Signature of antiferromagnetic long-range order in the optical spectrum of strongly correlated electron systems

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We show how the onset of a non-Slater antiferromagnetic ordering in a correlated material can be detected by optical spectroscopy. Using dynamical mean-field theory we identify two distinctive features: The antiferromagnetic ordering is associated with an enhanced spectral weight above the optical gap, and well-separated spin-polaron peaks emerge in the optical spectrum. Both features are indeed observed in LaSrMnO 4 [A. Gössling et al., Phys. Rev. B 77, 035109 (2008)].

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I. INTRODUCTION

This paper deals with the following issue: Is there a distinctive signature of the onset of antiferromagnetic (AF) long-range order in the optical spectrum of a strongly correlated insulating antiferromagnet, for example an antiferromagnetic transition-metal oxide?

AF ordering is a characteristic feature of strongly correlated Mott insulators where superexchange drives the ordering of localized magnetic moments. On the other hand, AF can also arise from Fermi-surface nesting in a weakly correlated material. From a theoretical point of view, a continuous crossover driven by the interaction strength connects weak- and strong-coupling antiferromagnets, and an unambiguous distinction between local-moment and Fermi-surface AF is lacking.

In this work we propose that, remarkably, optical spectroscopy can be used to infer the correlated nature of an AF state. Optical spectroscopy is an invaluable experimental probe of correlated materials, providing key physical information such as the optical gap, the relative weight of low-energy Drude excitations in metallic systems, and quite importantly the transfers of spectral weights often observed in correlated materials when temperature or composition is varied (for a recent review see Ref. 1). Spin degrees of freedom are however much less coupled to light than the charge so that the signatures of magnetism are expected to be comparatively weak. Zone-center magnons, for example, do not show up in the optical conductivity unless inversion symmetry is broken. 1 Furthermore, the key energy scale associated with magnetic ordering, the superexchange J (~D 2 /U with D being half the bandwidth), is much smaller than the scale U corresponding to the local matrix element of the screened Coulomb interaction, and hence generally significantly smaller than the optical gap itself (~U – 2D).

Despite these shortcomings, we show that clear signatures associated with antiferromagnetism are expected in the optical conductivity below the Néel temperature (T N ) when the system is in the strong-coupling (Mott insulating) regime. These signatures are twofold. First, there is a spectral weight transfer as one cools the system from above the Néel temperature down to low temperature. This transfer actually provides a diagnostics of the strong-coupling (superexchange) or weak-coupling (spin-density wave) nature of the antiferromagnetism. In the former case, spectral weight increase is observed, corresponding to a kinetic energy gain associated with the ordering. In the latter case, a spectral weight decrease takes place, corresponding to a kinetic energy loss (potential energy gain). For superconducting long-range order, similar diagnostics of weak vs strong-coupling pairing have been discussed and probed experimentally in the context of cuprate superconductors 2 and for models with attractive interaction. 3,4

Second, a multipeak structure develops above the gap in the ordered phase at strong coupling. These peaks correspond to the spin-polaron excitations associated with the motion of a hole in an antiferromagnetic background, a problem which has been addressed since the late sixties in a series of pivotal works. 5 In the context of infinite dimensions the physics of the spin polarons has also been extensively analyzed 6–9 and this work identifies their signature in the optical conductivity.

Finally, we consider the possibility of detecting these signatures experimentally and discuss in detail the case of LaSrMnO 4, where we argue that some of these effects may already have been observed. 10

The outline of the paper is as follows: In Sec. II we summarize the main aspects of our calculation of the optical conductivity for the Hubbard and t-J model. Details on the dynamical mean-field theory calculations for the two models are provided in Appendix 1 and 2, respectively. In Sec. III we discuss the main theoretical results before we compare them to experiment in Sec. IV. A summary and conclusion are given in Sec. V.

II. MODELS AND METHODS

In order to study the optical conductivity of strongly correlated antiferromagnets, we perform dynamical mean-field theory (DMFT) calculations for a half-filled Hubbard
model, described by the following Hamiltonian:

\[ H = -t \sum_{\langle ij \rangle \sigma} c_{i\sigma}^\dagger c_{j\sigma} + U \sum_i n_{i\uparrow} n_{i\downarrow}. \]  

(1)

Here, \( t \) denotes the hopping amplitude between nearest neighbors, \( U \) the Coulomb interaction, and \( c_{i\sigma}^\dagger \) (\( c_{i\sigma} \)) creates (annihilates) an electron with spin \( \sigma \) on site \( i \): \( n_{i\sigma} = c_{i\sigma}^\dagger c_{i\sigma} \). For simplicity, we use a semielliptical density of states of half width \( D \), corresponding to a Bethe lattice with infinite coordination. Our results however hardly depend on this specific choice (see also Appendix 1). We vary the temperature \( T \) from above to below the Néel temperature. We therefore need the paramagnetically and the antiferromagnetically ordered solutions, which can both be calculated within DMFT.\(^{11,12}\)

It is known that the Hubbard model for large values of the local repulsion \( U \) maps onto the \( t-J \) model. For the specific model under consideration, the relation between \( J \) and \( U \) is the following: For an individual pair of sites we have \( J_{\text{ind}} = 4t^2/U \) and summed over all nearest neighbors \( z \) we have \( J = z J_{\text{ind}} = D^z/U \) (coined \( J^* \) in Ref. 6). The quantities of interest in the \( t-J \) model in infinite dimensions can be expressed in the form of a simple continued fraction\(^{6,13}\) while the Hubbard model requires a more computationally expensive solver. Therefore we will often use the \( t-J \) solutions in the case of the antiferromagnetic phase. Indeed, the electron-removal part of the \( k \)-integrated spectral function of a half-filled Hubbard model below \( T_N \) coincides up to higher order terms in \( J \) with the spectral function of a single hole in the \( t-J \) model. The way we compute the optical conductivity for the \( t-J \) model, and in particular how we get across the absence of an electron addition part of the spectrum of the \( t-J \) model, is explained in detail in the Appendix.

Within DMFT the optical conductivity can be expressed exactly in terms of the convolution of two single-particle Green’s functions without vertex corrections.\(^{12,14,15}\) In the antiferromagnetic phase this reads\(^{16}\)

\[
\text{Re} \sigma(\Omega) = \frac{e^2 \hbar}{V \pi} \sum_{\sigma} \int d\omega \int_0^D d\epsilon \frac{f(\omega) - f(\omega + \Omega)}{\Omega} \frac{2}{\pi D^2} \times \left[ (D^2 - \epsilon)^{3/2} \text{Im} \epsilon_{\sigma}^{AA} (\epsilon, \omega) \text{Im} G_{\sigma}^{AA} (\epsilon, \omega + \Omega) + \text{Im} G_{\sigma}^{AB} (\epsilon, \omega) \text{Im} G_{\sigma}^{AB} (\epsilon, \omega + \Omega) \right],
\]

(2)

where \( f \) denotes the Fermi distribution function, \( G_{\sigma}^{AA} \) and \( G_{\sigma}^{AB} \) are respectively the Green’s function of an electron with spin \( \sigma \), \( \bar{\sigma} = -\sigma \) hopping between sites of the same sublattice and between sites of the two different sublattices A and B, in which the Bethe lattice is split in the AF phase.

In this expression, the quantity \( \frac{1}{\pi D^2} (D^2 - \epsilon)^{3/2} \) represents the product of the density of states of the Bethe lattice times the electric current vertex. This assumption insures that the \( f \)-sum rule associated with the optical conductivity holds, namely that its integral is proportional to the kinetic energy.\(^{14,17-20}\)

III. THEORETICAL RESULTS

In Fig. 1 we show our main result, namely the temperature dependence of the optical conductivity as one cools the system from close to the Néel temperature down to low temperature, in a regime of strong correlations, i.e., \( U/D = 5 \). For temperatures slightly lower than \( T_N \), the optical conductivity resembles that of the paramagnetic insulator. It has a gap of order \( U - 2D \), and the absorption peak above the gap is essentially featureless. Upon cooling, two remarkable features characterize the optical signal: (i) the optical absorption strongly increases and (ii) a multipeak structure emerges. Such a structure becomes more and more visible as one goes down in temperature. Note that these changes occur without a visible variation in the position of the absorption edge (size of the gap). As discussed below, they should nevertheless be detectable by accurate experiments as a function of temperature.

Later in this section we will explain the physical origin of the peaks in the optical conductivity by investigating the single-particle spectral function and its evolution with \( U \). At this stage, let us only stress that these peaks are by no means an artifact of the discretization required by our exact-diagonalization DMFT solver.\(^{9}\) To better compare with experiments, we used a Lorentzian broadening of 0.05\( D \) to plot the optical conductivity. This value is small enough to distinguish the first peaks but also large enough to mimic the smearing of the multipeak structure due to experimental resolution.

We now discuss in more detail the two key effects we mentioned before.

(i) Change of spectral weight through \( T_N \). The increase of spectral weight upon cooling is characteristic of a Mott antiferromagnet at strong coupling. Therefore it is the first gross feature one may look for in an optical experiment in order to establish whether the antiferromagnetism in the material is of strong-coupling nature, with preformed magnetic moments, or is of weak-coupling nature, with a collective Fermi-surface instability. To understand this, we can use the fact that the integral of the spectral weight is directly proportional to the kinetic energy. At strong coupling \( U \gg D \), the ordered phase is stabilized by a gain in kinetic energy, associated with the establishment of long-range coherence of the local magnetic moments. In simple words the system gains kinetic energy by increasing the staggered magnetization: Coherent hopping processes taking place in a staggered spin background are
favored if the spin pattern is as close as possible to the Néel one.\textsuperscript{3,9} Thus the lower the temperature, the larger the staggered magnetization and in turn the larger the spectral weight.

If we consider instead the weak-coupling regime $U \ll D$, a totally different mechanism for the stabilization of the ordered phase takes place. The AF phase is stabilized by a (small) potential energy gain, as the onset of a nonzero order parameter is correctly described by the static mean-field theory in this regime. These theoretical considerations are well confirmed by our numerical data shown in the bottom panel of Fig. 2, where we compare $K_{\text{AF}}$ and $K_{\text{PM}}$, the kinetic energy of the ordered and disordered phase, respectively. The complete evolution of the kinetic and potential energy as a function of $U$ is shown in Fig. 3. At strong coupling the antiferromagnetic order is clearly stabilized by the kinetic energy while at weak coupling ($U < 2D$) the ordered phase is driven by the gain in potential energy. At intermediate values of the interaction strength, corresponding to the crossover between weak and strong coupling, we have a mixed regime in which both energy differences are negative,\textsuperscript{3,4} i.e., where the onset of the AF is stabilized by both kinetic and potential energy.

(ii) Spin-polaron peaks. To clarify the origin of the multipeak structure, we compare in Fig. 2 the evolution from weak to strong coupling of the optical conductivity to that of the one-particle spectral function. It is evident that the multipeak structure of the spectral function\textsuperscript{5,5} survives in the optical conductivity, as already observed for the pure $t$-$J$ case by Cappelluti \textit{et al.}\textsuperscript{22} The peculiar behavior of the spectral function in infinite dimensions was first understood in the $t$-$J$ model in a pioneering work by Strack and Vollhardt\textsuperscript{6} and more recently reanalyzed for the case of the Hubbard model by some of us.\textsuperscript{9} The multiple peaks arise from stringlike excitations associated with a hole moving in a Néel background.\textsuperscript{5} In infinite dimensions the lifetime of such string excitations is infinite due to the absence of quantum spin-flip processes\textsuperscript{6,7,9} (which would act as “damage-repairing” processes), meaning that the Heisenberg interaction reduces to the Ising one (see Appendix 1 for further details). This results in the formation of coherent but heavy quasiparticles, “dressed” by a spinon cloud which is commonly called a “spin polaron.”

The formation of the spin polaron can be described only using methods (such as DMFT) that go beyond the static mean-field (Hartree-Fock) approximation of a spin-density wave. This is illustrated in the upper panels of Fig. 2, where we compare the evolution with $U$ of the dynamical (red line) and static mean-field (green line) results for the electron-removal spectrum. One clearly sees that the only feature of the static mean-field (Hartree-Fock) calculation is a Slater-like single peak centered around $-U/2$. The peak has weight $Z = 1$ and its width is given by $J$; i.e., the width shrinks to 0 as $1/U$ upon increasing $U$. In the dynamical mean-field calculation, instead, the weight of the lowest lying excitation is much smaller ($Z$ is proportional to $J/D$) and the remaining weight is transferred to the higher-energy peaks. This makes a total bandwidth of order $2D$ rather then $J$, resulting in a smaller gap compared to
the Hartree-Fock calculations. This is why the gap in Fig. 1 is of order $U - 2D$, while it would be exactly $U$ within static mean-field theory.

In the $t$-$J$ model in infinite dimensions (as well as in the Hubbard model within AF-DMFT, up to higher-order corrections in $t^2/U$) the position of the peaks can be analytically expressed through the zeros of the Airy functions and the spacing between them is proportional to $D^{1/3} J^{2/3}$. As we will also discuss in Sec. V these analytic expressions hold strictly only in infinite dimensions. Hence the applicability of our results to real materials, and in particular the emergence of spin-polaron peaks, strongly relies on the assumption that it is possible to neglect the quantum spin-flip processes. In principle this is not guaranteed for finite-dimensional systems; therefore one may expect some of the peaks to broaden or even to be washed out, due to the coupling to dispersive spin waves. On the other hand, recent sophisticated calculations for the two-dimensional $t$-$J$ model give clear evidence for the survival of the first one or two spin-polaron peaks with a similar separation as in the infinite-dimensional calculation. Furthermore the case of a double-layer antiferromagnet was recently considered in Ref. 24. There, it is shown that the interlayer excitation made of one hole and one doublon in each layer displays confinement effects when the interlayer coupling is smaller than the intralayer one. In such a situation, the spin-polaron excitations discussed here may become particularly relevant.

When considering three-dimensional cases, where DMFT is typically more accurate and finite-dimensionality effects are definitively weaker than in two dimensions, spin-polaron features are expected to be even more visible. In the following section, we discuss indeed the relevance of our results for the three-dimensional manganese LaSrMnO$_4$.

IV. COMPARISON WITH EXPERIMENTS

On the basis of the previous discussion, the ideal material for the observation of the temperature evolution predicted above would be a completely isotropic three-dimensional Mott antiferromagnet. To our knowledge the most relevant set of measurements in this respect is that by Gösslind et al., on LaSrMnO$_4$, a three-dimensional C-type layered antiferromagnetic material with $T_N = 133$ K, while we are not aware of similar studies as a function of temperature on G-type antiferromagnets, which would be the closest realization of our large-coordination results. In Fig. 4 we show the data of Ref. 10 for the optical conductivity of LaSrMnO$_4$ along the $a$ axis at different temperatures. The two main structures around 3.5 eV and 4.5 eV have been explained in terms of multiplet split transitions from a $d^4$ to a $d^5$ configuration of the Mn. Our main interest instead is focused on the fine structure arising upon cooling on top of both of these main structures.

In the light of our theoretical analysis, we can directly relate this observation with the physics of the spin polarons described in the previous section. Since the two structures are well separated, we shall make a simplifying assumption (not meant to be quantitative, but sufficient to reveal the main qualitative aspect of the phenomenon). Namely, we shall assume that we can adopt for each $d^4 \rightarrow d^5$ transition a simple single-band Hubbard model description.

Since the separation between the multiplet peaks is larger than the energy scale associated with the spin-polaron excitations, we expect to be able to detect the above-mentioned increase of spectral weight around each of the main absorption peaks and the consequent appearance of spin-polaron structures when going below the Néel temperature.

All curves below $T_N$ display, in the same region, a very similar low-frequency tail (below $\sim 3$ eV) while in that region the higher-temperature measurements show a gradual increase of spectral weight that we attribute to a more standard thermal broadening effect. It is likely that such thermal effects are visible also in the low-energy multiplet peak at 3.5 eV, which lies closer to this tail. This makes this spectral feature less visible with a first distinguishable excitation shifting with temperature unlike what happens for the second multiplet structure at 4.5 eV.

We focus therefore on the high-energy spectral structure around 4.5 eV, which seems to be unaffected by the temperature broadening of the low-frequency data, and we zoom the plot to focus on this feature. In order to highlight the temperature effect, we display in Fig. 5 the difference between the optical conductivity at a given temperature and the conductivity at the highest measured temperature (300 K).

Figure 5 and its theoretical counterpart (Fig. 6) endorse our physical interpretation of the peaks in the optical signal. Although the frequency grid in the experimental data is 0.05 eV, at least two well-spaced peak structures emerge below $T_N$ and, as expected within our spin-polaron picture, the lower the temperature the weaker the temperature dependence becomes. Indeed between 52 and 15 K there is hardly any difference in the optical conductivity in this frequency region, as one can expect from the behavior of the staggered magnetization (see inset to Fig. 1), which is almost saturated far below $T_N$. The appearance of the spin-polaron fine structures become clear below $T_N = 133$ K. Yet, the polaronic features start to be visible already in the 150 K curve. We attribute this to fluctuation effects above $T_N$: Even if long-range order is not yet fully developed, a sufficiently long coherence length of the spin-spin correlations is enough to sustain a non-fully-developed spin-polaron mechanism.
Features similar to the ones we have described are present in the optical absorption of another material: KCuF₃. For temperatures lower than the magnetic ordering temperature of this compound a multipeak structure is indeed observed [see Fig. 2(b) of Ref. 26], for which an interpretation in terms of spin waves was proposed.

We have therefore found in nature what we expect from our model calculation: a clear signature of antiferromagnetism arising from strong correlations in the optical conductivity of a strongly correlated material (LaMnSrO₄), which emerges as the system is cooled below $T_N$. If the material is sufficiently three-dimensional and if the resolution of the experiment is good enough, a clear signal with a multipeak structure can be observed. Another way of revealing the same effect would be to compare the increase of spectral weight to the measured staggered magnetization as a function of $T$.

In the strong-coupling regime, the sharp peaks associated with spin polarons originate from stringlike excitation created by the motion of a hole in an antiferromagnetic background, and we can safely distinguish them from artificially sharp peaks characteristic of exact-diagonalization solutions of DMFT.

Even if our formal derivation of the spin-polaron peak relies on the infinite-coordination limit in which DMFT becomes exact, we expect at least part of these features to survive in finite dimensions, and in particular in the three-dimensional case, in which DMFT typically proves more accurate. We notice that at least the lowest-energy excitation peak is found also in two dimensions, as shown by different approaches, testifying that the physics behind our DMFT results survives when the large-dimensionality limit is far.

If we look for experimental realizations of this scenario, we should take into account temperature effects and the finite experimental resolution, which can further smear out some of the fine structures. In this light, it is remarkable that an experimental study on LaMnSrO₄ reveals a multipeak structure whose form and temperature behavior are in very good agreement with our calculations on top of the multiplet structure.

Moreover, considering the recent work by Rademaker et al., optical spectroscopy performed along the $c$ axis in layered systems may be a promising tool to detect and investigate the occurrence of the spin-polaron physics.

Eventually, even stronger signatures of the spin-polaron effects are to be expected in three-dimensional correlated materials which present a three-dimensional $G$-type antiferromagnetism, such as SrMnO₃, CaMnO₃, BaMnO₃, calling for future ir-spectroscopy investigations of the PM-AF transitions in these compounds.

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APPENDIX

In this Appendix we give all the details on how to construct an approximated expression for the DMFT self-energy of the half-filled Hubbard model on the basis of the DMFT solution for the corresponding t-J model both in the antiferromagnetic and in the paramagnetic phase. We also discuss the validity of this procedure by exploiting the mapping of the (half-filled) Hubbard Hamiltonian onto the t-J one\(^{28}\) for \(U \gg D\) and by comparing our results for the self-energy and the optical spectra extracted from the DMFT solution of the t-J model with the corresponding ones for the Hubbard model.

1. DMFT for the t-J model

The main motivation to use the t-J results for approximating the DMFT self-energy stems, as mentioned in Sec. II, from the simplicity of the algorithm for solving the DMFT problem of one hole in the t-J model with (complete or partial) AF-Néel order\(^{48}\) as opposed to the bigger numerical effort needed for the Hubbard model. Specifically for one hole at \(T = 0\) one finds (for more details see Ref. 6) that for \(d \to \infty\)

1. the quantum spin-fluctuations associated with the term proportional to \(S_i^+ S_j^- + S_j^+ S_i^-\) of the SU(2)-symmetric Heisenberg interaction are completely frozen. In other words, the Heisenberg coupling of the t-J Hamiltonian is reduced to an Ising coupling, whose ground state is the AF-Néel state with a perfectly staggered spin configuration, i.e., with full staggered magnetization \([m = \frac{1}{N} \sum (n_i^+ - n_i^-)(-1)^i = 1,\text{ where } N\text{ represents the total number of sites}]\);

2. the "retraceable path" approximation becomes rigorously exact: In a perfect AF-Néel state, the local physics [e.g., the local Green’s function \(G(\omega)\)] is built only by hopping processes in which the one hole comes back to the starting site following precisely the same path. In fact, any non-retraceable path is associated with a higher-energy cost, as the hole movement would result in a path of misaligned spins in the AF-Néel background. This implies that the geometry of the underlying lattice becomes irrelevant in \(d \to \infty\): If only retraceable paths matter, every geometry becomes topologically equivalent to the Bethe lattice, a tree-structure which by construction only allows for retraceable paths.

As a consequence at \(T = 0\) the discrete energy levels characterizing the analysis of the one-hole dynamics in the AF-Néel background of the t-J model can be related to the simple quantum mechanical problem of one particle in a step potential \(V(n) = nJ\), being \(n\) the number of misaligned spin created by the hopping processes of the hole. This allows us in turn to obtain a compact continued-fraction expression for the retarded Green’s function \(G_{t-J}(\omega) = G_{t-J}(\omega + i0^+)\) (see Ref. 6)

\[
G_{t-J}(\omega) = \left[\omega - \frac{D^2}{4} G_{t-J} \left(\omega - \frac{1}{2} J\right)\right]^{-1}
\]

\[= \frac{1}{\omega - \frac{D^2}{4} \left[\frac{1}{\omega + \frac{1}{2} J - \frac{i\omega}{2}} + \frac{1}{\omega + \frac{1}{2} J - \frac{i\omega}{2}}\right]}, \tag{A1}
\]

which can be solved with high numerical accuracy by simply truncating the continued fraction to the desired level of precision, and then building up the full expression iteratively. Complications arise at finite \(T\): In particular point 2 is in principle no longer valid, as thermal fluctuations will reduce the value of the staggered magnetization, allowing for a contribution of non-retraceable paths to the local physics.\(^{7}\)

However, we can easily circumvent this problem, if we consider explicitly the Bethe lattice case for our DMFT calculations. This is, on the one hand, a reasonable choice, as the finite bandwidth of the Bethe lattice reflects more directly the realistic situation of finite bandwidths in real materials, and, on the other hand, it preserves by construction the validity of the “retraceable path” condition at all temperatures. Working with the Bethe lattice allows, in turn, for a straightforward generalization of the continued-fraction expression for \(G(\omega)\) to the finite-temperature case, along the lines of the exact derivation of Ref. 13. Specifically, one can easily generalize Eq. (A1) to finite temperatures by taking into account the probability \(P\) (or \(1 - P\)) of finding correctly aligned (or misaligned) spins along the retraceable path of the hole.

The probability \(P\) as a function of \(T\) is easily expressed as \(P = \frac{1}{2} [1 + m(T)] \) in terms of the staggered magnetization \(m(T)\). The latter, because of point 1 above, can be directly obtained by the simple mean-field Curie-Weiss self-consistent equation \(m(T) = \tanh(T S / 2 m)\), with \(T_s = J / 2\). The value of \(P\), when smaller than 1, will affect the self-consistent expression for the retarded Green’s function: \(^{13}\)

\[
G_{t-J}(\omega) = \left[\omega - \frac{D^2}{4} \times [P G_{t-J}(\omega - \omega_P) \right. \\
\left. + (1 - P) G_{t-J}(\omega + \omega_P))]^{-1}, \tag{A3}
\]

being \(\omega_P = \omega_P(T) = \frac{1}{2} m(T) J\), and \(\omega = \omega + i0^+\). Note that this expression is exactly reduced to Eq. (A1) in the limit \(T \to 0\), as \(m(T = 0) = P(T = 0) = 1\). The additional terms in Eq. (A3) at finite \(T\) result in an increasing proliferation of new (smaller) peaks no longer exactly matching with the original multipake structure of the continued-fraction expression at \(T = 0\). As a consequence, by increasing \(T\), one observes a gradual smearing out of the \(\delta\)-like peak structure of Eq. (A1), which is completely washed out at \(T \geq T_s\), when one has \(m = 0, P = \frac{1}{2}\) and the spectral representation \(G(\omega)\) is reduced to the characteristic semicircular shape of the noninteracting Bethe-lattice case. This result can be easily understood, because the Heisenberg interaction term, which is nonlocal in the site indices \(i, j\), becomes irrelevant due to the \(d \to \infty\) scaling in the paramagnetic phase (i.e., for \(m = 0\) and, therefore, for \(T \geq T_s\) one gets the same spectral function of the case \(J = 0\), independently from the actual value of \(J\).

2. Approximation for the DMFT self-energy of the Hubbard model

After having recalled how to solve exactly the DMFT problem of one hole in the t-J model, we will discuss here how to approximate the DMFT self-energy for the Hubbard model in the limit of intermediate-to-strong coupling by exploiting this solution. In fact, it is known\(^{28}\) that in the limit \(U \gg D\)
the Hubbard model can be mapped onto a corresponding $t$-$J$
model with $J = \frac{U}{2}$. When using the $t$-$J$ results to approximate
the Hubbard DMFT self-energy or the $k$-integrated spectral
and Green functions in the limit of $U \gg D$, one should not
forget that the exact “mapping” between the two models is
obtained by projecting out the doubly occupied sites. This
evidently means that the $t$-$J$ model cannot describe the whole
frequency range of the Hubbard model, even in the limit
of $U \gg D$, but, at most, only “half” of it. This becomes
quite obvious, when explicitly considering the $k$-integrated
spectral functions: The gap between the Hubbard bands is
originated by the energy difference ($\sim U$) between single
occupied and double occupied/empty sites, but this energy
difference is exactly what is projected out in the mapping
onto the $t$-$J$. Therefore, when computing the DMFT Green’s
function for one hole in the $t$-$J$ model, the associated spectral
function will be related to electron-removal ($\omega < 0$, i.e.,
photoemission) part of the $k$-integrated spectral function of
the Hubbard model: More precisely, taking into account the above-
mentioned energy shift of order $U$ (not explicitly included in
the $t$-$J$ Hamiltonian), it will correspond to the lower Hubbard
band.

These considerations are easily translated in a simple
practical implementation: In fact, considering the symmetry
with respect to the change $\omega \leftrightarrow -\omega$ for both the paramagnetic
and the antiferromagnetic case, one can obtain the $t$-$J$-like
approximation of the $k$-integrated retarded Hubbard Green’s
function $G_{\sigma}(\omega + i0^{+})$ on a given sublattice (for the sake of
definiteness we identify with $\sigma = \uparrow$ the majority spin
component on the AF order on the chosen sublattice) directly
from the retarded Green’s function (or the spectrum) $G_{\sigma,J}(\omega)$
($A_{\sigma,J}(\omega)$) of the one-hole $t$-$J$ problem as

$$G_{\uparrow}(\omega) = PG_{\uparrow,J}\left(\omega + \frac{U}{2}\right) + (1-P)G_{\downarrow,J}\left(-\omega + \frac{U}{2}\right),$$

(A4)

$$G_{\downarrow}(\omega) = (1-P)G_{\uparrow,J}\left(\omega + \frac{U}{2}\right) + PG_{\downarrow,J}\left(-\omega + \frac{U}{2}\right).$$

(A5)

The essential ingredients of Eqs. (A4) and (A5) are, hence, two:
(i) the spectral (Green’s) function of the one-hole problem of
the $t$-$J$ model, and (ii) the probability $P$ to find (on the chosen
site) the spin correctly aligned with the AF-Néel background.
A word of caution is due at this point: As the value of $P$
is defined in terms of the staggered magnetization $m(T)$, the latter
naturally represents the most appropriate “link” between
the DMFT $t$-$J$ and Hubbard Green’s functions. In fact, in using
the $t$-$J$ approximation for the DMFT spectra of Hubbard model,
one should keep in mind that the mapping is rigorously exact
only in the limit of $U \rightarrow \infty$; otherwise, corrections $\sim \frac{D}{U}$
have to be expected: From a more physical perspective, the presence
of doubly occupied/empty sites in the ground state of the
Hubbard model stays indeed finite (though becoming smaller
and smaller for increasing $U$) at any value of the Hubbard
interaction. As a consequence, for any given set of parameters
$U$ and $T$, the staggered magnetization of the Hubbard model
will be slightly lower than that for the $t$-$J$ model, as it is also
found in our DMFT data, plotted in Fig. 7.

It should be clear, hence, that the best agreement between
the Hubbard model (for a given $U$) and the corresponding $t$-$J$
(with $J = \frac{U}{2}$) approximation is found for sets of Hubbard and
$t$-$J$ Green’s or spectral functions with the same magnetization
values (which are achieved, according to Fig. 7, for slightly
different temperatures).

Having defined the equations (and the conditions) for best
approximating the Hubbard DMFT Green’s functions with
the $t$-$J$ model ones, it is straightforward to extract from these
the correspondent DMFT expressions for the (spin-dependent)
Hubbard self-energies $\Sigma_{\sigma}$, via the DMFT self-consistency
equations (generally written to include the AF order):^{12}

$$\Sigma_{\uparrow}(\omega) = \omega + \mu - \frac{D}{4} [G_{\uparrow}(\omega)]^{-1},$$

(A6)

$$\Sigma_{\downarrow}(\omega) = \omega + \mu - \frac{D}{4} [G_{\downarrow}(\omega)]^{-1},$$

(A7)

where the chemical potential $\mu$ is set fixed to $\frac{U}{2}$ (particle-
hole symmetric case). Representative results of our DMFT
$t$-$J$ approximation for the imaginary part of our Hubbard
self-energy $\text{Im} \Sigma_{\sigma}(\omega)$, computed according to Eqs. (A4)–(A7),
are reported in Fig. 8. The main feature of the $T$ dependence
of $\text{Im} \Sigma_{\downarrow}(\omega)$ well agree with the corresponding DMFT data
for the Hubbard model shown in Ref. 9, supporting the
validity of the $t$-$J$ approximation. In particular, if $U \gg D$,
both in Hubbard as well as in its $t$-$J$ approximation, one
finds that for $T \ll T_{N}$ (upper panel of Fig. 8) $\text{Im} \Sigma_{\downarrow}(\omega)$
is characterized by a multipeak structure for $\omega < 0$ which
reflects that of the spin-up spectral function, and, hence,
the above-mentioned spin-polaron physics. Remarkably the
“magnitude” of $\text{Im} \Sigma(\omega)$, e.g., if quantified by the value of its
frequency integral, is not particularly large for low $T$, implying
a certain degree of “coherence” of the spin-polaron
excitations in this regime.^{30} Note that except for the case of
full polarization, a much weaker multipeak structure is also
atomic limit \((D = 0)\), for which one can easily compute the Green’s function,\(^{12}\) e.g., for the spin \(\uparrow\) sector

\[
G_\uparrow(\omega) = \frac{1}{\omega + i0^+ + \frac{U}{2} + \frac{1}{\pi i} \ln \left( \frac{\omega + i0^+ - \frac{U}{2}}{\omega + i0^+ + \frac{U}{2}} \right)},
\]

(A8)

and, from this [and the corresponding \(G_\downarrow(\omega) = G_\uparrow(-\omega)\)], calculate the self-energy

\[
\Sigma(\omega) = U \left[ \frac{\omega - m\omega}{\omega + i0^+ + mU} \right],
\]

(A9)

which evidently displays a peak in its imaginary part located at \(\omega_{\text{peak}} = m\omega\) (i.e., a “softening” for increasing \(T\) or decreasing \(m\)) of weight \(w_{\text{peak}} = \pi \frac{U^2}{4} [1 - m^2]\) (i.e., of increasing strength for increasing \(T\) or decreasing \(m\)). At \(T = T_N\), one gets eventually \(\Sigma(\omega) = \Sigma_1(\omega) = \frac{U^2}{4\pi\omega+i0^+}\), whose divergence in \(\omega = 0\) directly corresponds to the spectral gap of (exact) size \(U\). In comparison with the atomic limit, however, beyond this huge central peak in \(\text{Im}\Sigma(\omega)\), one observes the formation of two almost semicircular structures at \(\omega = \pm \frac{U}{2}\) of width \(\sim D\), which can be interpreted as an hallmark of the incoherent nature of the electronic excitation of the Hubbard band in the paramagnetic phase.

3. Accuracy of the approximation

After having discussed the results obtained by approximating the Hubbard DMFT self-energy starting from the \(t-J\) one, and having noted that for \(U \gg D\) its structure is coincident with that of the exact Hubbard model within small corrections (roughly of order \(D^3/T^3\)), here we will explicitly compare DMFT data for the optical conductivity in order to show the accuracy of our \(t-J\) approximation for the quantity we are more interested in this paper.

We report in Fig. 9 our numerical data for the optical conductivity of the AF-phase, namely for \(U = 5D\) at \(T = 0\) calculated inserting the DMFT Hubbard self-energy and the correspondent \(t-J\) approximated one in Eq. (2). At \(T = 0\), in fact, the possibility to use the Lanczos as an impurity solver visible for \(\omega > 0\), roughly speaking where the upper Hubbard would be located in the PM phase.

By increasing \(T\), one observes a rapid strengthening of such “secondary” multiphase structure at \(\omega > 0\), which becomes quickly predominant (second and third panel of Fig. 8). More specifically, one notes that the lower energy peak of the “secondary” (\(\omega > 0\)) structure of \(\text{Im}\Sigma(\omega)\) is strongly enhanced with increasing \(T\), and at the same time undergoes a constant “softening,” as its position (inside the spectral gap) tends to go to zero frequency for \(T \to T_N\) (lower panel of Fig. 8). In the limit of \(T = T_N\), when \(\Sigma(\omega) = \Sigma_1(\omega)\), this (now extremely strong peak) would be located exactly in the center (that is at \(\omega = 0\)), ensuring the existence of the Mott-Hubbard gap \(\sim U - 2D\) also in the absence of any long-range magnetic order. This can be compared with the exact solution of the

FIG. 8. (Color online) Temperature evolution of the DMFT self-energy of the Hubbard model, calculated from the DMFT Green’s function of the \(t-J\) model, according to Eqs. (A5) and (A7) for the specific case \(U = 5D\) \((J = 0.2D)\). Shown is the spin-up self-energy on the sites with spin-up majority; the spin-down self-energy is the particle-hole counterpart (obtained by mirroring \(\omega \to -\omega\)). As for the requirement of an equal staggered magnetization between the Hubbard and the \(t-J\) model, the corresponding temperature of the \(t-J\) case is reported in the legend as \(T_{t-J}\). The dotted vertical lines are located at \(\omega = \pm \frac{U}{2} m\) and \(\omega = -\frac{U}{2} m\), which in a static mean-field context would represent the upper and lower limits of the mean-field spectral gap.

FIG. 9. (Color online) Comparison between the DMFT optical conductivity of Hubbard model (impurity solver: Lanczos with \(N_s = 25\)) and corresponding \(t-J\) model.
allows for a better resolution of the multiple-peak structure of the Hubbard self-energy and, therefore, for a better test case to evaluate the accuracy of the $t$-$J$ approximation for the optical conductivity of the Hubbard model. In fact, while Lanczos already allows a larger number of bath sites than the full exact diagonalization, in the AF case for conductivity of the Hubbard model.

In the AF phase we can reach with a reasonable computational effort a number of electrons as high as $N_s = 25$, while in the PM case we could reach only $N_s = 14$.

Going back to the analysis of Fig. 7, our data show that, at $U = 5D$, the overall agreement between the peaky structure of the optical conductivity of the $t$-$J$ approximation and that of the Hubbard model is rather good, as both the peak positions and intensity are well captured also in the $t$-$J$ scheme. This result justifies, for these values of $U$, the usage of the DMFT solution of the one-hole problem in the $t$-$J$ model as an accurate approximation, e.g., also when considering the finite-$T$ regime. In fact, when increasing $T$, Lanczos or full exact-diagonalization solution of DMFT is no longer accurate enough for the computation of multipeak spectral functions with sufficient resolution.
Note that within the $t$-$J$ model the temperature dependence of the optical conductivity above $T_N$ is just the one coming from the Fermi functions in Eq. (2). Within the Hubbard model we would instead get some temperature dependence, however only appreciable on a rather large energy scale.


For instance, at $U = 5D$, the maximal magnetization (at $T = 0$) for the Hubbard model is $m = 0.979$, which correspond to the $t$-$J$ solution at $\beta D = 46.6$. This slight difference in temperature is naturally reduced with increasing $U$.

In other words, this means that if the chemical potential were rigidly shifted to the value of the first (low-energy) AF magnetic peak, this would behave as a coherent quasiparticle excitation (though strongly renormalized by $Z \sim e^D$). A similar situation would occur, mutatis mutandis, in the case such an excitation were activated by an external perturbation of frequency $\omega$ of the size of the spectral gap, e.g., by an electromagnetic field, when measuring the optical conductivity $\sigma(\omega)$. 

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