I. INTRODUCTION

One of the outstanding problems in quantum many-body physics is to understand quasi-two-dimensional systems where both electron-electron interaction and geometric frustration are important. The triangular lattice is a prime example where the geometry frustrates near-neighbor antialignment of the spins that naturally tend to occur in the presence of short-range electron-electron interaction. Studying models of interacting electrons on such a lattice is thus certainly of fundamental interest, but it is also strongly motivated by the discovery of new materials. Prime examples of these materials are organic bis-(ethylenedithio) Cu3(CN)3 layers,5 triangular lattice antiferromagnets of the CuCrO2 family,3 and transition-metal oxide materials such as Na2CoO2 and Na1−xTiO2. The layered cobaltates have drawn much attention because of their unconventional properties. Sodium cobaltate shows an unusual strong thermopower4 at doping x = 2/3, which can be suppressed drastically by applying an in-plan magnetic field.5,6 The observation of Curie–Weiss behavior in the magnetic susceptibility while resistivity displays metallic behavior is another puzzle.7 The system also becomes superconductor when it is diluted by water.8–10 Various types of charge and spin orders also have been found in the system for various dopings.11–13 Na2CoO2 consists of two-dimensional CoO2 layers separated by insulating Na2+ layers. The CoO2 layers have Co atoms at the center of oxygen octahedra forming a 2D triangular lattice. The band structure calculation performed by Singh,14 revealed details of splitting of the 3d5 bands in Co atoms. With the help of this calculation and also of NMR experiments,7 one can find a rough estimate of hopping and exchange constants that would enter a two-dimensional Hubbard or t-J model for this system. However, the modeling is complicated by the fact that band structure calculations lead to hole pockets that are not observed experimentally, a question that is still debated by several groups using, for example, the Gutzwiller approximation,15 the local density approximation plus Hubbard U, and dynamical–mean-field theory17–19 In addition, the effect of long-range Coulomb interaction from the sodium leads to modifications to the simplest Hubbard Hamiltonian for the cobaltates.20,21 In this paper, we do not address the question of detailed modeling of the cobaltates or of other triangular lattice systems. Instead, we note that since several types of spin and charge-density waves are observed in these materials, it is quite likely that first-neighbor repulsion V and not only on-site repulsion U must be taken into account. U by itself tends to favor spin-density waves. We thus just focus on the simplest extended t-U-V one-band model Hubbard model on the triangular lattice and ask a few general questions: What types of phases are typical in different doping ranges, what type of interaction favors them, and should one expect pseudogap effects.

Previous theoretical and numerical works have obtained phase diagrams for the triangular lattice in the presence of competing interactions. There are, for example, variational Monte–Carlo calculations22,23 for the extended Hubbard model. That work focused mostly on the presence of the charge density wave (CDW) at filling n = 2/3 and RVB superconductivity at n = 1/3. Slave boson methods were used for the t-V and t-J models24,25 to study CDW, ferromag-
netism and also RVB superconductivity. Series expansion methods and cluster mean-field theory have also investigated CDW, Néel order, ferromagnetic order, dimer order, and phase separation in a $t$-$J$-$V$ model. We will comment further on some of these calculations in the context of our own results.

The results of this paper are obtained with the recently developed extended two-particle self-consistent approach (ETPSC) that is valid from weak to intermediate coupling. This method has been benchmarked against quantum Monte Carlo (QMC) simulations for the extended Hubbard model on a square lattice. The approach satisfies conservation laws and the Mermin–Wagner theorem, stating that no continuous symmetry can be broken at finite temperature in two dimensions. More traditional methods, such as the Random phase approximation, do not satisfy this requirement. With ETPSC, quantum renormalization of interactions (Kanamori–Brückner screening) is taken into account. Instability toward zero-temperature long-range order is signaled at finite temperature by crossover to the renormalized-classical regime where the correlation length grows exponentially. The wave vector of the instability is determined self-consistently by the approach and all wave vectors are in principle allowed. No a priori selection is necessary.

Within ETPSC, we can also compute the self-energy and other related quantities, such as the spectral weight that is measured in photoemission experiments. For the Hubbard model, it has been shown with the two-particle self-consistent approach (TPSC) that a pseudogap can appear as precursor induced by either antiferromagnetic or superconducting fluctuations. The former has been observed experimentally in electron-doped high-temperature superconductors. Our results demonstrate that CDW fluctuations can also induce a pseudogap. This is a relevant question experimentally given that CDW induced pseudogaps are observed and sometimes even show similarities with observations in high-temperature superconductors.34

We stress that our calculations begin at high-temperature. As the temperature is decreased we eventually reach a state without long-range order, which we call a Fermi liquid for simplicity, or we reach a crossover temperature below which a correlation length starts to grow exponentially at some wave vector. The diagrams for the crossover temperatures that we will present below specify whether the growing correlation length is in the charge or spin sector, as well as the wave vector of the growing correlations. For example, if that wave vector is not an integer fraction of the reciprocal lattice vectors and is in the spin channel, we say that we have an incommensurate spin-density wave. Our approach fails far below the crossover temperature, so one cannot be sure of the precise nature of the ground state. For example, we cannot tell if the incommensurate spin-density wave would be linear or spiral at zero temperature. In addition, in real systems, the precise nature of the coupling in the third direction will in general modify the precise details of the state that would be stabilized at finite temperature. Below the crossover temperatures that we find in two dimensions, the states generally have a pseudogap, so they are not Fermi liquid states. Also, at quantum critical points, non-Fermi liquid behavior extends to very high temperature.7

In the following, we first introduce the model and the ETPSC methodology. We next present our numerical results, discussing various physical effects in terms of the spin and charge structure factors. We display phase diagrams that help understand how microscopic parameters favor various phases. The CDW induced pseudogap and its effect on the Fermi surface are discussed before we present an overview and a conclusion.

II. MODEL AND METHOD

We write the extended Hubbard Hamiltonian in the following form,

$$H = -t \sum_{\langle ij \rangle \sigma} \sigma \langle c_{i\sigma}^\dagger c_{j\sigma} + c_{j\sigma}^\dagger c_{i\sigma} \rangle + U \sum_{\langle i \rangle} n_{i\uparrow} n_{i\downarrow} + V \sum_{\langle i \rangle \sigma} n_{i\sigma} n_{i\sigma'} - \mu \sum_{\langle i \rangle} n_{i\uparrow} + n_{i\downarrow} - \mu \sum_{\langle i \rangle} n_{i\sigma}$$

(1)

where $c_{i\sigma}$ ($c_{i\sigma}^\dagger$) is annihilation (creation) operator for electrons of spin $\sigma$ at site $i$ of a triangular lattice, $n_{i\sigma}$ is the density operator, and $t$ is the hopping matrix element. The quantities $U$ and $V$ are the on-site and nearest-neighbor interactions, respectively, and $\mu$ is the chemical potential.

For the Hubbard model ($V=0$), TPSC is a very reliable approach up to intermediate coupling limit. The functional derivative method is particularly convenient to obtain the TPSC equations. This is the method that was used to generalize TPSC to the extended Hubbard model.

The equations that need to be solved are the following. The charge and spin response functions take the form

$$\chi_{c\sigma}(q, \omega_n) = \frac{\chi^{0\sigma}(q, \omega_n)}{1 + \chi^{0\sigma}(q, \omega_n) U_{c\sigma}(q)}$$

(2)

$$\chi_{s\sigma}(q, \omega_n) = \frac{\chi^{0\sigma}(q, \omega_n)}{1 - \chi^{0\sigma}(q, \omega_n) U_{s\sigma}(q)}$$

(3)

where

$$U_{c\sigma}(q) = U \left[ \frac{\delta g_{\sigma\sigma}(0)}{\delta n} + n \frac{\delta g_{\sigma\sigma}(q)}{\delta n} + 4V \left[ g_{ss}(a) \gamma(a) + n \frac{\delta g_{ss}(a)}{\delta n} [3 + \gamma(a)] \right] \right]$$

(4)

and

$$U_{s\sigma}(q) = U g_{\sigma\sigma}(q) + 4V \left[ g_{ss}(a) \gamma(q) + n \frac{\delta g_{ss}(a)}{\delta n} [3 + \gamma(q)] \right]$$

are the charge and spin vertex functions and $\chi^{0\sigma}(q, \omega_n)$ is the free response function (noninteracting susceptibility) given by

$$\chi^{0\sigma}(q, \omega_n) = \int_{\text{BZ}} \frac{dp}{2\pi} \frac{\rho^\sigma(p + \frac{q}{2}) - \rho^\sigma(p - \frac{q}{2})}{i\omega_n - \epsilon_{p+q/2} + \epsilon_{p-q/2}}$$

(5)

with
\[ \epsilon_q = -2\left[ \cos(q,a) + 2 \cos(q,a/2)\cos(q,\sqrt{3}a/2) \right], \]

the noninteracting dispersion relation, and \( \gamma(q) = -\epsilon_q/2t \). In the above formula, \( v \) is the volume of the Brillouin zone, \( f^0(q) = 1/(1 + \exp[\epsilon_q - \mu_0]/T) \) is the Fermi function and \( \mu_0 \) is the noninteracting chemical potential. The pair correlation functions are related to the static structure factors by

\[ g_{sc}(r) = \int_{BZ} n \frac{d\mathbf{q}}{v} [S_{sc}(\mathbf{q}) - 1] \exp(i\mathbf{q} \cdot \mathbf{r}) \]  

and

\[ g_{ss}(r) = \int_{BZ} n \frac{d\mathbf{q}}{v} [S_{ss}(\mathbf{q}) - 1] \exp(i\mathbf{q} \cdot \mathbf{r}), \]

wherein \( S_{sc}(\mathbf{q}) = S_{ss}(\mathbf{q}) = S_{\sigma\sigma}(\mathbf{q}) \) are the charge and spin components of the static structure factor. The spin resolved static structure factor is defined by \( S_{\sigma\sigma}(\mathbf{q}) = (\langle n_{\sigma}(\mathbf{q}) n_{\sigma}(\mathbf{q}) \rangle)/n \) and \( n_{\sigma}(\mathbf{q}) \) is the Fourier transform of \( n_{\sigma}(\mathbf{a}) \). The quantities \( g_{sc}(\mathbf{a}) \) and \( g_{ss}(\mathbf{a}) \) entering the vertices in Eqs. (4) are simply the pair correlation functions at the first-neighbor distance.

Self-consistency is established by connecting the static structure factors to the response functions by the fluctuation-dissipation theorem

\[ S_{cc,ss}(\mathbf{q}) = \frac{T}{n} \sum_{\omega_n} \chi_{cc,ss}(\mathbf{q},\omega_n), \]

where \( \omega_n = 2n\pi T \) is the Bosonic Matsubara frequency. Substituting the expression for the susceptibilities in Eqs. (2) and (3) and the corresponding vertices in Eq. (4) on the right-hand side, one can use the result to obtain the pair correlation functions \( g_{ss} \) and \( g_{sc} \) entering the vertices using their relations in Eqs. (7) and (8) to the structure factors. Assuming that the functional derivatives of the pair correlation functions are known, as discussed below, we need only three equations to determine the pair correlation functions entering the vertices. This is because the Pauli principle imposes that \( g_{\sigma\sigma}(0) = 0 \). The equation that is dropped out is that for \( g_{sc}(0) = 0 \). This procedure and its impact on the Pauli principle is discussed in detail in Ref. 28.

Functional derivatives of the pair correlation functions with respect to density and magnetization enter the spin and charge vertices. The functional derivatives are obtained from the following equations:

\[ \frac{\partial g_{sc}(1,2)}{\partial n(1)} = [1 - g_{sc}(1,2)], \]

\[ \frac{\partial g_{ss}(1,1)}{\partial n(1)} = 2[1 - g_{ss}(1,1)], \]

and

\[ \frac{\partial g_{sc}(1,2)}{\partial n(1)} = [1 - g_{sc}(1,2)]. \]

These equations are strictly valid only when particle-hole symmetry is satisfied. On the square lattice, it has been checked by comparisons to QMC calculations that the results are satisfactory even in the absence of the full particle-hole symmetry.\(^\text{28}\) Apparently, particle-hole symmetry due to linearization of the dispersion relations near the Fermi surface suffices. We will make this assumption for the triangular lattice where strict particle-hole symmetry is not satisfied. This is justified \textit{a posteriori} by our results. Those that can be checked against variational QMC, for example, are in excellent agreement.

Finally, the self-energy needed to address the pseudogap problem is obtained following Ref. 27.

\[ \sum_{\sigma}(k,\omega_n) = (U n_{\sigma} + 6V n_{\sigma}) \left( \frac{T}{4} \sum_{\omega_n} \int_{BZ} \frac{d\mathbf{q}}{v} \left( U_{sc}(\mathbf{q}) \chi_{sc}(\mathbf{q},\omega_n) \right. \right. + \left. \left. U_{ss}(\mathbf{q}) \chi_{ss}(\mathbf{q},\omega_n) \right) \right), \]

where \( \omega_n = (2n + 1)\pi T \) is the Fermionic Matsubara frequency and \( \omega'_n \) is the Bosonic one. We can also find the spectral function \( A(\mathbf{q},\omega) = -\frac{1}{2\pi} \text{Im} \chi_{cc}(\mathbf{q},\omega) / G_0(\mathbf{k} + \mathbf{q}, \omega + \omega_n) \).

Finally, the interacting chemical potential is obtained from

\[ n = \frac{7}{2} \sum_{\omega_n} \int_{BZ} \frac{d\mathbf{q}}{v} \chi_{cc}(\mathbf{q},\omega_n). \]

### III. Response Functions

The “phase diagrams” that we present in the section that follows this one are determined from the growth of the spin and charge response functions as temperature decreases. When the interaction is local, \( U \), the wave vector of the instability, is determined entirely from the noninteracting susceptibility, in other words, from nesting properties of the Fermi surface. The introduction of near-neighbor repulsion \( V \) changes this since it introduces a wave vector dependence to the vertices. In order to disentangle the various effects, we present the noninteracting susceptibilities in the first subsection and the results with interactions in the second subsection. These numerical results are obtained from Eqs. (3)–(10).

#### A. Noninteracting susceptibility

The noninteracting susceptibility [Eq. (5)] is determined mostly by the shape of the noninteracting Fermi surface that is in turn determined by the dispersion relation given in Eq. (6). In Fig. 1, we present the Fermi surface for increasing...
values of the density, \( n = 0.5, 1, 1.25 \) and 1.5, respectively. The first Brillouin zone is plotted as a solid line. It is important to notice that the Fermi surface corresponding to \( n = 1.5 \) touches the first Brillouin zone boundary and has long parallel segments that lead to near nesting. We will show in more details that this causes a strong peak in the noninteracting response function. We also draw two wave vectors that are often found for the most important charge- or spin-density waves in the system. At these wave vectors, the charge or spin response functions often have a strong peak. The real-space modulations corresponding to these wave vectors are depicted in Fig. 2.

The noninteracting response function [Eq. (5)] is drawn in Fig. 3 for different values of densities \( n = 0.75, 1, 1.25, 1.5 \) and 1.75 at \( T = 0.2 \). The largest response is for \( n = 1.5 \). While one might have expected that parallel segments of the Fermi surface would have lead to a peak at a single dominant wave vector, it seems that the frustration imposes a less pronounced maximum. However, the height of the maximum at that density increases rapidly with decreasing temperature.

The position of all the peaks changes only slightly with temperature. The height of the peaks for the smaller values of density does not change drastically with decreasing temperature. That fact, in addition to quantum renormalization of the interactions, are the main reasons for the absence of any instability at low density up to intermediate coupling.

There is a deep minimum near the \( K \) point at higher values of the density. This is the main reason for absence of commensurate spin-density wave (SDW). The noninteracting response function has a peak at the commensurate wave vector \( Q_{c} \) (\( K \) point) at lower value of the density but, as we just mentioned, this peak does not grow enough to produce any sort of order including SDW up to intermediate coupling. That is not the case in the strong coupling limit but that is out of reach of our approach in the density regime where Mott physics is dominant.

It is quite remarkable that the free response function shows a strong peak at the origin for \( n = 1.75 \), a signature for ferromagnetism at nearby densities. It seems that, as we will see, frustration on the triangular lattice favors ferromagnetism at intermediate coupling, contrary to the square lattice case.

### B. Interacting response functions

In the presence of both types of interactions \( U \) and \( V \), the response functions are strongly modified; considering typical values of the interaction, \( U = 4 \) and \( V = 1.5 \). Using the same color code as in the figure for the noninteracting case, we show in Fig. 4 the spin [Eq. (3)] and in Fig. 5 the charge [Eq. (2)] response functions at \( T = 0.4 \) for the same densities as in the noninteracting case Fig. 3.

In the ordinary random phase approximation, the spin response function influenced only by the interaction \( U \) and the maxima are at the same location as in the noninteracting case. In the present approach, however, the nearest-neighbor interaction \( V \) also influences the spin response, introducing a
wave vector dependent vertex. Hence, some of the maxima of the spin response function in Fig. 4 are not at the same wave vector as in the noninteracting case. Nevertheless, the differences are much smaller than for the charge response function appearing in Fig. 5. In the latter case, the position of the maximum is near point $K$ (wave vector $Q_2$) for all densities, in other words, the charge response is dominated by the wave vector dependent vertex introduced by $V$.

One can summarize the results for the location of the maximum spin response function as follows. In the range $n \leq 1.5$, the tendency is toward an incommensurate spin-density wave (ISDW), while the response is maximum near zero wave vector (ferromagnetism) just above this density. Generally, the height of the peaks increases when $V$ is reduced, hence, nearest-neighbor repulsion does not favor spin order. For $n=0.75$, the maximum is near point $K$ corresponding to the same lattice structure as the CDW depicted in Fig. 2, except that one should replace the big or small points with up and down spins. This resembles the spin structure that would arise with ferrimagnetic order.

For the charge response in Fig. 5, tendency toward CDW order ($K$ point) is robust for these values of $U$ and $V$. The tendency is strongest at low values of the density because of a weaker effect of the frustration that leads to a dip in the noninteracting response function in Fig. 3. The effect of frustration is very strong for densities very close to $n=1.5$. Nevertheless, the height of the peak for densities around $n=1.5$ grows dramatically with decreasing temperature.

We verify the dominant effects of $U$ and $V$ discussed above, this time by fixing the filling at $n=1.5$ and changing the interactions. Figures 6 and 7 show, respectively, the effect on the charge and spin response functions at $T=0.4$.

For the charge response function in Fig. 6, we imply $U=4$ for those curves where the value of $U$ is not written. A simple comparison of the charge response function with the corresponding noninteracting susceptibility for $n=1.5$ in Fig. 3 shows the importance of the $V$ term in the vertex [Eq. (4)]. The noninteracting susceptibility has a deep minimum on the Brillouin zone boundary. The charge response function on the other hand shows two different maxima at wave vectors $Q_1$ and $Q_2$. The CDW modulation related to these wave vectors are illustrated in Fig. 2. We will see that these instabilities occur over a wide area in the $U-V$ plane. In fact, apart from the phase separation instability ($q=0$), which occurs for negative $V$, they are the only charge instabilities that can be found at this density. The situation changes as we change the density and one can expect to find incommensurate CDW instabilities in another region of the $U-V$ plane.

Moving on to the spin susceptibility, Fig. 7 shows that the presence of the $V$ term suppresses the spin response function
more and more as \( V \) increases, concomitant with the increase in the charge response function. In principle, one cannot find a strong maximum in both the spin and charge response functions. This is true in all one-band homogeneous paramagnetic systems as dictated by Eqs. (7) and (8) and \( g_{\alpha\beta}(0) = 0 \) (Pauli sum rule). Indeed, the Pauli sum rule (obtained from the Pauli principle \( \langle n_{\alpha}^2 \rangle = \langle n_{\beta} \rangle \)) connects \( g_{\alpha\beta}(0) \) to \( g_{\alpha\beta}(0) \) in such a way that an increase in one forces a decrease in the other.\(^{37}\)

**IV. CROSSOVER DIAGRAMS**

In mean-field theory, one normally finds finite temperature phase transitions, in contradiction with the Mermin–Wagner theorem. In ETNS, we obtain instead at a temperature \( T_X \) below which the correlation length begins to grow exponentially, diverging only at zero temperature. \( T_X \) is lower than the mean-field transition temperature because of the quantum Kanamori–Brückner renormalization of the vertices. In this low temperature regime, the characteristic frequency of the growing fluctuations becomes less than temperature in dimensionless units. This is the renormalized-classical regime. Either the spin or the charge correlations grow exponentially at some characteristic wave vector that suggests which long-range order will likely be stabilized at zero temperature. Since our approach is not valid deep in the renormalized-classical regime, one cannot be sure that the zero-temperature phase will be precisely that suggested by the behavior at \( T_X \).

The value of \( T_X \) depends on density, \( U \), and \( V \). We use \( \chi(q_x, q_y, 0) / \chi_0(q_x, q_y, 0) = \text{const} \) to estimate \( T_X \). For the sake of computational efficiency, we chose the constant to be 10 and checked that the general features do not change by choosing a larger value. This occurs because the exponential growth of the correlations is rather sudden. A detailed discussion of this issue can be found in previous publications.\(^{27,28}\)

**FIG. 7.** (Color online) The spin response function at fixed \( n = 1.5, U = 4, T = 0.4 \), and different values of \( V = 0, 1, 1.5, \) and 1.8.

**FIG. 8.** (Color online) Value of the crossover temperature \( T_X \) to the renormalized-classical regime as a function of \( U \) and \( V \) at filling \( n = 1.5 \). The wave vector and spin or charge character of the growing correlations is indicated by initials: Incommensurate spin-density waves, phase separation, or \( q = 0 \) charge instability (PS), and incommensurate charge-density waves. The wave vectors of the two special charge-density waves CDW1 and CDW2 (both \( \sqrt{3} \times \sqrt{3} \)) are shown in Figs. 1 and 2. The color scale (gray scale) appears on the right of the plot. Regions where either \( U \) or \( V \) are negative are shown for illustrative purposes only.

We present our results for \( T_X \) in Figs. 8–13 as color (gray scale) plots in various planes of parameter space. There are four plots that present the \( U-V \) dependence of \( T_X \) at four densities, and then two plots for the \( V-n \) dependence at fixed \( U \). We indicate by lines of various colors and types the boundaries between regions where there is either a change in the wave vector of the growing correlations or a change in the type of correlation, spin or charge. When we indicate a paramagnetic (PM) (Fermi liquid) region, we mean that correla-

**FIG. 9.** (Color online) Crossover temperature \( T_X \) to the renormalized-classical regime as a function of \( U \) and \( V \) at filling \( n = 4/3 \). Other symbols are defined in the caption of Fig. 8. Regions where either \( U \) or \( V \) are negative are shown for illustrative purposes only.
tions did not grow, in either the spin or charge sectors, at temperatures as low as $T=0.01$.

Figure 8 displays $T_X$ at $n=1.5$ as a function of $U$ and $V$ for both positive and negative values. At negative values of either $V$ or $U$, superconducting correlations will be competing. Since superconductivity has not been taken into account here, the results in all quadrants, except the first one, should be taken as just indicative of what may happen in the spin or charge sectors. When $V$ is negative, unless $U$ is large, there is a strong tendency to phase separation (PS), i.e., the static charge response function starts, at $T_X$, to grow exponentially for wave vector $q=0$. At positive $U$ and $V$, incommensurate spin-density waves are dominant, but $U$ and $V$ must be large enough, as expected from the absence of perfect nesting. At small $U$ and $V$, the system remains paramagnetic. Charge density waves appear at positive $U$ and $V$ only if $V$ is relatively large. Recall, however, that the effect of $V$ is amplified by the presence of several neighbors. The charge instability in this parameter range is of the CDW1 type illustrated in Fig. 2. Charge instabilities are further amplified at positive $V$ only if $U$ is allowed to become negative. The $\sqrt{3}\times\sqrt{3}$ CDW2 pattern is allowed only in extreme conditions of large positive $V$ and large negative $U$. This is not surprising since one can see from the noninteracting Fermi surface in Fig. 1 that the corresponding wave vector $Q_2$ is not particularly favored by nesting. The CDW2 phase is really governed by properties of the vertex $V$, not so much by the noninteracting Fermi surface.

When density is decreased to $n=4/3$, the noninteracting Fermi surface becomes almost circular so at positive $U$ and $V$
the tendency to order is strongly suppressed, as can be seen from Fig. 9. Compared to Fig. 8, the CDW2 vertex related instability is more robust while the CDW1 and ICDW instabilities occur in smaller regions, the ICDW existing over a larger region this time than CDW1. At larger fillings, $n = 5/3$, where again the Fermi surface becomes almost circular, similar features are observed.

As the filling decreases, the Fermi surface becomes more and more circular. Restricting ourselves to positive $U$ and $V$, and staying at weak to intermediate coupling where our theory is valid, nothing interesting occurs. The system remains paramagnetic down to $T = 0.01$. We thus also present, in Figs. 10 and 11, the results at large $U$ and $V$ where our theory is not strictly controlled. We feel these results are nevertheless interesting for two reasons. First, some of our “phase boundaries” compare favorably with results obtained from other methods. Second, the renormalized-classical regime occurs at such high temperature that $U/T$ and $V/T$ may begin to control the approach.

Figures 10 and 11 thus show the crossover diagram for, respectively, $n = 2/3$ and 0.5 over a wide range of positive $U$ and $V$. The CDW2 region now appears at positive $U$ and $V$, contrary to the results in the previous figures, as long as the stabilizing interaction $V$ is large enough. The boundary that separates CDW2 from PM in Fig. 10 is very close to QMC results, which gives us confidence in the validity of the results. The ferromagnetic phase is dominant when both $U$ and $V$ are large. The ICDW phase does not appear at filling $n = 2/3$ (Fig. 11). The CDW2 phase is influenced to some extent not only by the vertex, but also by commensurability, as can be seen from the fact that it is more important at $n = 2/3$ than at $n = 0.5$. There are competing tendencies for the CDW2 phase: 1) The density $n = 2/3$ is more favorable for the commensurate CDW2 as reflected in the free response function, and 2) The effect of off-site interaction is less important at lower value of the density. Based on these observations, one might surmise the presence of an optimal density where CDW2 appears over a larger area of $U$-$V$ space.

To explore in more details the density dependence, we present results as a function of $n$ and $V$ at fixed $U$. This was studied in particular by Motrunich and Lee. They calculated the phase diagram with different methods: (a) renormalized mean-field theory and variational quantum Monte–Carlo with a trial wave function and (b) slave boson mean-field theory. They suggest that the effect of the $V$ term is taken into the account more accurately in the first method than in the second one. In the first paper, they found that the CDW2 phase can be reached at smaller $V$ at the densities $n = 1/3$ and $2/3$ than at other densities. This is not the case in the second paper where these densities play no special role. Since their calculations are at $U = \infty$, this suggests that the general features of the phase diagram can be understood physically at low and high density; One needs a large $V$ to stabilize the CDW at low value of the density. Since at large value of the density, the hole plays the same role as the electron at low density, we expect at large $V$ to find the CDW as well. This argument is correct if other phases do not suppress the CDW.

We present the results of calculations at finite values of $U$ in Figs. 12 and 13. In Fig. 12, the results in the $V-n$ plane are for $U = 5$. It can easily be seen that CDW2 appears when both $V$ and $n$ are large. A ICDW region separates CDW2 from the paramagnetic or Fermi liquid phase. There is a small area on the left of the plane where the ferromagnetic phase is stable. Figure 13 shows the results for $U = 10$. CDW2 is still stable in the same region of the $V-n$ plane but the small FM region of the previous figure has now grown and pushed away slightly the CDW2 phase. In other words, at larger $U$, spin fluctuations are playing a more important role, as expected. This is also shown by the appearance of an ISDW regime. It is obvious from Figs. 12 and 13 that the densities $n = 1/3$ and $2/3$ do not play any special role, at least at these values of $U$. This is in agreement with the results of Ref. 24. However, at large $U$, we find more ferromagnetic spin fluctuations, a possibility that was not considered in Ref. 24. We also calculated the same phase diagram at higher value of $U$ where we find that the CDW2 region is completely swept away by ferromagnetism.

V. FERMI SURFACE AND PSEUDOGAP

It has already been shown that spin and superconducting thermal fluctuations can, in two dimensions, open up a pseudogap on the Fermi surface that reflects the wave vector of the fluctuations. The same study can be performed here to check the effect of charge-density wave fluctuations. We obtain the self-energy by substituting our results for the susceptibilities in Eq. (11), from which one can compute the spectral function.

We begin by the effect of spin fluctuations. In Fig. 14, we plot $A(q, \omega = 0)$ for $n = 1.5$, $U = 4$, $V = 0$, and $T = 0.4$. The dashed green line is the Brillouin zone. Following the largest intensity regions, one can recognize the shape of the noninteracting Fermi surface illustrated for $n = 1.5$ in Fig. 1. It is clear from this figure that the effect of the on-site interaction at this temperature is just to introduce damping. Since this is a region where ISDW appear at low temperature, the Fermi surface can be destroyed by lowering the temperature or increasing $U$.

For better understanding, we plot in Fig. 15 the spectral function as a function of frequency at different wave vectors for the same parameters as in the previous figure. These are energy dispersion curves (EDC). One can clearly observe the quasiparticle dispersion relation, the effect of $U$ appearing as damping. The extra features at higher frequency are precursors of extra bands that would appear in the ordered state. Indeed, the EDC are plotted in a regime where the correlation length associated with incommensurate fluctuations is
The spin-fluctuation induced precursor effects are also illustrated in Fig. 16. It is clear again from this figure that the pseudogap opens up around the M point where the suppression occurs in the MDC of Fig. 17. For other points, the peak positions change only slightly.

VI. DISCUSSION AND CONCLUSION

We have used ETPSC to clarify the leading instabilities of the extended Hubbard model on the triangular lattice. By instability we mean crossover to a renormalized-classical regime where there is generally a pseudogap and where Fermi liquid theory does not apply. The method is valid from weak to intermediate coupling. It satisfies the Mermin–Wagner theorem, includes quantum (Kanamori–Brückner) renormalization of the vertices, and does not assume a Migdal theorem for the self-energy. We interpret the entry into the renormalized-classical regime (exponential growth of the correlation length) as an indication of the phase that acquires long-range order at zero temperature. The actual order that is reached at zero temperature can in principle be at a different wave vector than the one that is dominant at the crossover temperature $T_X$. Also at $T_X$, one is still in a disordered state.
and it does not make sense to talk about the polarization of spin-density waves for example. Since we scan all wave vectors for both spin and charge instabilities, our method is not biased toward a restricted set of wave vectors, as most other studies. Superconducting fluctuations, however, have not been explored.

The range of possible fluctuations as a function of on-site interaction $U$, nearest-neighbor interaction $V$, and filling $n$ is quite rich. Not withstanding the superconducting instabilities, negative values of $U$ and $V$ favor charge instabilities either at zero wave vector (phase separation) when $U$ dominates, or at the CDW1 and CDW2 wave vectors when $V$ dominates. In the physically more relevant regime where $U$ and $V$ are both positive, spin instabilities are favored when $U$ dominates and they occur at wave vectors that are essentially determined by the Fermi surface. That is particularly clear at large fillings where the shape of the Fermi surface is non-trivial. Larger $V$, on the other hand, favors charge instabilities that are generally determined by the vertex itself instead of by details of the Fermi surface. This is particularly clear at small filling where the Fermi surface is essentially circular.

That predominance of the vertex is why the CDW2 ($\sqrt{3} \times \sqrt{3}$) region, for example, exists over a wide range of fillings and is not favored by commensurate fillings. We find that this CDW2 can compete with spin instabilities, especially ferromagnetism, when $U$ is large as well. Ferromagnetic fluctuations appear in a range of doping similar to that observed for the cobaltates. That competition with ferromagnetism has not been taken into account in earlier studies. All the crossover temperatures to the renormalized-classical regime appear at large values of the interactions when filling is small, somewhat outside the regime of validity of our approach. Nevertheless, agreement with other approaches suggests that ETPSC extrapolates in a reasonable way toward strong coupling. The fact that only short-range correlations (Fermi liquid) exist at small coupling and small fillings ($n = 2/3$) on the triangular lattice is a clear manifestation of the effects of frustration, as can be seen by contrasting with the square lattice case (where this does not occur).

Finally, we also showed that charge-density wave thermal fluctuations can also induce a pseudogap. A pseudogap associated with CDW is observed experimentally. If its origin is the one discussed in the present paper, it should disappear as temperature rises above that where the charge correlation length becomes of the order of the thermal de Broglie wavelength, as observed in the spin-fluctuation case in electron-doped cuprates.

Further studies should include the competition with superconducting fluctuations as well as more realistic modeling of the specific materials to which one wishes to apply our results. For example, the $n=1$ case is relevant for the layered organics.

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