Local moments versus itinerant antiferromagnetism: magnetic phase diagram and spectral properties of the anisotropic square lattice Hubbard model

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Using a cluster extension of the dynamical mean-field theory (CDMFT) we map out the magnetic phase diagram of the anisotropic square lattice Hubbard model with nearest-neighbor intrachain $t$ and interchain $t_\perp$ hopping amplitudes at half-filling. A fixed value of the next-nearest-neighbor hopping $t'=-t_\perp/2$ removes the nesting property of the Fermi surface and stabilizes a paramagnetic metal phase in the weak-coupling regime. In the isotropic and moderately anisotropic regions, a growing spin entropy in the metal phase is quenched out at a critical interaction strength by the onset of long-range antiferromagnetic (AF) order of preformed local moments. It gives rise to a first-order metal-insulator transition transition consistent with the Mott-Heisenberg picture. In contrast, a strongly anisotropic regime $t_\perp/t\lesssim 0.3$ displays a quantum critical behavior related to the continuous transition between an AF metal phase and the AF insulator. Hence, within the present framework of CDMFT, the opening of the charge gap is magnetically driven as advocated in the Slater picture. We also discuss how the lattice-anisotropy-induced evolution of the electronic structure on a metallic side of the phase diagram is tied to the emergence of quantum criticality.

I. INTRODUCTION

The Hubbard model at half-filling and finite doping has been a subject of numerous theoretical studies driven by its ability to account for a variety of collective behaviors in strongly correlated quantum systems such as a metal-insulator transition (MIT) [1, 2], antiferromagnetism with its precursors and competitors [3], dimensional crossover from the one-dimensional (1D) Tomonaga-Luttinger liquid physics to a higher-dimensional situation [4], pseudogap behavior in the single-particle spectral function [5], and high-$T_C$ superconductivity [6]. Even though the MIT is simply a consequence of the energy competition between Coulomb interactions which tend to localize electrons and the kinetic energy term which favors electron itineracy, its actual form is situation dependent with microscopic details of the electronic structure playing a leading role [7]. Accordingly, different mechanisms of the MIT have been proposed to explain the formation of an insulating phase.

In the Mott-Hubbard picture of a correlation-driven MIT, a strong on-site Coulomb repulsion $U$ splits the half-filled conduction band to open a gap in the electronic excitations between the resultant lower and upper Hubbard bands [8, 9]. Thus, the Mott-Hubbard MIT is driven solely by local electron correlations and does not involve any spontaneous symmetry breaking. Valuable insight into fundamental aspects of the Mott-Hubbard MIT has come from the dynamical mean-field theory (DMFT) [10]. Although DMFT neglects nonlocal correlations and thus becomes exact only in the limit of infinite dimensions [11, 12], it is able to capture the formation of local moments and resultant high frequency features in the single-particle spectrum — lower and upper Hubbard sidebands. In addition, DMFT predicts the first-order MIT line due to the coexistence regime of the metallic and insulating solutions at low temperature. The first-order MIT line terminates at a critical end point ($T_C, U_C$) similar to an ordinary liquid-gas transition. Consequently, one expects that the Mott critical end point belongs to the Ising universality class [13–15] with the double occupancy playing the role of a scalar order parameter of the transition. Subsequent studies based on quantum cluster techniques [16] revealed that the inclusion of short-range correlations on top of the local dynamics does not change the order of the Mott-Hubbard MIT which remains first order [17, 18].

Another issue when it comes to address the microscopic description of the MIT comes from the spin degrees of freedom which typically mask the Mott-Hubbard MIT by driving a magnetic instability of the metallic phase. This is for example known to happen in the Hubbard model on a two-dimensional (2D) square lattice at half-filling — due to the perfect nesting of the Fermi surface, the zero-temperature ground state displays long-range (AF) order for an arbitrary small on-site repulsion $U$. Essentially this type of localization is accounted for already at the single-particle level within a Slater formalism [19]: the onset of the antiferromagnetism leads to a doubling of the unit cell size which opens up an exponentially small gap $\Delta \propto t \exp(-2\pi \sqrt{t/U})$ even for small but nonzero $U$ at the Fermi level [20]. Increasing $U$ reduces the double...
occupancy and the Slater antiferromagnet progressively evolves into a Mott-Heisenberg insulator [2] with an AF gap of order of $U$ as verified within a variety of theoretical approaches [21–35]. In this regime, local moments form already on the temperature scale $T \sim U$ and thus the AF insulator at $T = 0$ is accounted for by a spin-1/2 Heisenberg model of localized spins with the superexchange constant $J = 4t^2/U$.

A prominent exception of the Mott insulator without any broken-symmetry-induced folding of the Brillouin zone down to $T = 0$ is found in the 1D Hubbard model at half-filling [36]. The essential difference in nature between 1D and higher-dimensional situations makes the studies of dimensional-crossover-driven phenomena very interesting [37–45]. Moreover, a combined effect of strong spatial anisotropy and geometrical frustration affects the strength of quantum fluctuations which in turn reduce the amplitude of broken-symmetry order parameters. This leaves a window for quantum critical behaviors and the emergence of novel ground states with anomalous single-particle spectra in their neighborhood [46–53]. Our previous works in this domain [54, 55] have established intriguing issues such as the unusual topology of the Fermi surface with dynamically generated pockets and fingerprints of Mott quantum criticality. Since the calculations in Refs. [54, 55] were carried out in the paramagnetic (PM) phase of the model, where by construction no long-range order is possible, it is conceivable that the onset of long-range order underlies the Mott transition. Thus, our aim is to revisit the phase diagram by adapting a cluster extension of DMFT (CDMFT) to handle standard Néel-type AF order [56].

Strictly speaking, long-range magnetic order on a 2D lattice with short-range interactions is destroyed by longwavelength fluctuations in the order parameter at any finite temperature [57]. However, at sufficiently low temperature, $T_{\text{DMFT}}$, captured relatively well already by a single-site DMFT [58], the correlation length grows exponentially with inverse temperature. This length scale defines an energy and time scale related to the fluctuations that are responsible for the destruction of the long-range order. Any experiment that is not able to resolve this energy or time scale will effectively perceive long-range order. Hence even at finite temperature, the fact that the correlation length diverges exponentially as a function of temperature has a very clear experimental signature observed in quantum simulations with ultracold atoms in optical lattices [59–62].

On the basis of CDMFT calculations, it was conceived in Ref. [63] that the position of maximum of $T_{\text{DMFT}}$ on the 2D square lattice is controlled by the critical end point $(T_c, U_c)$ of the MIT in the normal phase when the AF instability is artificially suppressed. Assuming that this (in general hidden) Mott-Hubbard MIT indeed marks the qualitative change in the microscopic mechanism behind the stability of AF order, an intriguing question arises: can one expect a profound influence on the underlying physics of the AF phase when the critical end point $T_c$ of the Mott-Hubbard MIT is driven down to zero upon increasing the degree of lattice anisotropy as in Ref. [55]? Most importantly, does this Mott quantum criticality affect the nature of the transition between a PM metal and the AF insulator? With this question in mind we proceed to discuss our findings.

The rest of the paper is organized as follows. In Sec. II we introduce the model, specify our implementation and technical details of the broken-spin-symmetry CDMFT algorithm, and define observables of interest. Our main results are discussed in Sec. III. We begin by presenting the anticipated finite-temperature phase diagram in the plane of interaction strength $U$ and hopping anisotropy $t_\perp/t$. Next, we elucidate the evolution of critical temperature and interaction strength $(T_c, U_c)$ terminating the first-order MIT upon varying the degree of lattice anisotropy. Subsequently, we turn our attention to the corresponding evolution of the electronic structure in the PM metal phase. Finally, we examine the reconstruction of low-energy quasiparticle excitations on going through the itinerant AF transition identified in the quasi-1D region of the phase diagram. We summarize our results and point out possible future directions in Sec. IV.

II. MODEL, METHOD, AND OBSERVABLES

To handle numerically crucial physical ingredients at play we use here CDMFT. Specifically, the $2 \times 2$ cluster is a minimal unit cell which allows one to capture the 1D umklapp scattering process opening a gap in the half-filled band [36, 64–66] and at the same time to treat short-range $x$- and $y$-axis AF spin fluctuations on an equal footing.

Our aim is to extend previous CDMFT studies of the influence of strong AF correlations on the nature of the MIT in the 2D Hubbard model [63, 67, 68] to the quasi-1D case. To this end, we consider the square lattice Hubbard model with an anisotropic hopping at half-filling,

$$H = -\sum_{ij,\sigma} t_{ij} c_{i\sigma}^\dagger c_{j\sigma} + U \sum_i n_{i\uparrow} n_{i\downarrow} - \mu \sum_i n_{i\sigma},$$

with a local Coulomb repulsion $U$, chemical potential $\mu$, and electron hopping amplitudes: $t_{ij} = t$ on the intrachain bonds, $t_{ij} = t_\perp$ on the interchain bonds, and $t_{ij} = t' = -t_\perp/2$ between next-nearest-neighbor sites on two adjacent chains. Thus, the energy dispersion for the non-interacting case reads

$$\epsilon_k = -2t \cos k_x - 2t_\perp \cos k_y - 4t' \cos k_x \cos k_y - \mu.$$
hopping $t'$ is strong enough to push the magnetic transition temperature below the critical end point temperature $T_c$ thus exposing the Mott-Hubbard MIT [69, 70]. A downside of the lifted perfect nesting condition is that one has to adjust the chemical potential $\mu$ as a function of $t_\perp, U$, and $T$ to keep the required condition of a half-filled band.

In CDMFT the original interacting lattice is mapped onto a cluster quantum impurity problem embedded in a self-consistent electronic bath. A predefined unit cell of volume given by the cluster size allows easily to study spin-symmetry-broken phases with a commensurate wavevector such as simple Néel AF order. To implement the CDMFT method, we decompose the lattice into $N_u$ supercells with $N_c$ atoms each. As a result, the non-interacting Green’s function $G_0(K, i\omega_n)$ and the spin-dependent self-energy $\Sigma_\sigma(K, i\omega_n)$ correspond to $N_c \times N_c$ matrices with wavevectors $K$ that span the reduced Brillouin zone of a supercell. In analogy to the DMFT approach, the CDMFT approximation neglects the $K$ dependency of the self-energy, $\Sigma_\sigma(K, i\omega_n) = \Sigma_\sigma(i\omega_n)$. The latter is extracted by solving the effective cluster Green’s function $G_\sigma(i\omega_n)$ and hence — via the Dyson equation — the cluster self-energy $\Sigma_\sigma(i\omega_n) = G_\sigma^{-1}(i\omega_n) - G_\sigma^{-1}(i\omega_n)$. The self-consistent loop is closed by requiring that the cluster Green’s function $G_\sigma(i\omega_n)$ matches the lattice Green’s function of the original model formulated in the cluster-site basis:

$$G_\sigma(i\omega_n) = \frac{1}{G_\sigma^{-1}(i\omega_n) - \Sigma_\sigma(i\omega_n)} = \frac{1}{N_u} \sum_K G_0^{-1}(K, i\omega_n) - \Sigma_\sigma(i\omega_n).$$

This allows us to compute a new bath Green’s function $G_{0,\sigma}(i\omega_n)$ which is fed back to the impurity solver and the procedure is repeated till convergence is reached.

While our preliminary results for the 2D case were obtained using the quantum Monte Carlo (QMC) method of Hirsch and Fye as cluster solver [71], we found it advantageous to switch to the continuous-time QMC (CTQMC) algorithm [72]. In particular we opted for its weak-coupling implementation based on a stochastic series expansion for the partition function in the interaction representation [73, 74]. It allowed us to reach temperatures as low as $T = t/100$ necessary to pin down the evolution of $T_c$ in the quasi-1D region. In addition, the CTQMC algorithm provides the possibility of Monte Carlo measurements directly on the Matsubara-frequency $\omega_n$ axis. Thus one avoids the cumbersome transformation from imaginary time to Matsubara frequencies necessary when the Hirsch-Fye solver is used instead.

To determine the domain of stability of the AF phase, in the actual simulation we explicitly break the SU(2) spin symmetry by introducing a small staggered magnetic field through the initial guess for the bath Green’s function $G_{0,\sigma}(i\omega_n)$. Hence, if the parameters of a simulation correspond to the regime with a thermodynamically stable AF phase, in the course of the CDMFT iterative process one converges to the solution with a finite staggered magnetization

$$m = \frac{1}{N_c} \sum_i (-1)^i (n_{i\uparrow} - n_{i\downarrow}).$$

We remark that allowing for a broken-spin-symmetry electronic bath substantially simplifies studies of otherwise a very intricate — in a generic nonrelativistic case — problem of a metal at the spin-density-wave quantum critical point involving coupling of gapless long-wavelength Goldstone modes to a Fermi sea [75–80].

Further insight into the underlying physics in different parts of the diagram is obtained from the following observables:

(i) Double occupancy

$$D = \frac{1}{N_c} \sum_i \langle n_{i\uparrow} n_{i\downarrow} \rangle.$$  

(ii) Lattice Green’s functions $g_\sigma(k, i\omega_n)$ in the original Brillouin zone with $k \in [-\pi, \pi]$. For consistency with our previous studies [54, 55], we extract it by periodizing the Green’s function in the cluster-site basis:

$$g_\sigma(k, i\omega_n) = \frac{1}{N_c} \sum_{\mu, \nu = 1}^{N_c} e^{i\mathbf{k}(\mathbf{a}_\mu - \mathbf{a}_\nu)} \times \left[ G_0^{-1}(k, i\omega_n) - \Sigma_\sigma(i\omega_n) \right]_{\mu, \nu},$$

where $\mathbf{a}_\mu, \mathbf{a}_\nu$ label cluster sites. It is however fair to remark that there are other periodization schemes such as the cumulant periodization which yield a faster convergence against the cluster size to the thermodynamic limit, an issue that becomes crucial in the proper description of doped 2D Mott insulators [81, 82]. Given that we are interested here in spectral properties of a moderately correlated metallic phase stabilized by strong magnetic frustration, one might still hope that the applied Green’s function periodization scheme reproduces the qualitative features of the thermodynamic solution.

(iii) Momentum-resolved spectral function at the Fermi level $A_k(\omega = 0)$. We estimate it from the behavior of lattice Green’s function at large imaginary time $\tau$ which gives the integrated spectral intensity in a frequency window of width $T$ around the Fermi level $\tilde{A}_k(\omega = 0) \propto \lim_{\beta \to \infty} \beta g(k, \tau = \beta/2)$ where $\beta = 1/T$ and

$$g(k, \tau = \beta/2) = \frac{1}{\beta} \sum_{\omega_n, \sigma} e^{-i\omega_n(\tau = \beta/2)} g_\sigma(k, i\omega_n).$$

We use $g(k, \tau = \beta/2)$ to analyze the evolution of the Fermi surface across the phase diagram.
\(\text{(iv)}\) Density of states at the Fermi level \(N(\omega = 0) = \frac{1}{N} \sum_k A_k(\omega = 0) \propto \lim_{\beta \to \infty} g_0(\tau = \beta/2)\) where
\[
g_0(\tau = \beta/2) = \frac{1}{\beta N} \sum_{k, \omega_m, \sigma} e^{-i \omega_m (\tau = \beta/2)} g_{\sigma}(k, i \omega_m). \quad (8)
\]

\(\text{(v)}\) Momentum-resolved spectral function \(A(k, \omega) = -\frac{i}{\pi} \text{Im} g(k, \omega)\). We have used the stochastic MaxEnt implementation \([84, 85]\) of the algorithms for Lattice Fermions (ALF) library \([86]\) to rotate the imaginary-time Green’s function \(g(k, \tau)\) to the real-frequency axis.

III. NUMERICAL RESULTS

A. Magnetic phase diagram

Our CDMFT results are summarized in the \((U, t_{\perp})\) phase diagram shown in Fig. 1. It was mapped out at a fixed temperature \(T = t/40\). On the one hand, a strong magnetic frustration introduced by the next-nearest-neighbor hopping \(t' = -t_{\perp}/2\) stabilizes in the weak-coupling region a PM metal phase which as shown in Fig. 1(a) extends to a fairly large interaction \(U_{\perp}/t = 5.17\) in the 2D limit. In this case, one finds a strong reduction of the double occupancy \(D\) at \(U_{\perp}\), indicative of the local moment \(\langle S_z^2 \rangle = 1 - 2D\) formation, see Fig. 1(b). On the other hand, we find that the Mott-Hubbard transition is preempted by AF order. Indeed, the \(U_{\perp}\) line in Fig. 1 corresponds to the transition from a PM metal phase to the AF insulator with increasing \(U\) while the \(U_{\perp}\) line to the transition from the AF insulator to the PM metal phase with decreasing \(U\). This hysteresis and jumps in both the staggered magnetization \(m\) [Fig. 1(a)] and double occupancy \(D\) [Fig. 1(b)] resolved for the moderately anisotropic region \(0.5 \leq t_{\perp}/t \leq 1\) clearly imply a first-order AF transition concurrent with the MIT. This contrasts with a static mean-field approximation where depending on the specific form of the band structure tuned by the magnitude of \(t'\), both the first-order and continuous transition between a PM metal and the AF insulator can be reproduced \([21, 87]\). In particular, within Hartree-Fock theory of the isotropic 2D Hubbard model one finds only a continuous transition for the particular choice \(t' = -t/2\) used here. In Sec. III C 1 we provide evidence that the first-order character of the MIT is actually a consequence of dominant local temporal fluctuations going beyond the mean-field approximation.

The situation in the strongly anisotropic part of the phase diagram is more subtle and requires more attention. In particular, a weak staggered magnetization in the vicinity of the AF transition (AFT) (dashed line in Fig. 1) accompanied by a relatively large double occupancy on the AF side are both suggestive of the itinerant magnetism.

In order to identify the character of the transition for \(t_{\perp}/t = 0.4\), we examine in Fig. 2 the behavior of the staggered magnetization \(m\) [Fig. 2(a)], double occupancy

![FIG. 1. Finite-temperature CDMFT phase diagram at \(T = t/40\) of the anisotropic 2D Hubbard model Eq. (1) at half-filling encompassing paramagnetic (PM) and antiferromagnetic (AF) phases. The shaded area between the \(U_{\perp}(U_{\perp})\) lines corresponding to a simulation with increasing (decreasing) the interaction strength \(U_{\perp}\) respectively, shows a coexistence region and implies a first-order metal-insulator transition (MIT). For interchain couplings \(t_{\perp}/t \leq 0.4\), a continuous AF transition (AFT) (dashed line) enables the onset of an AF metal phase which eventually undergoes a second-order MIT (dotted line) to the AF insulator. Color-coded circles display the behavior of: (a) staggered magnetization \(m\); (b) double occupancy \(D\), and (c) density of states at the Fermi level \(N(\omega = 0) \propto \lim_{\beta \to \infty} g_0(\tau = \beta/2)\).]
FIG. 2. Temperature dependence of the: (a) and (b) staggered magnetization $m$; (c) and (d) double occupancy $D$, and (e) and (f) density of states at the Fermi level $N(\omega = 0)$ [Fig. 2(e)] upon decreasing $U$ for various temperatures. One finds that the smooth increase of $m$ at the highest $T = t/20$ gets steeper at lower temperatures and is replaced by a small discontinuity at $T = t/40$. The latter is accompanied by a jump seen both in $D$ and $N(\omega = 0)$. This implies a first-order phase transition with a critical end point $T_c \simeq 1/40$ even though a slow convergence of the CDMFT self-consistency loop makes it hard to firmly assess the existence and range of the hysteresis at temperatures a little bit below $T_c$.

In contrast, simulations with a slightly smaller $t_{\perp}/t = 0.38$ yield a continuous onset of magnetism at $T = t/40$ and lower $T = t/50$ is required to resolve the discontinuity in all the three observables, see Figs. 2(b), 2(d), and 2(f). This result together with a shrinkage of the hysteretic region seen in Fig. 1 is suggestive of a systematic reduction of critical end point $T_c$ as a function of the growing lattice anisotropy. We analyze this issue in more detail in Sec. III B.

A continuous nature of the AF transition identified in the strongly anisotropic part of the phase diagram paves the way to an intermediate AF metal phase. Indeed, as shown in Fig. 1(c) there is a narrow region in the vicinity of the AF transition with small but finite $N(\omega = 0)$. Upon further increase of the interaction strength, a crossover from the AF metal to the AF insulator occurs once the staggered magnetic moment is sufficient to fully gap out hole and electron Fermi pockets of the AF metal phase.

Thus, in analogy with earlier studies restricted to PM solutions [54, 55], the continuous nature of the MIT stems from a smooth vanishing of the volume of Fermi pockets at the critical interaction. However, unlike in Ref. [55], where a dynamically generated breakup of the Fermi surface was the consequence of remnant 1D umklapp scattering, electron and hole Fermi pockets form here due to the doubling of the unit cell driven by AF order. As we discuss in Sec. III C 4, it results in a different topology of the Fermi surface in the vicinity of the MIT.

Finally, let us point out another important consequence of varying the ratio between inter- and intrachain hopping amplitudes — the existence of a certain critical point where the non-interacting 2D closed Fermi surface undergoes a topological change into an open surface. For the specific choice $t' = -t_{\perp}/2$ used in our studies, this is known to happen at $t_{\perp}/t \simeq 0.62$ [44]. As we demonstrate in Sec. III C 3, this topological (Lifshitz) transition is accompanied by a van Hove singularity in the single-particle density of states crossing the Fermi level. From a weak-coupling point of view, a large density of states might lead to divergent non-interacting susceptibilities in both particle-hole and particle-particle channels signaling enhanced ordering tendencies. Nevertheless, we find that phase diagram boundaries are insensitive to the van Hove singularity passing smoothly across the region $0.6 \leq t_{\perp}/t \leq 0.7$ with enhanced $N(\omega = 0)$, see Fig. 1(c). Together with a reduced double occupancy, this is yet another indication that the AF instability in this part of the phase diagram is not of a weak-coupling origin but
FIG. 3. Temperature dependence of the staggered magnetization $m$ (top) and double occupancy $D$ (bottom) obtained on decreasing $U$ in the case of: (a) and (b) weak anisotropy $t_\perp/t = 0.8$; (c) and (d) moderate anisotropy $t_\perp/t = 0.5$, and (e) and (f) strong anisotropy $t_\perp/t = 0.3$.

instead it should be considered as driven by the ordering tendency of preformed local moments.

B. Critical end point $T_c$

In this section we provide a detail analysis of the critical end point $(U_c, T_c)$ terminating the first-order MIT as a function of $t_\perp$. Identifying $(U_c, T_c)$ for a given $t_\perp$ requires numerous simulations at a variety of temperatures and as a function of $U$. We reduced this numerical effort by identifying first a crude estimate of the critical end point from simulations on a rough grid of temperatures. Next, we pinpointed $(U_c, T_c)$ to a better degree of accuracy by performing additional simulations on an appropriately refined grid restricted to the vicinity of the critical end point.

Figure 3 collects the resultant data for the AF order parameter $m$ and the double occupancy $D$ for three representative values of $t_\perp$ corresponding to different parts of the phase diagram. They range from $t_\perp/t = 0.8$ [Figs. 3(a) and 3(b)] through $t_\perp/t = 0.5$ [Figs. 3(c) and 3(d)] to $t_\perp/t = 0.3$ [Figs. 3(e) and 3(f)]. As apparent, for both $t_\perp/t = 0.8$ and $t_\perp/t = 0.5$ one finds a temperature range where the continuous behavior of $m(U)$ and $D(U)$ changes into a discontinuous jump. In contrast, for $t_\perp/t = 0.3$ we observe the persistence of a smooth behavior down to our lowest temperature $T = t/100$.

This motivated us to repeat the above analysis for other values of $t_\perp$ with the aim of elucidating the evolution of $T_c$ as a function of the interchain hopping. For each $t_\perp$ we define $T_c$ as a midpoint between the temperature at which $D(U)$ develops a singular behavior and the adjacent lower $T$ where $D(U)$ displays a clear jump. A striking outcome of this elaborate analysis is a nearly linear dependence of $T_c$ versus $t_\perp$, see Fig. 4. A linear fit to the data points in the $0.3 \leq t_\perp/t \leq 0.8$ range yields an estimate of a critical hopping $t_\perp^c/t = 0.31 \pm 0.03$ at which the MIT ceases to be first order, see Fig. 4(b).

This result can be rationalized by invoking a basic concept behind the DMFT algorithm [10]. It describes the formation of renormalized quasiparticles as a self-consistent Kondo screening of local moments by the electronic bath. In the case of a single-site DMFT, this screening involves only local spin-flip processes while in CDMFT also nonlocal spin-flips contribute. In addition, Kondo screening competes with the AF superexchange interaction between local moments. Given a substantial reduction of the double occupancy down to $D = 0.1348(4)$ in the PM metal on the verge of the MIT in Fig. 1(b), indicative of well formed local moments $S_0^z = 1 - 2D$, this competition is expected to be particularly strong in the isotropic 2D limit. In this limit we understand the Mott transition as a consequence of the jump to vanishingly small concentration of doubly occupied sites (doubloons) and empty sites (holons) triggered by the strong attraction of the doublon and holon scaled by $U$, which also drives the first-order transition due to the holon-doublon binding as described by the DMFT approximation. Magnetism is just a consequence of the
FIG. 4. (a) CDMFT estimate of the critical temperature and interaction strength ($T_c, U_c$) terminating the first-order MIT extracted from the vanishing of the double occupancy jump; for $t_{\perp}/t = 0.3$ we could not see resolve any signature of the discontinuous behavior down to our lowest temperature $T = t/100$. (b) Linear fit to the data points in the $0.3 \leq t_{\perp}/t \leq 0.8$ range which gives $t_{\perp}/t = 0.31 \pm 0.03$.

zero doublon and holon state that is thermodynamically unstable to magnetic ordering. In this sense magnetism rides on the MIT. This is very different from the MIT in semimetals (i.e. Hubbard model on a Honeycomb lattice) where the charge gap is a reflection of the magnetic ordering as in the case of Slater insulator [88, 89].

Upon growing lattice anisotropy, the observed increase in $D$ on the metallic side of the MIT indicates that the system crosses over to a weak-coupling regime. This diminishes the impact of local moments on the nature of the MIT and reduces continuously the magnitude of a jump in $D$, and thus $T_c$, down to zero.

C. Electronic properties

As discussed in Sec. IIIA, lattice anisotropy controls the behavior of the doubly occupancy and hence the degree of localization in the metallic phase. This shall have a strong impact on electronic properties of the metal. Below we systematically analyze the evolution of both Fermi surface and momentum-resolved single-particle (A($k$, $\omega$). It allows us to reveal dynamical effects arising from quantum fluctuations and to identify the underlying physics in different parts of the phase diagram.

1. Quasi-2D region: local moment formation

We begin with a weakly anisotropic case with $t_{\perp}/t = 0.8$. Figure 5 displays the evolution of the Fermi surface segment across two quadrants of the Brillouin zone with increasing $U$ in the PM phase at constant temperature $T = t/40$. The dynamical contribution to the self-energy in CDMFT yields a finite lifetime of quasiparticles. As a result, one finds a relatively sharp Fermi surface only for the smallest value $U/t = 1$, see Fig. 5(a), while dynamical effects become already discernible at $U/t = 2$ as Fermi surface blurring, see Fig. 5(b). As shown in Figs. 5(c) and 5(d), further increase of the interaction strength $U$...
induces substantial transfer of spectral weight from the Fermi level to finite-frequency parts of the single-particle spectrum indicative of a correlated metal.

A few additional comments are in order:

(i) It is known that in small cluster the effects of periodic boundary conditions are particularly strong which typically results in some artificial features in the single-particle spectra. For example, in Fig. 5(d) there is a noticeable depletion of weight at $k_x = \pi/2$. However, as we show later, its position is $t_{\perp}$ independent and pinned to $k_x = \pi/2$, and thus we consider it merely as a spurious feature of the $2 \times 2$ cluster.

(ii) Since the $2 \times 2$ CDMFT captures short-range AF spin fluctuations, the imaginary part of the self-energy can acquire a strong momentum dependence. If that is the case, the disappearance of the Fermi surface starts near the so-called hot spots, i.e., regions with an enhanced quasiparticle scattering rate. This gives rise to a pseudogap in the single-particle spectrum that precedes the Mott-Hubbard MIT [17, 90–97]. One could argue that the absence of momentum selective opening of the charge gap up to $U/t = 4.15$, see Fig. 5(d), where the system is on the verge of the transition to the AF insulator, is simply because the critical interaction strength is smaller when AF spin order is allowed. However, we believe that it is the consequence of a large next-nearest-neighbor hopping $|t'| = t_{\perp}/2$ used in the present studies which brings about a strong frustration of the AF spin correlations. Because of the first-order nature of the transition, the transition takes place before an appreciable momentum dependence manifested by the momentum differentiation gets started when $U$ is increased. The growth of the local moment indicated by the suppressed double occupancy without the growth of the AF spin correlations in the metallic side does not allow the strong momentum dependence in the spectral weight as well. This conclusion is supported by previous studies of the 2D half-filled Hubbard model within the eight-site dynamical cluster approximation which reported the suppression of momentum-space differentiation as the magnitude of $t'$ increases [98].

(iii) Dynamical correlations can also renormalize the Fermi surface topology via the real part of the self-energy [99]. Nevertheless, the inspection of Fig. 5 indicates that increasing $U$ does not modify noticeably the Fermi surface shape. Furthermore, anisotropic hopping amplitudes $t_{\perp} \neq t$ breaks the fourfold rotational ($C_4$) symmetry. This can lead to a surprisingly large directional anisotropy in the spectral intensity reflecting a large dynamically generated anisotropy in the self-energy close to the Mott-Hubbard MIT [100]. We do not observe here such anomalies possibly due to a combined effect of

FIG. 6. Single-particle spectrum $A(k, \omega)$ with increasing $U$ at $t_{\perp}/t = 0.8$ in the PM phase at $T = t/40$. The solid white line shows the free dispersion. Dashed blue (green) line with weaker (steeper) slope in (d) denotes the Fermi velocity for the interacting (free) case, respectively.
a relatively small critical interaction sufficient to trigger the transition from a PM metal to the AF insulator and a strong magnetic frustration.

In order to gain further insight into the onset of a correlated metal, we plot in Fig. 6 the corresponding momentum-resolved single-particle spectra $A(k, \omega)$. On the one hand, the spectral function in a weak-coupling regime $U/t = 1$ follows essentially the non-interacting dispersion shown as the solid white line in Fig. 6(a). On the other hand, qualitative changes in the spectrum produced by dynamical correlations are already found at $U/t = 2$: apart from the overall broadening, weak renormalization effects near the $k = (0, 0)$ momentum become visible as a reduced bandwidth of the coherent quasiparticle dispersion with respect to the non-interacting one, see Fig. 6(b). As shown in Figs. 6(c) and 6(d), further increase in $U$ leads to the transfer of the zero-frequency spectral weight into higher frequency regions as already anticipated in Fig. 5. As a result, one observes the formation of the incoherent lower and upper Hubbard bands: the former appears predominantly in the region of Brillouin zone around the $k = (0, 0)$ momentum while the latter emerges as $k$ moves towards the $(\pi, \pi)$ point. At the same time, the flattening of the quasiparticle dispersion near the Fermi level signals an increased effective mass of the quasiparticles and thus growing localization tendency. To quantify this effect we have plotted in Fig. 6(d) the Fermi velocity for the non-interacting and interacting cases. We see a reduction of approximately 30% around $(\pi, 0)$ and 40% around $(\pi/2, \pi/2)$ momenta.

2. **Quasi-2D region: thermal melting of local moments**

A jump in the double occupancy across the MIT implies its first-order nature. It is thus natural to expect that the physical mechanism that underlies the opening of the charge gap, i.e., the local moment formation, is robust in temperature.

We confirm this in Fig. 7 by examining the temperature evolution of $A(k, \omega)$ for a fixed $t_\perp/t = 0.8$ and at the corresponding critical interaction $U \lesssim U_c$ in the PM phase. Indeed, one finds the persistence of the coherent quasiparticle band near the Fermi level coexisting with incoherent high-frequency parts of the spectrum up to $T = t/10$, see Fig. 7(a). This is in accordance with a sizeable double-occupancy jump at this temperature illustrated in Fig. 3(b). At even higher $T = t/7.6$ [Fig. 7(b)], i.e., the highest $T$ at which one observes a jump in the double occupancy, a remnant quasiparticle band is still resolved in a narrow frequency window near the Fermi level. It is only at our highest $T = t/6.7$ [Fig. 7(c)] that the low-frequency quasiparticle band is washed out and the overall spectrum becomes blurred. At this temperature $T > T_c$, we could only detect a smooth transition from the PM to AF phase, see Figs. 3(a) and 3(b).

![FIG. 7. Evolution of the spectral function $A(k, \omega)$ with increasing temperature $T$ at $t_\perp/t = 0.8$ and at the critical interaction $U \lesssim U_c$ in the PM phase.](image)

3. **Moderate anisotropy: Fermi-surface topology change**

The presence of saddle points of energy dispersion yields a van Hove singularity in the single-particle density of states. Typically the access to study the behavior of a system near a van Hove singularity is achieved by the fine tuning of the electron density such that the Fermi surface approaches the singularity. Here we provide evidence that the lattice anisotropy is yet another control parameter that allows one to drive the van Hove singularity to cross the Fermi level.

We illustrate it in Fig. 8 which shows $A(k, \omega)$ at the critical interaction $U \lesssim U_c$ in the PM phase at $T = t/40$:
In accordance with a continuous nature of the transition for $t_{\perp}/t = 0.7$ [Fig. 11(a)] and $t_{\perp}/t = 0.3$ differs at $T = 0.03$ as established in Fig. 4(b).

Another signature of a reduced quasiparticle scattering off local moments at $t_{\perp}/t = 0.4$ is substantially restored coherence of low-energy quasiparticle excitations. Indeed, the resultant distinct Fermi surface found for both $t_{\perp}/t = 0.38$ [Fig. 11(a)] and $t_{\perp}/t = 0.4$ [Fig. 12(a)] contrasts sharply with that found at $t_{\perp}/t = 0.8$ where a clear loss of spectral weight is apparent, cf. Fig. 5(d). However, a precise physical mechanism of the magnetic phase transition for $t_{\perp}/t = 0.38$ differs at $T = t/40$ from that for $t_{\perp}/t = 0.4$. We discuss now both cases separately.

In accordance with a continuous nature of the transi-
FIG. 10. Spectral function $A(k, \omega)$ at the critical interaction $U < U_c$ in the PM phase at $T = t/40$ in the strongly anisotropic region with: (a) $t_{\perp}/t = 0.4$ and (b) $t_{\perp}/t = 0.3$. For comparison, panel (c) shows $A(k, \omega)$ for $t_{\perp}/t = 0.3$ at our lowest $T = t/100$.

At $t_{\perp}/t = 0.38$, one observes a smooth disappearance of the Fermi surface near "hot" regions corresponding to the intersection of the AF Brillouin zone with the Fermi surface, see Fig. 11(b). A more detailed inspection of the low-frequency part of $A(k, \omega)$ on the AF side of transition shows the backfolding of the quasiparticle dispersion due to the broken translation symmetry and the resultant depletion of spectral weight just below (above) the Fermi level in Figs. 11(c) and 11(e) [Fig. 11(d)], respectively. This depletion should be considered as a precursive feature of electron and hole pockets that open up at larger $U$. Indeed, one finds that the backfolded quasiparticle band crosses the Fermi level at two momenta — the spectral weight at the second crossing is much weaker than the original quasiparticle dispersion and causes a faint "ghost" side of the pockets. In fact, the observed gradual reconstruction of the low-energy quasiparticle dispersion

FIG. 11. Single-particle properties in the proximity of magnetic transition for $t_{\perp}/t = 0.38$ at $T = t/40$. Fermi surface cuts on the: (a) PM and (b) AF side of the transition. In (b) the measured value of the staggered magnetization $m = 0.0749(6)$. As a consequence of relatively weak magnetic order, the whole Fermi surface arc continues to exist on the AF side with a partial suppression of the spectral weight at the hot spots due to folding of Brillouin zone in the AF phase. (c)-(e) Low-energy part of the spectral function $A(k, \omega)$ along $(0,0) \to (\pi,0)$, $(0,0) \to (\pi,\pi)$, and $(0,\pi) \to (\pi,\pi)$ paths in the AF metal at $U/t = 2.28$ and (f)-(h) at $U/t = 2.3$ with the magnetization $m = 0.1419(4)$.
FIG. 12. Single-particle properties in the proximity of magnetic transition for \( t_\perp/t = 0.4 \) at \( T = t/40 \): Fermi surface cuts on the: (a) PM and (b) AF side of the transition. In (b) the measured value of the staggered magnetization \( m = 0.1275(8) \). Due to strong AF order, the Fermi surface breaks into disconnected pockets. (c)-(e) Low-energy part of \( A(k,\omega) \) along the Brillouin zone paths as in Fig. 11 in the AF metal at \( U/t = 2.365 \) and \( t_\perp/t = 0.4 \). (f)-(h) in the AF insulator at \( U/t = 2.6 \). Upon further increasing \( U \), the bands shift away from the Fermi level, gradually reducing the size of Fermi pockets in the intermediate AF metal phase which ultimately brings about the transition into the AF insulator, see Figs. 12(f)-12(h). Hence, we find aspects of both local and nonlocal correlation participating in the emergence of the insulating phase.

FIG. 13. Low-energy part of the spectral function \( A(k,\omega) \) at \( t_\perp/t = 0.3 \) and \( T = t/40 \) along \((0,0) \rightarrow (\pi,0)\) (a) and (d), \((0,0) \rightarrow (\pi,\pi)\) (b) and (e), and \((0,\pi) \rightarrow (\pi,\pi)\) (c) and (f) paths in the Brillouin zone at \( U/t = 2.2 \) (top) and \( U/t = 2.4 \) (bottom) across the transition from an AF metal to the AF insulator.

as Slater-like with quasiparticle scattering off essentially static staggered moment whose growing magnitude controls the size of the pockets as \( U \) grows, see Figs. 11(f)-11(h).

In contrast, the magnetic transition at \( t_\perp/t = 0.4 \) is weakly first order and significant AF order builds up right away at the critical interaction. This rapid develop is a consequence of a remnant local moment physics which enhances localization effects. This is reflected in disconnected Fermi surface segments on the AF side of the magnetic transition, see Fig. 12(b), accompanied by a definite gap between the lower and upper quasiparticle bands illustrated in Figs. 12(c)-12(e). Upon further increasing \( U \), the bands shift away from the Fermi level, gradually reducing the size of Fermi pockets in the intermediate AF metal phase which ultimately brings about the transition into the AF insulator, see Figs. 12(f)-12(h). Hence, we find aspects of both local and nonlocal correlation participating in the emergence of the insulating phase.

We turn now to the analysis of the low-frequency part of \( A(k,\omega) \) as a function of \( U \) for smaller hoppings \( t_\perp/t = 0.3 \) and \( t_\perp/t = 0.2 \) with the goal of elucidating the location of the corresponding MITs. In each case, a continuous loss of metalicity upon increasing \( U \) is ac-

\[
E_k^\pm = \frac{\epsilon_k + \epsilon_{k+Q}}{2} \pm \left( \frac{\epsilon_k - \epsilon_{k+Q}}{2} \right)^2 + \Delta^2, \quad (9)
\]

where \( \Delta = Um/2 \), consistent with that of the mean-field band structure in the spin-density-wave state. Hence, we identify the origin of the itinerant antiferromagnetism can be reproduced by a functional form
companied by the occurrence of more pronounced backfolded quasiparticle features such that they are visible in a broader momentum range, see Figs. 13 and 14. Let us also point out that the identified redistribution of spectral weight restricted to a narrow frequency region on the scale of the charge gap of the AF insulator is a generic feature of the magnetic instability in a weakly correlated metal driven predominately by the Fermi surface. This contrasts with the strong coupling regime where the onset of AF order in the Mott insulator involves the spectral weight transfer within the two Hubbard bands such that it accumulates at their lower edges \[68,102\]. Hence, the emergent magnetic ordering modifies the spectral properties over a broad energy range that is much larger than the AF charge gap itself.

The resultant MIT phase boundary anticipated from Figs. 12, 13, and 14 complements the CDMFT phase diagram shown in Fig. 1. Together with the line indicating the onset of the staggered magnetization, they both delimit the domain of stability of itinerant antiferromagnetism. It matches qualitatively the region of phase space with strongly reduced but nevertheless finite \[N(\omega=0)\], see Fig. 1(c).

Let us conclude the discussion of the established AF metal phase by contrasting its Fermi surface topology with that obtained in the previous studies in Refs. \[54,55\] restricted to the normal phase of the model Eq. (1). In this case the destruction of the Fermi surface starts at momenta \[k = (\pi/2, \pm \pi/2)\] where the interchain hopping matrix elements vanish and is driven by the remnant 1D umklapp scattering \[47,51\]. The resultant broken Fermi surface of the compensated metal displays elliptic electron and hole pockets around the \(k = (\pi/2, 0)\) and \((\pi/2, \pm \pi)\) points, see Fig. 5 of Ref. \[55\].

IV. SUMMARY AND CONCLUSIONS

Understanding the nature and factors controlling the degree of electron localization is crucial for exploring functional applications of quantum materials such as transition metal oxides. In this work we have made a contribution to this issue by studying the interplay between electron correlation, frustration, and dimensionality effects in the anisotropic Hubbard model at half-filling. To this end, we have adapted CDMFT to handle long-range AF order. An important outcome from our study is that the quasi-1D region of the magnetic phase diagram harbors an AF quantum critical metal. Consequently, in the CDMFT scenario one finds a crossover from a local momentum physics of a correlated isotropic 2D metal to the itinerant AF behavior in the strongly anisotropic case.

It is very interesting that, independently of a specific implementation of CDMFT, i.e., paramagnetic or broken spin symmetry, one observes a full suppression of the critical end point \(T_c\) of the MIT upon approaching the quasi-1D region. In both cases, the emergent quantum criticality can be traced back to a growing relative importance of spatial versus local fluctuations. Indeed, strong quasiparticle scattering off local moments along the whole Fermi surface explains its sudden disappearance and the resultant first-order character of the MIT in the 2D case. On the contrary, damping of low-energy quasiparticles in the quasi-1D region begins near "hot" regions of the Fermi surface. This leads to the formation of hole and electron Fermi surface pockets in a resultant compensated metal which ultimately undergoes a continuous MIT.

Specifically, when the CDMFT loop is constrained to converge to the PM solution, the MIT is driven by remnant 1D umklapp scattering and corresponds to the vanishing of the Fermi pockets driven by their continuous shift away from the Fermi level \[54,55\]. Likewise, when AF spin order is allowed, imperfect nesting of the model band structure paves the way to the itinerant AF transition followed by a MIT whose continuous nature is again the consequence of a smooth disappearance of Fermi pockets. In the actual simulations we explicitly broke the translational symmetry of the lattice by allowing for a nonvanishing staggered magnetic moment. The latter is equivalent to divergence of the static spin susceptibility at the AF wavevector \(Q = (\pi, \pi)\) measured on a sufficiently large cluster size capturing correctly the correlation length scale of spin fluctuations. From this point of view it becomes clear that introducing the lattice anisotropy tips the balance between local temporal fluc-
tions responsible for the Mott-Hubbard physics and thus—the first-order MIT—and spatial AF spin fluctuations playing a key role in the established continuous transition from an AF metal to the AF insulator.

The established CDMFT phase diagram with PM, AF metal, and AF insulator phases bears similarity with that of the extended Hubbard model featuring at the mean-field level PM, charge-ordered metal, and charge-order insulator phases [103, 104]. In the latter case one finds a tricritical point where all the three phases coexist. This very special point terminates also the continuous transition between the PM and charge-ordered metal phases. In our analysis we were unable to locate such a tricritical point. Instead as a function of the anisotropy $\frac{t'}{t}$ we find three situations: (i) first-order transition from a PM metal to the AF insulator; (ii) a crossover region where the dynamical correlations trigger a weakly first-order AF transition but they are not strong enough to fully localize charge carriers, and (iii) continuous transition between the PM and AF metal phases. Still, an attempt to induce the tricritical point could be made by fine tuning of the non-interacting band structure, i.e, the ratio of $\frac{t'}{t}$.

Another possible scenario in the strongly anisotropic limit would be the interaction-driven emergence of the genuine Mott insulator without any symmetry breaking characterized as the quantum spin liquid and sandwiched by the PM metal and the AF insulator. Such an intervening Mott insulating quantum spin liquid phase is beyond the scope of the CDMFT approximation. Since the auxiliary-field quantum Monte Carlo algorithm is hindered in the presence of geometrical frustration by the negative sign problem [86], a many-variable variational Monte Carlo method is an appealing option [105] to clarify this point of view.

Our second important result is evidence that the next-nearest-neighbor hopping $t'$ brings about an efficient mechanism to suppress the AF instability. In particular, we found that in the isotropic 2D situation with $t' = -\frac{t}{2}$, the PM metal extends to a fairly large interaction $U/t = 5.17$ below which previous CDMFT studies of the half-filled $t'$-Hubbard model reported a finite $d$-wave superconducting order parameter [106]. Since those studies did not consider long range AF order, it was possible that the latter prevails and leads to the insulating behavior at $T = 0$ instead. Our findings corroborate the scenario that the frustration of AF spin interactions by finite $t'$ shifts the onset of antiferromagnetism to a critical interaction which is large enough to expose $d$-wave superconductivity [107, 108]. Moreover, given our evidence for itinerant antiferromagnetism with Fermi surface pockets, it would be worth examining, e.g., using the spin fluctuation approach [109], the leading superconducting pairing instabilities in the spatially anisotropic situation.

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