I. INTRODUCTION

The calculation of non-zero-temperature and dynamical properties, such as magnetic transition temperatures and conductivities, is a long-standing and difficult problem in materials theory, but one for which the recent theoretical development of the ‘dynamical mean field’ method offers a promising avenue for progress. This method, which allows an exact (in principle) treatment of local correlations has been recently used in combination with ‘ab initio’ band calculations to estimate the magnetic transition temperature and optical conductivities are found to depend sensitively on the band structure. For parameters consistent with local spin density approximation band calculations, the computed transition temperatures are lower than observed, and in particular decrease dramatically as band filling is increased, in contradiction to experiment. Band parameters which would increase the transition temperature are identified.

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In this paper we present a model for the low (less than, say 3eV) lying electronic states, which are important for transport and magnetism. Our model consists of a tight binding description of the bands, which we derive from previously published first principles calculations and general arguments, and a local interaction (namely a Hunds coupling on the Fe site). The key physical assumption made in our model is that the magnetism is driven by the strong Hunds coupling on the Fe site. Important technical issues include the two dimensional nature of the underlying band structure and the multi-orbital nature of the material. We solve the model in the dynamical mean field approximation, and from our solution determine the magnetic transition temperature and optical conductivity, and attempt to determine the general materials aspects which control $T_c$. This paper supercedes a previous paper in which the model Hamiltonian used did not provide an adequate approximation to the underlying band structure.
II. MATERIAL AND MODEL

A. Material

Double perovskite systems form in the $AB\prime B'O_6$ crystal structure which generalizes the $ABO_3$ perovskite structure familiar from ferroelectrics, high temperature superconductors and the ’colossal’ magnetoresistance rare earth manganites by having two different $B$ site ions. In the double perovskite materials of interest here, $A$ is an alkaline earth such as $Sr$, $Ca$ or $Ba$ and the $B',B''$ sites form a rocksalt structure, i.e. a simple cubic lattice with a doubled unit cell and one sublattice occupied by $Fe$ and the other by a transition metal from the 4$d$ or 5$d$ series such as $Mo$ or $Re$. The crystal fields and atomic energetics are such that the formal valences correspond to $Fe$ with a half filled, maximally polarized $d$-shell while the $Mo/Re$ has one or two $d$ electrons distributed over the $t_{2g}$ levels. We will focus on electronic states arising from the transition metal $d$-levels.

B. Hamiltonian

1. Overview

The Hamiltonian describing the low lying, electronically active degrees of freedom may be written as the sum of a ’hopping’ part arising from the band structure and an interaction part:

$$H = H_{\text{band}} + H_{\text{int}}$$  \hspace{1cm} (1)

The relevant portions of the calculated band structure involve three bands (degenerate in the ideal double perovskite structure) arising from the three transition metal $t_{2g}$ levels $d_{xy,yz,xz}$. To a high degree of accuracy these three bands do not hybridize with each other and the physics is therefore described by a three-fold degenerate tight binding model. The planar character of the $t_{2g}$ levels implies that the tight binding model has an interesting two dimensionality, which may be summarized as follows. The $d_{xy}$ orbital on a $Fe$ site hybridizes via a matrix element $t_1$ with the $d_{xy}$ levels on the four nearest neighbor ($Mo/Re$), sites in the same plane and via a much smaller matrix element $t_3$ to the four nearest $Fe$ ions also in the same plane. The hopping in the third direction is negligible, because of the planar character and $xy$ orbital symmetry of the $d_{xy}$ wave function. The $d_{xy}$ orbital on a $Mo/Re$ site hybridizes with the four in-plane near neighbor $Fe$ sites via the same hopping matrix element $t_1$ and with the four in-plane second nearest neighbor ($Mo/Re$) sites, via another matrix element $t_2$, which is not particularly small, because of the more spatially extended character of the $d$-electrons in 4$d$/5$d$ orbitals. Further neighbor hoppings are also found to be important in other $t_{2g}$-based 4$d$ systems such as $Sr_2RuO_4$.

It is natural to assume that the magnetic character of the material derives from the strongly magentic nature of the $Fe$ ion and we therefore assume that the dominant interaction arises from the strong atomic Hunds coupling of the $Fe$.

2. Hopping Hamiltonian

To write the Hamiltonian explicitly we focus the cubic lattice of $B,B'$ sites in the underlying single perovskite structure, labelling these sites by $i$ and the operator creating an electron into orbital $a(= xy,yz,xz)$ and spin $\sigma$ by $c_{a,i,\sigma}^+$. Although we refer to this orbital as a ’$d$-orbital’ it in fact represents a hybrid, composed mainly of transition metal $d$ and oxygen $p$ orbitals, of the correct local symmetry. We introduce a nearest neighbor ($Fe \leftrightarrow Mo/Re$) hopping $t_1$ and two second neighbor (same sublattice) hoppings $t_2$ and $t_3$ representing $Mo - Mo$ or $Fe - Fe$ hoppings respectively. As noted above we expect that $t_2$ corresponding to to $Mo - Mo$ hopping is relatively large, because of the larger spatial extent of the $4d/5d$ orbitals while $t_3$ is essentially negligible. To obtain the conductivity we couple in the electric field by using a vector potential and the Peierls phase ansatz. This approximation has been shown to be accurate in other transition metal oxide contexts. Thus the hopping portion of the Hamiltonian is the sum of three identical tight binding models, one for each orbital. The Hamiltonians take the general form (note that the first sum runs over all lattice sites, the second over the $B'$ (non-$Fe$) and the third over the $B$ ($Fe$) sites, while $\delta_a$ labels the in-plane direction relevant to orbital $a$ and we have set the electric charge $e$ and the speed of light $c$ equal to unity)

$$H_{\text{band}} = - \sum_{a,i,\delta_a,\sigma} (t_{1,a} e^{iA\delta_a} c_{a,i,\sigma}^+ c_{a,i+\delta_a,\sigma} + H.c.)$$

$$- \sum_{a,i \in B',\delta_a} \left( t_{2,a} e^{iA\delta_a} c_{a,i,\sigma}^+ c_{i+\delta_a,a,\sigma} + H.c. \right)$$

$$- \sum_{a,i \in B,\delta_a} \left( t_{3,a} e^{iA\delta_a} c_{a,i,\sigma}^+ c_{i-\delta_a,a,\sigma} + H.c. \right)$$ \hspace{1cm} (2)

$H_{\text{band}}$ implies an interesting band structure, which is most plainly revealed by writing $H_{\text{band}}$ in momentum space in a matrix notation where the upper left entry corresponds to $Fe$ and the lower right to $Mo$, thus if $A = 0$ we have, for the $xy$ orbitals

$$H_{\text{band,}xy}[A = 0] =$$

$$\begin{pmatrix}
0 & -2t_1 (\cos p_x + \cos p_y) \\
-2t_1 (\cos p_x + \cos p_y) & -4t_2 (\cos p_x \cos p_y)
\end{pmatrix}$$

where we have set the $Fe - Mo$ distance to unity and the momenta are restricted to the reduced Brillouin zone $|p_x| + |p_y| < \pi$. 

3. Interaction

The most important interaction effect constrains the occupancy of the \( B (F e) \) site. The formal valence of \( F e \) is \( d^5 \) and the strong Hunds coupling characteristic of \( F e \) (and found in the local spin density approximation to band theory) means that in the \( d^5 \) configuration all of the \( F e \) \( d \)-electrons are aligned, leading to a filled, completely spin-polarized \( d \)-shell. Two charge fluctuation processes are possible: \( F e d^5 \leftrightarrow F e d^4 \) or \( F e d^5 \leftrightarrow F e d^6 \). The strongly stable nature of the filled \( d \)-shell implies that the \( d^5 - d^6 \) process is dominant. To express this physics we introduce a strong Hunds coupling on the \( B (F e) \) site, expressing the fact that in the ground state the \( F e \) is in the \( d^5 \) maximal spin configuration, and an energy splitting parameter \( \Delta \) expressing the differing electronegativities of the \( B \) and \( B' \) sites. Thus we write

\[
H_{\text{int}} = -J \sum_{a,i \in B, \alpha \beta} \vec{s}_i \cdot \vec{c}_{a,i,\alpha}^{\dagger} \vec{c}_{a,i,\beta} + \sum_{a,i \in B', \sigma} \Delta \sigma \vec{c}_{a,i,\sigma}^{\dagger} \vec{c}_{a,i,\sigma} \tag{4}
\]

For the calculations presented in this paper we will specialize to cubic symmetry, so the \( \Delta \) are the same for all three orbitals, but this restriction may easily be lifted. The energy scale relevant for the \( F e d^5 \leftrightarrow F e d^6 \) valence fluctuation is \( J - \Delta \). Examination of published band structures indicates that \( J - \Delta \sim 1 \text{eV} \) while the \( d^5 \leftrightarrow d^4 \) process has a much larger energy of \(|J + \Delta| \gtrsim 5 \text{eV}|. In a fully spin polarized ground state, the interaction terms simply become level shifts, \( \Delta_{\text{maj}} = J + \Delta \) and \( \Delta_{\text{min}} = -J + \Delta \) for the majority and minority spin bands. Transitions onto the majority-spin \( F e \) orbital involve very large energies, so to simplify the calculations at \( T > T_c \) we will take the limit \( J + \Delta \to \infty \) with \( J - \Delta \) fixed. We henceforth refer to the quantity \( J - \Delta \) as \( \Delta \). Because the local spin density approximation may not be accurate for strongly interacting systems such as the double perovskites, we will consider a range of \( \Delta \) here. Other authors have argued that an additional Hunds-type coupling on the \( B' \) site is important. Technical limitations prevent us from treating such an interaction accurately, so we do not include it here.

4. Discussion: \( T=0 \) Band structure, ferromagnetic case

Eq.\(^3\) may be thought of as describing two bands of electrons: one on the \( F e \) sites, with ‘intrinsic’ bandwidth set by \( t_3 \) and one on the non-\( F e \) sites, with ‘intrinsic’ bandwidth set by \( t_2 \). The two bands hybridize via the overlap \(-2t_1 (\cos p_x + \cos p_y)\). We see immediately that the hybridization vanishes along the line \( \cos p_x + \cos p_y = 0 \) which also contains the van Hove points \( 0, \pi \) and \( \pi, 0 \) at which the density of states of the two individual bands diverges. Near these points a complicated structure including divergences in the density of states is expected.

FIG. 1: Total (solid line) and \( F e \)-projected (dashed line) densities of states computed using tight binding parameters \( t_1 = 0.25 \text{eV}, t_2 = 0.15 \text{eV}, t_3 = 0.03 \text{eV} \). Top panel: \( T = 0 \) majority spin density of states using \( \Delta_{\text{maj}} = -2.5 \text{eV} \) Middle panel: \( T = 0 \) minority spin density of states computed from Eq.\(^3\) with \( \Delta_{\text{min}} = 0.3 \text{eV} \). Lowest panel: total (both spins) density of states at \( T > T_c \) computed as described in section III.
The full density of states and the projection of this
density of states onto the Fe orbitals are shown in the
upper panels of Fig. 1 for parameters $t_1 = 0.25$ eV,
$t_2 = 0.15eV$, $t_3 = 0.03eV$ (note that most of the ma-
jority spin Fe density of states is at a low energy out-
side the range of this plot). Comparison of this den-
sity of states to the published band theory results
shows that these parameters reproduce the band density
of states accurately. The main difference is that if the $J$
and $\Delta$ are adjusted to correctly reproduce the minority
spin band, then the upper (non-Fe-portion of the major-
ity spin band is positioned about 0.5eV too low in energy.
The extra shift in the majority spin Mo orbitals must be
attributed to a Hunds coupling on the non-Fe site, not
included here.

The two features seen in our calculated density of
states near 0.5eV arise from states in the vicinity of
the van Hove points $(0, \pi)$ and $(\pi, 0)$ where the hy-
bridization vanishes and the $B$ and $B'$ sites have en-
ergy $\Delta + 4t_3 \approx 0.4eV$ and $4t_2 \approx 0.6eV$ re-
spectively, whereas the peak at $\omega = 0$ arises from the van Hove
point ($\pi/2, \pi/2$) of the $B'$ (non-Fe) band, where as noted
above the hybridization to the Fe vanishes.

Formal valence arguments indicate that the material
contains one or two d-electrons beyond the filled shell
Fe $-d^6$ $Re/Mo - d^6$ configuration. Fig 1 shows that
for the band theory parameters, these carriers go into
states with only a small admixture of Fe. The physics
behind this result is that for this sign of $t_2$ the strongly
hybridized states near $p_x = p_y = 0$ are at the bottom of
the band described by the $t_2$–only term in $H_{band}$, and
are pushed further away from the Fe states by usual
level repulsion, leading to a mainly non-Fe character of
the lowest states. For $n = 1$ (Re) only the minority spin
band is occupied (the majority spins occupy low-lying Fe
states off of the plotted scale). However, for $n = 2$ within
this approximation, chemical potential is $\mu_2 \approx 0$ and the
majority spin band is somewhat occupied, so the mate-
rial is not a ‘half-metal’ in this approximation. These
features will be seen to be of importance for the calcu-
lated transition temperature and optical conductivity.

It is interesting to consider a contrasting set of param-
eters, for which the level repulsion argument works in the
opposite manner. If $t_2$ has an unphysical (negative) sign
and $\Delta$ is near 0 then the low lying states of the $t_2$-band
do not mix with the Fe states, which are pushed down-
wards by hybridization with the higher-lying Re/Mo lev-
eles, leading to low-lying states of mainly Fe character, as
shown in Fig 2.

C. Conductivity

The current operator $\hat{J} = \delta H/\delta \delta^{23}$. For electric field
in the $x$ direction the $xy$ and $xz$ orbitals contribute, thus
$J_x = J_{x,xy} + J_{x,xz}$ with $\delta$ the lattice vector in the $x$
direction and $\delta'$ labelling the four ‘second neighbor’
lattice vectors $(\delta' = \pm (\delta_x \pm \delta_y))$ so by expanding Eq. 3
in

\[ \mu_2 \approx 0 \text{ eV}. \]

\[ \Delta \min = 0.3eV. \]

\[ \Delta \max = -2.5eV. \]

\[ T \approx 0 \text{ majority spin density of states computed from Eq. 3 with} \]

\[ \mu_2 \approx 0 \text{ with} \]

\[ \Delta \min = 0.3eV. \]

\[ \Delta \max = -2.5eV. \]

\[ T \approx 0 \text{ minority spin density of states computed from Eq. 3 with} \]

\[ \mu_2 \approx 0 \text{ with} \]

\[ \Delta \min = 0.3eV. \]

\[ \Delta \max = -2.5eV. \]

\[ T \approx 0 \text{ total (both spins) density of states at} \]

\[ T \approx 0 \text{ computed as described in section III.} \]
powers of $A$ we obtain

$$J_{xy}(A) = - \sum_{i, \pm \delta_x, \sigma} (i \delta t_{1, \alpha} e^{i A \delta_x} c_{xy, i, \alpha} e_{xy, i + \delta_x, \sigma} - H.c.) - \sum_{i \in B', \delta'_x} (i t_{2, \alpha} (\delta' \cdot \vec{x}) e^{i A \delta'_x} c_{i, \alpha} c_{i + \delta'_x, \alpha} - H.c.)$$

(5)

The expectation value of the term in $J$ proportional to $A$ gives the total oscillator strength, $S(\infty) = \frac{2}{a} \int_0^{\infty} d\omega \sigma(\omega)$ in the conduction band contribution to the optical conductivity (see Appendix A for details). Restoring units we have $(a$ is the Fe–Mo/Re distance, the sum rule is conventionally defined in terms of the conductivity per unit volume and the factor of two comes from the $xy$ and $xz$ orbitals, whose contributions to the expectation values are identical in cubic symmetry)

$$S(\infty) = \frac{2e^2}{a} \left\langle \sum_{i, \pm \delta_x, \sigma} (t_{1, \alpha} c_{xy, i, \alpha} e_{xy, i + \delta_x, \sigma} + H.c.) \right\rangle$$

(6)

$$\sigma(\Omega) = \frac{S(\infty) - 2 \chi_{ij}(\Omega)}{i\Omega}$$

(7)

with $\chi_{ij}$ the usual Kubo formula current-current correlation function evaluated using $J_{xy}(A = 0)$ (Eq 3) and again the factor of two represents the orbital degeneracy.

III. METHOD OF EVALUATION

A. Overview

To evaluate the properties of $H$ we use the dynamical mean field method [13]. This method is extensively described and justified elsewhere, and is relevant here because the principal interactions are local. In brief the central approximation is that the electron self energy, $\Sigma$, is momentum independent. In this circumstance the physics may be derived from a local theory which may be viewed as a quantum impurity model combined with a self consistency condition. The application to the double perovskite systems requires some discussion. In these systems the unit cell contains two sites, each site contains three orbitals and there are two choices of spin, so the local theory has twelve degrees of freedom. However, the problem may be simplified. First, the three orbitals ($d_{xy}$ etc) are coupled only via the interaction. Second, the interaction exists only on the Fe site, so that we may formally integrate out the electrons on the non-Fe ($B'$) site, defining a single-orbital model with the inverse Fe ($B$)-site Green function for e.g. the $xy$ orbitals viz.

$$G_{BB}^{xy,band}(p, \omega) = \omega - \frac{4t_1^2 (\cos(p_x) + \cos(p_y))^2}{\omega + \Delta^{xy} - 4t_2 (\cos(p_x) \cos(p_y))}$$

(8)

We measure momenta in units of $\pi/a$ where $a \approx 4\AA$ is the distance from a $B$ to a nearest neighbor $B'$ site. The two dimensional Brillouin zone is defined by $|p_x + p_y| < \pi$.

The physics is then described by a three-orbital local theory given by the partition function $Z_{loc} = \int Dc^+ e^{\text{exp}[S_{loc}]}$ with an action $S_{loc}$ which we write in the Matsubara frequency representation as

$$S_{loc} = T \sum_{\omega} Tr \left[ c^+_{\alpha \alpha}(\omega) \left( a_{\alpha \beta}^{ab}(\omega) - JS \cdot \sigma_{\alpha \beta} c_{\beta \beta}(\omega) \right) \right]$$

(9)

involving fields $c_{\alpha \alpha}$ and specified by a tensor mean field function $a$ which has orbital ($ab$) and spin ($\alpha \beta$) indices (the trace is over the spin and orbital indices). The mean field function is fixed by the condition that the Green function defined from $S_{loc}$.

$$G_{loc}(\tau) = \frac{\delta \ln Z_{loc}}{\delta a(\tau)} = (a - \Sigma)^{-1}$$

(10)

is equal to the local Green function defined by integrating Eq 8 over momenta using the self energy defined by Eq 10 i.e.

$$G_{loc}^{xy}(\omega) = \int \frac{d^2p}{(2\pi)^2} G_{BB}^{xy,band}(p, \omega - \Sigma)$$

(11)

and the integral is over the Brillouin zone defined above.

Substitution of Eqs 8,9,10 into Eq 11 yields explicit equations which are solved numerically by iteration.

B. Calculation of $T_c$

We calculate the ferromagnetic transition temperature by decomposing the mean field function $a$ into nonmagnetic ($a_0$) and magnetic ($a_1$) parts

$$a = a_0 + a_1 \mathbf{m} \cdot \sigma$$

(12)

and linearizing in $a_1$. We take the magnetization direction $\mathbf{m}$ to be parallel to $z$ and take the limit $J \to \infty$ so that after integrating out the fermions and redefining $a \to a + J$ we obtain ($\cos(\theta)$ is the dot product between the direction of the core spin and of the magnetization)

$$S_{imp} = Tr \ln [a_0^\dagger (\omega) + a_1^\dagger (\omega) \cos(\theta)]$$

(13)

where the $Tr$ is over the frequency index and the orbital degree of freedom.

The Green function of the impurity model becomes

$$G_{imp}^\dagger(\omega) = \frac{1}{2} \left\langle \frac{1 - \hat{S} \cdot \sigma}{a_0 - a_1 \cos(\theta)} \right\rangle$$

(14)
where the expectation value is over the directions of the 'core spin' $S$.

In the paramagnetic phase $a_1 = 0$. Expanding near the magnetic transition (assumed second order) yields

$$G^{\text{imp}}(\omega) = \frac{1}{2a_0} \left( 1 - \left( m + \frac{a_1}{3a_0} \right) \sigma_z \right)$$

(15)

with $m = < \cos \theta > $ so that

$$\Sigma(\omega) = -a_0 - \left( 2a_0m - \frac{a_1}{3} \right) \sigma_z$$

(16)

At $T > T_c$ $m = a_1 = 0$ and substitution of Eqs. (4) into Eq. (11) yields

$$\frac{1}{2a_0(\omega)} = I_1(\omega, a_0(\omega))$$

(17)

where the $n^{th}$ order integral $I_n$ is given by

$$I_n = \int \frac{d^3p}{(2\pi)^3} (G_{22}(\omega))^n$$

(18)

This equation is solved numerically by iteration for a sufficiently dense set of frequency points (typically frequency spacing 0.04$t_1$). Once a solution for $a_0$ is obtained we may linearize Eq. (11) in the magnetic part of the self energy and local Green function, obtaining

$$\frac{m}{2a_0} - \frac{a_1}{6a_0^2} = I_2(\omega, a_0(\omega)) \left( 2a_0m + \frac{a_1}{3} \right)$$

(19)

where

$$m = < \cos \theta > = \sum_{\omega,\alpha} \frac{a_1}{3a_0^2}$$

(20)

Solving for $a_1$ and then using this to obtain an expression for $m$ yields a self consistent equation for $T_c$ which in the limit of cubic symmetry becomes

$$1 = \sum_n \frac{1 - 4a_0^2 I_2}{1 + 2a_0^2 I_2} = \sum_n \left[ -2 + \frac{9}{1 + 2a_0^2 I_2} \right]$$

(21)

It turns out that the transition temperatures are low compared to the other scales of the model so that one may recast this equation via analytical continuation to the real axis as ($\mu$ is the chemical potential corresponding to the desired carrier density)

$$T_c = \int_{-\infty}^{\mu} d\omega \frac{\text{Im} \left[ \frac{9}{1 + 2a_0^2 I_2} \right]}{\pi}$$

(22)

IV. RESULTS AND DISCUSSION

We have calculated the magnetic transition temperature from Eq. (22), finding that for the parameters used to construct Fig. 1 (and which are the ones following from band theory) $n = 1$ (Re case) $T_c \approx 110K$ and that for $n = 2$ (Mo case) the ground state is not ferromagnetic. These calculated values are in sharp disagreement with the experimental values $T_c \gtrsim 400K$ for both $n = 1$ and $n = 2$. The relatively small values of $T_c$ found for $n = 1$ in this calculation may be understood from the density of states, which shows that the low-lying states lie mainly on the non – Fe sites, which are far displaced in energy from the magnetic site and therefore do not hybridize strongly with it, so the effective carrier-spin interaction is not strong. That the $n = 2$ is non-magnetic may be understood by combining the results of with the observation that the band structure is effectively two dimensional. In the extreme weak coupling limit, the nature of the magnetic ground state is determined by the wave vector at which the susceptibility is maximal. For the two dimensional band structures considered here this maximum is not at $q = 0$. Ref. showed that in the DMFT approximation, increasing the carrier-spin coupling increased the range in which ferromagnetism existed, but that as band filling is increased, a transition to an antiferromagnetic state generically occurs, and gets pushed to the half-filled band only for $J$ of the order of the bandwidth. These effects are more pronounced for the two dimensional band structure we consider.

The relative weakness of the virtual Re/Mo $\leftrightarrow$ Fe transitions is reflected in the temperature dependence of the many-body density of states, shown for the 'band' parameters in the lower panel of Fig. 1. Comparing these we see that disordering the Fe spins leads to a slight narrowing of the bands, but the larger (30%) band narrowing effects found in CMR manganese are not observed for these parameters.

FIG. 3: Left panel: Contour plots of calculated transition temperatures for range of model parameters and conduction band density $n = 1$. Right panel: Contour plots of calculated transition temperature for $n = 2$. In each figure, contours are spaced approximately 100 Kelvin apart and the white areas correspond to transition temperatures in excess of 400K.

To understand the behavior of the model in more detail we have evaluated the predicted ferromagnetic transition temperatures for wide range of model parameters. The resulting are summarized in the two panels of Fig. 3 which
show via contour plots the values of $T_c$ predicted by the method. The contours are spaced approximately 100K apart, and the black regions indicate the areas in which the calculated $T_c$ vanishes. It is seen that in order to obtain a reasonably high transition temperature, especially for the $n = 2$ band filling, one must choose the parameter $t_2$ to have the opposite sign from the physical one. The reason for this behavior is reveal by Fig. 2, which shows the density of states for parameters which maximize the $n = 2$ $T_c$. The low-lying states for this case are seen to be of mainly Fe character, because the level repulsion argument which pushed down the non-Fe states for the LSDA parameters is not operative here.

![Fig. 4: $T = 0$ (dashed) and $T > T_c$ conductivities for $n = 1$ using best-fit band parameters $t_1 = 0.25 eV$, $t_2 = 0.15 eV$, $t_3 = 0.03 eV$ and $\Delta = 0.3 eV$ used in Fig. 1. The $T = 0$ conductivity was computed using an artificial broadening of 0.1 eV applied to the Fe site.](image1)

We have also calculated the optical conductivity for various model parameters. Results obtained using the LSDA parameters are shown in Figs. 4 (Re-case, $n = 1$) and Fig. 5 (Mo-case, $n = 2$). One would in principle expect two classes of transitions: a 'Drude' peak centered at $\omega = 0$ involving motion of electrons near the Fermi surface and an interband transition involving moving an electron from a Re/Mo to a Fe. Our calculations indicate that for the tight binding parameters corresponding to the LSDA calculation, the interband feature is very weak, indeed not visible in the Figure again demonstrating the weakness of the Fe–Mo coupling for these parameters. We observe that the 'Drude' part has a distinctly non-Drude form, which arises because in our calculation the scattering processes couple to the 'B' (Fe) site only; although the regions of momentum space where the hybridization vanishes are of measure zero, they do lead to a frequency dependence of the scattering rate which explains the peculiar form. We also note that the main cause of the changes in conductivity and oscillator strength between $T = 0$ and $T = T_c$ is the change in band filling, which leads to a change in optical matrix element. In the paramagnetic state one has three bands, each with a two-fold spin degeneracy, corresponding to a filling of $n/6$ ($n=1,2$ is the particle density), so the relevant states are quite close to the bottom of the band where the optical matrix element is small. In the ferrimagnetic state for $n = 1$ one loses the spin degeneracy, so one has three bands each filled to a higher level, so with a correspondingly higher fermi velocity and optical matrix element, whereas for the $n = 2$ case the temperature induced shift corresponds to a change from 1/3 to 2/3 filled band, with much smaller change in optical matrix elements. The temperature dependent change in the oscillator strength is therefore much less. As noted above, in the ferrimagnetic case for $n = 2$ one has a small filling of the majority spin band, leading to a small additional contribution to $\sigma$, shown as the dot-dashed line in Fig. 5.

The conductivity corresponding to the parameters which maximize $T_c$ (Fig 2) is shown in Fig. 6. We see that the different electronic structure leads to a different optical conductivity: the Drude absorption is weaker, and a peak corresponding to excitation of carriers from Fe to Re/Mo is evident.

![Fig. 5: Upper panel: $T = 0$ (dashed) and $T > T_c$ (solid) conductivities for $n = 2$, using best-fit band parameters $t_1 = 0.25 eV$, $t_2 = 0.15 eV$, $t_3 = 0.03 eV$ and $\Delta = 0.3 eV$ used in Fig. 1. The $T = 0$ conductivity was computed using an artificial broadening of 0.1 eV applied to the Fe site. The small contribution to the $T = 0$ conductivity arising from the minority spin band is shown as the dash-dot line.](image2)

The conductivity of $Sr_2FeMoO_6$ has been measured by Jung and co-workers. These authors found a conductivity which was of roughly the Drude form, (albeit with a rather larger scattering rate than we have used) but additionally has a weak kink at a frequency of approximately 0.6 eV. It is interesting to speculate that this kink is a signature of the 'interband' feature which we found only for the 'antiphysical parameters. A more detailed experimental investigation of the band structure...
may be warranted.

V. CONCLUSIONS

We have used the dynamical mean field method to determine the ferromagnetic transition temperature, density of states and optical conductivity of a model representing key physics (two dimensionality of band structure and strong on-site interaction on Fe site) of the double perovskite ferrimagnets Sr$_2$Fe(Mo/Re)O$_6$. Our method can easily be generalized to include the effects of mis-site disorder, or lattice distortions which split the $t_{2g}$ levels. However, such generalization is not immediately warranted because the calculated transition temperatures are, at least for the parameters following from band structure calculations, in qualitative disagreement with experimental data—in particular, the calculation predicts that Sr$_2$FeMoO$_6$ is not ferrimagnetic, whereas experiment indicates that it is with a $T_c$ excess of 400K, and underpredicts the $T_c$ of Sr$_2$FeReO$_6$ by a factor of almost 4.

The essential reason for this was found to be that the band theory parameters imply that the mobile carriers reside mainly on the non-Fe sites, and hybridize weakly with these sites. Optical conductivity signatures of the weak hybridization were demonstrated. The calculation indicates that transition temperatures would be substantially raised if parameters are used for which the added carriers are largely on the Fe sites. An alternative possibility is that an interaction omitted from the model is crucially important; in particular that the magnetism should not be regarded as arising from correlations on the Fe site, but should instead be thought of more as a Stoner instability of the band arising from the Re/Mo states. The additional interaction proposed in Ref. 3 would tend to produce this physics and an important next step would be to extend the methods developed here to the treatment of this case.

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