Charge Order of Strongly Bounded Electron Pairs on the Triangular Lattice: the Zero-Bandwidth Limit of the Extended Hubbard Model with Strong Onsite Attraction

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Abstract

The inhomogeneous distribution of charged particles is known as charge-order phenomena. Recent experimental studies of (quasi-) two-dimensional materials found different ordered patterns on the triangular lattice. In this work, the preliminary studies of the extended Hubbard model (with the intersite Coulomb density-density repulsion) in the atomic limit on the triangular lattice are presented. The model is investigated within the mean-field approach that treats the onsite term exactly. The finite temperature phase diagrams including metastable phases and some characteristics of the model are determined for the limit of strong onsite attraction. It is shown that in the limit considered all transitions between stable phases are discontinuous (for fixed chemical potential), which is totally different from the model on the alternate lattices. Continuous transitions between metastable phases are also found.

Keywords

Charge order · Triangular lattice · Extended Hubbard model · Atomic limit · Metastable phases · Mean-field theory · Phase diagram

1 Introduction

The charge-order phenomenon is associated with inhomogeneous distribution of charged particles on a lattice [1, 2]. The particles can form different patterns depending, among others, on relative values of interactions in the system as well as on a type of the lattice on which they interact. In this paper, the paradigmatic model for strongly correlated fermion (electron) system is studied on the triangular lattice. The motivation of this work is recent experimental [3–11] and theoretical [12–19] studies of quasi-two-dimensional systems, e.g., NbSe2 or organic conductors, where various charge-ordered patterns have been observed on such a type of the lattice.

The extended Hubbard model in the zero-bandwidth (or equivalently, atomic) limit analyzed in this work (e.g., Refs. [20–33]) has the following form

\[ \hat{H} = U \sum_i \hat{n}_i^\uparrow \hat{n}_i^\downarrow + \frac{W}{2z} \sum_{\langle i,j \rangle} \hat{n}_i \hat{n}_j - \mu \sum_i \hat{n}_i, \]  

where \( \hat{n}_i = \hat{n}_i^\uparrow + \hat{n}_i^\downarrow \) is a operator of particle number at site \( i \), \( \hat{n}_i^\sigma = \hat{c}_i^\dagger \hat{c}_i^\sigma \) is a operator of number of particles with spin \( \sigma \) (\( \sigma \in \{ \uparrow, \downarrow \} \)) at site \( i \), and \( \hat{c}_i^\dagger \) (\( \hat{c}_i^\sigma \)) is a operator of annihilation (creation) of a fermion (electron) with spin \( \sigma \) at site \( i \). \( \langle i, j \rangle \) denotes the summation over the nearest-neighbor sites, independently. \( U \) denotes the Hubbard onsite interaction, which is assumed to be attractive (i.e., \( U < 0 \)), \( W \) is the intersite density-density Coulomb repulsion (i.e., \( W > 0 \)), whereas \( \mu \) is the chemical potential. \( z \) denotes the coordination number (the number of nearest neighbors).

The model is analyzed using the variational approach that treats the onsite term exactly and the intersite term within the standard mean-field approximation (including only the Hartree terms) and in the thermodynamic limit (in the grand canonical ensemble) [26–30]. This approach is an exact one in the limit of large coordination number (\( z \to +\infty \)) [26, 34–36], but even that it can be a good starting point for the triangular lattice with \( z = 6 \) (Fig. 1). For the ground state of the model on this lattice, it is an exact theory. For finite temperatures, it obviously overestimates the critical temperatures, at which the long-range order vanishes.

The model is studied on the triangular lattice. This lattice can be divided into three equivalent sublattices labeled by \( \alpha = A, B, C \) (cf. Fig. 1) (the tri-sublattice assumption). Various homogeneous phases occurring on phase diagrams...
of the model are characterized by average occupations of the sublattices: $n_\alpha = \frac{1}{L} \sum_{i \in \alpha} \langle \hat{n}_i \rangle$, where $L$ is a number of lattice sites (within the approach used $n_\alpha = \langle \hat{n}_i \rangle$ for each $i \in \alpha$).

2 Numerical Results for Large Onsite Attraction

For the limit studied (i.e., $U/W = -10$), only two different phases are found: (i) a nonordered (NO) phase with $n_A = n_B = n_C$ and (ii) a charge-ordered (CO) phase, where the concentrations in two of three sublattices are equal to each other and the concentration in the third sublattice is different (in a general case, one has three possibilities: $n_A \neq n_B = n_C$, $n_B \neq n_A = n_C$, or $n_C \neq n_A = n_B$). The concentration which occurs only in one of the sublattices is denoted as $n_1$ (e.g., $n_1 = n_A$), whereas the other one (occurring in two of the sublattices) is indicated as $n_2$ (e.g., $n_2 = n_B = n_C$).

One can imagine the existence of the ordered phase where all three $n_\alpha$’s are different (i.e., $n_A \neq n_B \neq n_C$, and $n_C \neq n_A$; cf., e.g., Ref. [33]), but such a phase has not been found in the studied range of model parameters. To distinguish the CO and NO phases, charge-order parameter $\Delta$ defined as $\Delta = (1/2)(n_1 - n_2)$ is introduced. In the CO phase, $\Delta \neq 0$, whereas in the NO phase, $\Delta = 0$. Notice that such defined $\Delta$ is different from order parameter $\Delta_Q$ for checker-board-ordered phases occurring on alternate lattices [27–30].

The expresions for $\Delta$ and $n_\alpha$ for the triangular lattice can be derived from Ref. [28], where the equations for a general case were derived within a site-dependent approach.

2.1 The Phase Diagrams

The phase diagram of the model for the fixed chemical potential (as a function of chemical potential $\tilde{\mu} = \mu - U/2 - W$) and for $U/W = -10$ is shown in Fig. 2. First, the occurrence of stable phases, i.e., phases with the lowest grand canonical potential, is discussed. On the diagram, two distinguishable
CO phases are present: (i) a CO1 phase with $\Delta > 0$ stable for $\bar{\mu} < 0$ and (ii) a CO2 phase with $\Delta < 0$ occurring for $\bar{\mu} > 0$. These two regions of the CO phase are separated by the first-order CO1–CO2 transition at $\bar{\mu} = 0$. The order-disorder transitions, i.e., NO–CO1 and CO2–NO transitions, are also discontinuous ones. The maximal temperature of the transitions is $k_B T_M/W \approx 0.5$ and it is located at $\bar{\mu} = 0$. For $|\bar{\mu}|/W > 1$, a re-entrant behavior is found. Each of these discontinuous transitions is associated with an occurrence of the metastable phases in their neighborhood. Thus, as a consequence of the CO1–CO2 transition, the CO1 (CO2) phase is metastable in the CO2 (CO1, respectively) phase region of stability (cf. also Fig. 4). Similarly, next to the NO–CO1 (CO2–NO) line in the region of the NO phase stability, the CO1 (CO2, respectively) phase is metastable for some define range of model parameters (very narrow region, not visible in the figure), whereas the NO phase is metastable in the CO1 (CO2, respectively) region (cf. also Figs. 5 and 6). It turns out that inside the CO1 (CO2) region the continuous NO–CO2 (CO1–NO, respectively) transition between metastable phases occurs (dashed line in Fig. 2).

Fig. 4 Dependencies of order parameters: a particle concentrations in sublattices: $n_1 = n_A$ and $n_2 = n_B = n_C$, b total particle concentration $n$, and c charge-order parameter $\Delta = \frac{1}{2}(n_1 - n_2)$ as a function of chemical potential $\bar{\mu}$ for $k_B T/W = 0.2$, $W > 0$, and $U/W = -10$. Solid and dash-dotted lines correspond to stable solutions, whereas dotted and dashed lines correspond to metastable solutions. The metastable solutions for the CO1 and CO2 phases inside the NO region are not shown (cf. also Fig. 5). Labels as in Fig. 2.

Fig. 5 Dependencies of a total particle concentration $n$ and b charge-order parameter $\Delta$ as a function of chemical potential $\bar{\mu}$ for $k_B T/W = 0.2$ (i.e., $T < T_C$), $W > 0$, and $U/W = -10$ in the neighborhood of the first-order NO–CO1 transition. Solid and dotted lines correspond to stable and metastable solutions, respectively. Labels as in Fig. 2.
Figure 3 shows the phase diagram of the system considered for $U/W = -10$ as a function of total electron concentration $n = (1/3)(n_1 + 2n_2)$. Due to the fact that the CO1–CO2, NO–CO1, and CO2–NO transitions are first order for fixed chemical potential, three regions of phase separations, PS1:NO/CO1, PS2:CO2/NO, and PS3:CO1/CO2, exist on the diagram. In these regions, phase-separated states have the lowest free energy and two homogeneous phases can coexist in a defined range of $n$. In particular, in the PS3 region, two charge-ordered phases, i.e., the CO1 and CO2 phases, coexist. The fraction of the system occupied by each phase depends on $n$ (for details, see, e.g., Refs. [37–39]).

In the regions of the phase separations, one or two homogeneous phases may be metastable. In the whole region of the PS3 state occurrence, both CO1 and CO2 phases are metastable (they have higher free energy than the PS3 state), which free energy $f_{PS3}$ can be calculated by Maxwell’s construction [37–39]). But for $n < 1$, the free energy of the CO1 phase is lower than that of the CO2 phase ($f_{PS3} < f_{CO1} < f_{CO2}$), whereas for $n > 1$, the CO1 phase has higher free energy than the CO2 phase ($f_{PS3} < f_{CO2} < f_{CO1}$). At $n = 1$, a first-order transition between the CO1 and CO2 (metastable) phases occurs for any $T < T_M$.

The situation is more complex in a case of regions of the PS1 and PS2 states occurrence. For fixed temperature $k_BT/W$, there are two possible sequences of phase transitions with increasing $n$ (for $n < 1$). The first one occurring at low temperatures (for $T < T_C$, where $k_BT_C/W \approx 0.33$) is NO–PS1(NO)–PS1(NO;CO1)–PS1(CO1;NO)–PS1(CO1;CO2)–CO1(CO2). The denotation, e.g., PS1(NO;CO1), denotes that in the PS1 region, two phases, NO and CO1, are metastable with $f_{PS1} < f_{NO} < f_{CO1}$, whereas, e.g., PS1(NO) indicates that the NO phase is the only metastable solution with $f_{NO} > f_{PS1}$. At the beginning, the NO phase is only metastable phase in the PS1 region. Next, inside the PS1 region, the metastable CO1 solution appears [$n_\mu(T)$]. With increasing $n$, a first-order NO–CO1 transition between metastable phases occurs at $n_{\nu}(T)$ (before it: $f_{PS1} < f_{NO} < f_{CO1}$, whereas after it: $f_{PS1} < f_{CO1} < f_{NO}$, cf. Fig. 5), analogously to the CO1–CO2 transition inside the PS3 region discussed previously. After that, but still in the PS1 region, the second-order NO–CO2 transition between metastable phases occurs at $n_{\eta}(T)$ and $f_{PS1} < f_{CO1} < f_{CO2}$ for $n > n_{\nu}(T)$. The second possible sequence occurring for $T_C < T < T_M$ is NO–PS1(NO)–PS1(NO;CO1)–PS1(CO1;NO)–CO1(NO)–CO1(CO2). In such a case, the metastable CO2 phase does not occur in the PS1 region and the second-order NO–CO2 transition between metastable phases occurs in the CO1 region, not in the PS1 region as previously (cf. Fig. 6). At $T = T_C$, the NO–PS1(NO)–PS1(NO;CO1)–PS1(CO1;NO)–CO1(CO2) sequence is present. Note that in Fig. 3, two lines inside the region of the PS1 state occurrence are not denoted: a line, where the CO1 solution appears as the second metastable phase [$PS1(NO)–PS1(NO;CO1)$ boundary, $n_m(T)$], and a line of the first order NO–CO1 transition between metastable phases [$PS1(NO;CO1)–PS1(CO1;NO)$ boundary, $n_\nu(T)$].

Due to the particle-hole symmetry of the model discussion for $n > 1$ is similar, one should change PS1 to PS2, exchange CO1 and CO2 with each other, and discuss the sequences with decreasing $n$ (or equivalently change $n$ into $2-n$ and $\bar{\mu}$ into $-\bar{\mu}$). As it is shown before, near $n = 1$, a CO1(CO2)–PS3(CO1;CO2)–PS3(CO2;CO1)–CO2(CO1) sequence occurs for any $T < T_M$.

2.2 Dependencies of the Order Parameters

To give deeper insight into properties of the system, Figs. 4, 5, and 6 present dependencies of order parameters as a function of chemical potential for fixed $k_BT/W$. In Fig. 4a, concentrations $n_1$ and $n_2$ are presented, whereas in Fig. 4b and c, dependencies of total electron concentration...
n and charge-order parameter $\Delta$ are shown, respectively. It is clearly seen that at $\tilde{\mu} = 0$, the first-order CO1–CO2 transition occurs, namely $\Delta$ and $n$ change discontinuously from $\Delta_- > 0$ and $n_- < 1$ (for $\tilde{\mu} = 0^-$) to $\Delta_+ = -\Delta_- < 0$ and $n_+ = 2 - n_- > 1$ (for $\tilde{\mu} = 0^+$). The metastable solutions exist on both sides of the transition. Notice that for fixed $n$ between $n_-$ and $n_+$, the PS3:CO1/CO2 state has the lowest free energy, but in this region, both homogeneous phases are metastable. The comparison of free energies of both metastable phases reveals that indeed for $n = 1$, there is the discontinuous CO1–CO2 transition between metastable phases.

Now, behavior of the system in the neighborhood of the discontinuous order-disorder transition is considered. Only a case of $\tilde{\mu} < 0$ ($n < 1$) is discussed here, but discussion of a case of $\tilde{\mu} > 0$ ($n > 1$) is analogous due to the particle-hole symmetry (as indicated at the end of the previous section). At the NO–CO1 transition for fixed $\tilde{\mu}$ concentration, $n$ changes discontinuously from $n_{NO}$ (in the NO phase) to $n_{CO1}$ (in the CO1 phase), and thus for $n_{NO} < n < n_{CO1}$, the PS1:NO/CO1 state occurs for fixed $n$. In Figs. 5 and 6, one can notice that the CO1 phase is the metastable on the left side from a point of the first-order NO–CO1 transition. One can also see that in the range between $n_{NO}$ and $n_{CO1}$, only the NO phase is metastable (for $n < n_m$) or two (CO1 and NO for $n_m < n < \text{Min}[n_c,n_{CO1}]$), or, CO1 and CO2 only for $T < T_C$ and $n_c < n < n_{CO1}$ are metastable phases. The location $n_t$ of the first-order NO–CO1 transition between metastable phases for fixed $n$ can be determined by the comparison of their free energies (not indicated in the figures, and $n_t > n_m$). Moreover, in all figures, the continuous NO–CO2 transition between metastable phases occurring at $n_t$ is presented. For fixed $n$ and for $T < T_C$, the NO–CO2 transition between metastable phases occurs inside the PS1 region, i.e., $n_{NO} < n_c < n_{CO1}$ (cf. Fig. 5a). Moreover, $n_c$ is also located in the region, where the CO1 phase is the metastable phase with lower free energy (i.e., $n_m < n_t < n_c$ and $f_{PS1} < f_{CO1} < f_{NO} = f_{CO2}$ at $n_t$, respectively). In contrary, for $T_C < T < T_M$, the NO–CO2 transition between metastable phases occurs inside the CO1 region, i.e., $n_c > n_{CO1}$ (cf. Fig. 6a).

At the end of this section, one should stress that for $T_C < T < T_M$, the following relations between all critical concentrations are satisfied: $n_{NO} < n_m < n_t < n_{CO1} < n_c$, whereas for $T > T_C$, one gets $n_{NO} < n_m < n_t < n_c < n_{CO1}$. To be strict, at low temperatures (smaller than $k_B T/W \approx 0.1$), $n_{NO} < n_c < n_m < n_{CO1}$ could be also possible, which would correspond to NO–PS(NO)–PS(CO2)–PS(CO1;CO2)–CO1(CO2) sequence of transitions ($f_{CO1} < f_{CO2}$ for any $T$ and $n < 1$). This is the only possible sequence not discussed previously. However, its existence cannot be unambiguously adjudged numerically due to very small difference $|n_c - n_m|$, which is of the order of a few thousandths. Nevertheless, for $T \to 0$, all above critical concentrations goes to zero.

Note that generally homogeneous phases can be metastable on both sides of a PS region (such as for the PS3 region above or the PS region in the model of Ref. [40]) or any homogeneous phase cannot be metastable in some part of a PS region (cf. the model with longer-range interactions on alternate lattice [29]). In the present work, none of these cases has been found for the PS1 and PS2 regions.

### 3 Summary and Final Remarks

In this work, the extended Hubbard model in the atomic limit on the triangular lattice for strong onsite attraction was discussed in the context of charge-order phenomena and the analysis including metastable phases has been performed. The phase diagram of the model including metastable phases was derived for $U/W = -10$. All transitions between stable phases for fixed chemical potential were found to be discontinuous and thus phase separations occur for fixed concentration. It is different from a case of bipartite (alternate) lattices (with only nearest-neighbor interaction), where only a continuous order-disorder transition occurs for $U < 0$ [27–30].

Notice that the discussed model in the limit of $U \to -\infty$ is equivalent with the antiferromagnetic $S = 1/2$ Ising model on the triangular lattice [41–43]. The results obtained in these works do not predict long-range order at zero field (i.e., $\tilde{\mu} = 0$) [41]; however, longer-range interactions [43] or weak interactions between different (triangular) layers occurring in realistic systems could stabilize such an order (cf. also Ref. [33]).

Increasing $U$ should decrease the temperatures of the order-disorder transitions in similar way as for alternate lattices, at least for $U < 0$ [27, 30]. The analyses of this issue will be presented in the future work.

Additionally, one can note that the mean-field results for the atomic limit of the extended Hubbard model with attractive $W < 0$ are the same for both two-sublattice and tri-sublattice assumptions. In such a case, two different nonordered phases exist with the discontinuous first-order transition at $\tilde{\mu} = 0$, and thus for fixed $n$, the electron-droplet state (phase separation NO/NO) exists (cf. Refs. [26, 29, 30] for large attractive $U$ and Ref. [41] for the ferromagnetic Ising model).

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References

1. Micnas, R., Ranninger, J., Robaszkiewicz, S.: Superconductivity in narrow-band systems with local nonretarded attractive interactions. Rev. Mod. Phys. 62, 113 (1990). https://doi.org/10.1103/RevModPhys.62.113

2. Imada, M., Fujimori, A., Tokura, Y.: Metal-insulator transitions. Rev. Mod. Phys. 70, 1039 (1998). https://doi.org/10.1103/RevModPhys.70.1039

3. Zhou, S., Wang, Z.: Charge and spin order on the triangular lattice: Na$_x$CoO$_2$ at $x = 0.5$. Phys. Rev. Lett. 98, 226402 (2007). https://doi.org/10.1103/PhysRevLett.98.226402

4. Soumyaranayan, A., Yeea, M.M., Era, Y., van Wezel, J., Chatterjee, U.: Orbital selectivity causing anisotropy and particle-hole asymmetry in the charge density wave gap of 2H-TaS$_2$. Phys. Rev. B. 96, 245103 (2017). https://doi.org/10.1103/PhysRevB.96.245103

5. Ptok, A., Głodzik, S.Z., Domański, T.: Yu-Shiba-Rusinov states of impurities in a triangular lattice of NbSe$_2$ with spin-orbit couplings. Phys. Rev. B. 96, 184425 (2017). https://doi.org/10.1103/PhysRevB.96.184425

6. Xi, X., Wang, Z., Zhao, W., Park, J.H., Law, K.T., Berger, H., Forro, L., Shan, J., Mak, K.F.: Ising pairing in superconducting NbSe$_2$ atomic layers. Nat. Phys. 12, 139 (2016). https://doi.org/10.1038/nphys3538

7. Lian, C.-S., Si, C., Duan, W.: Unveiling charge-density wave, superconductivity, and their competitive nature in two-dimensional NbSe$_2$. Nano Lett. 18, 2924 (2018). https://doi.org/10.1021/acs.nanolett.8b02037

8. Pätztor, A., Scarfato, A., Barreteau, C., Giannini, E., Renner, C.: Dimensional crossover of the charge density wave transition in thin exfoliated VSe$_2$. 2D Mater 4, 041005 (2017). https://doi.org/10.1088/2053-1583/aa86de

9. Zhao, J., Wijayaratne, K., Butler, A., Yang, J., Milliakas, C.D., Chung, D.Y., Louca, D., Kanatzidis, M.G., van Wezel, J., Chatterjee, U.: Orbital selectivity causing anisotropy and particle-hole asymmetry in the charge density wave gap of 2H-TaS$_2$. Phys. Rev. B. 96, 125103 (2017). https://doi.org/10.1103/PhysRevB.96.125103

10. Kolekar, S., Bonilla, M., Ma, Y., Díaz, H.C., Batzill, M.: Layer- and substrate-dependent charge density wave criticality in FeTiSe$_2$. 2D Mater 5, 015006 (2018). https://doi.org/10.1088/2053-1583/aa866f

11. Ryu, H., Chen, Y., Kim, H., Tsai, H.-Z., Tang, S., Jiang, J., Liou, F., Kahn, S., Jia, C., Omranif, A.A., Shim, J.-H., Hussain, Z., Shen, Z.-X., Kim, K., Min, B.-I., Hwang, C., Crommie, M.F., Mo, S.-K.: Persistent charge-density-wave order in single-layer TaSe$_2$. Nano Lett. 18, 689 (2018). https://doi.org/10.1021/acs.nanolett.7b03264

12. Cano-Cortés, L., Ralko, A., Février, C., Merino, J., Fratini, S.: Geometrical frustration effects on charge-driven quantum phase transitions. Phys. Rev. B. 84, 155115 (2011). https://doi.org/10.1103/PhysRevB.84.155115

13. Merino, J., Ralko, A., Fratini, S.: Emergent heavy fermion behavior at the Wigner-Mott transition. Phys. Rev. Lett. 111, 126403 (2013). https://doi.org/10.1103/PhysRevLett.111.126403

14. Tocchio, L.F., Gros, C., Zhang, X.-F., Eggert, S.: Phase diagram of the triangular extended Hubbard model. Phys. Rev. Lett. 113, 246405 (2014). https://doi.org/10.1103/PhysRevLett.113.246405

15. Kaneko, R., Tocchio, L.F., Valenti, R., Gros, C.: Emergent lattices with geometrical frustration in doped extended Hubbard models. Phys. Rev. B. 94, 195111 (2016). https://doi.org/10.1103/PhysRevB.94.195111

16. Łitak, G., Wysokiński, K.I.: Evolution of the charge density wave order on the two-dimensional hexagonal lattice. J. Magn. Magn. Mater. 440, 104 (2017). https://doi.org/10.1016/j.jmmm.2016.12.042

17. Kaneko, R., Tocchio, L.F., Valenti, R., Becca, F.: Charge orders in organic charge-transfer salts. New J. Phys. 19, 103033 (2017). https://doi.org/10.1088/1367-2630/aa887b

18. Ptok, A., Głowidz, S.Z., Domanski, T.: Yu-Shiba-Rusinov states of impurities in a triangular lattice of NbSe$_2$ with spin-orbit couplings. Phys. Rev. B. 96, 184425 (2017). https://doi.org/10.1103/PhysRevB.96.184425

19. Cichy, A., Ptok, A.: Reentrant Fulde-Ferrell-Larkin-Ovchinnikov superfluidity in the honeycomb lattice. Phys. Rev. A. 97, 053619 (2018). https://doi.org/10.1103/PhysRevA.97.053619

20. Jędrzejewski, J.: Phase diagrams of extended Hubbard models in the atomic limit. Phys. A. 295, 125052 (2004). https://doi.org/10.1016/j.physa.2004.12.002

21. Bursill, R.J., Tompson, C.J.: Variational bounds for lattice fermion models. Phys. Rev. B. 94, 125103 (2016). https://doi.org/10.1103/PhysRevB.94.125103

22. Micnas, R., Robaszkiewicz, S., Chao, K.A.: Multicritical behavior of impurities in wide band organic charge-transfer salts. New J. Phys. 19, 103033 (2017). https://doi.org/10.1088/1367-2630/aa887b
30. Kapcia, K.J., Robaszkiewicz, S.: On the phase diagram of the extended Hubbard model with intersite density-density interactions in the atomic limit. Phys. A. 461, 487 (2016). https://doi.org/10.1016/j.physa.2016.05.056
31. Rademaker, L., Pramudya, Y., Zaanen, J., Dobrosavljević, V.: Influence of long-range interactions on charge ordering phenomena on a square lattice. Phys. Rev. E. 88, 032121 (2013). https://doi.org/10.1103/PhysRevE.88.032121
32. Kapcia, K.J., Barański, J., Ptok, A.: Diversity of charge orderings in correlated systems. Phys. Rev. E. 96, 042104 (2017). https://doi.org/10.1103/PhysRevE.96.042104
33. Kaneko, R., Nonomura, Y., Kohno, M.: Thermal algebraic-decay charge liquid driven by competing short-range Coulomb repulsion. Phys. Rev. B. 97, 205125 (2018). https://doi.org/10.1103/PhysRevB.97.205125
34. Müller-Hartmann, E.: Correlated fermions on a lattice in high dimensions. Z. Phys. B. 74, 507 (1989). https://doi.org/10.1007/BF01311397
35. Pearce, P.A., Thompson, C.J.: The anisotropic Heisenberg model in the long-range interaction limit. Comm. Math. Phys. 41, 191 (1975). https://doi.org/10.1007/BF01608757
36. Pearce, P.A., Thompson, C.J.: The high density limit for lattice spin models. Comm. Math. Phys. 58, 131 (1978). https://doi.org/10.1007/BF01609416
37. Arrigoni, E., Strinati, G.C.: Doping-induced incommensurate antiferromagnetism in a Mott-Hubbard insulator. Phys. Rev. B. 44, 7455 (1991). https://doi.org/10.1103/PhysRevB.44.7455
38. Bąk, M.: Mixed phase and bound states in the phase diagram of the extended Hubbard model. Acta Phys. Pol. A. 106, 637 (2004). https://doi.org/10.12693/APhysPolA.106.637
39. Kapcia, K.J., Czart, W.R., Ptok, A.: Phase separation of superconducting phases in the Penson–Kolb–Hubbard model. J. Phys. Soc. Jpn. 85, 044708 (2016). https://doi.org/10.7566/JPSJ.85.044708
40. Kapcia, K.: Metastability and phase separation in a simple model of a superconductor with extremely short coherence length. J. Supercond. Nov. Magn. 27, 913 (2014). https://doi.org/10.1007/s10948-013-2409-8
41. Houtappel, R.M.F.: Order-disorder in hexagonal lattices. Physica 16, 425 (1950). https://doi.org/10.1016/0031-8914(50)90130-3
42. Campbell, C.E., Schick, M.: Triangular lattice gas. Phys. Rev. A 5, 1919 (1972). https://doi.org/10.1103/PhysRevA.5.1919
43. Mihura, B., Landau, D.P.: New type of multicritical behavior in a triangular lattice gas model. Phys. Rev. Lett. 38, 977 (1977). https://doi.org/10.1103/PhysRevLett.38.977