Site-selective electronic structure of pure and doped Ca$_2$O$_3$Fe$_3$S$_2$

L. Craco,$^{1,2}$ B. Freelon,$^3$ and S. Leoni$^4$

$^1$Instituto de Física, Universidade Federal de Mato Grosso, 78060-900, Cuiabá, MT, Brazil  
$^2$IFW Dresden, Institute for Solid State Research, P.O. Box 270116, D-01171 Dresden, Germany  
$^3$Department of Physics, Massachusetts Institute of Technology, Cambridge, Massachusetts, 02139, USA  
$^4$School of Chemistry, Cardiff University, Cardiff, CF10 3AT, UK

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Using density functional dynamical mean-field theory we investigate the site-selective electronic structure of Ca$_2$O$_3$Fe$_3$S$_2$. We confirm that the parent compound with two distinct iron sites is a multi-orbital Mott insulator similar to La$_2$O$_3$Fe$_3$S$_2$. Electron/hole doping, carrier localization is found to persist on the two active iron channels because the chemical potential lies in a gap structure with anisotropic and almost vanishing states near the Fermi energy. This emergent behavior stems from large electronic reconstruction caused by dynamical spectral weight transfer involving states with distinct d-shell occupancies and orbital character at low energies. We detail the implications of our microscopic analysis and discuss the underlying physics which will emerge in future experiments on Ca$_2$O$_3$Fe$_3$S$_2$.

I. INTRODUCTION

In recent years the large family of Fe-based compounds$^{1,2}$ has been the subject of intensive research after the discovery of high temperature superconductivity (high-TC) in F-doped LaOFeAs.$^3$ It has emerged that these unconventional superconductors (both pnictides and chalcogenides) increasingly fall into the bad-metal category. In SmFeAsO$_{1-x}$F$_x$,$^4$ for example, suppression of superconductivity by high magnetic fields reveals a low temperature insulating behavior, similar to underdoped high-TC cuprates.$^5$ Interestingly, in some cases superconductivity arises directly from a normal state with insulating-like$^6$ resistivity above $T_c$. Given telling similarities, along with notable differences, between the iron- and cuprate-based superconductors,$^8$ an especially important issue has been to confirm whether the parent compounds are itinerant metals or if they lie in close proximity to a Mott metal-to-insulator transition.$^7,9$ particularly interesting for the latter view are Fe-oxycgalogenides, which have been synthesized with various compositions,$^{2,10-14}$ showing different degrees of electron localization.

A Mott insulator undergoes an electronic phase transition to a correlated metal at low temperatures.$^{15}$ The metallic behavior can vary from canonical Fermi to non-Fermi liquid depending on the degree of electronic correlation, disorder and structural-induced changes in orbital polarization patterns. In all cases, the insulator-to-metal phenomenon is due to the development of a severely renormalized lattice coherence scale, driven by increasing relevance of electronic delocalization upon external perturbations like pressure and chemical substitution.$^{15}$ Optical and spectroscopic studies in 122-iron selenide superconductors,$^{16}$ for example, show large-scale spectral weight transfer as function of temperature across the magnetic and superconducting phase instabilities, which is a fingerprint of Mottness.$^{17}$

In this context, finding insulating and antiferromagnetically ordered ground state in layered Fe-oxycgalogenides$^{11,12,14,18,19}$ is relevant, since it confirms theoretical suggestions that Mott-insulating parent compounds in the Fe-superconducting systems could be found by increasing the ratio of the interaction-to-hopping beyond a critical value for a Mott transition.$^{20}$ The Fe-oxycgalogenides$^2$ thus help to sharpen the fundamental debate$^{21}$ on the degree of electronic correlations in Fe-based superconductors in general. However, finding Mott electrons with different degrees of activation energies$^{11-14,18}$ suggests strong multi-orbital (MO) electronic localization in this material class. In contrast to Fe-based superconductors, the consequences of Mottness (or the proximity to a insulator-metal transition) upon external perturbations in the Fe-oxycgalogenides has not received the theoretical attention it deserves. This is even more important, when the Mott transition is unconventional in nature, as shown here in Ca$_2$O$_3$Fe$_3$S$_2$ where carrier localization persists on different Fe-channels upon doping. Thus, following earlier studies on Fe-oxycgalogenides$^{22,23}$ we use the local density approximation plus dynamical mean-field theory (LDA+DMFT)$^{24}$ to study the role played by sizable MO electron-electron interactions in the Fe 3d shells of Ca$_2$O$_3$Fe$_3$S$_2$. As shown below, all five 3d-bands of two distinct Fe-sites (Site 1, Site 2) must be kept in order to satisfactorily resolve the site-selective Mott localization and bad metallic regimes obtained, respectively, for pure and doped Ca$_2$O$_3$Fe$_3$S$_2$. In MO Mott insulators, sizable electronic correlations drives new physical effects from (electron, hole) doping. They can induce a pseudogap regime, where the chemical potential lies in an energy region of vanishing density-of-states (DOS) near the Fermi energy ($E_F$),$^{25}$ or orbital-selective incoherent states, naturally yielding co-existent insulating and bad-metallic states as in cuprates$^{26}$ and Fe-pnictide superconductors.$^{27}$ In this work, we also use our results to discuss the influence of Mottness in lightly doped Ca$_2$O$_3$Fe$_3$S$_2$ providing specific predictions which can be tested in future experiments.
The crystal structure of Ca$_2$O$_3$Fe$_3$S$_2$ (space group $P4/mmm$) is built from alternate stacking of antiperovskite-like [Ca$_2$FeO$_2$]$^{2+}$ layers and [Fe$_2$OS$_2$]$^{2-}$ building blocks along the $c$-axis, as shown in Fig. 1. Similar to Na$_2$Fe$_2$OS$_2$ oxychalcogenide Mott insulator, in the Fe$_2$OS$_2$ unit, the Fe$^{2+}$ ($d^6$ electronic configuration) ions are located between oxygen atoms, forming a square-planar layer, which is an anti-configuration with respect to the CaO$_2$ layer of high-$T_c$ cuprates. As seen in Fig. 1, the two-dimensional [CaFeO$_2$]$^{2+}$ layers contain FeO$_2$ planar sheets separated from the [Fe$_2$OS$_2$]$^{2-}$ layers by Ca ions. The Fe ions (Site 2) in the [Fe$_2$OS$_2$]$^{2-}$ blocks are sixfold-coordinated by two O ions and four S ions, forming a FeO$_2$S$_2$ octahedron. Extant refinement of structural data for the individual sites of Ca$_2$O$_3$Fe$_2$O$_4$S$_2$ revealed that the Fe sites at the planar layer (Site 1) are fully occupied, while at Site 2 the Fe ions are about 20% deficient in this vacancy ordered compound.

To explore the site-selective electronic structure of Ca$_2$O$_3$Fe$_3$S$_2$ parent compound, self-consistent band structure calculations were performed using the linear muffin-tin orbital (LMTO) method$^{29}$ in the atomic sphere approximation as implemented in PY LMTO computer code.$^{30}$ The Perdew-Wang parameterization$^{31}$ was used to construct the exchange correlation potential within LDA. The total density was converged on a grid of 819 irreducible $k$ points, and the radii of the atomic spheres were chosen as $r_{Ca}=$3.57 a.u., $r_O=$1.97 a.u., $r_{Fe}=$2.45 a.u., $r_S=$3.12 a.u. in order to minimize their overlap. The LDA calculation we performed using the cell parameter values reported elsewhere for Ca$_2$O$_3$Fe$_3$S$_2$ parent compound.$^{32}$

Our results in Figs. 2 and 3 confirm that similar to Fe-based superconductors the active electronic states involve Fe 3$d$ carriers. A sizable enhancement of the average LDA bandwidth ($W$) relative to that of La$_2$O$_3$Fe$_2$S$_2$ due to short Fe-Fe bonds is obtained. As shown previously, La$_2$O$_3$Fe$_2$S$_2$ is a narrow band-gap Mott insulator and the significantly larger on-particle bandwidth for Ca$_2$O$_3$Fe$_3$S$_2$ naturally implies proximity to metallization. Site- and orbital-driven anisotropies in the LDA band structure are also clearly manifested in Fig. 3. At Site 2 the orbital-resolved DOS show similar features as found for La$_2$O$_3$Fe$_2$S$_2$, a result consistent with the Fe$^{2+}$ oxidation state with six electrons in the Fe 3$d$ shell. On the other hand, interesting deviations are seen for the $xz, yz, xy$ and the $x^2 − y^2$ orbitals of Site 1, albeit less pronounced in the latter case. Due to changes in the crystal-field splittings the electronic states at Site 1 are transferred to energies about $E_F$ severely renormalizing the LDA orbital occupancies compared to Site 2, promoting a Fe$^{3+}$ valence state ($d^5$ electronic configuration) with enhanced orbital polarization. Thus, we attempt to address the following question: How do electroncorrelations change from $d^6$ to $d^5$ electronic configuration? Typically going from partially to half-filled 3$d$ electron shell electron correlation effects are expected to increase. Here it is shown that this scenario holds true for Ca$_2$O$_3$Fe$_2$S$_2$. Additionally, as common to Fe-oxychalcogenides$^{22,23}$ and peculiar to the Fe-based superconductors, strong crystal-field splitting effects lift the $xz, yz$ orbitals degeneracy, leaving an antiferromagnetically ordered state at low temperatures without tetragonal-to-orthorhombic structural phase transition or orbital nematic instabilities. It thus follows that the novelties found in Fe-based superconductors, relating to electronic nematic instabilities in...
the tetragonal phase near the borderline of structural and magnetic transitions, will not play an active role in Ca$_2$O$_2$Fe$_3$S$_2$: Only spin nematic phase fluctuations are expected to be seen across the magnetic phase transition of Ca$_2$O$_2$Fe$_3$S$_2$ as compared to La$_2$O$_2$Fe$_3$S$_2$.

Currently, the theoretical microscopic understanding of Ca$_2$O$_2$Fe$_3$S$_2$ is restricted to LDA and LDA+U calculations. Specifically, these ab initio density functional calculations demonstrate that the chalcogen and oxygen $p$ states lie well below $E_F$ and are weakly hybridized with Fe 3$d$ states. Hence, the most relevant electronic states near the Fermi energy for Ca$_2$O$_2$Fe$_3$S$_2$ derive from the Fe layers with almost direct Fe-Fe hopping. In previous works we undertook systematic LDA+DMFT studies of Fe-oxyselenides, showing that narrow-band Kondo-Mott localization and orbital-selective localization-delocalization transition can be understood within a single theoretical picture. Good semiquantitative agreement with experimental data of La$_2$O$_2$Fe$_2$M$_2$ ($M=$Se,S) systems serves as support to explore intrinsic, dynamical correlation effects in Ca$_2$O$_2$Fe$_3$S$_2$.

Although LDA+U provides reliable structural and magnetic ground state information of strongly correlated electron systems, it generically fails to capture the ubiquitous dynamical correlation effects in $d$-band compounds, and so cannot access normal state incoherence and the emergence of Hubbard satellites (local moments) at low and high energies. Combining LDA with DMFT is the state-of-the-art prescription for overcoming this problem. Thus, as common to Fe-based superconducting materials within LDA the one-electron part of the Hamiltonian for Ca$_2$O$_2$Fe$_3$S$_2$ reads $H_0 = \sum_{k,a,\sigma} c_{\sigma}(k)c_{\sigma}^\dagger(k)$, where $a = 3z^2-r^2, xx, yz, x^2-y^2, yz$ label the (diagonalized in orbital basis) five Fe 3$d$ bands, which are the only ones we retain, since the non-3$d$-orbital DOS at Sites 1 and 2 have negligible or no weight at $E_F$. Here, the relevant inputs to our LDA+DMFT treatment for Ca$_2$O$_2$Fe$_3$S$_2$ are the LDA DOS for the five 3$d$ orbitals at Sites 1 and 2 shown in Fig. 3, the on-site Coulomb interaction $U$, the inter-orbital term $U' = U - 2J_H$, and the Hund’s coupling $J_H$. Based on this, at each Fe-channel the correlated part of the many-body Hamiltonian for Ca$_2$O$_2$Fe$_3$S$_2$ is $H_{\text{int}} = U\sum_{i,a} n_{i,a}n_{i,a} + \sum_{i,a\neq b}[U' n_{i,a}n_{i,b} - J_H S_{i,a} \cdot S_{i,b}]$. We choose values of $U = 5.0$ eV, $J_H = 0.7$ eV as employed in our earlier works for the Fe-oxyselenides. Our pa-
Carefully examining the orbital-resolved spectral function \( s \) is orbital-dependent, i.e., intrinsically anisotropic. (i) Strong dynamical MO correlations originating from \( U, U' \) lead to sizable spectral weight redistribution over large energy scales and the formation of a severely reconstructed (compared to LDA) correlated electronic structure, with concomitant reduction of the orbital polarization patterns found in LDA. This behavior is characteristic of multiband Mott systems, where emergent upper and lower Hubbard bands are expected at high-energies. These latter features are known to be related to coupled spin-orbital local moments defining a Mott insulator without long-range charge, orbital or magnetic order.

Since there is no particle-hole symmetry in the bare electronic states of \( \text{Ca}_2\text{O}_2\text{Fe}_3\text{S}_2 \) (see Fig. 3), it is interesting to inquire what happens to the parent compound upon carrier (electron/hole) doping. In particular, does selective localization characteristic of MO systems in the verge of Mott transition could be found in future studies on doped \( \text{Ca}_2\text{O}_2\text{Fe}_3\text{S}_2 \)? Indeed, the generic appearance of incoherent metallic states with low energy pseudogaps, and the instabilities of such states to unconventional orders in a wide variety of correlated electron systems makes this an important question to inquire about. Our aim here is to build upon the strengths of MO correlated electronic structure calculations to predict the effect of carrier doping on the two Fe-sites of \( \text{Ca}_2\text{O}_2\text{Fe}_3\text{S}_2 \). In particular, we will discuss a set of predictions which could be tested in future experimental work.

Fig. 5 shows the changes in the correlated electronic structure upon electron and hole doping \(( n \equiv 6 \pm \delta \) at Site 2). Similar to \( \text{Na}_2\text{Fe}_2\text{OSe}_2 \), when \( \delta \) increases to small negative values a selective-Mott state develops, in which the \( 3z^2 - r^2 \), \( xz \), \( yz \) and \( x^2 - y^2 \) spectral functions show behavior of a Mott insulator with vanishing DOS at \( E_F \). While the \( xy \) orbital shows metallic behavior, characterized by the presence of incoherent in-gap states at \( E_F \). On the other hand, enhanced metallicity is predicted for electron doped \(( n = 6.2 \) \( \text{Ca}_2\text{O}_2\text{Fe}_3\text{S}_2 \) where, in addition to the \( xy \) orbital, the \( xz \), \( yz \) bands will also contribute to orbital-selective electronic transport and Fermi surface reconstruction within LDA+DMFT. Although the Mott gap in the \( 3z^2 - r^2 \) and \( x^2 - y^2 \) band is considerably reduced upon electron doping, strong localization is clearly visible in this orbital sector, as seen in Fig. 5. From the above observation, it can be suggested that strong Mottness and orbital-selective Mott physics coexist in the \( \text{Fe}^{2+} \) electronic channel. Since there is no particle-hole symmetry in \( \text{Fe}^{2+} \) channel, it is interesting to inquire as to the effects of electron doping \(( n \equiv 5 \pm \delta \) in this half-filled electronic reservoir. It is worth noticing here that electron-electron interactions are expected to reach their maximum at half-filled

![FIG. 5: (Color online) Effect of electron/hole doping on the LDA+DMFT \(( U = 5.0 \text{ eV}, J_H = 0.7 \text{ eV}) \) spectral functions for the Fe 3d orbitals of \( \text{Ca}_2\text{O}_2\text{Fe}_3\text{S}_2 \) at Site 2. Notice the electron delocalization and the appearance of incoherent, pseudogapped shoulders quasiparticle resonances on different orbitals near the Fermi energy.]
In a MO system like the Fe-oxychalcogenides, strong (indoped counterpart. 

To orbital-selective metallicity compared to its electron hole doping spectral functions now show pseudogap behavior. Hence, emergent spectral weight at $E_h$ hole doping, and the other hand, metallicity is found to be favored upon orbitals at $E_f$.

Fig. 6 exhibits the answer to this fundamental questions. As seen, Mott localization persists in all orbitals at Site 1 for electron doped Ca$_2$O$_3$Fe$_3$S$_2$. On the other hand, metallicity is found to be favored upon hole doping, and the $x^2-y^2$, $yz$ spectral functions show emergent spectral weight at $E_F$, while the $3z^2-r^2$, $xz$, $yz$ spectral functions now show pseudogap behavior. Hence, hole doping Site 1 of Ca$_2$O$_3$Fe$_3$S$_2$ is predicted to lead to orbital-selective metallicity compared to its electron doped counterpart.

What is the origin of these orbital-selective features? In a MO system like the Fe-oxychalcogenides, strong (incoherent) scattering between different carriers in orbital states split relative to each other due to the specific crystal field and one-electron band fillings leads to two main effects: (i) It leads, via static-Hartree contributions (from the static part of the orbital-dependent self-energies) to orbital-dependent shifts of the $d$-bands relative to each other, and (ii) strong dynamical correlations due to sizable $U,U'$, $J_H$ cause appreciable spectral weight transfer over large energy scales, from high to low-energy, upon carrier doping. This second feature leads to a drastic modification of the spectral lineshape as shown in Figs. 5 and 6. Microscopically, strong incoherent scattering, arising from co-existence of Mott-localized and bad metallic states, leads to an almost complete suppression of the Fermi liquid quasiparticles and the emergence of an incoherent state, reminiscent of what is seen in cuprate oxides. Our results imply that doped Ca$_2$O$_3$Fe$_3$S$_2$ should be located in pseudogap regime and the metallic state obtained by the filling-controlled Mott transition would be unconventional both at Sites 1 and 2. The origin of this emergent electronic state is the lattice orthogonality catastrophe$^{39}$ that occurs due to orbital-selective blocking of the coherent motion within LDA+DMFT due to sizable $U'$ in the MO Hubbard model.

Armed with the above results, let us now discuss the implications of our results to future nuclear magnetic resonance (NMR) experiments. Here we recall that the spin-lattice relaxation rate ($1/T_1$) or the nuclear spin relaxation rate ($1/T_1T$) are directly computable using the DMFT propagators. $^{40-42}$ While $1/T_1$ quantifies the...
decay of the nuclear magnetic moments providing valuable information about the fluctuations of the electronic magnetic moments in the solid, $1/T_1 T$ measures the slope of the imaginary part of the local spin susceptibility in the zero-frequency limit.\textsuperscript{42} With this caveat in mind, in Fig. 7 we display the temperature dependence of local dynamical spin fluctuations by computing the NMR spin-relaxation rate $1/T_1$ as a function of $T$ within the paramagnetic phase of Ca$_2$O$_3$Fe$_3$S$_2$ as done earlier for the LaO$_{1-x}$FeAsF$_x$ system.\textsuperscript{40} Though experimental data does not exists yet, the $T$-dependence of $1/T_1$ for Ca$_2$O$_3$Fe$_3$S$_2$ parent compound is predicted to show similar Site-dependent features in the non-magnetically ordered state of Ca$_2$O$_3$Fe$_3$S$_2$. By assuming the hyperfine coupling constants for the two Fe-sites to be identical, a particularly interesting feature to be seen is the power-law behavior and the change in slope at low temperatures, albeit less pronounced as compared to La$_{1-x}$FeAsF$_x$, which marks the changes of MO spin fluctuation modes as $T$ is reduced. If the nuclear spin relaxation at both Fe sites is driven by a single spin degree of freedom, one would expect that the scaled relaxation rate be equal.\textsuperscript{43} However, the different relaxation rates we obtain for Ca$_2$O$_3$Fe$_3$S$_2$ indicates that both the magnitude and the $T$-dependence of the two sites is not driven by the single relaxation mechanism, and that this relaxation mechanism is electronic state dependent. Enhanced Mott localization within Fe$^{3+}$ electronic channel tends to suppress local spin fluctuations due the formation of stronger localized moments in the half-filled shell. On the other hand the similarities in the relaxation rates seems to suggest that the Fe 3$d$ spins fluctuate incoherently on both Fe Sites, so that both of them contribute to the relaxation almost similarly in future site-selective NMR experiments.

III. CONCLUSION

To summarize, we have used LDA+DMFT for a realistic two-channel Hubbard model with five active 3$d$ bands at each channel to perform a detailed study of the paramagnetic Mott insulating state of Ca$_2$O$_3$Fe$_3$S$_2$. In particular, considering the Fe$^{2+}$ ($S = 2$) and Fe$^{3+}$ ($S = 5/2$) valence (spin) states intrinsic to this Fe-oxoxychalcogenide, we predict orbital-selective blocking of electrons and strong correlation fingerprints in the half-filled (Fe$^{3+}$) channel as an effect of multi-orbital Hubbard correlations. Compared to La$_{1-x}$FeAsF$_x$ short Fe-Fe bonds intrinsic to Ca$_2$O$_3$Fe$_3$S$_2$ lead to decreased electron correlation through increasing bandwidth, an undoubtedly significant feature also present in the high-$T_c$ FeSe/SrTiO$_3$ system.\textsuperscript{44} Interesting as well is the emergent electronic behavior when we consider the effect of electron/hole doping in this two-channel system. We predict that localization or low-energy incoherence persists in doped Ca$_2$O$_3$Fe$_3$S$_2$ due to a lattice orthogonality catastrophe induced by orbital-selective Mottness. This orbital blocking phenomenon can be directly tested by a combination of future spectral, transport and nuclear spin-lattice relaxation measurements. Such studies are called for, and should confirm or refute our predictions.

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