A General Mechanism for Orbital Selective Phase Transitions

Yu-Zhong Zhang, Huypo Lee, Hai-Qing Lin, Chang-Qin Wu, Harald O. Jeschke, and Roser Valentí

1Shanghai Key Laboratory of Special Artificial Microstructure Materials and Technology, Physics Department, Tongji University, Shanghai 200092, P.R. China
2Max-Planck-Institut für Physik komplexer Systeme, D-01187 Dresden, Germany
3Beijing Computational Science Research Center, Beijing 100084, China
4Department of Physics and State Key Laboratory of Surface Physics, Fudan University, Shanghai 200433, China

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I. INTRODUCTION

Orbital selective phase transitions (OSPTs) leading to phases where localized and itinerant electrons coexist have attracted extensive interest from both experimentalists and theoreticians, since the observation in the metallic phase of Ca$_{2-x}$Sr$_x$RuO$_4$ ($0.2 \leq x \leq 0.5$) of an anomalous behavior with a Curie-Weiss-like local spin.

In spite of the controversies regarding the applicability of such a proposal to real compounds, various mechanisms for OSPTs have been investigated, such as two orbitals with different bandwidth at half-filling away from half filling with crystal field splitting, coexistence of different orbital degeneracies with crystal field splitting at any filling, or different magnetic states in different orbitals at half-filling.

Recently, various models based on an assumption of the coexistence of localized and itinerant electrons have been proposed in order to describe the magnetism in the new iron-based superconductors. Less work has been done on understanding the origin of such orbital selective phases (OSPs). As one possible mechanism for the OSPTs in the pnictides, bands with similar bandwidths suffering from different intra-band Coulomb repulsion were suggested in analogy to the mechanism of a difference in intra-band, inter-band Coulomb interaction and Hund’s coupling, respectively, which fulfill the rotational invariance of the magnetism observed in the parent compounds of most iron pnictides.

In this paper, a possible OSPT mechanism is proposed based on a simple two-dimensional (2D) two-orbital Hubbard model with both orbitals having different band dispersions. We solve the model in the context of mean-field theory (Hartree-Fock approximation (HFA)). A comparison of our results to those obtained from the dynamical mean-field approximation (DMFA) at and near half filling shows that OSPT can be qualitatively captured already at the mean-field level without taking dynamical fluctuations into account. The advantage of working with the mean-field approximation is that we will be able to investigate a large variety of possible cases not easily accessible within the DMFA. We will show that even in the absence of crystal field splittings or when bandwidths, orbital degeneracies and magnetic states are equal for both orbitals the orbitals have different band dispersions. Such conditions should be generally satisfied in many materials. We find that this OSPT is not sensitive to the strength of Hund’s rule coupling and that heavy doping favors the collinear antiferromagnetic state over the OSPT. We discuss our results in relation to the iron pnictides.

II. MODEL AND METHOD

The 2D two-orbital Hubbard model is defined as

\[
H = - \sum_{\langle ij \rangle, (\gamma \gamma \sigma)} t_{ij, \gamma} \epsilon_{\gamma \sigma} c_{\gamma \sigma}^\dagger c_{\gamma \sigma} + U \sum_{\langle \gamma \rangle} n_{\gamma \uparrow} n_{\gamma \downarrow} + \left( U' - \frac{J}{2} \right) \sum_{\langle \gamma \gamma' \rangle} n_{\gamma \uparrow} n_{\gamma' \uparrow} - 2J \sum_{\langle \gamma \gamma' \rangle} S_{\gamma \uparrow} \cdot S_{\gamma' \downarrow},
\]

where $t_{ij, \gamma} = t_s$ ($t'_s$) is the intra-orbital hopping integral between NN (NNN) sites denoted by $\langle ij \rangle$ ($\langle\langle ij \rangle\rangle$) with band indices $\gamma = \alpha, \beta$ in units of $t$. $U$, $U'$ and $J$ are the intra-band, inter-band Coulomb interaction and Hund’s coupling, respectively, which fulfill the rotational invariance condition $U = U' + 2J$. The pair-hopping term is ignored as it does not affect our mean-field results. $c_{\gamma \sigma}^\dagger$ ($c_{\gamma \sigma}$) creates (annihilates) an electron in orbital $\gamma$. A General Mechanism for Orbital Selective Phase Transitions

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of site $i$ with spin $\sigma$. $n_{i\gamma\sigma}$ is the occupation operator, while $n_{i\gamma} = n_{i\gamma\uparrow} + n_{i\gamma\downarrow}$, and $S_{i\gamma}$ the spin operator.

In order to access the true ground state in a 2D system with hoppings up to NNN sites, the original lattice is divided into two sublattices A and B (see Fig. 1 (a)), allowing us to consider various magnetic states in uniform form, such as the PM state, ferromagnetic (FM) state with momentum $Q_{A/B,\gamma} = (0, 0)$ and magnetization $m_{x/y} = m_{B,\gamma}^{x/y}$, NAF state with $Q_{A/B,\gamma} = (0, 0)$ and $m_{x/y} = -m_{B,\gamma}^{x/y}$, collinear AF (CAF) state with $Q_{A/B,\gamma} = (0, 0)$ and $m_{x/y} = m_{B,\gamma}^{x/y}$, non-collinear AF (NCAF) state with $Q_{A/B,\gamma} = (\pi, \pi)$ and $m_{x/y} = m_{B,\gamma}^{x/y}$, bi-collinear AF (BAF) state with $Q_{A/B,\gamma} = (0, \pi)$ and $m_{x/y} = m_{B,\gamma}^{x/y}$, collinear AF (CAF) state with $Q_{A/B,\gamma} = (\pi, \pi)$ and $m_{x/y} = m_{B,\gamma}^{x/y}$. The correspond-}

FIG. 1: (Color online) Cartoons for the different magnetically ordered states we use in our calculations. (a) Paramagnetic state. The choices of sublattice and coordinate system are shown. (b) Ferromagnetic, (c) Neel, (d) collinear, (e) non-collinear, (f) bi-collinear antiferromagnetic states.

III. COMPARISONS BETWEEN THE RESULTS FROM HFA AND DMFA

In order to check the validity of our mean-field calculations, we first compare our results with those obtained using the DMFA. For this comparison, the chemical potential rather than the filling is fixed as is usually done in DMFA studies, and only the NAF state is allowed as required by a two-sublattice calculation within the DMFA. Fig. 2 (a) shows the sublattice magnetization as a function of interaction $U/t$ for the case $t_\alpha = 1$, $t_\beta = 0.6$, and $t'_\beta = 0$. We find that while the magnetic phase transition obtained from the HFA happens earlier than that from the DMFA and higher magnetization is detected in the HFA -indicating that dynamical fluctuations ignored in the HFA strongly suppress the magnetically ordered states- the variation of the magnetization with $U/t$ obtained from the DMFA can be qualitatively reproduced by the results from the HFA. Furthermore, all the phases given from the DMFA can be qualitatively captured by the HFA as shown in Figs. 2 (b)-(d) and (e)-(g) which depict the density of states (DOS) in the different phases obtained from DMFA and HFA, respectively. The OSPT, where one orbital becomes localized while the other remains metallic, is clearly detected by both DMFA (Fig. 2(c)) and HFA (Fig. 2(f)). The resulting OSP is sandwiched between the PM metallic state and the NAF insulating state as seen in Fig. 2 (b), (d) for DMFA and Fig. 2 (e), (g) for HFA. The qualitative consistency between the results from HFA and DMFA imply...
the validity of our following discussion on the OSP as well as on other phases in our model at the mean-field level. In fact, it is already known from the DMFA - where spatial fluctuations are absent - that the PM metal-insulator transition which is inaccessible to the HFA is precluded by a magnetic phase transition in the half-filled case at zero temperature [3], which may be qualitatively described by the HFA. Comparing DMFA and HFA at other hopping parameters with different \( t'_\beta \) (not shown here), we concluded that dynamical fluctuations play a minor role in the OSPT.

IV. THE GENERAL MECHANISM FOR OSPT

In the following we shall investigate the case of fixed filling at 1/2 in contrast to the case of fixed chemical potential where the filling is changed as a function of interaction \( U/t \). All the magnetically ordered states shown in Fig. [1] are taken into account and the ground state is the one with lowest total energy. Fig. [2] (a) shows the phase transitions happening at \( t_\alpha = 1, t_\alpha' = 0.8 \) and \( t_\beta = 1, t'_\beta = 0 \) as a function of \( U/t \). As long as \( U/t < 2.88 \), the ground state is a PM metal with orbital order. In a small interaction region of \( 2.88 < U/t < 3.08 \), an NAF metal with orbital order appears. Further increasing \( U/t \) from 3.08 up to 4.02, the \( \alpha \) orbital exhibits NAF insulating behavior while the \( \beta \) orbital keeps the NAF metallic state, indicating an OSP. Orbital order disappears in this interaction region. At \( U/t > 4.02 \), both orbitals display NAF insulating behavior.

Though we have demonstrated that the OSPT is still present at fixed filling in finite dimension, the mechanism for it has not yet been identified. After analyzing the noninteracting DOS, we find that several possible mechanisms coexist, such as (i) two orbitals having different bandwidth with the ratio of \( W_\alpha/W_\beta \) which can be viewed as the existence of an effective crystal field splitting, and (ii) two orbitals having distinct band dispersions which leads to different shapes of the noninteracting DOS. The last effect was not considered in previous DMFA studies where a semicircular DOS with particle-hole symmetry is usually employed for all the orbitals. In the following, we will reveal that orbitals having distinct band dispersions play a crucial role in the OSPT with magnetic order.

In order to figure out the essential mechanism responsible for the OSPT observed above, we will study separately three cases. (i) We first eliminate the effect of different bandwidths by rescaling the hopping parameters of the \( \alpha \) orbital from \( t_\alpha = 1, t'_\alpha = 0.8 \) to \( t_\alpha = 0.769, t'_\alpha = 0.615 \) so that the ratio of \( t'_\alpha/t_\alpha = 0.8 \) is retained while the ratio of bandwidths becomes \( W_\alpha/W_\beta = 1 \). Fig. [3] (b) presents the various phases as a function of \( U/t \) after rescaling. Though the critical points are changed due to the change of the total bandwidths, all the phases involving OSP are preserved, indicating that such an OSPT exists in the absence of bandwidth differences between orbitals.

As a second case (ii), we remove the orbital order by adding an effective crystal field splitting, by which the half-filling condition is simultaneously satisfied at \( U/t = 0 \) in both orbitals. Fig. [3] (d) shows that the OSPT is still present in the absence of orbital order. However, the states with metallic behavior in both orbitals vanish since the Fermi level is located right at the van Hove singularity in the \( \beta \) orbital at \( U/t = 0 \). We have checked that a small \( t'_\beta \) which shifts the van Hove singularity away from the Fermi level leads to the appearance of metallic phases in both orbitals at finite \( U/t \).

As a third option (iii), we eliminate the effect of orbitals having distinct band dispersions but retain the difference in bandwidth by choosing \( t_\alpha = 1, t'_\alpha = 0.8 \) and \( t_\beta = 0.769, t'_\beta = 0.615 \) which leads to \( t'_\alpha/t_\alpha = t'_\beta/t_\beta = 0.8 \) and \( W_\alpha/W_\beta = 1.3 \). As shown in Fig. [3] (c), an OSPT is precluded by NCAF states, resulting in only two successive phase transitions from PM metals to NCAF insulators through NCAF metals in both orbitals. Clearly, the OSPT will be replaced by NAF insulating states in both orbitals at any finite \( U/t \) if we take \( t_\alpha = 1, t'_\alpha = 0 \) and \( t_\beta = 1.3, t'_\beta = 0 \), which means a similar dispersion relation \( t'_\alpha/t_\alpha = t'_\beta/t_\beta = 0 \) but different bandwidth \( W_\alpha/W_\beta = 1.3 \), since the Fermi level crosses the van Hove singularities in both orbitals.

Our results so far strongly point to the fact that orbitals with distinct band dispersions are crucial for the OSPT since it is present even though all the other mechanisms mentioned above are absent while different bandwidth alone will not support the existence of OSPT when magnetic order is considered. Fig. [3] (e) presents a phase diagram in the \( U/t-t'_\alpha \) plane at \( t_\alpha = 1, t_\beta = 1 \), and \( t'_\beta = 0 \). An OSP exists in a wide region of the phase diagram. The phase transitions from both NAF states to the NCAF state and from PM metal to NAF insulator are of first order (solid line), otherwise second order (dotted line). The NAF metallic state has also been detected in the one-band Hubbard model with NN and NNN hopping [15].

V. VARIOUS EFFECTS ON THE OBSERVED OSPT

Finally, we investigate various effects on the observed OSPT. From the phase diagram, it is obvious that we should discuss two cases separately: 1) \( t'_\alpha/t_\alpha > 1 \) where different effective orders like NAF and NCAF orders compete with each other; 2) \( t'_\alpha/t_\alpha < 1 \) where only NAF order occurs. In Fig. [4] we show the results at \( t_\beta = 1, t_\alpha = 1, t'_\alpha = 2 \). We first present the effect of adding NNN hopping \( t'_\beta \). It is found that increasing \( t'_\beta \) favors the NCAF state, which squeezes the region of the NAF OSP. As seen in Fig. [4] (b), at \( t'_\beta = 0.4 \), the region of OSP is smaller than that at \( t'_\beta = 0 \) (Fig. [4] (a)) and at \( t'_\beta = 0.8 \).
FIG. 3: (Color online) Variation of the magnetization as a function of $U/t$ (a) at $t_\alpha = 1$, $t'_\alpha = 0.8$ and $t_\beta = 1$, $t'_\beta = 0$, (b) at $t_\alpha = 0.769$, $t'_\alpha = 0.615$ and $t_\beta = 1$, $t'_\beta = 0$, (c) at $t_\alpha = 1$, $t'_\alpha = 0.8$ and $t_\beta = 0.769$, $t'_\beta = 0.615$, (d) after eliminating the orbital order by adding an effective crystal field splitting of $\Delta = 0.798$ at $t_\alpha = 0.769$, $t'_\alpha = 0.615$ and $t_\beta = 1$, $t'_\beta = 0$, where the term for crystal field splitting is written as $\sum_i \Delta(n_i^\beta - n_i^\alpha)$. (e) Phase diagram in $U/t-t'_\alpha$ at $t_\alpha = 1$, $t_\beta = 1$, $t'_\beta = 0$. Here $J/U = 0.25$ and filling is $1/2$. Regions of different phases are indicated by the abbreviations defined in the text. M (I) denotes metal (insulator). Solid and dotted lines represent first and second order phase transitions, respectively.

the OSP completely vanishes (see Fig. 4 (c)). However, for the case of $t'_\alpha = 0.8$ (not shown), the region of the OSP remains unchanged at $t'_\beta = 0.4$, while it is reasonably replaced by NCAF state at $t'_\beta = 0.8$. The effect of Hund’s rule coupling is presented in Fig. 4 (d). Compared to Fig. 4 (a) where $J/U = 0.25$, the region of OSP is enlarged at $J/U = 0.0625$ and a direct first-order phase transition from PM metals in both orbitals to the NAF OSP is observed instead of two successive second-order phase transitions through an intermediate NAF metallic state at $J/U = 0.25$. For the case of $t'_\alpha = 0.8$, $t'_\beta = 0$ (not shown), a similar effect of the Hund’s rule coupling is found.

Fig. 4 (e) shows that at a small concentration of electronic doping of 2.5%, the OSP with NAF order is slightly moved to higher values of $U/t$ and the NCAF insulating states existing in the undoped case is replaced by a small region of OSP with NCAF order which eventually become CAF metallic states at larger $U/t$. At large doping of 20%, only two phases with PM and CAF metallic states
remain and the OSP vanishes as seen in Fig. 4(f). The critical value of doping concentration where the OSP disappears is around 13.6%. For the case of $t'_\alpha = 0.8$, $t'_\beta = 0$ (not shown), the OSPT also exists at 2.5% doping but is excluded by PM and CAF metallic states at 20% doping. It is interesting to notice that CAF metallic states only appear when the system is doped. 20% electronic doping is related to the filling factor in the pnictides where 6 3d electrons occupy 5 3d orbitals. However, after examining various sets of model parameters, including those for the pnictides, we should emphasize that the OSPT disappears whenever CAF order occurs.

VI. DISCUSSIONS AND CONCLUSIONS

Recently, various efforts have been made in order to reconcile the controversies about the origin of the CAF phases observed in the iron pnictides. Models containing coupled local spins and itinerant electrons have been proposed. Experimental data has been also interpreted in terms of a coexistence of local and itinerant electrons. However, such a compromise doesn’t seem to be supported by the present study of OSPT with magnetic order. We find that OSPT and CAF order tend to avoid each other. Also, involving the inter-orbital hoppings do not favor the OSP with CAF order. On the other hand, it is not to be expected that increasing the orbital degrees of freedom will dramatically change the situation. Furthermore, existing mechanisms proposed within the PM state are in conflict with the fact that the low temperature phases of most pnictides are magnetically ordered, and the bandwidth of different orbitals are almost the same. However, quantum fluctuations, especially spatial rather than dynamical fluctuations, which favors paramagnetic states, may be responsible for possible OSPT in the pnictides.

In summary, we propose a general mechanism for
an OSPT in magnetically ordered states. Different orbitals with different band dispersions should be quite widespread in real materials. Importantly, the OSPT according to the presented mechanism occurs in a wide range of model parameters, suggesting that this mechanism could be realized in nature.

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* Corresponding author. E-mail: yzzhang@tongji.edu.cn

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