT result giving the quite large in-plane resistivity anisotropy. The anisotropy difference between DFT and eDMFT calculations comes from the correlation effect on iron $3d$ orbitals. The resistivity along $b$-axis ($\rho_b$) is almost dominated by the non-correlated zig-zag As chain. However, the resistivity along $a$-axis ($\rho_a$) is enhanced by the correlated FeAs layer in the systems. Therefore the correlation effect enhances the in-plane resistivity anisotropy $\rho_a/\rho_b$ induced by the structural anisotropy exhibited in the systems. Compared eDMFT with DFT optical conductivities, the eDMFT calculations show higher spectral weights at low energy. It is due to the incoherent spectral weight induced by the local correlation effect, which is well described by eDMFT, however is not present in DFT. The incoherent spectral weight is also clearly shown in inset of Fig. 8(c), where we have compared the eDMFT and DFT calculations with the measured in-plane average conductivity. Based on the high spectral weight at low energy ($\sim$0.4 eV) in experiment, eDMFT calculations give better description rather than DFT. But still, eDMFT and experiment seem to have rather different positions of maxima, at 1.2 eV and at 0.4 eV, respectively. Note that when CaFeAs$_2$ is doped, its crystal structure could change as well. However, we do not take this effect into account, and the change in the anisotropy we observe is purely due to changes in the electronic structure, and not due to changes in the crystal structure.

The mass enhancement $m^*/m_e$ of the iron $3d$ orbitals is reduced upon doping as shown in Fig. 10(a). This means that the band becomes more coherent upon doping and this leads to enhance the Drude peak in the optical conductivity upon doping in Fig. 8(c). Note that the mass enhancement is not equal in all orbitals, and $t_{2g}$ orbitals have larger enhancement than $e_g$ orbitals. Among the $t_{2g}$ orbitals, the $xy$ orbital has the largest mass enhancement. These behaviors are quite consistent with the previous eDMFT calculation in iron-based superconductors [65–69]. We also show the mass enhancement extracted from angle-resolved photoemission spectroscopy (ARPES) experiments [19, 20]. (With help of polarization-dependent ARPES experiments [20], it is possible to extract the band dispersion having iron $d_{xz}$ and $d_{yz}$ orbital characters near the Fermi level and calculate the corresponding mass enhancement for individual orbitals.) A good agreement between the eDMFT and experiment is shown in Fig. 10(a).

Figure 10(b) shows the orbital occupation of iron $3d$ orbitals. Upon doping the $x^2 − y^2$ orbital has almost constant occupation, and $xz/yz$ and $z^2$ orbitals have the similar increment in the orbital occupation. (The increment from $x = 0$ to $x = 0.3$ compound is 0.030 for $z^2$, and 0.035 for $xz/yz$ orbitals.) The largest increment in the orbital occupation of iron $3d$ orbitals is the $xy$ orbital and is 0.075 from $x = 0.0$ to $x = 0.3$ compound. Since this system has the metallic As spacer, there is some additional charge in the spacer As $4p$ orbitals upon doping. The change in the charge on the As $4p$ orbitals between $x = 0$ and $x = 0.3$ is about 0.05. This shows, again, that the metallic As spacer has an important role in the doping process.

Note that two orbitals $z^2$ and $xy$ have very different mass enhancements among other $3d$ orbitals and their enhancements change a lot as a function of doping. The orbital occupation of $z^2$ is the largest among other $3d$ orbitals over the doping ratio up to 30 % and that of $xy$ is significantly increased upon doping. These are effects beyond DFT and are very important for the resistivity anisotropy realized in eDMFT calculations.

**LANDAU FREE ENERGY**

In this section, we outline the basics of a Landau free energy relevant to $\text{Ca}_1-x\text{La}_x\text{FeAs}_2$, as well as other ternary iron pnictide superconductors. The reason for such a study is that the absence of the $C_4$ symmetry in the high-temperature phase of CaFeAs$_2$ might lead a misconception that the nematic transition in this compound has to be significantly different from its counterpart in other tetragonal iron pnictides. Below, we show that this is not the case. Even though there have been various studies which wrote down the Landau theory for these systems, such as references [70–72], to the best of our knowledge there are no studies which emphasize the difference between ferro- and antiferro-orbital orders and their connection with the nematicity.

We start by writing a Landau free energy expansion around a high symmetry tetragonal phase of these compounds. (A tetragonal phase only serves as a reference structure.) The point we would like to emphasize is that the primitive unit cell of this high symmetry phase con-
tains two formula units. As a result, even though many model tight-binding studies usually consider a single Fe atom, we need to consider a two-Fe primitive cell (inset of Fig. 9) when building a Landau theory. We label the two Fe atoms with numbers 1 and 2, and choose cartesian axes such that \( x \) and \( y \) directions point towards nearest neighbors (inset of Fig. 9). We define the orbital polarization for each atom, \( \phi_i \) \((i = 1, 2)\), as the difference between the occupations of \( d_{xz} \) and \( d_{yz} \) orbitals:

\[
\phi_i = n_{i,xz} - n_{i,yz}.
\]

The two cases where the signs of the orbital polarization of the two atoms are the same or opposite correspond to the ferro- and antiferro-orbital orders (Fig. 11). We label the corresponding order parameters by \( \phi^+ \) and \( \phi^- \) as follows:

\[
\phi^+ = \frac{\phi_1 + \phi_2}{2},
\]

\[
\phi^- = \frac{\phi_1 - \phi_2}{2}.
\]

While the onset of either \( \phi^+ \) or \( \phi^- \) would break the symmetry between \( d_{xz} \) and \( d_{yz} \) orbitals on an individual Fe atom, these two order parameters break different space group symmetries of the crystal. The nematic transition commonly observed in iron pnictides involve the ferro orbital order \( \phi^+ \). This order parameter breaks the four-fold rotational symmetry in a specific way - it chooses one of the \( x \) or \( y \) cartesian axes over the other, and hence can couple bilinearly with the shear strain, which we denote as \( \eta \) (Fig. 12(b)), so the free energy has a term \( \sim \eta_\phi \).

At this point, we would like to note that the effect of the shear strain \( \eta \) on the iron sublattice is to convert it from a square to a tetragonal one. As a result, for models considering a single-Fe unit cell, the strain relevant to the nematic transition is a normal strain, but not a shear strain. However, the normal strain in the actual crystallographic unit cell (Fig. 12(c)) is totally different. It is this normal strain \( \eta_n \) that is present at high temperature in \( \text{Ca}_{1-x}\text{La}_x\text{FeAs}_2 \). While \( \eta_n \) also breaks the four-fold rotational symmetry, it does in a different way than \( \eta \) and does not differentiate between \( d_{xz} \) and \( d_{yz} \) orbitals on the Fe atoms. As a result, there is no bilinear coupling \( \sim \eta_\phi \) in the Landau free energy.

Another way to think about the shear and normal strains is to consider the mirror planes present in the reference structure. In the tetragonal structure of iron pnictides, there are 4 mirror planes that are perpendicular to the Fe layers (Fig. 12(d)). The presence of \( \text{m}_{100} \) and \( \text{m}_{010} \) (shown in red in the figure) flips the \( x \) and \( y \) axes on a Fe atom and as a result imposes the condition that \( \phi_i = 0 \). The normal strain breaks the \( \text{m}_{110} \) and \( \text{m}_{110} \) mirror symmetries but preserves \( \text{m}_{100} \) and \( \text{m}_{010} \). So, even though it breaks the \( C_4 \) symmetry, it does not create an orbital order. The shear strain does the opposite, it breaks the \( \text{m}_{100} \) and \( \text{m}_{010} \) mirror symmetries but preserves \( \text{m}_{110} \) and \( \text{m}_{110} \). This gives rise to not only a nonzero \( \phi \) but also a nonzero \( \phi^- \).

The stripe antiferromagnetic (AFM) order observed in iron pnictides has a \( k \)-vector along the \( \{110\} \) family of directions, and involves a doubling of the unit cell. We denote the AFM order parameters with different wave vectors as \( L_x \) and \( L_y \) (Fig. 13) [74]. At the lowest order, there is no \( \sim L_\alpha \) term in the free energy since there is no other parameter that breaks the time reversal symmetry. However, the onset of the AFM order breaks the four-fold rotational symmetry and differentiates between \( x \) and \( y \) axes. It also breaks the \( \text{m}_{100} \) and \( \text{m}_{010} \) mirror symmetries. (Whether the AFM order breaks the other mirror
ensures that $\phi$ is zero. However, there is no symmetry operation that takes into account an additional $\sim$ each other and flips the $x$ pound: It connects the two Fe atoms in the unit cell to a crucial effect for the nematic transition in this compound.

apart from identity and translations is the screw rotation does not include any terms linearly coupled to superconductors, and showed that the free energy expression does not include any terms linearly coupled to $\phi_+$ in the compound under study, $\text{Ca}_{1-x}\text{La}_x\text{FeAs}_2$. Therefore there will be a sharp phase transition between a phase with $\phi_+ = 0$ at high $T$ and a low $T$ phase with $\phi_+ \neq 0$ like in all other iron pnictide materials.

**SUMMARY AND CONCLUSIONS**

We have checked the phase stability of $\text{CaFeAs}_2$ and $\text{Ca}_{1-x}\text{La}_x\text{FeAs}_2$ compounds. The spacer, zig-zag As chain, nearly has the $1\overline{1}0$ valence state, so that it does not form the As square-net, which is reminiscent of the Peierls type instability. According to the convex-hull construction, $\text{CaFeAs}_2$ is above the hull with 13 meV/atom. Further stabilization is possible with rare earth doping in $\text{CaFeAs}_2$ material. We have also calculated the optical conductivity of $\text{Ca}_{1-x}\text{La}_x\text{FeAs}_2$ based on the DFT + eDMFT method, and found a large in-plane resistivity anisotropy. This large anisotropy does not originate from electronic nematicity, but from the structural anisotropy arising from the zig-zag As chain. The electronic correlations do not induce but nevertheless enhance this anisotropy, as seen from the difference of DFT and eDMFT results. For $\text{Ca}_{0.7}\text{La}_{0.3}\text{FeAs}_2$ compound, we found a low-frequency peak of 0.34 eV in the in-plane $yy$ component of the optical conductivity. This peak corresponds to the coherent interband transition within the
FIG. 14: (Color Online) $T_c$ vs. Anion height in iron pnictide and chalcogenide superconductors. (Reproduced with permission from Ref. [73].) The three data points we added (black) for the 112 superconductors display a similar trend as the other iron based high $T_c$ superconductors. This suggests that the metallic spacer layer in the 112 family does not influence superconductivity.

zig-zag As chain.

In Fig. 14 we reproduce the plot of the distance between the anion and Fe layers and $T_c$ in iron pnictide superconductors. The three data points we add from published data falls perfectly on the lines that were drawn to underline the striking correlation with this single structural parameter and the superconducting critical temperature. This suggests that despite being noncentrosymmetric and having a metallic spacer layer, the superconductivity in the rare-earth doped CaFeAs$_2$ is no different that in other iron pnictide superconductors. This supports the spin-fluctuation based theories as opposed to theories that rely on charge fluctuations.

ACKNOWLEDGMENTS

C.-J. Kang and G. Kotliar were supported by the DOE EFRC CES program. T. Birol acknowledges the support of the Center for Materials Theory at Rutgers University.

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