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Published in:
PHYSICAL REVIEW B

DOI:
10.1103/PhysRevB.91.144510

Published: 01/04/2015

Please cite the original version:
Superfluidity and density order in a bilayer extended Hubbard model

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(Received 13 November 2014; revised manuscript received 23 March 2015; published 23 April 2015)

We use cellular dynamical mean-field theory to study the phase diagram of the square lattice bilayer Hubbard model with an interlayer interaction. The layers are populated by two-component fermions, and the densities in both layers and the strength of the interactions are varied. We find that an attractive interlayer interaction can induce a checkerboard density-ordered phase and superfluid phases, with either interlayer or intralayer pairing. Remarkably, the latter phase does not require an intralayer interaction to be present: it can be attributed to an induced attractive interaction caused by density fluctuations in the other layer.

DOI: 10.1103/PhysRevB.91.144510 PACS number(s): 71.10.Fd, 03.75.Ss, 67.85.Fg, 67.85.Hj

I. INTRODUCTION

Layered lattice systems may exhibit interesting physics since they allow effects typical to two and three dimensions to interplay. For instance, the study of superfluidity in layered systems has been an active area of research since the discovery of high-temperature superconductivity in the cuprates [1].

Another intriguing phenomenon in layered systems is the condensation of interlayer electron-hole pairs, or excitons, induced by repulsive interactions between electrons [2]. Ultracold gases provide one more system where various types of layered systems can be realized. There, the possibility to control inter- and intralayer tunnelings (hoppings) has been available for some time [3,4] but the interactions have been limited to on-site intralayer type. The interesting new development are dipolar atoms and molecules [5–10] that provide long-range interaction, since there is an exact particle-hole condensation in the continuum, they show that in one-dimensional systems, they create a particle of spin component \(\sigma\) at site \(i\) of layer \(l\), the corresponding density operators are defined by

\begin{equation}
\label{eq:1}
H_l = -t \sum_{\sigma,l,(i,j)} c_{\sigma l i}^\dagger c_{\sigma l j} - t_{\perp} \sum_{\sigma,l} (c_{\sigma l i}^\dagger c_{\sigma l 2i} + \text{H.c.})
- \epsilon \sum_{\sigma,l} (n_{\sigma l 1i} - n_{\sigma l 2i}) - \mu \sum_{\sigma,l} (n_{\sigma l 1i} + n_{\sigma l 2i}).
\end{equation}
so that half-filling corresponds to \( \epsilon = \mu = 0 \). We focus on the case where the on-site interaction \( U \) is tuned to zero (e.g., using a Feshbach resonance), and set \( V = -3 \), corresponding to an attractive interaction between the layers. These parameters can be realized in experiment, for instance, with \(^{161}\text{Dy} \) atoms in a bilayer with intralayer lattice spacing \( d = 225 \text{ nm} \) and the interlayer spacing \( d_{12} = 0.5 \text{ nm} \). The dipolar interaction \( V_{\text{dip}} \) is perpendicular to the layers, and we take the intralayer hopping as our unit of energy, \( t = 1 \).

The interaction Hamiltonian is written in the particle-hole symmetric form

\[
H_I = V \sum_{i,\sigma,\sigma'} \left( n_{\sigma i} - \frac{1}{2} \right) \left( n_{\sigma' j} - \frac{1}{2} \right) + U \sum_{i,l} \left( n_{\uparrow li} - \frac{1}{2} \right) \left( n_{\downarrow li} - \frac{1}{2} \right),
\]

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\]

III. PHASE DIAGRAM

The intralayer and interlayer superfluid order parameters are shown as a function of the external fields \( \mu \) and \( \epsilon \) in Fig. 1. These results were obtained within two-site DMFT \((L=1)\) that excludes translational symmetry breaking. We find that the system prefers interlayer superfluidity for low values of the interlayer density polarizing field \( \epsilon \), while higher polarizations drive it into the intralayer superfluid state. The interlayer superfluid region is larger for the more dense layer, i.e., the layer further away from half-filling, and therefore the intralayer order parameter is plotted for that layer.

We have thus predicted that, when the symmetry of the layers is distorted by the interlayer density polarizing field, an intralayer superfluid emerges that does not require intralayer interactions. An explanation of this intriguing finding could be that the superfluidity is induced by second-order effects mediated by the interlayer interaction. To test this hypothesis, we calculate the interaction strength in layer \( l' \) mediated by density fluctuations in layer \( l \) using a simple mean-field theory [38],

\[
U_{l'}^{\text{ind}} \propto 2V^2 \sum_{k,n} G_{knl}^0 G_{knl}^0 \propto 2V^2 \frac{1}{k_BT} \sum_k n_F(\epsilon_k - \mu_l)[n_F(\epsilon_k - \mu_l) - 1],
\]

where in the first line \( G_{knl}^0 \) is the noninteracting Green’s function for the particles in layer \( l \) and the summation runs over all energy and momentum states, while the factor 2 results from summing over spin states. In the second line, \( n_F \) are the Fermi

![FIG. 1. (Color online) The interlayer order parameter \( \Delta_\perp \) (yellow or light grey) and interlayer order parameter \( \Delta_\parallel \) for the more dense layer (red or dark grey) from translation invariant two site DMFT, as functions of the interlayer density polarizing field \( \epsilon \) and the chemical potential \( \mu \). Half-filling is given by \( \mu = 0 \). The intralayer order parameter is zero where the interlayer order parameter is nonzero, and vice versa. The inverse temperature is \( \frac{1}{k_BT} = 15 \) and the interaction strength \( V = -3 \), in units of the hopping \( t = 1 \). The data points are marked by black dots, and the surfaces interpolate between them. The Cooper pairs forming the interlayer superfluid are directly bound by the interlayer interaction, while the attraction necessary for the intralayer superfluid is indirectly induced by the other layer.]
distribution functions, where $\epsilon_k$ are the particle dispersions and $\mu_\sigma$ determines the density in layer $l$. The strength of $U^{\text{ind}}$ in one layer thus depends on the density in the other layer and turns out to be stronger if mediated by the layer closer to half-filling. This explains why the superfluid is more easily formed in the more dense layer, i.e., in the layer further away from half-filling, while intuition gained from, e.g., the single-band attractive Hubbard model would suggest the opposite.

Now, the crucial question is whether the superfluids, we predict survive the competition with density order, highly typical for lattice systems. To investigate this, we have obtained the phase diagram of the system also using the $L = 2$ cluster, allowing the possibility of a checkerboard density-ordered phase, which can be identified by a non-zero value of the order parameter

$$D = \frac{1}{N} \sum_{\sigma \sigma' l} \text{sgn}(i)(n_{\sigma l}) \right|_l,$$

where \text{sgn}(i) is an alternating sign that has the opposite values for any pair of neighboring sites inside one layer, and $N$ is the total number of sites on the lattice. The density order is present close to half-filling and zero polarization, as can be seen in the phase diagrams in Figs. 2(a) and 2(b). Compared to Fig. 1, it can be seen that in quite a considerable region where we first found a superfluid phase, actually the density-ordered phase turns out to be the ground state of the system. However, we still find a region where the interlayer superfluid phase is the most favorable and, strikingly, the intralayer superfluid phase induced by density fluctuations in the opposite layer remains present in the phase diagram.

Above the critical temperatures of the superfluids, the transition from the density-ordered phase to the normal phase is of a small intralayer repulsion on the phase diagram. This explains why the superfluid is more easily formed in the more dense layer, i.e., in the layer further away from half-filling. The rapid variation of the density and polarization as a function of $\epsilon$ and $\mu$ suggests phase separation, in particular between the density-ordered phase and the intralayer superfluid, for which DMFT solutions can be obtained at the same external polarizing field $\epsilon$ but with large differences in polarization $P$.

The translation invariant solution is in itself an interesting model for two-band superfluidity where the interband coupling is the dominant one. The order parameters and densities near the intralayer-interlayer transition are shown in Fig. 3. It can be seen that the transition between these two superfluid states is a sharp, first-order transition. The interlayer superfluid order parameter $\Delta_\perp$ is zero when the intralayer superfluid order parameter $\Delta_l$ is nonzero and vice versa. The discontinuity in the polarization and the interlayer order parameter gradually decreases with increasing density, and the transition turns into a second-order one when the density is high enough so that the intralayer superfluid is not present. Figure 3 can be compared to Fig. 2 of Ref. [37], where analogous behavior is found in a single-band attractive Hubbard model as the spin polarization within the single band is increased by a magnetic field. In their manifest in the fact that the density-ordered region is slightly larger above the superfluid critical temperatures, although it never totally covers the superfluid regions present in Fig. 2.
case, the transition is always a second-order one, since the competition between two superfluid phases as in our case is absent.

The intralayer-interlayer superfluid transition is caused by a competition between the different modes of pairing. For instance, if we force the intralayer pairing fields to be zero, the interlayer superfluid region is extended and exhibits a second-order transition to the normal state along the whole phase boundary. Similarly, the intralayer superfluid is also present at zero polarization if the interlayer pairing is excluded. Nevertheless, we did not find any hysteretic behavior in the simulations near the intralayer-interlayer transition, which supports the reliability of the numerics.

**IV. EFFECT OF THE CLUSTER SIZE AND COMPARISON TO MEAN-FIELD RESULTS**

The critical temperatures of the superfluids and the density-ordered phase at selected points in the \((\epsilon,\mu)\) plane are listed in Table I. There is only little variation between different cluster sizes, which is a sign that neglecting nonlocal quantum fluctuations beyond two sites has only a small quantitative effect on the results. This is in contrast to the case of exotic superfluidity in quasi-1D systems \([39]\) or \(d\)-wave superfluidity in 2D \([40]\) where nonlocal correlations play a crucial role.

It is possible to allow translational symmetry breaking also within the two-site CDMFT by treating the system in a partial real space formalism \([41,42]\) by including two inequivalent impurity problems representing the high-density and low-density sublattices of the checkerboard density order. A comparison of the phase diagrams obtained within this approximation and using the \(L = 2\) cluster is presented in Figs. 2(b) and 2(c). We find that the results are in good agreement, which further supports the conclusion that the effect of the cluster size is not very important.

The effects of different approximations on the transition from the normal phase to the density order as a function of the inverse temperature are illustrated in Fig. 4. As expected, the critical temperature is slightly lower for larger clusters, but the variation is small and the effect of the cluster size is insignificant when the system is not very close to the critical point. In contrast, the mean-field approximation gives considerably higher critical temperatures, emphasizing the importance of more accurate methods. For a weaker interaction strength \(V = -3\), the critical temperature from two-site DMFT \((L = 1)\) is \(T_c = 0.13 \pm 0.01\). In this case, the system is well in the perturbative region, and while the mean-field treatment still gives considerable errors, including second-order corrections (see Appendix C) already yields a good approximation. For \(V = -3\), however, even with the second-order corrections, mean-field theory is clearly inadequate.

**V. EFFECT OF ON-SITE REPULSION AND INTERLAYER HOPPING**

The presence of a small repulsive on-site interaction \(U\) does not destroy the intralayer superfluid. For example, the superfluidity persists up to \(U \approx 0.1\) at the point \((\epsilon = 1.1,\mu = 0.4)\), when the other parameters are as in Fig. 2.

The \(\mu-\epsilon\) phase diagram for \(U = 0.1\) is also shown in Fig. 2(d). The on-site interaction has only a small effect on the density-ordered and interlayer superfluid phases, and the intralayer superfluid also remains present in the phase diagram despite the suppression of intralayer Cooper pairing. On the other hand, we find that an attractive on-site interaction (not shown) favors the intralayer superfluid phase as expected.

Finally, we study the effect of a finite interlayer hopping amplitude in the translation invariant case. A comparison of the density, polarization and the superfluid order parameters for different values of \(t_{\perp}\) is shown in Fig. 5. The possibility to tunnel between the layers turns the sharp first-order transition between the interlayer and intralayer superfluid states into a smooth crossover. This is expected, since in this case an intralayer Cooper pair can turn into an interlayer one and vice versa thus mixing the two types of superfluids. Note that the smoothening is not very large for \(t_{\perp} = 0.1\), and for \(t_{\perp} = 0.01\) it is negligible.

**VI. CONCLUSIONS**

In conclusion, we have studied the bilayer extended Hubbard model for attractive interlayer interactions using a beyond mean-field approach. We found density order starting from half-filling and, when the density is increased, an interlayer superfluid that appears before reaching the normal state. This superfluid, where the Cooper pairs are formed by particles from different layers, is conceptually related to bilayer exciton condensates where pairing happens between particles in one layer and holes in the other. Notably, the freedom to polarize a density difference between the layers revealed also a novel type of superfluid with intralayer pairing that is induced by density fluctuations in the opposite layer. The predicted phases could be experimentally realized, for instance, with
ultracold dipolar atoms or molecules. A prospect for future research is to study if some other type of density order than the checkerboard considered here, possibly incommensurate, would be present near the density order phase boundary. This would be particularly interesting as there is a possibility that a supersolid phase.

**ACKNOWLEDGMENTS**

This work was supported by the Academy of Finland through its Centers of Excellence Programme (2012-2017) and under project Nos. 263347, 251748, and 272490, and by the European Research Council (ERC-2013-AdG-340748-CODE and ERC-2011-AdG-290464-SIMCOFE). T.I.V. is grateful for the support from the Vilho, Yrjö and Kalle Väisälä Foundation. Computing resources were provided by CSC – the Finnish IT Centre for Science and the Triton cluster at Aalto University.

**APPENDIX A: HOPPING PARAMETERS AND ENERGIES IN THE EXTENDED HUBBARD MODEL**

In the Hubbard model, the hopping parameter \( t \) describing hopping from site \( j \) to the nearest-neighbor site \( j' \) is given by

\[
t = -\int d^3 r \phi_j^*(\mathbf{r}) \left[ -\frac{\hbar^2}{2m} \nabla^2 + V_{\text{lat}}(\mathbf{r}) \right] \phi_{j'}(\mathbf{r}),
\]

where \( V_{\text{lat}}(\mathbf{r}) \) is the lattice potential. To calculate the interlayer hopping \( t_{\perp} \) in our model, \( j \) and \( j' \) are taken on different layers, whereas for the intralayer hopping \( t \) the sites \( j \) and \( j' \) are in the same layer. The interaction parameters are given by

\[
U = \frac{4\pi \hbar^2 a_S}{m} \int d^3 r |\phi_j(\mathbf{r})|^4 + \int d^3 r d^3 r' |\phi_j(\mathbf{r})|^2 |\phi_{j'}(\mathbf{r}')|^2 U(\mathbf{r} - \mathbf{r}')
\]

\[
V_{j,j'} = \int d^3 r d^3 r' |\phi_j(\mathbf{r})|^2 |\phi_{j'}(\mathbf{r}')|^2 U(\mathbf{r} - \mathbf{r}').
\]

where \( a_S \) is the s-wave scattering length and \( U(\mathbf{r}) \) is the dipole-dipole interaction [9], which in the case of all dipoles being aligned reads

\[
U(\mathbf{r}) = \frac{C_{dd}}{r^3} \left[ 1 - \frac{3}{4\pi} \cos^2 \theta \right],
\]

where \( \theta \) is the angle between \( \mathbf{r} \) and the direction of the polarization. Here, \( C_{dd} \) is the strength of the dipolar interaction, which in case of dysprosium, is \( C_{dd} = \mu_0 \mu^2 \) with the magnetic dipole moment \( \mu = 10 \mu_B \). The on-site interaction \( U \) results from the
contact interaction and the dipole-dipole interaction, whereas the intersite interactions $V_{ij,j'}$ result from the dipole-dipole interactions only. To make an order of magnitude estimation for the hoppings and interactions we use in this paper, we calculated the above coefficients using the tight binding approximation (the on-site wave functions $\phi_0(r)$ were taken to be harmonic oscillator ground state wave functions). We found that for $^{164}$Dy in a layer with lattice spacing $d = 225$ nm [20] and lattice height $V_0 = 3.5E_R$, with $E_R$ the recoil energy, the on-site interaction can be tuned to zero, $U = 0$, since there are several Feshbach resonances available to control $a_S$. An interlayer interaction $V = -1t$ can be reached when the spacing between the layers is $d_{\perp} = d/3$ and the lattice height in the perpendicular direction is $V_{0,\perp} = 20V_0$. In that case, the interlayer hopping $t_{\perp}$ and intralayer nearest-neighbor interaction $V_{l}$ are small enough (less than 10%) in comparison to $t$ and $V$, respectively, to be taken to zero in our model. A stronger interaction, compared to the hopping, can be reached by taking larger lattice heights in all directions. The stronger interaction we consider in this article can be reached when the spacing between the layers is $d_{\perp} = d/3$ and the lattice heights $V_0 = 7E_R$ and $V_{0,\perp} = 20V_0$ in the parallel and perpendicular directions, respectively.

APPENDIX B: CHOICE OF THE SUPERFLUID ORDER PARAMETERS

In this Appendix, we provide details for the choice of the superfluid order parameters taking into account the symmetries of the system. We first discuss the case $t_{\perp} = 0$ and comment on the general case in the end.

Within the $L = 1$ cluster, it is possible to have a six component pairing order parameter which consists of the intralayer order parameter $\Delta_{l} = \langle c_{1l}c_{1l}^\dagger \rangle$ in each layer $l$ and of the four component matrix $M$ defined by

$$\psi_{il} = \begin{pmatrix} c_{1l}^\dagger \\ c_{1l} \end{pmatrix}, \quad M = \langle \psi_{i1}^\dagger \psi_{i2}^T \rangle. \quad (B1)$$

As the interlayer hopping vanishes, and the interlayer interaction only couples the total densities of the layers, the Hamiltonian is invariant under rotations $\psi_{i} = U_{l} \psi_{il} \in \text{SU(2)}$ in both layers $l$ where $U_{l}$ is a unitary matrix. The SU(2) part of this symmetry corresponds to conservation of spin, and the U(1) part corresponds to conservation of particle number, in both layers separately. Letting

$$U_{l} = \begin{pmatrix} a & b \\ c & d \end{pmatrix}, \quad (B2)$$

the transformation for $\Delta_{l}$ is

$$\Delta'_{l} = \langle c_{1l}^\dagger c_{1l} \rangle = \langle (ac_{1l} + bc_{1l})(ac_{1l} + dc_{1l}) \rangle = \langle (ad - bc)c_{1l}c_{1l}^\dagger \rangle = \text{det}(U_{l})\Delta_{l}. \quad (B3)$$

Thus the $\Delta_{l}$ are invariant in pure SU(2) spin rotations and only gain a phase $\text{det}(U_{l})$ in general unitary transformations.

The transformation for the interlayer order parameters is given by

$$M' = \langle \psi_{i1}^\dagger (\psi_{i2}^\dagger)^T \rangle = \langle U_{l1}^\dagger U_{l2}^\dagger \psi_{i1}^\dagger \psi_{i2}^T \rangle = U_{l1}^\dagger \langle \psi_{i1}^\dagger \psi_{i2}^T \rangle U_{l2}^\dagger = U_{l1}^\dagger MU_{l2}^\dagger. \quad (B4)$$

By the singular value decomposition, it is always possible to choose $U_{1}$ and $U_{2}$ so that $M'$ is diagonal and only has nonnegative real elements. Thus, without losing generality, the order parameter reduces to two nonnegative real valued fields and the two, in general complex valued, $\Delta_{l}$. By doing a further transformation with the matrices

$$U_{1} = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}, \quad U_{2} = I, \quad (B5)$$

$\Delta_{1}$ gets a minus sign and the interlayer order parameter matrix $M'$ becomes purely off diagonal, the nonzero components being $\langle c_{11}^\dagger c_{12} \rangle$ and $\langle c_{11}^\dagger c_{12} \rangle$.

This is the representation where we perform our numerics. Because of technical limitations, we also consider the intralayer order parameters to be real valued. As we have fixed the interlayer order parameters to be nonnegative, this still leaves the signs of the interlayer order parameters as nontrivial factors that cannot be rotated away by symmetry. However, from the simulations we find that the interlayer order parameters (the matrix $M'$) are always zero, when the intralayer order parameters are nonzero, and vice versa. Under this assumption, it is always possible to find a transformation that flips the sign of one of the nonzero components, leaving the other one invariant. Thus all sign configurations are equivalent, and we are free to choose the order parameters to be positive, including the intralayer $\Delta_{l}$. Furthermore, we find that the two nonzero components of $M'$ are always equal, and thus we just call this value the interlayer order parameter $\Delta_{1} \equiv \langle c_{11}^\dagger c_{12} \rangle = \langle c_{11}^\dagger c_{12} \rangle$.

Also, within larger clusters, our CDMFT implementation includes all anomalous Green’s functions between opposite spin components within the cluster. The $t_{\perp} = 0$ case can be thought of as a limiting case of a model where the particles have a small probability to tunnel between the layers. This tunneling breaks the conservation of particle number in both layers separately, thus leaving only two exactly conserved spin components. Thus it is expected that the superfluidity can be treated in a picture where the Cooper pairs are only formed between particles of opposite spin, and not between particles of the same spin in different layers. In the case of nonzero $t_{\perp}$, the intralayer and interlayer order parameters take finite values simultaneously, and the relative signs of the order parameters become important. We find that the preferred sign configuration is such that $\Delta_{1} \equiv \langle c_{11}^\dagger c_{12} \rangle = -\langle c_{11}^\dagger c_{12} \rangle = \langle c_{12}^\dagger c_{11} \rangle$ and $\Delta_{2} = 0$.

APPENDIX C: MEAN-FIELD TREATMENT OF THE DENSITY-ORDERED PHASE

To make comparisons with the CDMFT solution of the model, we have also considered a mean-field type approximation for the density-ordered phase. The lowest-order mean-field approximation can be obtained by decomposing the interlayer density-density interaction as

$$(n_{A} - \frac{1}{2})(n_{B} - \frac{1}{2}) = n_{A}n_{B} - \frac{1}{2}(n_{A} + n_{B}) + \frac{1}{4} = (n_{A} + \delta_{A})(n_{B} + \delta_{B}) - \frac{1}{2}(n_{A} + n_{B}) + \frac{1}{4}$$

where $\delta_{A} = (n_{A} - \langle n_{A} \rangle)$ is the deviation of the particle number from its expectation value. The deviation from the equilibrium value is thus

$$\delta_{A} = \langle n_{A} \rangle = \frac{\mu}{E_{R}},$$

where $\mu$ is the chemical potential.

By using this approximation, the interlayer density density interaction becomes

$$(n_{A} - \frac{1}{2})(n_{B} - \frac{1}{2}) = (n_{A} + \delta_{A})(n_{B} + \delta_{B}) - \frac{1}{2}(n_{A} + n_{B}) + \frac{1}{4}$$

which results in a deviation from the equilibrium value of

$$\delta_{A} = \langle n_{A} \rangle = \frac{\mu}{E_{R}}.$$
\[ H = H_0 + \sum_{i,\sigma,\sigma'} \left( n_{\sigma li} - \frac{1}{2} \right) n_{\sigma' li} - V \sum_{i,\sigma,\sigma'} \langle n_{\sigma li} \rangle \langle n_{\sigma' li} \rangle, \] (C1)

where \( l' = 1 \) when \( l = 2 \) and vice versa. The resulting self-energy is diagonal in spin, site, and layer indices, and is given in Matsubara frequency space by

\[ \Sigma^{(1)}_\sigma(i\omega_n) = -V \sum_{\sigma'} \left( \langle n_{\sigma' li} \rangle - \frac{1}{2} \right). \] (C2)

Subsequently, the propagator \( G_{\sigma li}(\tau) = \langle c_{\sigma li}(\tau)c^\dagger_{\sigma li}(0) \rangle \) can be calculated as

\[ G_{\sigma li}(\tau) = \frac{1}{\beta} \sum_{\omega_n} \exp(-i\omega_n\tau)(-i\omega_n + T - \Sigma_\sigma(i\omega_n))^{-1}, \] (C3)

where \( G, T, \) and \( \Sigma \) are matrices in the site indices, \( T \) is the hopping matrix of the square lattice including the chemical potential contributions, and \( \beta \) is the unitless inverse temperature. In practice the matrix inversion is done in Fourier space with a unit cell that allows the breaking of translation invariance. The self-consistency condition is that the density \( \langle n_{\sigma li} \rangle = 1 - G_{\sigma li}(\tau = 0^+) \) agrees with the density in Eq. (C2).

It is possible to derive the mean-field theory from the Baym-Kadanoff (or Luttinger-Ward) functional formalism, which can also be used to systematically include higher-order corrections to the approximation [43–45]. In this formulation, the self-energy is expressed as a diagrammatic series expansion in terms of the interaction vertices and interacting propagator lines. As only the two-particle irreducible diagrams enter this series, the contributions up to second order in \( V \) consist of only two diagrams, which are depicted in Fig. 6.

Including only the first-order diagram gives the mean-field self-energy (C2). The second-order diagram includes contributions, which are not diagonal in the site indices, see Fig. 6. However, we neglect these contributions and apply a local approximation where both vertices of the diagram are on the same site \( i = j \). Evaluating the second-order diagram gives the correction term

\[ \Sigma^{(2)}_\sigma(\tau_1 - \tau_2) = -4V^2 G_{\sigma li}(\tau_1 - \tau_2) \times \sum_{\sigma'} G_{\sigma' li}(\tau_1 - \tau_2) G_{\sigma'i li}(\tau_2 - \tau_1), \] (C4)

where again \( l' \neq l \). We calculate this term directly in an imaginary time grid and then perform a numerical Fourier transformation to Matsubara frequencies. This yields a self-energy \( \Sigma_\sigma(i\omega) = \Sigma^{(1)}_\sigma(i\omega) + \Sigma^{(2)}_\sigma(i\omega) \), which can be used to calculate the propagator according to Eq. (C3). The then obtained propagator is used to calculate the self-energy again, and the process is iterated until a converged, self-consistent solution is found.


