One-dimensional Hubbard model at quarter filling on periodic potentials

C. Schuster and U. Schwingenschlögl
Institut für Physik, Universität Augsburg, 86135 Augsburg, Germany
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Using the Hubbard chain at quarter filling as a model system, we study the ground state properties of highly doped antiferromagnets. In particular, the Hubbard chain at quarter filling is unstable against 2\(k_F\)- and 4\(k_F\)-periodic potentials, leading to a large variety of charge and spin ordered ground states. Employing the density matrix renormalization group method, we compare the energy gain of the ground state induced by different periodic potentials. For interacting systems the lowest energy is found for a 2\(k_F\)-periodic magnetic field, resulting in a band insulator with spin gap. For strong interaction, the 4\(k_F\)-periodic potential leads to a half-filled Heisenberg chain and thus to a Mott insulating state without spin gap. This ground state is more stable than the band insulating state caused by any non-magnetic 2\(k_F\)-periodic potential. Adding more electrons, a cluster-like ordering is preferred.

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I. HOLE DOPED ANTIFERROMAGNETS

The study of low-dimensional doped antiferromagnets was triggered by investigations of the phase diagram of high temperature superconductors. Upon doping, they show a transition from an antiferromagnetic to a superconducting state. On the other hand, the investigation of one-dimensional compounds is also an active field. A simple form of low doping in a Cu-chain is the replacement of a few Cu ions by non-magnetic ions like Zn. In this case, the system is well described by the Heisenberg model with some spins removed [1]. The charge and spin order of highly doped compounds, on the contrary, are not understood by far. A prototypical material revealing strongly doping dependent charge order and antiferromagnetism in a less than half-filled band is the spin chain compound \((\text{Ca,Sr})_{14}\text{Cu}_{24}\text{O}_{41}\). This composite crystal contains two different structural components along the c-axis: \(\text{Cu}_2\text{O}_3\) ladders and \(\text{Cu}_2\text{O}_2\) chains. Whereas the ladders contain only \(\text{Cu}^{2+}\)-ions, nearly all \(\text{Cu}^{3+}\)-ions are located on the chains [2]. Substitution of calcium on the strontium sites leads to a transfer of holes from the chains into the ladders. For the Ca-rich compounds one generally assumes one hole per formula unit on the ladders and five on the chains. Near the quarter-filled band antiferromagnetic order is established [2].

DFT-LDA calculations [3], using the ASW method [5], reveal that the ladder and chain substructures can be treated individually. In particular, the density of states of the chains shows a partially filled valence band, as illustrated in Fig 1. The origin of antiferromagnetism in these compounds is still unclear, and the doping dependence of the charge order and spin gap is discussed controversially [6]. Recent studies on the basis of the Heisenberg model show that a simple spin model is not appropriate to describe the magnetization [7], indicating the necessity of studying more advanced models, as e.g. the Hubbard model.

In this article we address the stability of the proposed charge order (cluster-like versus staggered) [8] in \((\text{Ca,Sr})_{14}\text{Cu}_{24}\text{O}_{41}\), considering the competition between kinetic energy, interaction and periodic distortions, on the basis of the Hubbard model. Since no periodic lattice distortion – neither with two-site nor with four-site periodicity – is found in the crystal structure [9] we consider only periodic potentials. Nevertheless, our results are relevant for all correlated materials with quarter filled bands. Thus we aim at understanding a whole class of materials on a common basis.
II. PERIODIC DISTORTIONS IN INTERACTING CHAINS

Charge order – independent of magnetic ordering – in less than half-filled bands is found, for example, in organic charge transfer salts, like the Bechgaard salts (TMTSF)$_2$X, with X=PF$_6$, AsF$_6$, ClO$_4$, ReO$_4$, or Br, or their sulfur analogs (TMTTF)$_2$X. These systems, which consist of stacks of organic molecules forming weakly coupled one-dimensional chains, exhibit a large variety of low temperature phases [10, 11]. They have been studied using a purely electronic model, namely the extended Hubbard model with next-nearest-neighbor interaction. Recent results have been obtained by mean-field approaches [12] and exact diagonalization [13]. On the other hand, charge order is often connected to a structural transition and hence to electron-phonon coupling. The coupling of a one-dimensional metal to an elastic lattice results in an instability towards a periodic lattice distortion commensurate to the band-filling, known as Peierls transition. Within a lattice model, the Peierls transition is captured by a modulated hopping term in the Hamiltonian of the non-interacting system [14]. A periodic potential, instead of a modulated hopping, has similar effects on the electronic structure of the one-dimensional metal, leading to a band insulator.

The interplay of electron-lattice and electron-electron interaction is conveniently studied in the framework of the Hubbard model. With interaction, the Peierls and the so-called ionic model show different ground-state phase diagrams [10]. Many results are available for the half-filled Hubbard model with periodic hopping or potential, for which the transition from a band insulator to a Mott insulator has been studied in [20, 21, 22, 23, 24]. On the other hand, away from half filling no insulator-insulator-transition is present, but a variety of ordering processes have to be taken into account [22]. The stability of the different ground states of the quarter-filled Hubbard model with respect to a periodic potential has been investigated [26, 28] less extensively as compared to the half-filled model. The phase diagram of the quarter-filled Peierls-Hubbard model is discussed in [29]. Hence, we restrict our calculations to the ionic potentials.

We access the stability regions of the ionic Hubbard model by introducing several potentials connected with different ordering processes of charge and spin degrees of freedom into the Hamiltonian. In particular, we investigate which potential leads to the largest energy gain, and analyze against which potential or order the system is most unstable, indicating which order will be established in nature. In our calculations, we concentrate on the ionic Hubbard model at quarter filling, for both weak and strong interaction. We determine the ground state and the ground state energy as a function of interaction, doping and potential strength, using the density matrix renormalization group method [30]. Thus, our calculations give detailed insights into the properties of hole-doped antiferromagnets.

III. MODELS

In the following section we first discuss the properties of non-interacting electrons on a chain with different periodic potentials and the phase diagram of the homogeneous Hubbard chain. In the last subsection we discuss some basic features of the periodic Hubbard model.

A. Peierls model

The Peierls model is the prototypical model to study the coupling of non-interacting electrons to the lattice. The Hamiltonian is given by

$$H_{\text{Peierls}} = -\sum_{i,\sigma} t_i \left( c_{i,\sigma}^+ c_{i+1,\sigma} + \text{h. c.} \right) , \quad (1)$$

where we consider a modulated hopping term with $t_i = t[1 + u \cos(Qa \cdot i)]$. $N$ denotes the number of lattice sites and $a$ the lattice constant. In addition, $N_e$ is the number of electrons on the chain and $Q$ the wave vector of the periodic distortion. In case of a commensurate distortion we have $Q = 2mk_F$, $m = 1, 2, \ldots$. The dimerized chain, $Q = \pi/a$, at half filling, $n = N_e/N = 1$, was investigated by Su, Schrieffer, and Heeger (SSH) [14]. Due to the periodic distortion a band gap, $\Delta = 4ut$, opens at $k_\Delta = Q/2$. Since at half filling we have $k_\Delta = k_F$, the system is insulating. As is characteristic for a band insulator, the gaps for charge and spin excitations are identical. In the following, we concentrate on the quarter-filled band and 4a-periodic distortions, because in case of the 2a-periodic distortion the quarter-filled system is metallic, since $k_\Delta \neq k_F$.

To begin, we determine the energy bands and the energy gain due to a periodic modulation of the hopping parameter with $Q = \pi/2a$. For quarter filling, $k_F = \pi/4a$, this system shows the Peierls transition, since $Q = \pi/2a = 2k_F$. The Fourier transformation of the hopping term yields, in analogy with the SSH-model, the expression

$$H_{\text{Peierls}} = \sum_k \left( c_k^+ c_{k+Q}^+ c_{k+2Q}^+ c_{k+3Q}^+ \right) \tilde{h}_{\alpha\beta} \begin{pmatrix} c_k \\ c_{k+Q} \\ c_{k+2Q} \\ c_{k+3Q} \end{pmatrix} , \quad (2)$$

where the matrix $\tilde{h}_{\alpha\beta}$ is given by
The eigenvalues of this matrix are

\[
\epsilon(k) = \pm t \sqrt{2 \left( 1 + 2u^2 \pm \sqrt{\cos^2 2ka + 4u^2(1 + \sin^2 2k)} \right)}, \tag{4}
\]

compare \[31\]. The four energy bands are separated by the energy gaps

\[
\Delta_1 = 4ut, \quad \Delta_2 = 2u^2t, \quad \text{and} \quad \Delta_3 = \Delta_1, \tag{5}
\]

where \(\Delta_1 = \varepsilon_2(k = k_F) - \varepsilon_1(k = k_F)\) and \(\Delta_2 = \varepsilon_3(k = 2k_F) - \varepsilon_2(k = 2k_F)\). The energy gain of the fermions, for the quarter-filled case, obtained by summing up the contributions of all occupied states, is found to be \(\propto u^2t \ln u\), as for half filling.

### B. Ionic model

Another type of periodic distortion is given by local potentials, resulting in the so-called ionic model

\[
H_{\text{ionic}} = -t \sum_{i,\sigma}^{N} (c_{i,\sigma}^{+} c_{i+1,\sigma} + \text{h. c.}) + \sum_{i}^{N} W_{i} n_{i} - \sum_{i}^{N} h_{i} S_{i}^{z}. \tag{6}
\]

Again, for the half-filled band the same calculation as before can be performed for a periodic potential, with \(\Delta_i = \Delta_3 = 2W\). In case of the ionic model for a quarter-filled system we compare several potentials, which correspond to different charge and spin patterns:

a) First, we study a simple potential with \(2k_F(4a)\)-period, given by \(W_{i} = W_{\cos(\pi i/2)}\). Here, the charge order for zero hopping is given by \(1/0.5/0.5\) electrons per site, periodically continued. Spin order is not established.

b) A modification of a) given by \(W_{i} = W_{\cos(\pi i/2)} + W_{\cos(\pi (i+1)/2)}\), which leads to two occupied and two unoccupied sites (“cluster”) \[33\], i. e. a \(1/1/0/0\) charge order.

c) A charge order with a \(1/0/1/0\) pattern, i. e. a \(4k_F(2a)\)-periodic order, is forced by a potential with \(W_{i} = W_{\cos(\pi i)}\). With interaction we likewise expect magnetic ordering.

d) In addition, a local magnetic field with \(h_{i} = W_{\cos(\pi i/2)(n_{i,\uparrow} - n_{i,\downarrow})}\) enforces a \(\uparrow/0/\downarrow/0\) pattern. As in c), we expect a \(2k_F(4a)\)-period for the spins and correspondingly a \(4k_F(2a)\)-period for the charges.

Diagonalizing the resulting Hamilton matrix, as described above, yields the dispersions

\[
a) \text{d)} \quad \epsilon(k) = \pm t \sqrt{2 + w^2/2 \pm \sqrt{4 \cos^2(2k) + 2w^2 + w^4/4}},
\]

\[
b) \epsilon(k) = \pm t \sqrt{2 + w^2/2 \pm \sqrt{4 \cos^2(2k) + 4w^4}},
\]

\[
c) \epsilon(k) = \pm t \sqrt{w^2 + 4 \cos^2 k},
\]

with \(w = W/t\). Fig. 2 shows the dispersion for potentials a) and c). In comparison to the Peierls model not only a gap opens at \(k = Q/2\) but the lower and upper bands are shifted down and up, respectively. In case of a small perturbation, the dispersion is similar for the Peierls and the ionic model, as well as for potentials a) and b). Potential a) and d) are equivalent in the non-interacting case.

The energy gain in the quarter filled band is found to be about \(-W^{1.65}\) in cases a), b), and d). For small \(W\), the energy gain here is nearly the same in case a) and b), whereas it becomes weaker in case b) than in case a) at \(W \approx 0.5t\). In case c), the energy gain is quadratic, \(E(W) - E(0) = \frac{8}{7}W^2\). It mainly traces back to the band shift.

Accordingly, the gaps are given by

\[
a) \text{d)} \quad \Delta_1 = W, \quad \Delta_2 = \sqrt{2}W, \quad \Delta_3 = W;
\]
b) $\Delta_1 \sim W$ for small $W$, $\Delta_1 \to \Delta_1^c$ for strong $W$, $\Delta_2 = \sqrt{2} - W/2$, $\Delta_3 = \Delta_1$;

c) $\Delta_1 = 0$, $\Delta_2 = W$, $\Delta_3 = 0$.

C. Hubbard model

The Hubbard model is known to capture the interplay between kinetic energy (delocalization) and interaction (localization) in electronic systems. The Hamiltonian is given by

$$H_{\text{Hubb}} = -\sum_{i,\sigma} t_i \left(c_{i,\sigma}^+ c_{i+1,\sigma} + \text{h.c.}\right) + U \sum_i n_{i,\uparrow} n_{i,\downarrow}. \quad (7)$$

The Hubbard model in one dimension is exactly solvable by means of the Bethe ansatz [15]. Note that in one dimension another useful formulation of the Hubbard model is available on the basis of the bosonization technique. The low-lying excitations of the non-interacting as well as the interacting fermions system are sound waves, i.e., the Fermi system can be described as a non-interacting Bose system, called a Luttinger liquid, showing spin-charge separation. In the clean case, the Hubbard model has three phases. For $U < 0$, the spin excitation spectrum has a gap and the low-lying charge excitations can be described by those of a Luttinger liquid. For $U > 0$ and away from half filling, spin and charge excitations are those of a Luttinger liquid. The last phase occurs for $U > 0$ and half filling, where the charge excitations have a gap and the spin excitations are of Luttinger type. A relevant $4k_F$-Umklapp scattering term, only present for half filling, is responsible for the Mott gap in the latter phase. The bosonization technique is adequate for metallic systems or in the weak coupling regime. It is useful to determine the phase boundary between metals and insulators but it is not suitable for distinguishing different insulating phases for intermediate or strong perturbations.

D. Periodic Hubbard model

A commensurate periodic distortion – i.e., commensurate to the band filling – introduces an additional nonlinear term in the bosonized Hamiltonian, which couples spin and charge degrees of freedom and destroys the integrability of the clean Hubbard model. In the half-filled case we therefore find a transition between two possible insulating phases, i.e., the band (Peierls) insulator ($W, ut \gg U$) with charge and spin gap of the same order and the Mott insulator ($W, ut \ll U$) with a charge gap proportional to the interaction, but vanishing spin gap, where periodic distortion and interaction oppose each other. However, the transition is fundamentally different for the Peierls and the ionic Hubbard model [10, 17]. In the Peierls model the transition between the two phases is smooth and continuous [18], whereas a third phase – a spontaneous dimerized insulating phase – is found in the ionic Hubbard model. Within this phase, charge and spin gap are non-zero, and the dimerization operator has a non-zero expectation value. In fact, the extension of this phase is discussed controversially.

The situation in the quarter filled band is completely different. With interaction, no insulating phase is established, except for very strong interaction, where higher order terms of the Umklapp scattering can cause a Mott insulator on its own [26]. In the quarter-filled case with $2k_F$-periodic potential we expect from bosonization a transition from the Luttinger liquid to the band insulator in the non-interacting as well as in the interacting system. The interaction modifies the exponents of the gaps and the energy gain. Since they become smaller with increasing interaction, interaction stabilizes the band insulator. Here, the interplay between periodic distortion and interaction is weak, since the quarter filled model with interaction is still a metal. A similar situation is found for half filled spinless fermions for weak interaction, where the Umklapp scattering induces the insulator at finite interaction strength [25]. In case of a $4k_F$-periodic potential Umklapp scattering becomes relevant [28] due to the doubling of the unit cell or, equivalently, due to the reduction of the Brillouin zone, see the straight line in Fig. 1. In this case, the system undergoes a transition from the Luttinger liquid to a Mott insulating phase when the potential and the interaction are present. Potential and interaction hence cooperate rather than oppose each other as in the half filled ionic model.

An analysis of the ionic Hubbard model with another $2k_F$-periodic potential is given in [19]. The authors find the band insulator for weak interaction but strong potential. Our $4k_F$-periodic pattern, hence the Mott insulator, is obtained in their model in the limit of strong interaction. Both phases are separated by a bond ordered phase. Thus, the model potential used in [19] reveals the features of the half filled ionic Hubbard model. The analysis, however, concentrates on large $|U - W|$. In our case, the $4k_F$-periodic pattern is not a limit of the $2k_F$-periodic patterns.

In the following, we concentrate on the ionic Hubbard model at weak to intermediate potential strength and intermediate to strong interaction. In addition, we study the region of stability for each potential.

IV. NUMERICAL RESULTS FOR THE IONIC HUBBARD MODEL AT QUARTER FILLING

In the following we discuss the numerical data for the ground state energy, the charge and spin gap, and the spin-spin correlation function. The numerical results are obtained by the density matrix renormalization group method (DMRG), as implemented by Brune [32]. The DMRG is a quasi-exact numerical method to determine the ground state properties, i.e., the ground state and
the ground state energy, of long one-dimensional (non-integrable) systems with reasonable accuracy \[30\]. Regarding different boundary conditions, it is useful to take into account the equivalence of fermion and spin models and to implement the spinless-fermion model in terms of an equivalent spin chain, and the Hubbard model as two coupled spin chains without perpendicular XY-coupling. Using the DMRG, it is possible to extend the tractable system lengths for the Hubbard chain to \(N = 60 \ldots 100\) sites. In our simulations we perform five lattice sweeps and keep 300 to 500 states per block. Correlation functions can be obtained within an error of \(10^{-6}\) in the Hubbard model when using open boundary conditions. A memory of about 700 MB is required.

A. Energy

First, we determine the ground state energy as a function of \(W\) and \(U\). The energy gain depends almost quadratically on the potential strength for all potential types, and is dominated by the band shift. The numerical data are depicted in Fig. 3. In case of a coupling to the charge density, \(W_1 n_i\), the \(2k_F\)-periodic potential is stabilized for small \(U\) (case a)), and the \(4k_F\)-periodic potential (case c)) for large \(U\), as found for the Hubbard-Holstein model \[33\]. This energy gain is mainly due to the prefactors. A more detailed analysis of the algebraic behavior for \(W \to 0\) shows that \(E_a \propto W^{1.68}\) and \(E_c \propto W^2\) if \(U = 2t\), but \(E_a \propto W^{1.76}\) and \(E_c \propto W^{1.76}\) if \(U = 10t\). In case a) the exponent increases with interaction, in case c) it decreases. For all interactions, however, term d), \(h_iS_i^z\), is dominant. Of course, in a real system, a potential coupled to the density is much bigger than a magnetic field. It leads – like potential c) – to \(4k_F\)-oscillations in the charge density and \(2k_F\)-oscillations in the magnetization. In the non-interacting system the data points for potentials a) and d) lie nearly on top of each other (by use of open boundary conditions the exact equivalence of a) and d) is waived), but in the interacting system the energy gain due to potential d) grows faster with interaction than in all other cases. To be more specific, we find \(E_d \propto W^{1.62}\) for \(U = 2t\) and \(E_d \propto W^{1.44}\) for \(U = 6t\).

B. Charge and spin gap

We note that the excitation gaps

\[ \Delta_e = \frac{1}{2} [E(N_e + 1) + E(N_e - 1) - 2E(N_e)]\]

and

\[ \Delta_s = \frac{1}{2} [E(S^z = 1) + E(S^z = -1) - 2E(S^z = 0)]\]

are calculated for finite systems only. An extrapolation to the thermodynamic limit, \(\Delta = \Delta_0 + f(N)\) with \(f(N) \to 0\) for \(N \to \infty\), is not performed, for the following reasons:

for small gap \(\Delta_0\) the accuracy of the DMRG away from half filling is not accurate enough and for large \(\Delta_0\) we have \(f(N) \approx 0\) for the considered system sizes. The dependence of the excitation gaps on the potential strength is shown in Fig. 3. In cases a), b) and d) we see that \(\Delta_s \sim \Delta_e\), according to the band insulating state. An almost linear dependence on \(W\), as for non-interacting particles, is recovered in the interacting system for the charge gap in cases a), b), and c). The saturation of the gaps in case b), as discussed in Sec. 3B, is obtained for the spin gap in the considered parameter regime. The influence of the interaction in the band insulating state can be summarized as follows. In case a) the charge gap becomes smaller with interaction, the spin gap even
the interaction strength is $U$ more. In case b) the charge gap increases with interaction and the Parzen filter function $\text{[34]}$ to reduce the Friedel oscillations. In the Mott insulating regime, the Friedel oscillations in the spin sector are long ranged, but decay fast in spin gap systems $\text{[32]}$. The main question concerns the correspondence of the spin-spin correlation functions of the $2a$-periodic potential (case c)) and the Heisenberg model. For comparison, we calculate the spin-spin correlation function of the half-filled Hubbard model, which can be mapped onto a Heisenberg chain in case of strong interaction. In the Heisenberg model we have

$$\langle S_i^z S_j^z \rangle = \frac{1}{4\pi(a \cdot i)^2} + \frac{a_{2k_F} \cos(2k_Fa \cdot i)}{a \cdot i}. $$

In gapped systems, the spin-spin correlation functions reflect the behavior of the energy gaps. Thus, the spin-spin correlation function decreases exponentially in cases a) and b), which show a spin gap without spin order. In case d) the spins are fixed by the potential $h_i$. The exponentially decaying part is hard to extract from the mean magnetization in the numerical data due to the incomplete suppression of the boundary oscillations.

In the Mott insulating regime, however, the linear decrease of the spin-spin correlation function with distance is obtained already for small potential $W$. For better comparison with the data of the half-filled Hubbard chain, we show data for $W = t$ in Fig. 5. Obviously, the quarter-filled Hubbard chain with $4k_F$-periodic distortion can be mapped onto the Heisenberg model, too. We note that in case d) already a weak potential—connected with a small spin gap—leads to significant structures in $\langle S_i^z S_j^z \rangle$, whereas in case c) the potential has to be much stronger to yield effects of similar size. The increase of $S(q) \propto q$ appears both in the half-filled Hubbard chain and in the quarter-filled Hubbard chain with $4k_F$-periodic potential. $S(q)$ has a sharp maximum at $q = \pi/2$ in case d).

V. SUMMARY

In summary, we have studied the effects of periodic potentials on a chain. In particular, we have considered the one-dimensional Hubbard model at quarter filling. We have compared periodic potentials yielding different types of behavior: a) a $2k_F$-periodic potential leading to a band insulator, b) a $2k_F$-periodic potential leading to a band insulator with cluster-like arrangements of the charges, c) a $4k_F$-periodic potential leading to a Mott insulator with antiferromagnetic alignment of Heisenberg type spins on next-nearest neighbor sites, and d) a $2k_F$-periodic magnetic field leading to a band insulator with antiferromagnetic alignment of Ising type spins. Cases c) and d) reveal the same charge and spin distribution but

![FIG. 4: (Color online) Charge and spin gap versus potential strength (both in units of $t$). In the upper plot we show $\Delta_c$ and in the lower $\Delta_s$, comparing potentials a), b), c), and d). The number of sites varies between $N = 28$ and $N = 60$ and the interaction strength is $U = 6t$.](image-url)
different spin excitations. In the Heisenberg case the spin excitations are gapless, whereas in the Ising case they are gapped.

In non-interacting systems, we find $E_a = E_d > E_b > E_c$, thus the band insulator. The charge distribution is quite homogeneous, and the cluster like arrangement is not minimal in energy. Turning on the interaction, potential d) results in the largest energy gain, while the order of the remaining potentials depends on both the potential strength and the interaction. The interaction strongly supports the spin order, where double occupancy is suppressed.

In the following discussion we rely on the potentials a) to c), where the spin-order, if present, is a result of the charge order. For weak interaction and weak potential, we recover the behavior of the non-interacting system, with $E_a > E_b > E_c$. Turning to intermediate values ($U \approx 3t$, $W \approx t$ ), the cluster-like arrangement of potential b) gains more energy than the homogeneous distribution of case a), $E_b > E_a > E_c$. For strong $W \rightarrow t$, also a $4k_F$ pattern is favored against the homogeneous case, $E_b > E_c > E_a$. Thus, for small or intermediate interaction and potential, the gain of energy due to the hopping is stronger than the effects of the repulsive interaction or the underlying potential. With increasing potential, however, the effects of the potential dominates.

On the other hand, regarding a weak potential but increasing interaction, we find that a pattern related to potential a) is established only at small interaction. The cluster-like arrangement is formed at intermediate interaction ($U \approx 3 - 5t$), but at strong interaction the $4k_F$-periodic pattern dominates. Concerning the spin chain compound (Ca,Sr)$_{14}$Cu$_{24}$O$_{41}$, we have applied the potential only in order to clarify the leading instability of the system. For comparison with a real material which shows no lattice effects, the limit $W \rightarrow 0$ is relevant. In addition, for copper oxides usually a correlation parameter of $U \approx 8\text{eV}$ is assumed. Together with the bandwidth of about 1 eV, see Fig. 1, we have $U > 15t$.

In this parameter region, we find – neglecting the magnetic field – the $4k_F$-periodic charge pattern with antiferromagnetism to be the most probable ground state. The energetic order found for weak potentials extends to strong potentials. For strong interaction, we always have $E_c > E_b > E_a$. The tendency of the repulsive interaction to separate the charges dominates the phase diagram in this region. The influence of the potential is weaker, leading to a preference of the cluster over the homogeneous distribution.

Finally, we remark that adding electrons stabilizes the cluster formation. This observation likewise agrees with experimental results, obtained for the (Ca,La)$_{14}$Cu$_{24}$O$_{41}$ series, where the cluster size grows on additional charge. Therefore we conclude, that the antiferromagnetism is due to the band filling, thus a commensurability effect.

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