Correlation effects in the tetragonal and collapsed tetragonal phase of CaFe$_2$As$_2$

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We investigate the role of correlations in the tetragonal and collapsed tetragonal phases of CaFe$_2$As$_2$ by performing charge self-consistent DFT+DMFT (density functional theory combined with dynamical mean-field theory) calculations. While the topology of the Fermi surface is basically unaffected by the inclusion of correlation effects, we find important orbital-dependent mass renormalizations which show good agreement with recent angle-resolved photoemission (ARPES) experiments. Moreover, we observe a markedly different behavior of these quantities between the low-pressure tetragonal and the high-pressure collapsed tetragonal phase. We attribute these effects to the increased hybridization between the iron- and arsenic orbitals as one enters the collapsed tetragonal phase.

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I. INTRODUCTION
The iron-pnictide superconductor CaFe$_2$As$_2$ belongs to the so-called 122 family, AFe$_2$As$_2$ (e.g. A = Ba, Sr, Ca) which crystallizes in the ThCr$_2$Si$_2$ structure with the I 4/mmm space group. CaFe$_2$As$_2$ is tetragonal (TET) at room temperature and ambient pressure and undergoes a structural phase transition to an orthorhombic (ORT) phase upon cooling below 170 K. Whereas the tetragonal phase is non-magnetic, the orthorhombic phase shows a stripe-like magnetic order. Upon application of pressure, the appearance of a collapsed tetragonal (CT) phase characterized by a collapse of the c lattice parameter and a volume shrinkage of about 5% with respect to the tetragonal phase was observed. First principles studies have shown that for increasing pressures at low temperature the system goes from the orthorhombic phase directly into the collapsed tetragonal phase at 0.36 GPa, whereas for higher temperatures at high pressure the tetragonal phase is energetically more favorable than the collapsed tetragonal phase. Moreover, the ORT→CT structural transition coincides with the disappearance of the magnetic moment. Also in BaFe$_2$As$_2$ such a collapsed tetragonal phase has been theoretically predicted and experimentally observed, though at much higher pressures of 27 GPa under hydrostatic pressure conditions.

The appearance of a superconducting phase under pressure was reported in CaFe$_2$As$_2$ with a critical temperature of 10 K at 0.69 GPa. However, it was recently established that the superconducting region is disjoint from the non-magnetic collapsed tetragonal phase and it is still not entirely clear if superconductivity appears in the orthorhombic phase or in a low temperature tetragonal phase that is stabilized by special non-hydrostatic pressure conditions. In order to understand this behaviour, a lot of effort has been devoted in the last years to investigate the electronic properties of the collapsed tetragonal phase and its main differences compared to the orthorhombic and tetragonal phases. Angle-resolved photoemission (ARPES) measurements for the orthorhombic and tetragonal phases in CaFe$_2$As$_2$ at ambient pressure were performed by Liu et al., where a two- to three-dimensional transition in the Fermi surface was observed, corresponding to the transition from the tetragonal to the orthorhombic phase at low temperatures. Measurements have also been performed for isostructural materials which are in the collapsed tetragonal phase at ambient pressure: CaFe$_2$P$_2$ and Ca(Fe$_{1-x}$Rh$_x$)$_2$As$_2$. In both cases hole pockets around the zone center Γ present in the tetragonal phase disappear in the collapsed tetragonal phase.

Only very recently CaFe$_2$As$_2$ samples could be grown in the collapsed tetragonal phase at ambient pressure by introducing internal strain. In the same work, the authors performed detailed ARPES measurements and found that collapsed tetragonal CaFe$_2$As$_2$ shows a similar behavior to CaFe$_2$P$_2$ and Ca(Fe$_{1-x}$Rh$_x$)$_2$As$_2$, namely the disappearance of the hole pockets at the Γ point. While density functional theory (DFT) calculations correctly predict this feature, ARPES measurements show a strong band renormalization compared to the DFT calculations.

In order to investigate the origin of this discrepancy, we present in this work an analysis of the electronic structure of tetragonal and collapsed tetragonal phases of CaFe$_2$As$_2$ by combining DFT in the GGA approximation with dynamical mean-field theory (GGA+DMFT). This method has been proven to provide a good description of correlation effects in a few families of Fe-based superconductors. While the 122 family has been argued to be less correlated than the so-called 111 or 11 families, we will show that also in CaFe$_2$As$_2$ correlations are necessary to understand the renormalization of
TABLE I: Lattice parameters for the experimentally measured tetragonal and collapsed tetragonal structure from Ref. 5

|          | TET       | CT      |
|----------|-----------|---------|
| $T$ (K)  | 250       | 50      |
| $p$ (GPa)| 0.0       | 0.35    |
| $a$ ($\text{Å}$) | 3.8915    | 3.9792  |
| $c$ ($\text{Å}$)  | 11.690    | 10.6073 |
| $z_{\text{As}}$ | 0.372     | 0.3663  |
| $V$ ($\text{Å}^3$) | 177.03    | 167.96  |

The bands, where we find a distinct change of orbital-dependent mass enhancements in the transition from the tetragonal to the collapsed tetragonal phase.

II. METHODS

For our fully charge self-consistent GGA+DMFT calculations we consider the tetragonal and collapsed tetragonal structures obtained by neutron diffraction experiment.\cite{5} Lattice parameters and As $z$ position are shown in Table I. The DFT calculations were performed with the WIEN2k implementation of the full-potential linear augmented plane wave (FLAPW) method.\cite{29} As exchange-correlation functional we considered the generalized gradient approximation\cite{29} (GGA). The self-consistency cycle employed 726 $k$-points in the irreducible Brillouin zone, resulting in a 21 $\times$ 21 $k$ mesh in the conventional Brillouin zone, and a $R_{\text{int}}k_{\text{max}} = 7.0$. For the projection of the Bloch wave functions to the localized Fe 3$d$ orbitals we used our own implementation of the method described in Ref.\cite{29}.\cite{29} The energy window for the projection was chosen to be in the range from $-5.9$ to $16.0$ eV ($-6.3$ to $16.0$ eV) for the tetragonal and collapsed tetragonal structures. We were able to set the lower energy boundary in a gap in the density of states (DOS). The impurity problem was solved with a continuous-time quantum Monte Carlo method in the hybridization expansion\cite{32} as implemented in the ALPS\cite{32} project. Calculations were done at $\beta = 40$ eV$^{-1}$ with $2 \times 10^6$ Monte Carlo sweeps. For the double counting correction the fully localized limit\cite{33} (FLL) scheme was used, although the around mean field (AMF) scheme led to comparable results with only slightly less renormalized masses. The interaction parameters are used in the definition of the Slater integrals\cite{34} $F^\alpha$ with $U = F^\alpha$ and $J = (F^2 + F^\alpha)/14$. For the onsite correlation we consider a value of $U = 4$ eV and for Hund’s rule coupling $J = 0.8$ eV and we analyze the dependency of our results on variations of these parameters. For the analytic continuation of the Monte Carlo data on the imaginary time axis we used a combination of Padé-approximation and a fourth order polynomial fit to the first eight Matsubara frequencies to obtain real frequency data.

In the projection of the Fe 3$d$ orbital character, we use a coordinate system which is rotated by 45° around the $z$-axis with respect to the conventional $I4/mmm$ unit cell so that $x$- and $y$-axis point towards neighboring Fe atoms as shown in Fig. (a). In the band structure and Fermi surface plots we choose the usual high symmetry points $X$, $M$ and $Z$ of the $I4/mmm$ space group to facilitate comparison with the other families of iron pnictides.

III. RESULTS

A. Band structure and spectral function

In Fig. 2 we show a comparison of the DFT (GGA) band structure calculations and the spectral function obtained with GGA+DMFT. We find that correlations mostly renormalize bands in both structures without introducing significant band shifts or altering the topology of the Fermi surface. In the tetragonal phase we observe in both DFT (GGA) as well as GGA+DMFT calculations the presence of three hole bands crossing the Fermi level at the zone center $\Gamma$, two electron pockets at
X and three well-defined hole pockets at the zone corner M formed by strongly dispersive hole bands with a large outer pocket and two smaller inner pockets almost identical in size. In the collapsed tetragonal phase the bands at Γ are pushed below the Fermi level in agreement with experiments\textsuperscript{10,18–20} the inner electron pocket at X is pushed up to positive energies leaving only the slightly enlarged outer electron pocket present. At M the bands forming the inner two hole pockets are pushed onto the Fermi level, leaving two extremely shallow bands of which only one just barely crosses $E_F$. GGA+DMFT introduces a significant separation between the two bands not observed in the DFT(GGA) calculations. This is a result of the orbital dependent correlations introduced by DMFT.

In Fig. 3 we show the GGA+DMFT results for the same energy range and along the same path in the Brillouin zone as in Ref.\textsuperscript{20} in order to allow a better comparison to the ARPES measurements. We find a good agreement between ARPES and our GGA+DMFT calculation in both the tetragonal and the collapsed tetragonal phases albeit GGA+DMFT finds a smaller band renormalization than the value extracted from ARPES. Our band renormalizations are about a factor of $1.7$ compared to GGA masses while the ARPES measurements report a factor of $5$. This suggests that other possible contributions not considered in DMFT may also be important for the description of the electronic behavior of CaFe$_2$As$_2$ like non-local correlations and electron-phonon interactions.

The GGA+DMFT Fermi surface for CaFe$_2$As$_2$ shows only slight changes compared to DFT(GGA) (see Fig. 4) and agrees reasonably well with ARPES measurements\textsuperscript{20}. The main features of the collapsed tetragonal phase are the disappearance of the hole pockets at Γ as well as a change from a more two-dimensional shape in the tetragonal phase to a three-dimensional shape in the collapsed tetragonal phase (compare the cuts along a plane parallel to the z direction in Figs. 3(a) and (b)) due to increasing Fe 3$d$-As 4$p$ hybridizations.

### B. Mass enhancements and sensitivity to interaction parameters

We calculate the effective masses directly from the impurity self-energy via

$$m^* = 1 - \frac{\partial \text{Im} \Sigma(i\omega)}{\partial \omega} \bigg|_{\omega \rightarrow 0}. \quad (1)$$

For the interaction parameters set to $U = 4$ eV and $J = 0.8$ eV we obtain mass renormalizations between 1.2 and 1.7 as shown in Tab.\textsuperscript{11} for the different orbital characters. Mass renormalizations are strongest for the $t_{2g}$ orbitals Fe 3$d_{xy}$ and 3$d_{xz/yz}$ in the tetragonal phase while the $e_g$ orbitals 3$d_{z^2}$ and 3$d_{x^2-y^2}$ are less renormalized both in the tetragonal and collapsed tetragonal phases.
As shown in Table II and Fig. 5 we observe a change in the strengths of the mass renormalizations. Interestingly, the iron Fe 3d orbital undergoes a change from being the most strongly renormalized orbital in the tetragonal phase to the least renormalized orbital in the collapsed tetragonal phase. This can be understood in terms of increased hybridization in the collapsed tetragonal phase. The structural collapse in this phase is assisted by a formation of As 4p–As 4p bonds between the Fe-As layers as shown in Fig. (b) and (c) with a strong bonding-antibonding splitting of the As 4p bands. The Fe 3d orbitals, which are pointing in the direction of the As atoms, become less localized in the collapsed tetragonal phase due to increased Fe 3d–Fe 3d as well as Fe 3d–As 4p x and 4p y hybridizations. This higher degree of delocalization leads to less mass renormalization upon inclusion of correlations.

By varying the interaction parameters $U$ and $J$ we have investigated their influence on the effective masses. The effective masses show stronger dependencies on the Hund’s rule coupling $J$ than the Hubbard $U$ as shown in Fig. and as already reported for other members of the iron pnictides. Our results are stable for all values of the chosen interaction parameters and, except for the stronger band renormalization, we observe only very small qualitative changes in the Fermi surface.

In Fig. 6 we show a comparison of the occupation numbers between the GGA and the GGA+DMFT results for the tetragonal and the collapsed tetragonal phases.

![Diagram](image)

**FIG. 4:** (Color online) Comparison of the Fermi surface from GGA (left) and GGA+DMFT (right) along a plane at $k_z = 0$ and a vertical cut through the $\Gamma$ and $X$ point.

**FIG. 5:** (Color online) Sensitivity of effective masses $m^*/m_{GGA}$ with respect to changes in the interaction parameters. (a), (c) show variations in $U$ in the tetragonal and the collapsed tetragonal phases respectively, and (b), (d) show variations in $J$.

**TABLE II:** Mass renormalizations calculated with GGA+DMFT for the Fe 3d orbitals.

|          | $d_{x^2}$ | $d_{x^2-y^2}$ | $d_{xy}$ | $d_{3z^2-r^2}$ |
|----------|-----------|---------------|----------|----------------|
| Tetragonal| 1.45      | 1.44          | 1.72     | 1.62           |
| Collapsed tetragonal | 1.39      | 1.42          | 1.36     | 1.57           |
The $3d_{xy}$ and $3d_{x^2−y^2}$ show the largest occupation with respect to $3d_{z^2}$ and $3d_{x^2−y^2}$ reflecting the crystal field splitting in $t_{2g}$ and $e_g$ orbitals. At the GGA level the transition from tetragonal to collapsed tetragonal phase implies a pronounced increase of charge occupation of the $3d_{xy}$ orbital and to a lesser extent of the $3d_{x^2−y^2}$, while the occupation for the $e_g$ states decreases. This can also be understood in terms of the change in hybridizations as explained above, where due to the enhanced delocalization of the $3d_{xy}$ electrons in the collapsed tetragonal phase the $3d_{xy}$ orbital becomes less correlated. Regarding the GGA versus GGA+DMFT occupations we observe only little changes and a general trend of electronic charge being shifted from the most correlated orbitals to the less correlated orbitals, as expected, with the total charge on the Fe $3d$ orbitals staying basically identical to the DFT calculation.

Recently, we became aware of the ARPES investigations by Golfry et al.\textsuperscript{119} who reported a distinct increase of the effective masses of the bands around the $\Gamma$-point when entering the collapsed tetragonal phase. In order to understand this, we calculated the effective masses $m^*_{\text{GGA}}/m_e$ of the three hole bands around the $\Gamma$-point according to the method we described in a previous article.\textsuperscript{11} In the tetragonal phase we obtained 1.11, 1.62 and 1.71$m_e$, while in the CT phase we obtained 1.53, 2.00, 2.89$m_e$, with the bands ordered from higher to lower binding energies. Thus, already at the GGA level the trend of increasing renormalization of the bands around $\Gamma$ in the CT phase is correctly described, albeit the absolute values are lower compared to what was reported from experiment.\textsuperscript{29} Therefore, we conclude that the observed increase in band renormalizations from the tetragonal to the CT phase around $\Gamma$ is mostly due to stronger hybridizations in the collapsed tetragonal phase, as discussed in this section, leading to a shift of the hole bands below the Fermi level. Electronic correlations contribute further only to a minor degree to the effective electronic mass of the bands around $\Gamma$, which we attribute to the fact that CaFe$_2$As$_2$ is a weakly to moderately correlated metal.

**IV. CONCLUSIONS**

We have performed charge self-consistent GGA+DMFT calculations for CaFe$_2$As$_2$ in the tetragonal and collapsed tetragonal phases. We observe that while the topology of the Fermi surface in both phases remains nearly unaffected, the orbital-selective mass renormalizations of a factor 1.3 to 1.7 introduced by GGA+DMFT improve the agreement of the calculations with ARPES experiments. The analysis of the influence of the tetragonal to collapsed tetragonal transition on the orbital-dependent effective masses shows that Fe $3d_{xy}$ changes from being the most strongly correlated orbital in the tetragonal phase to being the least correlated one in the collapsed tetragonal phase. We attribute this to the change in hybridization of the Fe 3d orbitals in the collapsed tetragonal phase, where due to the decreased distance of the Fe-As layers the hybridization for the Fe $3d_{xy}$-Fe $3d_{xy}$ as well as Fe $3d_{xy}$-As 4$p_x$ and 4$p_y$ orbitals increases, rendering the Fe $3d_{xy}$ less localized and thus less correlated. The orbital occupations confirm this trend and show a higher occupation for the Fe $3d_{xy}$ orbital in the collapsed tetragonal phase.

With these observations we conclude that correlation effects beyond DFT(GGA) as introduced by GGA+DMFT are needed even for weakly correlated pnictides like CaFe$_2$As$_2$ in order to understand the orbital-selective mass renormalizations observed in ARPES. However, we also observe that such a description is, nevertheless, still insufficient for explaining the large mass renormalizations observed experimentally. We attribute this discrepancy to possible non-local correlations as well as phononic effects and this will be a subject of future investigations.

During finalization of this manuscript we became aware of another preprint of a DFT+DMFT study of CaFe$_2$As$_2$\textsuperscript{242} where the authors also find the same trend of reduced renormalization in the CT phase and their results agree, except for minor quantitative differences, with our findings.

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