Metal-insulator transition in the quarter- filled frustrated checkerboard lattice

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We study the electronic structure and correlations in the geometrically frustrated two dimensional checkerboard lattice. In the large $U$ limit considered here we start from an extended Hubbard model of spinless fermions at half-filling. We investigate the model within two distinct Green’s function approaches: In the first approach a single-site representation decoupling scheme is used that includes the effect of nearest neighbor charge fluctuations. In the second approach a cluster representation leading to a ‘multi-orbital’ model is investigated which includes intra-cluster correlations exactly and those between clusters on a mean field basis. It is demonstrated that with increasing nearest-neighbor Coulomb interaction $V$ both approaches lead to a metal-insulator transition with an associated ‘Mott-Hubbard’ like gap caused by $V$. Within the single site approach we also explore the possibility of charge order. Furthermore we investigate the evolution of the quasiparticle bands as function of $V$.

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I. INTRODUCTION

To treat the problem of a metal-insulator transition (MIT) driven by electron-electron interactions in lattice models with a fractional electron site occupancy, both the on-site and long-range Coulomb repulsions are equally important. A lattice with geometrical frustration imposes additional complications and may produce new features of MIT. The well known example is the Verwey transition in magnetite ($Fe_3O_4$) at $T_V \approx 120 K$. Magnetite has a spinel structure and $A$ sites are occupied regularly with $Fe^{3+}$ ions. On the $B$ sites forming a pyrochlore lattice, the $Fe$ ions are in the mixed valent state of $Fe^{2+}:Fe^{3+}=1:1$. Therefore, in the high-temperature charge disordered state, $T > T_V$, a band structure calculation would predict a quarter-filled conduction band, i.e., a metallic state with one itinerant electron per two sites. Below the Curie temperature $T_C \approx 850 K$ of magnetite the itinerant electrons are (almost) fully ferromagnetically polarized. Therefore they can be treated as a system of interacting spinless fermions. In terms of spinless fermions, the charge occupancy of one electron per two sites corresponds to a half band-filling. Below the Verwey temperature $T_V$, the charge degrees of freedom are ordered and an insulating state occurs. As was first pointed out by Anderson \cite{4}, properties of the Verwey transition can not be understood without taking into account the geometrical frustration and the resulting huge degeneracy of the ground-state charge configurations in the pyrochlore lattice. Two more recent examples of electronic structure changes connected to charge ordering in spinels are found in $AlV_2O_4$ with temperature decreasing at ambient pressure \cite{5,6} and in $LiV_2O_4$ under the external pressure \cite{7,8}. An interesting observation is that charge ordering in these three spinel systems is accompanied with a lattice structural change, i.e., the system tries to avoid the geometrical frustration of the pyrochlore structure. At the same time, no MIT and charge ordering were observed in $LiV_2O_4$ at ambient pressure down to very low temperatures, but instead the metallic compound $LiV_2O_4$ exhibits a heavy-fermion behavior below $30 K$ \cite{9,10}. Most of the work until now has been devoted to an understanding of the magnetic properties of geometrically frustrated lattices such as pyrochlore structure, since in the presence of antiferromagnetic interaction frustration acts against a conventional long-range order and may stabilize a spin-liquid state \cite{10}. Charge degrees of freedom have in contrast been studied much less \cite{11}. In fact, charge ordering in geometrically frustrated systems has been an intriguing and unsettled problems \cite{12}. For example, numerical diagonalizations of a Hamiltonian for spinless fermion with strong nearest-neighbor repulsion on a checkerboard lattice have given evidence that at half filling (number of spinless fermion equals one half the number of sites) the ground state is two-fold degenerate but a liquid \cite{13}. This must be kept in mind when that system is treated within different approximations leading to a charge ordered ground state. In other words, the observed charge order in frustrated structures could crucially depend on associated lattice deformations, i.e., the involvement of lattice degree of freedom.

The extended Hubbard model on non-frustrated lattices has been extensively investigated in one dimension at quarter-\cite{14} or half-filling\cite{17}, for two-leg ladders at quarter-filling\cite{16}, for two-dimensional square lattices at half-filling\cite{17} and in the limit of infinite dimensions at quarter-\cite{18} or half-filling\cite{19}. A variety of techniques, such as Hartree-Fock approximation, pertubation theory, dynamical mean-field theory, as well as numerical methods, e.g., quantum Monte Carlo and density-matrix renormalization group have been employed. However all these investigations were based on the non-frustrated lattice.
In the present paper, we apply a Green’s function approach to study the possible phase transitions of a half-filled spinless fermion model, i.e., a system with one electron per two sites, on the frustrated checkerboard lattice (see the inset of Fig. 1). The lattice can be viewed as a two-dimensional projection of a pyrochlore lattice. The spinless fermion model arises naturally for ferromagnetic materials in which one of the spin-split bands is completely occupied or completely empty as in magnetite \[20\]. Also this model can be viewed as a quarter-filled extended Hubbard model in the large \( U \) limit. Then double occupancy of a site is forbidden and the nearest-neighbour Coulomb repulsion \( V \) plays a crucial role. Because for (spinless) half filling every second site is unoccupied on the average one would naively expect a metallic state. Our main goal in this work is to show that inter-site correlations \( V \) can lead to a MI transition even for a case with less than one electron per site. We are using two different methods to study this model.

Firstly we employ a single site approach within a Hartree-Fock as well as Hubbard-I type approximations \( \text{Sect. III} \). By using the ‘Hubbard-I’-type decoupling scheme which includes the effect of nearest-neighbor charge fluctuations, we find that with increasing value of \( V \) first a metal-insulator transition occurs with a gap in the excitation spectrum, while at even larger values of \( V \) charge ordering appears. This is opposite to the result in the simple Hartree-Fock approximation, i.e. inter-site correlations favor the MI transition and suppress the CO. This observation may indicate that CO is indeed not present for a rigid checkerboard or pyrochlore lattice for any \( V/t \) ratio (it certainly is not in the limit \( V/t \to \infty \)). Indeed in the compounds \( \text{AlV}_2\text{O}_4 \) and \( \text{LiV}_2\text{O}_4 \) (under pressure) where CO has been found it is accompanied by a lattice distortion.

Secondly we start from a cluster representation of the model where the intra-cluster correlations are taken into account exactly and the inter-cluster terms are treated in Hartree-Fock approximation \( \text{Sect. III} \). This transformation leads to an effective multi-orbital extended Hubbard model. Again we find a M-I transition at a value similar to the first approach. In view of the suggestion above we do not consider the possibility of CO in this case, although within the Hartree-Fock approximation for the inter-cluster interactions it would presumably be present in the ground state.

Furthermore, we investigated the evolution of quasiparticle bands in the various phases in the single-site and cluster approaches. Following the selfconsistently determined chemical potential and the formation of interband gaps allows to determine the critical value for the MI transition.

\[ \text{II. SINGLE SITE APPROXIMATION} \]

The Hamiltonian for the spinless fermion model is given by \[11\]

\[
H = -t \sum_{\langle ij \rangle} c_{i}^\dagger c_{j} + \frac{V}{2} \sum_{\langle ij \rangle} n_{i} n_{j}
\]

where \( V \) is the Coulomb repulsion between nearest neighbors denoted by \( \langle ij \rangle \). We refer to this spinless Hamiltonian as the \( t-V \) model.

First, let us briefly review previous work on the \( t-V \) model at half-filling and the closely related quarter-filled extended Hubbard model in the large \( U \) limit for different lattices. In one dimension, the \( t-V \) model can be mapped onto an anisotropic Heisenberg model \[21\] and solved exactly via Bethe ansatz. In this case, one finds a gapless metallic phase for \( V < 2t \) and a gapped charge-ordered state for larger values of \( V \). For \( t-V \) model on a two-leg ladder, it was found from renomalization group calculations that contrary to a single chain, the ladder becomes a Mott insulator for arbitrarily small repulsive interactions \( V \) \[22\]. By using the density matrix renormalization group method, Vojta \[16\] et. al. studied the extended Hubbard model for two-leg ladders in the large \( U \) limit. They found that the charge-ordered phase vanishes for \( V < 2.5t \) but claimed that there will be a charge gap for all values of \( V/t \). For a square lattice, McKenzie \[22\] et al. argued from slave-boson theory that the insulating phase with charge order is destroyed below a critical value \( V \) of order \( t \) and the system becomes metallic.

Here we will study the \( t-V \) model on the checkerboard lattice at half-filling (one electron per two sites) with the aim to find out about possible MI and CO transitions in this frustrated lattice which is a 2D model for the pyrochlore lattice. At first we will employ the Green’s function method and various decoupling schemes within the single-site representation. There are two sites per unit cell. For convenience, we rewrite the Hamiltonian (1) in the following form

\[
H = H_t + H_{V,MF} + H_{t'}
\]

where

\[
H_t = -t \sum_{l=1}^{4} \left[ c_{l,1}^\dagger (c_{l-1,x,y,1} + c_{l,y,1} + c_{l+1,x,1}) + c_{l,2}^\dagger (c_{l,2} + c_{l+1,x,2} + c_{l-1,y,1}) + h.c. \right],
\]

and

\[
H_{V,MF} = \frac{V}{2} \sum_{\langle ij \rangle} n_{i} n_{j}
\]

and

\[
H_{t'} = \frac{t'}{2} \sum_{\langle ij \rangle} n_{i} n_{j}
\]
By introducing the Fourier transformation the equation (8) can be now written explicitly as

$$H_{V,MF} = V \sum \delta n_{1,2} \left( \langle n_{1-x,y,1} + n_{1-y,1} + n_{1+x,2} \rangle + \langle n_{1,2} \rangle \right) + \langle n_{1,2} \rangle \left( n_{1-x,y,1} + n_{1-y,1} + n_{1+x,2} \right)$$

$$+ \delta n_{1,1} \left( \langle n_{1+2,2} + n_{1-1,1} \rangle + \langle n_{1,1} \rangle \right) \left( n_{1+2,2} + n_{1-1,1} \right)$$

$$- \delta n_{1,2} \left( \langle n_{1-x,y,1} + n_{1-y,1} + n_{1+x,2} \rangle \right) - \langle n_{1,1} \rangle \left( \langle n_{1+2,2} + n_{1-1,1} \rangle \right)$$

$$H_V = V \sum \left[ \delta n_{1,2} \left( \delta n_{1-x,y,1} + \delta n_{1-y,1} + \delta n_{1+x,2} \right) + \delta n_{1,1} \left( \delta n_{1+2,2} + \delta n_{1-1,1} \right) \right].$$

Here we have introduced a charge fluctuation operator \( \delta n_{1,i} = n_{1,i} - \langle n_{1,i} \rangle \) on sites \( i = 1, 2 \) of the \( 1 \)-th cell. \( H_I \) is the kinetic energy term, \( H_{V,MF} \) is mean-field part of the interaction term while \( H_V \) is the residual interaction part. In the following we only consider the simplest charge ordered pattern, namely a staggered checkerboard pattern with wave vector \( Q = (0,0) \) illustrated in the inset of Fig. 1. The average charge density on different sites in the cell is \( \langle n_{1,i} \rangle = \frac{1}{2} - (-1)^i n \), where the order parameter \( n \) means a charge disproportionation within each unit cell.

The electron propagation is described by a retarded Green’s function (for simplicity the conventional symbol \( R \) is omitted)

$$G_{i,j} (1 - 1', \omega) = \langle \langle c^\dagger_{i} | c^\dagger_{j} \rangle \rangle_\omega = \int \langle \langle c^\dagger_{i}(t) | c^\dagger_{j}(0) \rangle \rangle e^{i\omega t} dt.$$  

(6)

where

$$\langle \langle c^\dagger_{i}(t) | c^\dagger_{j}(0) \rangle \rangle = i \theta (t) \langle \{ c^\dagger_{i}(t) , c^\dagger_{j}(0) \} \rangle.$$  

(7)

and \( \{ , \} \) denotes the anticommutator. The above \( (2 \times 2) \)-matrix Green’s function must satisfy the equation

$$\omega \langle \langle A | B \rangle \rangle_\omega = \langle \{ A , B \} \rangle + \langle \left[ A , H \right] | B \rangle \rangle_\omega.$$  

(8)

By introducing the Fourier transformation

$$G_{i,j} (k, \omega) = \frac{1}{N} \sum_{k} e^{i k (R_i - R_j)} G_{i,j} (1 - 1', \omega),$$  

(9)

the equation (8) can be now written explicitly as

$$\left( \omega \hat{1} + \hat{\Lambda} (k) \right) \hat{G} (k, \omega) = \hat{1} + V \hat{1} \hat{G} (k, \omega).$$  

(10)

with

$$\hat{\Lambda} (k) = \begin{pmatrix} (3 - 2n) V + 2 t \cos k_y & 4 t e^{-i k_x k_y} \cos \frac{k_y}{2} \cos \frac{k_x}{2} \\ 4 t e^{i k_x k_y} \cos \frac{k_y}{2} \cos \frac{k_x}{2} & -(3 + 2n) V + 2 t \cos k_x \end{pmatrix}.$$  

(11)
FIG. 2: Quasiparticle dispersion within mean-field theory for different value of $V/t$. The chemical potential $\mu$ is fixed at zero energy. (a) is the noninteracting case without charge order. These two bands touch at $(\pi, \pi)$ and the upper band is flat. (b) is in the metallic phase with small charge order. The two bands still touch while the upper band become dispersive due to the inequivalence of diagonal hopping. (c) is exactly at the critical point. The two bands touch at $(0, \pi)$. (d) is in the insulating phase. The two bands separate.

where $\hat{1}$ is the unit matrix and $\hat{\Gamma}(k, \omega)$ is the Fourier transformation of the $(2 \times 2)$-matrix $\Gamma_{i,j}(1 - \Gamma', \omega)$ $(i, j = 1, 2)$ which is defined as

$$
\Gamma_{i,j}(1 - \Gamma', \omega) = \left\langle \left\langle \sum_{(n.n.) \in (l, i)} \delta n_{(n.n.)} |c^\dagger_{l', j}\rangle \right\rangle \right\rangle_{\omega}.
$$

Here the summation is over six nearest-neighbor sites (n.n.) surrounding the site $(l, i)$

A. Mean-field approximation

In the first step we may decouple

$$
\Gamma_{i,j}(1 - \Gamma', \omega) \approx \left\langle \sum_{(n.n.) \in (l, i)} \delta n_{(n.n.)} \right\rangle \left\langle \left\langle c_{l,i} |c^\dagger_{l', j}\rangle \right\rangle \right\rangle_{\omega} = 0,
$$

which leads to a mean-field approximation and the quasiparticle dispersions are given by

$$
E_{\pm}^k/t = - (\cos k_x + \cos k_y - 3V/t) \\
\pm [4(1 + \cos k_x)(1 + \cos k_y) + (2nV/t - \cos k_x + \cos k_y)^2]^{1/2}.
$$

Now the intra-cell charge disproportionation $n$ has to be determined from the following self-consistent equation

$$
n = \frac{1}{2N} \sum_k \frac{(2nV/t - \cos k_x + \cos k_y)}{[4(1 + \cos k_x)(1 + \cos k_y) + (2nV/t - \cos k_x + \cos k_y)^2]^{1/2}}.
$$

The calculated phase diagram is shown in Fig. 1. In the present mean-field approximation ($\hat{\Gamma} \equiv 0$) the metallic phase is already charge ordered before a charge-transfer-type metal-insulator phase transition takes place at $V = 2.8t$. In the non-interacting case $V = 0$ (Fig. (2a)), the upper band is flat while the lower band is dispersive. It touches the flat band at $(\pi, \pi)$. As shown in Fig. (2b, 2c) with increasing intersite repulsion $V$ the flat band becomes increasingly dispersive with increasing charge order and the touching point moves from $(\pi, \pi)$ towards $(0, \pi)$. The dispersion of the previously flat band is due to the inequivalence of diagonal hopping (see inset of Fig. 1) induced by charge order. Finally, for $V > 2.8t$ the two bands separate as shown in Fig. (2d). In the half-filled case, the lower band is fully occupied and the upper band is empty resulting in a charge-transfer-type insulator.
B. ‘Hubbard I’-approximation for the inter-site correlations

The neglect of correlations overestimates the tendency to CO symmetry breaking. Therefore, in the strongly correlated case \(V \gg t\), the mean-field results are unreliable and it is necessary to consider equations of motion of higher-order Green’s functions \(\Gamma_{i,j}(1 - \Gamma, \omega)\) which can be written as

\[
\begin{align*}
\left( \omega - V \sum_{(n.n.) \in \{l,1\}} \langle n(n.n.) \rangle \right) \left\langle \left( \sum_{(n.n.) \in \{l,1\}} \delta n(n.n.) | c_{l,i}^\dagger \right) \right\rangle_{\omega} \\
+ V \left\langle \left( \sum_{(n.n.) \in \{l,1\}} \delta n(n.n.) \right)^2 | c_{l,i}^\dagger \right\rangle_{\omega}
\end{align*}
\]

(16)

Here we adopt the following approximations, which consist in an extension of the on-site Hubbard I decoupling scheme \[24\] to spinless fermion on a checkerboard lattice with inter-site Coulomb interaction:

\[
\left\langle \left( c_{l,i}^\dagger c_{l,i}^\dagger c_{l,i}^\dagger c_{l,i}^\dagger \right) \right\rangle_{\omega} \approx 0.
\]

(17)

In equation (16), the fourth term involves both the nearest- and next-nearest-neighbor charge correlations and we neglect the latter. When treating the last term in equation (16) we also neglect the more distant charge correlations and approximate as

\[
\begin{align*}
\left\langle \left( c_{l,i}^\dagger \left( \sum_{(n.n.) \in \{l,1\}} \delta n(n.n.) \right)^2 c_{l,j}^\dagger \right) \right\rangle_{\omega} \\
\approx \left\langle \left( \sum_{(n.n.) \in \{l,1\}} \delta n(n.n.) \right)^2 \left\langle c_{l,i}^\dagger c_{l,j}^\dagger \right\rangle \right\rangle_{\omega} \\
\approx \left( \frac{3}{2} - 6n^2 + 2 \sum_{(n.n.) \in \{l,2\}} \delta n_{l,2} \delta n(n.n.) \right) \left\langle \left\langle c_{l,i}^\dagger c_{l,j}^\dagger \right\rangle \right\rangle_{\omega}.
\end{align*}
\]

(18)

Finally the equation of motion for the higher-order Green’s function \(\Gamma_{i,j}(k, \omega)\) can be written as

\[
\left( \omega \mathcal{\Gamma} + \hat{\Lambda}(k) \right) \hat{\Gamma}(k, \omega) = \hat{B}(k) + \hat{M}(k) \hat{G}(k, \omega).
\]

(19)

where

\[
\hat{M}(k) = \left( \begin{array}{cc}
V K_2 & -8n t e^{-i \frac{k_x + k_y}{2}} \cos \frac{k_x}{2} \cos \frac{k_y}{2} \\
8n t e^{i \frac{k_x + k_y}{2}} \cos \frac{k_x}{2} \cos \frac{k_y}{2} & V K_1
\end{array} \right),
\]

(20)
FIG. 3: Results within Hubbard I-like approximation. (a) charge order (CO) parameter \( n \) versus \( V/t \). The dotted line indicates the phase boundary between the metallic and insulating phase without charge ordering. Contrast to the mean-field approximation, the charge-ordered phase transition is of first order and occurs after a Mott-Hubbard-like metal-insulator transition takes place. The dashed line shows the quasiparticle gap versus \( V/t \) in both insulating phases. (b) Hopping amplitude (solid line) from site 2 to site 1 \( \langle \hat{c}^\dagger_{l1} \hat{c}^{\phantom{\dagger}}_{l2} \rangle \) and nearest-neighbor charge fluctuation correlations \( \sum_{\text{n.n.} \in (l,i)} \delta n_{\text{n.n.}} \) (dashed line) versus \( V/t \). In the insulating phase without charge order, the expectation value \( \langle \hat{c}^\dagger_{l1} \hat{c}^{\phantom{\dagger}}_{l2} \rangle \) is equal to 0 while \( \sum_{\text{n.n.} \in (l,i)} \delta n_{\text{n.n.}} \) keeps constant and maximum. Note that \( \sum_{\text{n.n.} \in (l,i)} \delta n_{\text{n.n.}} \) is site-independent for all value of \( V/t \). (c) Energy difference between ordered and disordered state versus \( V/t \). For \( V > V_\text{c2} \), the charge-ordered state has lower energy.

and

\[
\tilde{B}(\mathbf{k}) = \left( \begin{array}{c} 
2 \langle \hat{c}^\dagger_{11} \hat{c}^\dagger_{1-\mathbf{k},1} \rangle \cos k_y \\
4 \langle \hat{c}^\dagger_{11} \hat{c}^\dagger_{12} \rangle e^{i k_x + i k_y} \cos \frac{k_x}{2} \cos \frac{k_y}{2} \\
4 \langle \hat{c}^\dagger_{11} \hat{c}^\dagger_{12} \rangle e^{-i k_x + i k_y} \cos \frac{k_x}{2} \cos \frac{k_y}{2} \\
2 \langle \hat{c}^\dagger_{12} \hat{c}^\dagger_{1\mathbf{k},2} \rangle \cos k_x 
\end{array} \right).
\]  

(21)

Here we used the definition

\[
K_i = \left( \frac{3}{2} - 6n^2 + 2 \left( \delta n_{1,i} \sum_{\text{n.n.} \in (l,i)} \delta n_{\text{n.n.}} \right) \right).
\]  

(22)

By solving equation (19) with respect to \( \hat{\Gamma} \) and after substituting \( \hat{\Gamma} \) into equation (10) one obtains the final solution for the Green's function \( \hat{G} \). For the electron concentration 1/2 (one electron per unit cell), the chemical potential \( \mu \), the charge disproportionation \( n \) and the hopping amplitudes \( \langle \hat{c}^\dagger_{11} \hat{c}^\dagger_{12} \rangle \) are calculated from the following set of self-consistent equations:

\[
1 = \frac{1}{N/2} \sum_{\mathbf{k}} \int_{-\infty}^{\mu} d\omega \left( -\frac{1}{\pi} \right) \text{Im} [G_{11}(\mathbf{k},\omega) + G_{22}(\mathbf{k},\omega)],
\]  

(23)
The behavior of $V > V_c$ shown that a first order MI transition occurs at $V = V_c$ and a decoupling scheme that includes the effect of nearest-neighbor charge fluctuations, however, it was observed that the tendency to symmetry breaking, CO appears before the system becomes an insulator. By using a Green’s function lattice within mean field theory and a ‘Hubbard-I type’ approach. In the former approximation which overestimates the increase, charge order becomes more and more pronounced and can no longer distinguish between bonding and antibonding states.

As a consequence, four bands are obtained. For $V < V_c$ these bands split into four when $0 < V < V_c$ and $E^+_k|_{V=0} = -2t(\cos k_x + \cos k_y) + 1$ and $E^+_k|_{V=0} = 2t$ as expected. When $t = 0$, two poles $\omega = 2V$ and $\omega = 4V$ are obtained from above equations, which describes one hole or one particle excitation.

In Figure (3a) we show the phase diagram within the previously used decoupling scheme. First, a metal-insulator transition occurs at $V_c = 4.86t$ and then charge ordering appears at $V > V_c$ and $V < V_c$ a gap opens and increases with increasing value of $V$. At $V_c$, it drops to a smaller value. For $V > V_c$, the gap increases again monotonously. In the charge ordered phase, there exist always two self-consistent solutions, i.e., a charge ordered and a disordered one. By comparing the energy of these two states, (see Fig. 3 (c)), we find for $V > V_c$ that the charge ordered state has lower energy. The phase transition is of first order. Fig. 3 (b) shows that at the metal-insulator transition the hopping amplitude $\langle c_{11}^c c_{12}^\dagger \rangle$ vanishes.

In the following we want to explain this feature in more detail. For that purpose we study the poles of the retarded Green’s function exhibiting four poles given by the roots of $D = 0$ where $D$ is defined as

$$D = (\omega - E^+_1)^2 (\omega - E^-_1)^2 + V^4 K_1 K_2 + 16n^2 V^2 t^2 (1 + \cos k_x) (1 + \cos k_y)$$

$$+ V^2 K_1 [(\omega - (3 - 2n) V + 2t \cos k_y)^2 - 4t^2 (1 + \cos k_x) (1 + \cos k_y)]$$

$$+ V^2 K_2 [(\omega - (3 - 2n) V + 2t \cos k_x)^2 - 4t^2 (1 + \cos k_y) (1 + \cos k_y)]$$

As a consequence, four bands are obtained. For $V = 0$, two branches of momentum-dependent dispersions are obtained from equation (27): $E^-_k|_{V=0} = -2t(\cos k_x + \cos k_y) + 1$ and $E^+_k|_{V=0} = 2t$ as expected. When $t = 0$, two poles $\omega = 2V$ and $\omega = 4V$ are obtained from above equations, which describes one hole or one particle excitation.

In the regime $0 < V < V_c$, states in the antibonding flat band become more and more occupied as $V$ increases. Therefore the expectation value $\langle c_{11}^c c_{12}^\dagger \rangle$ decreases since an equal occupational probability of a bonding and antibonding state implies that $\langle c_{11}^c c_{12}^\dagger \rangle = 0$. This is the case in the regime $V_c < V < V_{c2}$ shown in Figs. 4 (d), (e). Note that a correlation gap has opened above the two occupied lower bands which implies a Mott-Hubbard type insulating state and the nearest-neighbor charge fluctuation correlations $\delta n_{i,i} \sum_{(n.n.) \in \{1,1\}} \delta n_{(n,n.)}$ remain constant. When $V > V_{c2}$, then due to symmetry breaking one can no longer distinguish between bonding and antibonding state and $\langle c_{11}^c c_{12}^\dagger \rangle \neq 0$ (see Figs. 4 (f)). As $V$ continues to increase, charge order becomes more and more pronounced and $\langle c_{11}^c c_{12}^\dagger \rangle$ decreases again monotonously. This explains the behavior of $\langle c_{11}^c c_{12}^\dagger \rangle$ shown in Fig. 3. The inflexion point at $V/t = 2.2$ shown in Fig. 3 (b) is due to the lower flat band crossing the inflexion points (maximum of DOS) of the upper dispersive band shown in Fig. 4 (b). For $V > V_{c2}$, CO strongly suppresses the nearest-neighbor charge fluctuation correlations $\delta n_{i,i} \sum_{(n.n.) \in \{1,1\}} \delta n_{(n,n.)}$ which is site-independent for all value of $V/t$ as expected.

In this section we have studied the half-filled (one electron per two sites) spinless $t – V$ model on a checkerboard lattice within mean field theory and a ‘Hubbard-I type’ approach. In the former approximation which overestimates the tendency to symmetry breaking, CO appears before the system becomes an insulator. By using a Green’s function approach and a decoupling scheme that includes the effect of nearest-neighbor charge fluctuations, however, it was shown that first a MI transition into an insulating state without CO takes place and only for larger $V$ eventually
FIG. 4: evolution of the quasiparticle spectrum within a Hubbard I-like approximation for different value of \( V/t \). The chemical potential \( \mu \) is fixed at zero energy. \( V/t = 0 \) is already shown in Fig. 2 (a). Figs. (a), (b) and (c) are in the metallic phase \((0 < V < V_{c1} = 4.86t)\). Figs. (d) and (e) are in the insulating phase without CO symmetry breaking \((V_{c1} < V < V_{c2} = 6.47t)\). Reminiscent of the upper and lower Hubbard band splitting, the metal-insulator phase transition at \( V_{c1} \) is of Mott-Hubbard type. Fig. (f) is in the insulating phase with CO \((V > V_{c2})\).

CO appears. It is not entirely clear, however, how realistic the appearance of CO is. In the limit \((V/t \to \infty)\) when the hopping vanishes, the ground state for half filling is macroscopically degenerate [4, 11] and no CO is present [25]. It is therefore possible that CO obtained here for the rigid checkerboard lattice above \( V_{c2} \) may be due to the employed Hubbard-I type approximation. In fact, the charge ordered state obtained here is one of the macroscopically degenerate states which must obey the tetrahedron rule (Anderson rule) [4]. If there would exist corresponding lattice distortion, for example, compressed along the diagonal direction [12], the system would select such a CO state shown in the inset of Fig. 1 out of the macroscopically degenerate states. Indeed in some 3D pyrochlore compounds which exhibit CO as mentioned in the introduction, it is always accompanied by symmetry lowering lattice distortions which remove the frustration by introducing inequivalent bond lengths in the tetrahedrons of the corner sharing lattice.

III. CLUSTER APPROXIMATION

We want to supplement the previous calculation based on a Hubbard I type of approximation by another one where the strong electron correlations are treated exactly within a cluster but within mean-field approximation outside the cluster. Within this scheme we want to determine the critical interaction \( V_{c1} \) at which a gap opens in the excitation spectrum when the case of half-filling is considered. We do not care here about a possible charge order at large value of \( V \) because, as pointed out before, that may turn out to be an artefact of the involved approximations.

We divide the checkerboard lattice into sublattice A and B of the plaquette so that each sublattice contains \( N/4 \) units where \( N \) is the number of sites. Fig. 5 shows black (basic) clusters linked across white square. The Hamiltonian (1), \( H = H_t + H_V \), is decomposed into

\[
H_t = H_t^{(\text{intra})} + H_t^{(\text{inter})}, \quad H_V = H_V^{(\text{intra})} + H_V^{(\text{inter})}
\]

Here each of the terms, \( H_t^{(\text{intra})} \) and \( H_V^{(\text{intra})} \), is a sum over \( l = 1, \ldots, N/4 \) decoupled basic clusters while \( H_t^{(\text{inter})} \) and
\(H^{(\text{inter})}_{V}\) are due to the intercluster coupling. Specifically the decoupled clusters are treated by the Hamiltonian

\[
H^{(\text{intra})}_t + H^{(\text{intra})}_V = \sum_1 H^{(\text{intra})}_{t-V} (1)
\]

(29)

\[
H^{(\text{intra})}_{t-V} (1) = \sum_{i,j} M_{ij} (0) \left( -tc^\dagger_{i\alpha} c_{j\beta} + \frac{V}{2} n_{i\alpha} n_{j\beta} \right).
\]

(30)

Here \(i, j = 1, \ldots, 4\) denote the sites within a cluster with lattice vector \(1\) and \(M_{ij} (0)\) is a \((4 \times 4)\)-matrix with elements \(M_{ij} (0) = 1\), if \(i \neq j\), and \(M_{ii} (0) = 0\). Furthermore \(n_{i\alpha} = c^\dagger_{i\alpha} c_{i\alpha}\). Thus, \(H^{(\text{intra})}_{t-V} (1)\) represents an intracluster Hamiltonian to be solved below.

To write down in a systematic way the intercluster coupling given by \(H^{(\text{inter})}_t\) and \(H^{(\text{inter})}_V\), we refer to Fig. 5 where a fragment of the checkerboard lattice is shown with the origin located at the center of a basic cluster. There are eight neighboring basic clusters connected to a given one with the translation vectors \(\mathbf{\tau}\} = \mathbf{\tau}_1, \mathbf{\tau}_2, \ldots, \mathbf{\tau}_8\). Three of them are depicted in Fig. 5. The intercluster electron hopping and Coulomb repulsion terms can be written as

\[
H^{(\text{inter})}_t = -t \sum_1 \sum_{\{\mathbf{\tau}\}} \sum_{i,j} \mathbf{M}_{ij} (\mathbf{\tau}) c^\dagger_{i,1+\mathbf{\tau}} c_{j,1+\mathbf{\tau}}
\]

(31)

\[
H^{(\text{inter})}_V = \frac{V}{2} \sum_1 \sum_{\{\mathbf{\tau}\}} \sum_{i,j} \mathbf{M}_{ij} (\mathbf{\tau}) n_{i,1+\mathbf{\tau}} n_{j,1+\mathbf{\tau}}
\]

(32)

where the eight \((4 \times 4)\)-matrices \(\mathbf{M}_{ij} (\mathbf{\tau})\) are specified by referring to Fig. 5. Consider first the matrix \(\mathbf{M}_{ij} (\mathbf{\tau}_1)\) that connects two neighboring clusters by \(\mathbf{\tau}_1\)-translation. According to Fig. 5, only two individual two-site bonds contribute to this connection (reading from the right to the left): \((ij) = (14), (23)\). Therefore, we define \(M_{14} (\mathbf{\tau}_1) = M_{23} (\mathbf{\tau}_1) = 1\) and \(M_{ij} (\mathbf{\tau}_1) = 0\) otherwise. In a similar way, one finds \(M_{13} (\mathbf{\tau}_2) = 1, M_{12} (\mathbf{\tau}_3) = M_{43} (\mathbf{\tau}_3) = 1\) and \(M_{ij} (\mathbf{\tau}_2) = M_{ij} (\mathbf{\tau}_3) = 0\) otherwise. With this procedure, the other five matrices \(\mathbf{M}_{ij} (\mathbf{\tau})\) can be easily found as well.

Now we solve the intra-cluster eigenvalue problem by diagonalizing the Hamiltonian \(H^{(\text{intra})}_{t-V} (1)\) from 30 separately in each \(n\)-particle \((n=0,1,\ldots,4)\) sector

\[
H^{(\text{intra})}_{t-V} (1) |\Psi^{(n)} > > 1 = E^{(n)}_\nu |\Psi^{(n)} > > 1
\]

(33)

The empty state \(|\Psi^{(0)}_\nu > > 1\) with energy \(E^{(0)}_\nu > > 1 = 0\) defines the cluster vacuum state \(|0 > > 1\). There are four singly occupied cluster states \(|\Psi^{(1)}_\nu > > 1\) with \(\nu = 1, \ldots, 4\) which are found to be

\[
|\Psi^{(1)}_\nu > > 1 = \sum_{i=1}^4 \beta^{(1)}_{\nu i} c^\dagger_{i1} |0 > > 1 = f^\dagger_{\nu i} |0 > > 1
\]

(34)

where \(\beta_{\nu i}\) is a \((4 \times 4)\)-matrix

\[
\beta_{\nu i} = \frac{1}{2} \begin{pmatrix}
1 & 1 & 1 & 1 \\
1 & -1 & 1 & -1 \\
\sqrt{2} & 0 & -\sqrt{2} & 0 \\
0 & \sqrt{2} & 0 & -\sqrt{2}
\end{pmatrix}
\]

(35)
and the corresponding eigenvalues are

\[ E_1^{(1)} = -3t; \quad E_2^{(1)} = E_3^{(1)} = E_4^{(1)} = t \]  

(36)

The eigenvectors \( \mathbf{C} \) are basis vectors of different irreducible representations of \( C_{4h} \) point group: the case \( \nu = 1 \) describes the lowest energy fully symmetric \((a_1)\) solution, \( \nu = 2 \) belongs to \( b_g \) and \( \nu = 3, 4 \) to \( e_u \) representation. Below we will refer to these 1-particle solutions as cluster 'orbitals'. According to \( \mathbf{C} \), for a given cluster 1 the transformation from the original site operators \( c_{i1}^l \) \((i=1,...,4)\) to the cluster 'orbital' operators \( f_{i1}^l \) is given by the matrix \( \beta_{\nu \nu'} \) defined in \( \mathbf{C} \). Note that the \( f_{i1}^l \) \((f_{i1})\) operators anticommute. The cluster states of higher occupancy \( n=2,3,4 \) can be understood as a result of successive filling of cluster 'orbitals'. For instance, for \( n=2 \) one obtains six eigenstates (the cluster index 1 is dropped):

\[ |\Psi_{1,2,3}^{(2)}\rangle = f_2^1 f_1^1 f_1^1 f_1^1 |0\rangle, \quad |\Psi_{4,5,6}^{(2)}\rangle = f_4^1 f_2^1 f_1^1 f_1^1 |0\rangle \]  

(37)

with corresponding energies

\[ E_{1,2,3}^{(2)} = -2t + V; \quad E_{4,5,6}^{(2)} = 2t + V \]  

(38)

In the cluster 'orbital' representation, the terms \( H_1^{(\text{intra})} \) and \( H_2^{(\text{intra})} \) in \( \mathbf{C} \) can be now written as follows

\[ H_1^{(\text{intra})} = \sum_{i} \left(-3t f_{i1}^1 f_{i1}^1 + t \sum_{\nu} f_{i\nu}^1 f_{i\nu}^1 \right) \]  

(39)

\[ H_2^{(\text{intra})} = \frac{V}{2} \sum_{\nu} \sum_{\nu'} \sum_{\nu''} \sum_{\nu'''(\nu'') n_{\nu1} n_{\nu1}'}. \]  

(40)

From the result \( \mathbf{C} \), one can see that \( H_2^{(\text{intra})} \) is the Hubbard term in an effective, 'multi-orbital' electronic model. Such a model is derived below by adding to \( \mathbf{C} \) and \( \mathbf{C} \) the intercluster hopping \( H_1^{(\text{inter})} \) and the Coulomb \( H_2^{(\text{inter})} \) terms and by using for the latter a mean-field approximation, i.e., \( n_{i,1+\tau} n_{j,1} \simeq \langle n_{i,1+\tau} \rangle n_{j,1} + n_{i,1+\tau} \langle n_{j,1} \rangle - \langle n_{i,1+\tau} \rangle \langle n_{j,1} \rangle \). The approximated term \( H_1^{(\text{inter})} \) reads

\[ H_1^{(\text{inter})} = \sum_{\{\tau\}} \sum_{\nu,\nu'} \sum_{\nu'',\nu'''} T_{\nu\nu'}^{\{\tau\}}(\nu') f_{\nu+\tau}^1 f_{\nu'1}. \]  

(42)

The \((4 \times 4)\)-matrices \( T_{\nu\nu'}^{\{\tau\}} \) are related to the \( M_{ij}^{\{\tau\}} \) by a rotation

\[ T_{\nu\nu'}^{\{\tau\}}(\nu') = -t \sum_{ij} \beta_{\nu i} M_{ij}^{\{\tau\}} \beta_{\nu' j}, \]  

(43)

because \( (\beta^{-1})_{\nu'} = \beta_{\nu' j} \). Finally, by collecting the contributions \( \mathbf{C} \), we obtain an effective 'multi-orbital' Hubbard-like Hamiltonian

\[ H_{\text{HUB}} = H_1^{(\text{inter})} + H_2^{(\text{intra})} + H_1^{(\text{inter})} + H_2^{(\text{intra})}, \]  

(44)

The effective Hamiltonian \( \mathbf{C} \) has a lower symmetry compared to that of the original Hamiltonian (1). This may lead to an artificial low-symmetry ground-state solution, for instance, to a long-range charge/bond ordering. Here we suggest, however, that the geometrical frustration of the checkerboard lattice prevents this kind of long-range ordering, as was discussed already in the beginning of this section. Therefore we supplement the effective Hamiltonian \( \mathbf{C} \) with additional restrictions, which prevent an artificial low-symmetry solution of \( \mathbf{C} \). We require: (a) a homogeneous
electron distribution over the lattice sites, \( \langle c_i^\dagger c_i \rangle = n \), with \( n \) being the electron concentration, and (b) a bond-independence of hopping amplitudes, \( \langle c_i^\dagger c_j \rangle = k \), where a pair of sites \( i, j \) denotes a bond (there are six bonds within each plaquette). Note that \( k \) is a real \( n \)-dependent quantity to be calculated self-consistently. On general grounds, one expects that \( k > 0 \) in a metallic phase and \( k = 0 \) in an insulating phase. Both requirements (a) and (b) together impose restrictions on cluster ‘orbital’ averages

\[
\langle f_{\nu 1}^\dagger f_{\nu 1} \rangle = \sum_{ij} \beta_{\nu i} \beta_{\nu j} \langle c_i^\dagger c_j \rangle = n \sum_i \beta_{\nu i} \beta_{\nu i} + k \left( \sum_i \beta_{\nu i} \right) \left( \sum_j \beta_{\nu j} \right) - \sum_i \beta_{\nu i} \beta_{\nu i} \tag{45}\]

By using that \( \sum_i \beta_{\nu i} \beta_{\nu i} = \delta_{\nu \nu'} \) and

\[
\sum_i \beta_{\nu i} = \begin{cases} 2; & \nu = 1 \\ 0; & \nu = 2, 3, 4 \end{cases} \tag{46}\]

one obtains for \( \nu = \nu' \) the following relations for cluster ‘orbital’ occupancies

\[
\langle f_{\nu 1}^\dagger f_{\nu 1} \rangle = n + 3k, \ (\nu = 1); \ \langle f_{\nu 1}^\dagger f_{\nu 1} \rangle = n - k, \ (\nu = 2, 3, 4). \tag{47}\]

If \( \nu \neq \nu' \), the expression (45) leads to

\[
\langle f_{\nu 1}^\dagger f_{\nu 1} \rangle = 0, \ (\nu 
eq \nu'). \tag{48}\]

For an isolated cluster with an integer electron occupancy equation (48) is obviously fulfilled because of symmetry arguments.

Based on the effective model (44) we calculate the electronic band structure from the Fourier transformation \( G_{\nu \nu'} (q, \omega) \) of the retarded matrix Green’s function:

\[
G_{\nu \nu'} (1 - 1', t - t') = \left\langle \{ f_{\nu 1}(t) , f_{\nu 1}^\dagger (t') \} \right\rangle = -i \theta (t - t') \left\langle \{ f_{\nu 1}(t) , f_{\nu 1}(t') \} \right\rangle. \tag{49}\]

To obtain the equation of motion for \( G_{\nu \nu'} (q, \omega) \), we use the method of the two-time ‘irreducible’ Green’s function. A successive differentiation of (49) with respect to both times \( t \) and \( t' \) with the use of properly defined projection procedure lead to the Dyson’s equation:

\[
\left[ \omega \hat{1} - \hat{\Omega} (q) - \hat{\Sigma} (q, \omega) \right] \hat{G} (q, \omega) = \hat{1}. \tag{50}\]

Here \( \hat{1}, \hat{\Omega} (q) \) and \( \hat{\Sigma} (q, \omega) \) are the unit, a frequency and a self-energy \((4 \times 4)\)-matrices, respectively. To define \( \hat{\Omega} \) and \( \hat{\Sigma} \) explicitly, it is convenient to introduce a notion of a scalar product of two fermionic operators \( A \) and \( B \) as \( \langle A | B^\dagger \rangle = \langle \{ A , B^\dagger \} \rangle \). In this notation, the matrix elements of \( \hat{\Omega} \) are

\[
\Omega_{\nu \nu'} (q) = \left\langle i f_{\nu q} | f_{\nu' q}^\dagger \right\rangle = \left\langle \{ f_{\nu q} , H_{1 - v} \} , f_{\nu' q}^\dagger \right\rangle, \tag{51}\]

resulting in

\[
\Omega_{\nu \nu'} (q) = \delta_{\nu \nu'} E_{\nu}^{(1)} + T_{\nu \nu'} (q) + \delta_{\nu \nu'} V \left( \sum_{\nu_i (\neq \nu)} \langle n_{\nu_i} \rangle + \frac{3}{4} \langle N_c \rangle \right). \tag{52}\]

Here the \( E_{\nu}^{(1)} \) are given by (30) and

\[
T_{\nu \nu'} (q) = \sum_{\{\tau\}} e^{i q \cdot \tau} T_{\nu \nu'} (\tau). \tag{53}\]

The frequency matrix \( \hat{\Omega} \) provides for a mean-field description of the electronic band structure. In order to include effects of electron correlations, the self-energy part \( \hat{\Sigma} \) must be calculated.
For this purpose, the set of basis operators \( f_{\nu q} \) is complemented with a new set of operators \( F_{\nu q} (\nu = 1, \ldots, 4) \):

\[
F_{\nu q} = i \dot{f}_{\nu q} - \sum_{\nu'} \Omega_{\nu \nu'} (q) f_{\nu' q},
\]

which are orthogonal to \( f_{\nu q} \), i.e. \( \langle f_{\nu q} | f_{\nu' q}^\dagger \rangle = 0 \). Then, the self-energy \( \Sigma \) has the form of an 'irreducible' matrix Green’s function \( \Sigma \) whose elements are defined by

\[
\Sigma_{\nu \nu'} (q, \omega) = \langle \langle F_{\nu q} F_{\nu' q}^\dagger \rangle \rangle_\omega - \sum_{\nu_1 \nu_2} \langle \langle F_{\nu q} F_{\nu_1 q}^\dagger \rangle \rangle_\omega \frac{1}{\langle \langle f_{\nu_1 q} f_{\nu_2 q}^\dagger \rangle \rangle_\omega} \langle \langle f_{\nu_2 q} F_{\nu' q}^\dagger \rangle \rangle_\omega.
\]

The irreducible Green’s function matrix \( \langle \langle F_{\nu q} F_{\nu' q}^\dagger \rangle \rangle_\omega \) obeys the following equation (the lower indices are omitted for brevity):

\[
\omega \langle \langle F | F^\dagger \rangle \rangle_\omega = \langle F | F^\dagger \rangle + \left\{ \langle \dot{F} | F^\dagger \rangle + \langle i F | -i F^\dagger \rangle \right\}_\omega \frac{1}{\langle \langle F | F^\dagger \rangle \rangle_\omega} \langle \langle F | F^\dagger \rangle \rangle_\omega.
\]

We truncate this equation by neglecting in (56) the next-order 'irreducible' matrix Green’s function \( \langle \langle F | F^\dagger \rangle \rangle_\omega \). For the effective 'multi-orbital' model \( [4] \), the truncation corresponds to the first step of the Hubbard-I approximation and results in the following form of \( \Sigma \):

\[
\Sigma_{\nu \nu'} (q, \omega) = \left[ \omega \hat{\Lambda} (q) \right]_{\nu \nu'}^{-1} \langle \langle F_{\nu q} F_{\nu' q}^\dagger \rangle \rangle_\omega,
\]

where the frequency matrix \( \hat{\Lambda} (q) \) is given by

\[
\Lambda_{\nu \nu'} (q) = \frac{\langle i F_{\nu q} | F_{\nu' q}^\dagger \rangle}{\langle F_{\nu q} F_{\nu' q}^\dagger \rangle}.
\]

In the Hubbard-I approximation applied to the standard Hubbard model, the corresponding self-energy is \( q \)-independent. We shall apply a similar approximation here by dropping the intercluster hopping term \( H_{12}^{\text{inter}} \) which appears in the equation \( i F_{\nu q} = [F_{\nu q}, H_{12}] \). In the following, \( \hat{\Lambda} \) and \( \bar{\Sigma} \) are \( q \)-independent matrices and the latter is of the form

\[
\Sigma_{\nu \nu'} (\omega) = \delta_{\nu \nu'} \frac{\langle F_{\nu 1} | F_{\nu 1}^\dagger \rangle}{\omega - \langle i F_{\nu 1} | F_{\nu 1}^\dagger \rangle / \langle F_{\nu 1} | F_{\nu 1}^\dagger \rangle}.
\]

With the definition (57), one obtains explicitly

\[
F_{\nu 1} = V \delta N_{1}^\nu f_{\nu 1},
\]

where

\[
\delta N_{1}^\nu = N_{1}^\nu - \langle N_{1}^\nu \rangle, \quad N_{1}^\nu = N_{1} - n_{\nu 1} = \sum_{\nu_1 (\neq \nu)} n_{\nu_1 1}.
\]

Then the diagonal elements of the frequency matrix entering into the denominator of (59) are found to be

\[
\Lambda_{\nu \nu} = \frac{\langle i F_{\nu 1} | F_{\nu 1}^\dagger \rangle}{\langle F_{\nu 1} | F_{\nu 1}^\dagger \rangle} = E_{\nu}^{(1)} + \frac{3}{4} V \langle N_{c} \rangle + V \frac{\langle N_{1}^\nu (\delta N_{1}^\nu)^2 \rangle}{\langle (\delta N_{1}^\nu)^2 \rangle}.
\]
FIG. 6: Energy bands of spinless fermions on the checkerboard lattice calculated in the cluster approach for different values of inter-site repulsion $V/t = 0, 0.3 < V_c/t$ and $V/t = 10 > V_c/t \simeq 4$. The bands are unfolded along the highly symmetrical directions in the square Brillouin zone; energy unit is $t \equiv 1$. The electron concentration is chosen as $n = \frac{1}{2}$ (equivalent to $\langle N_c \rangle = 2$) and the chemical potential $\mu$ is fixed at zero energy.

Both averages, $\langle (\delta N^\nu) \rangle$ and $\langle N^\nu_1 (\delta N^\nu_1) \rangle$, are approximated in a mean-field manner. For instance, in this approximation, the mean value $\langle (\delta N^\nu) \rangle$ which describes the intracluster charge correlations, reads

$$
\langle (\delta N^\nu) \rangle = \sum_{\nu_1 (\neq \nu)} \langle n_{\nu_1} \rangle (1 - \langle n_{\nu_1} \rangle).
$$

We summarize the results for the self-energy as follow ($\langle n_{\nu} \rangle = \langle n_{\nu} \rangle$):

$$
\Sigma_{\nu\nu} (\omega) = \frac{\sum_{\nu_1 (\neq \nu)} \langle n_{\nu_1} \rangle (1 - \langle n_{\nu_1} \rangle)}{\omega - \Lambda_{\nu\nu}},
$$

(63)

$$
\Lambda (\omega) = E^{(1)}_\nu + \frac{3}{4} V \langle N_c \rangle + V \frac{\sum_{\nu_1 (\neq \nu)} \langle n_{\nu_1} \rangle (1 - \langle n_{\nu_1} \rangle)^2}{\sum_{\nu_1 (\neq \nu)} \langle n_{\nu_1} \rangle (1 - \langle n_{\nu_1} \rangle)} + V \frac{\sum_{\nu_1 (\neq \nu)} \sum_{\nu_2 (\neq \nu, \nu_1)} \langle n_{\nu_2} \rangle (1 - \langle n_{\nu_2} \rangle) \langle n_{\nu_1} \rangle (1 - \langle n_{\nu_1} \rangle) \langle n_{\nu_2} \rangle}{\sum_{\nu_1 (\neq \nu)} \langle n_{\nu_1} \rangle (1 - \langle n_{\nu_1} \rangle)}.
$$

(64)

In this expression the second term is the inter-cluster Hartree-Fock correction. The third and fourth terms are intra-cluster Hubbard-I type correlation corrections, where the latter appears only for the present ‘multi-orbital’ Hubbard Hamiltonian. In the common Hubbard model (with just one ‘orbital’) only the third correction would be present. To perform self-consistent band-structure calculations, a chemical potential $\mu$ is introduced in a standard way. For
a given spinless-fermion concentration $n$ varying within the range $0 \leq n \leq 1$, the value of $\mu$ is determined from the equation

$$n = \frac{1}{4} \langle N_c \rangle = \frac{1}{4} \sum_{\nu=1}^{4} \langle n_{\nu} \rangle,$$

(65)

where $\langle n_{\nu} \rangle$ is an average 'orbital' cluster occupancy

$$\langle n_{\nu} \rangle = \frac{1}{M} \sum_{q} \int_{-\infty}^{\mu} d\omega \left( -1/\pi \right) \text{Im} G_{\nu\nu}(q, \omega).$$

(66)

Here the summation is over $M = L/4$ $\mathbf{q}$-vectors in the first Brillouin zone of the square lattice formed by the basic clusters in the checkerboard lattice. The 'orbital' occupancies should obey the relations [17].

The most representative results of the self-consistent band structure calculations are shown in Figs. 6 and 7. In Fig. 6, band spectra for different values of coupling $V$ are given along highly symmetrical directions in the square Brillouin zone. To show the insulating gap opening in the band spectrum with increasing $V$, the electron concentration $n=1/2$ corresponding to an integer mean cluster occupancy $\langle N_c \rangle=2$ is chosen; the chemical potential $\mu$ is located at zero energy. From the upper panel, a doubly degenerate completely flat branch on the top of two highly dispersive bands is seen for the non-interacting case, $V=0$. For finite coupling $V$, but less than some critical value $V < V_c$, these four branches are split and the chemical potential intersects one of the dispersive branches as shown in the middle panel of Fig. 6. In this weakly correlated regime of the model one expects a metallic state of the system. We found that starting from $V = V_c \approx 4t$, the spectrum is clearly split into the low- and high-energy Hubbard subbands separated with a gap growing as $V$ increases above $V_c$. For a given concentration $\langle N_c \rangle=2$ and $V > V_c$, the four lowest Hubbard subbands are filled completely and located below the chemical potential $\mu$ as shown for $V = 10$ in the lower panel of Fig. 6. If one includes a weak next neighbor hopping $t'$, the former flat energy branches acquire a small dispersion, the picture of the insulating gap opening still retains at slightly changed value of the critical coupling $V_c$.

In this paper, however, we restrict ourselves to the study of the limit, $t'=0$. In this limit, to avoid discussing a rather special case of the partially filled upper flat band, the electron concentration is chosen to vary in the range $n \leq 1/2$. Within this range, for strong coupling $V$ the model displays an insulating state at the integer cluster occupancy $\langle N_c \rangle=1$ as well. This can be easily seen from Fig. 7, where the calculated mean cluster occupancy $\langle N_c \rangle$ is depicted as a function of the chemical potential $\mu$ for $V=10$. The charge compressibility $d \langle N_c \rangle / d\mu$ is found to be

![FIG. 7: Mean cluster occupancy $\langle N_c \rangle$ calculated as a function of the chemical potential $\mu$ at $V/t = 10$. Two wide flat regions with $d\langle N_c \rangle/d\mu = 0$ at $\langle N_c \rangle =1,2$ indicate an insulating state of the system.](image-url)
zero in a wide range of varying $\mu$ at both integer occupancies $\langle N_c \rangle = 1, 2$. Near the occupancy value $\langle N_c \rangle = 1.5$, the sharp change of $\langle N_c \rangle$ is connected to the fact that $\mu$ intersects the flat band peculiarity in the density of states.

IV. CONCLUSION

In this work we have studied an extended Hubbard type spinless fermion model for the frustrated checkerboard lattice which is a 2D analogue for pyrochlore or spinel-type lattices. We have studied the possibility of a metal-insulator phase transition as function of the inter-site Coulomb interaction and the band filling. We have used single-site and cluster representations of the model. In the latter the intra-cluster correlations are accounted for exactly by transforming to a 'cluster orbital' basis. Within a Green’s function approach decouplings on Hartree-Fock and 'Hubbard-I' type level for the inter-site Coulomb term have been employed. In both approaches we find a metal-insulator transition for increasing $V$ for half filling (that is one fermion per two sites) and in the cluster approach even for quarter filling. The MI transition is of the Mott Hubbard-type and is associated with a gap opening in the quasiparticle excitations. The critical interaction for the MI transition is $V_c = \frac{4.86}{t}$ in the single site approach and $V_c \approx 4$ in the cluster approach are in reasonable agreement. We also consider the possibility of charge ordering within the single-site approach where we find a transition to a staggered CO state with $Q = (0,0)$ at $V_{c2}/t = 6.47$. This state may be a result of the approximations employed since for $V/t \to \infty$ CO has to vanish due to the macroscopic degeneracy of the ground state. Stabilisation of CO may indeed require the coupling to the lattice to lift this degeneracy. Further progress in understanding the nature of the insulating state caused by the inter-site Coulomb interaction may require the use of more advanced methods like cluster dynamical mean field methods which has sofar not been achieved.

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