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Is the nature of magnetic order in copper-oxides and in iron-pnictides different?

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Abstract. - We use the results of first-principles electronic structure calculations and a strong coupling perturbation approach, together with general theoretical arguments, to illustrate the differences in super-exchange interactions between the copper-oxides and iron-pnictides. We show that the two magnetic ground states can be understood in a simple manner within the same theoretical foundation. Contrary to the emerging view that magnetic order in the iron-pnictides is of itinerant nature, we argue that the observed magnetic moment is small because of frustration introduced by the electrons of the Fe orbitals as they compete to impose their preferred magnetic ordering.

The copper-oxide layers present in the high-Tc superconducting families are turned into superconductors by introducing dopants that create electrons or holes in these otherwise antiferromagnetic (AF) insulating layers [1]. The proximity of antiferromagnetism to superconductivity has led to the general view that this form of magnetic order is intimately related to the mechanism of superconductivity in these materials [2]. In the recently discovered iron-pnictide based superconductors [3–8], which exhibit superconductivity at relatively high-Tc, the copper-oxide layer is replaced by an iron-pnictide layer. Interest in the new materials, reminiscent of that seen when the cuprates were discovered more than two decades ago, is due to the fact that many unsuccessful attempts were made to replace the copper-oxide layer in high-Tc materials to facilitate practical applications. The parent compounds of these iron-pnictide materials, like the copper-oxide parent materials, show a spin-density-wave order [9, 10] illustrated in Fig. 1(a). Unlike the copper-oxides, the parent compounds in the iron-pnictides, such as the pure LaOFeAs, are metallic, but are magnetically ordered and non-superconducting and they become superconductors by doping with electrons or holes. The copper-oxide parent compounds are well described as spin-1/2 Heisenberg quantum antiferromagnets [2]. Furthermore, it is widely believed that superconductivity in copper-oxides arises when, by doping the quantum antiferromagnet, the carriers (holes or electrons) form a strongly correlated Anderson-Mott type system with the spin-spin correlations playing a fundamental role in the superconductivity mechanism. After the discovery of the iron-pnictide superconductors, there is an emerging view that the magnetism in these compounds is of itinerant type [11–14] and that these systems are different from the cuprates and in fact weakly correlated. This is an important issue to settle, because any further theoretical analysis of other properties of this new family of materials, including the still unknown superconductivity mechanism, depends on it.

Here we focus on this issue and seek a broader framework to reconcile the different forms of magnetic order and to explain the magnetic properties in both families of materials with the same approach. Our arguments are inspired by the results of Ref. [15], where first-principles electronic structure calculations based on density functional theory (DFT) were combined with a strong coupling expansion to obtain an effective low-energy Hamiltonian which describes the electrons occupying the five Fe orbitals. In the present work, we address the nature of the magnetic order in the iron-pnictide and the copper-oxide based materials using a more general scheme, which is
Fig. 1: (a) The spin-density-wave order (columnar antiferromagnetic) as observed by neutron diffraction. The Fe magnetic moments along the (1,1) direction are aligned, while two nearest neighbouring such chains are antiferromagnetically aligned. (b) The familiar antiferromagnetic ordering of the cuprous oxides. The shaded square denotes the unit cell. With the mapping Fe → Cu, As → O, the CuO₂ unit cell differs from the Fe₂As₂ one by a magnetic atom missing from the centre.

Fig. 2: Super-exchange between two nearest neighbour (NN) or next nearest neighbour (NNN) Fe atoms through electron-hopping (hybridization) process between any Fe d-orbital and any p or s orbital of an intervening As atom. One of the As spin-up electrons (shown as red) hops to a singly occupied Fe d-orbital (process 2), then the spin-down electron (shown as blue) from the same As orbital hops to a NN or NNN Fe d-orbital (process 3). With two successive hops (processes 4 and 5) the other electrons that were initially occupying the same two Fe d states, hop to the same As site, restoring its doubly occupied status. The initial configuration 1 differs from the final configuration 5 by the spin orientation on the two Fe orbitals. Such processes lead to AF spin-spin interaction between electron spins occupying any two d orbitals belonging to NN or NNN Fe atoms.

The case in which an orbital is occupied by two electrons is assumed that the particular Fe d-orbital is singly occupied. The initial configuration 1 differs from the final configuration 5 by the spin orientation on the two Fe orbitals. Such processes lead to AF spin-spin interaction between electron spins occupying any two d orbitals belonging to NN or NNN Fe atoms.

for NN and NNN pairs of magnetic ion sites. Here we have assumed that the particular Fe d-orbital is singly occupied. The case in which an orbital is occupied by two electrons is discussed separately. Other models in the literature, as in Refs. [16–18], are similar to ours with the very important difference that our Hamiltonian, Eq. (1), treats differently the various Fe d-orbitals labelled by ν. The condition for quantum- Néel (checkerboard) ordering (shown in Fig. 5(b)) is

\[ J^{(1)}_{\nu} > 2J^{(2)}_{\nu}; \]

if the coupling between spins of electrons occupying the orbital ν do not satisfy the above condition, the SDW order of Fig. 5(a) is energetically favourable against the checkerboard order (Fig. 5(b)).

By fitting the first-principles electronic structure results to a tight-binding model, we have established that, with the exception of the \(d_{x^2-y^2}\) orbital, for any given Fe d-orbital, there is only one orbital of the intervening As (of s or p mixed character) with which the hybridization amplitude is most significant. Therefore, the most significant
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![Diagram showing magnetic interactions](image)

Fig. 3: (a) An example of the paths that contribute to the spin-exchange interactions $J^{(1)}_v$ and $J^{(2)}_v$ between NN and NNN Fe atoms for $v = d_{xz}$ and through the $p_x$ or $p_y$ orbital of the intervening non-magnetic As atoms. (i) Contributions to $J^{(1)}_v$ of the NN Fe atoms indicated as M1 and M2: First there is a contribution from direct hopping $t$ between the magnetic ions as shown by the red path. Second there is the super-exchange through the $p_x$ or $p_y$ orbital (only the $p_x$ is drawn) of the intervening As atom N1 or N2. One of these is shown by the green-blue path. (ii) Contributions to $J^{(2)}_v$ for the NNN magnetic atoms M1 and M3 through the $p_x$ or $p_y$ orbital of the intervening As atom N1 shown as a green-yellow path. (b) The corresponding situation for the copper-oxide layer: There is no corresponding M2 magnetic atom and the spin-spin interaction between the NN Cu atoms M1 and M3 for this case occurs through the hybridization of the $d_{xz}$ orbital of the Cu atom M1 or M3 and the $p_x$ orbital of the intervening O atom N1.

Super-exchange processes involve the same intervening As orbital for both of the exchanged electrons. The super-exchange process illustrated in Fig. 2 requires four steps because two electrons, each occupying one of the two NN or NNN magnetic (Fe) ions, need to move through hopping to the intervening non-magnetic ion (As) and, thus, each electron has to make two steps. To obtain the total spin-spin interactions we need to consider also the direct hopping of electrons between these two neighbouring Fe or Cu d-orbitals. The results of Ref. [15] show that the direct hopping matrix element between two NN Fe d-orbitals, indicated as $t$ in Fig. 3, is smaller but not negligible. In the case of copper-oxides this direct process is non-applicable because two NN Cu atoms correspond to NNN Fe atoms in the Fe-pnictides.

Fig. 3 illustrates paths that give rise to the most significant contribution to $J^{(1)}_v$ and $J^{(2)}_v$, and compares the super-exchange interactions between the iron-pnictide layer and the copper-oxide layer through the intervening non-magnetic As or O atom respectively. For example, for iron-pnictides, $J^{(1)}_{xz}$ has two contributions: (i) one from the direct hopping $t_{xz}$ between two NN Fe $d_{xz}$ orbitals, (shown as the red path in Fig. 3), and (ii) a super-exchange contribution due to processes such as the one illustrated by the combined green-blue path in Fig. 3. The hybridization $V_{xz}^x$ of the $d_{xz}$ orbital of the Fe atom M1 and the $p_x$ orbital of the As atom N1 is very large ($\sim 1.4$ eV), while its hybridization $V_{yz}^y$ with the $p_y$ orbital of the same atom N1 is negligible ($\sim 0.1$ eV). The reverse is true for the hybridization of the same Fe $d$-orbital with the $p_x$ and $p_y$ orbitals of atom in position N2 because of the 90° relative orientation. The contribution to $J^{(1)}_{xz}$ due to the green-red process illustrated in Fig. 3 is proportional to

$$J^{(1)}_{xz} \sim (V_{xz}^x/V_{xz}^y)^2.$$  

(3)

On the other hand, the contribution to $J^{(2)}_{xz}$ due to the green-yellow process illustrated in Fig. 3 is proportional to

$$J^{(2)}_{xz} \sim (V_{xz}^y)^4.$$  

(4)

Therefore, $J^{(2)}_{xz}/J^{(1)}_{xz} \sim (V_{xz}^y/V_{xz}^x)^2 \sim 0.01$ and the super-exchange contribution to $J^{(1)}_{xz}$ is negligible as compared to that of $J^{(2)}_{xz}$ between NNN Fe atoms. Hence, we can conclude that in the subspace formed by $d_{xz}$ and $d_{yz}$, $J^{(1)}_{xy}$ has contributions mainly through direct hopping while $J^{(2)}_{xy}$ has significant super-exchange contributions similar to the copper-oxide case, as shown in Fig. 3(b).

Fig. 4: The Hilbert space of the five Fe d-orbitals is divided into three subspaces ($S_1$ spanned by $d_{x^2-y^2}$, $S_2$ spanned by $d_{xz}$, $d_{yz}$, and $S_3$ spanned by $d_{xy}$, $d_{xz}$). These subspaces are coupled through the Hund’s rule coupling $J^H$ which is indicated by the blue arrows.

On the other hand, the NN spin-spin interaction $J^{(1)}_v$, involving the other three Fe d-orbitals ($yz, xy, z^2$), has contributions from super-exchange processes in which the intervening As orbital is the $sp_z$ (a linear combination of the As 4s and 4p$_z$). In these processes the magnitude of the hybridization of the $yz, xy, z^2$ Fe d-orbitals with the $sp_z$-As orbital does not change with a 90° rotation which necessarily occurs in the red-green path. Therefore, for these three orbitals, $J^{(1)}_v$ is significantly larger than $J^{(2)}_v$, which implies that the condition (2) for checkerboard order is fulfilled.
Fig. 5: (a) Illustration of the observed columnar AF order in the undoped iron-pnictides. This state is strongly favoured by the $S_2$ subspace (spanned by $d_{xz},d_{yz}$). (b) Illustration of the familiar checkerboard Néel ordering which is preferred by the $S_1$ subspace (spanned by $d_{x^2−y^2}$ and by the $S_3$ subspace (spanned by $d_{xy},d_{z^2}$). Notice that the magnetic moments of the Fe atoms along the green lines are frustrated because their orientations in case (a) are opposite to their orientations in case (b).

The NN AF couplings $J^{(1)}_d$ in the subspaces $S_1$ and $S_3$ are greater than the NNN AF couplings $J^{(2)}_d$; however, in the subspace $S_2$ the NNN AF coupling $J^{(2)}_{xz}$ is large, that is, the condition (2) is not satisfied for the arguments presented above. As a result the subspace $S_2$ alone prefers the observed SDW state illustrated in Fig. 5(a), while the other two subspaces prefer the checkerboard-type quantum-Néel order shown in Fig. 5(b). These two competing orders create a magnetic frustration illustrated in Fig. 5. Namely, in the two ordered states illustrated in Fig. 5, the magnetic moments of the electrons on the iron atoms along the alternating green lines in Fig. 5 are opposite. The Hund’s rule coupling, which is significantly stronger than the AF coupling between NN and NNN atoms, forces the electrons on these different subspaces to choose a common spin orientation. However, any such global choice within the same atom will minimize the energy in one subspace and at the same time frustrate the other subspaces as discussed in the caption of Fig. 5. In the case of the undoped material where there are six electrons occupying the five Fe $d$-orbitals, the $S_1$ subspace has zero spin; hence, the $S_2$ subspace competes with the $S_3$ subspace (which prefers the checkerboard AF order) only and because $J^{(2)}_{xz}$ is larger than the NNN AF couplings in $S_3$, it imposes the observed SDW state. The local magnetic moment is expected to be small due to this frustration. If we neglect the influence of the other subspaces, and we restrict ourselves to $S_2$, the maximum expected order should be less than $2\mu_B$ because of the reduction of the magnitude of the order parameter due to zero-point spin fluctuations. If we turn on the interaction with the other subspaces, this has a frustrating effect of the magnetic moment. Therefore, the present analysis gives a natural explanation for the observed small magnitude of the magnetic moment [9], while the calculations based on the itinerant picture [11,12,17–19] produce values for the Fe magnetic moment greater than $2\mu_B$.

From this analysis, we have shown that the different ordering in the iron-pnictides and copper-oxides can be understood within the same theoretical foundation. The observed small Fe magnetic moment arises from the fact that only the electrons occupying the subspace spanned by $d_{xz},d_{yz}$ prefer the observed SDW order (Fig. 5(a)), while the electrons occupying the other three Fe orbitals prefer a checkerboard ordering (Fig. 5(b)), which create large zero-point spin fluctuations. Including these frustrating effects, we expect a significantly reduced local moment in agreement with neutron scattering experiments.

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