Spin- and charge-density waves in the Hartree–Fock ground state of the two-dimensional Hubbard model

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Abstract
The ground states of the two-dimensional repulsive Hubbard model are studied within the unrestricted Hartree–Fock (UHF) theory. Magnetic and charge properties are determined by systematic, large-scale, exact numerical calculations, and quantified as a function of electron doping, h. In the solution of the self-consistent UHF equations, multiple initial configurations and simulated annealing are used to facilitate convergence to the global minimum. New approaches are employed to minimize finite-size effects in order to reach the thermodynamic limit. At low to moderate interacting strengths and low doping, the UHF ground state is a linear spin-density wave (l-SDW), with antiferromagnetic order and a modulating wave. The wavelength of the modulating wave is 2/h. Corresponding charge order exists but is substantially weaker than the spin order, hence holes are mobile. As the interaction is increased, the l-SDW states evolve into several different phases, with the holes eventually becoming localized. A simple pairing model is presented with analytic calculations for low interaction strength and small doping, to help understand the numerical results and provide a physical picture for the properties of the SDW ground state. By comparison with recent many-body calculations, it is shown that, for intermediate interactions, the UHF solution provides a good description of the magnetic correlations in the true ground state of the Hubbard model.

(Some figures may appear in colour only in the online journal)

1. Introduction
The Hubbard model is one of the most fundamental models in quantum physics. Despite numerous analytic and numerical investigations [1–6] key questions still remain about the properties of this model [7–11]. Surprisingly, even at the mean-field level, its phase diagram has not yet been fully determined, and the ground state magnetic properties are not completely known.

The Hubbard model was originally proposed to describe correlations between d-electrons in transition metals [12]. At half-filling (one electron per lattice site), it gives a simple description of the so-called Mott insulator, with antiferromagnetic (AFM) order. Soon after the discovery of high-\(T_c\) cuprate superconductors, it was pointed out that the two-dimensional (2D) Hubbard model might be an appropriate minimal model for high-\(T_c\) cuprates [13], because of the copper–oxygen plane geometry and the proximity of the superconducting transition to the AFM phase of undoped mother compounds. The 2D Hubbard model has since become a focal point of research in condensed matter and quantum many-body physics.

Recently, rapid experimental progress in optical lattice emulators [14] has promised a new way to approach Hubbard-like models. Using ultra-cold fermionic atoms trapped in periodic laser-field potentials, these highly controllable experiments are capable of potentially ‘simulating’ the
Hubbard model directly. Thus the properties of the Hubbard model are not only of importance theoretically but can also be of direct experimental relevance.

The Hartree–Fock (HF) method is the simplest paradigm to describe a quantum many-fermion system. The method finds the single Slater determinant wavefunction which minimizes the variational energy. As is well known, the mean-field approximation involved can turn out to be very severe. Nevertheless the HF method has often provided the foundation for our qualitative understanding of many systems in condensed matter and quantum chemistry. For example, HF correctly predicts an AFM order in the ground state of the Hubbard model at half-filling, even though the strength of the AFM order is overestimated and translational symmetry is (necessarily) broken. In quantum chemistry, HF is the starting point for most calculations and serves as the basis for understanding the electronic structure of many systems.

Because correlation effects (e.g. the correlation energy, which is a fundamental concept in the framework of density functional theory) are often defined using the HF solution as a reference, qualitative and quantitative understanding of the HF state is of key importance. This has not always been easy to achieve. For example, the nature of the unrestricted HF (UHF) state in the electron gas at high and intermediate densities was only recently determined [16].

In the Hubbard model, HF calculations are in principle straightforward. The 2D Hubbard model has been studied within the HF approximation in some of the pioneering works on high-$T_c$ superconductors. Inhomogeneous states have been found at small dopings, such as spin polarons [17], domain walls [18–20], and spin-density waves [21–23] (SDW), and phase diagrams have been proposed [1, 23, 24]. Due to computing power limitations, however, these studies have either done exact numerical calculations at only a few doping and interaction parameters [19, 20, 22], or have scanned parameters with restricted forms of the solution [23, 19, 24]. Furthermore, finite-size effects were difficult to remove, as we discuss below, which can mask the true solution. Furthermore, finite-size effects were difficult to remove, as we discuss below, which can mask the true solution. Due to particle–hole symmetry, we confine ourselves to denote the number of holes in the $\sigma$ channel. The total number of spin-$\sigma$ electrons is denoted by $N_{\sigma}$, and we assume that the system has no spin polarization, i.e. $N_{\uparrow} = N_{\downarrow}$. Under this assumption, the model has only two parameters, namely, the onsite repulsion $U$ and the doping $h \equiv 1 - (N_{\uparrow} + N_{\downarrow})/N \equiv N_{\text{hole}}/N$.

The remainder of the paper is organized as follows. In section 2, the self-consistent scheme used for solving the mean-field Hubbard model is summarized. The numerical results are presented in section 3, and analytic calculations are described in section 4. In section 5 the results are discussed and summarized in a phase diagram, and we conclude the paper in section 6.

2. Method

The Hamiltonian of the single-band repulsive Hubbard model reads

$$H = -t \sum_{\langle rr' \rangle, \sigma} (c^\dagger_{r\sigma} c_{r'\sigma} + c^\dagger_{r'\sigma} c_{r\sigma}) + U \sum_r n_{r\uparrow} n_{r\downarrow},$$

(1)

where $U > 0$ is the interaction strength and $t$ is the hopping amplitude between nearest neighbor sites (denoted by $\{rr\}$ in the summation). Throughout this work, energy is quoted in units of $t$ and we set $t = 1$. The operator $c_{r\sigma}^\dagger$ ($c_{r\sigma}$) creates (annihilates) an electron with spin $\sigma$ ($\uparrow$, $\downarrow$) at site index $r$, which runs through the lattice of size $N = L_x \times L_y$. The total number of spin-$\sigma$ electrons is denoted by $N_{\sigma}$, and we assume that the system has no spin polarization, i.e. $N_{\uparrow} = N_{\downarrow}$. Under this assumption, the model has only two parameters, namely, the onsite repulsion $U$ and the doping $h \equiv 1 - (N_{\uparrow} + N_{\downarrow})/N \equiv N_{\text{hole}}/N$.

$$h \equiv 1 - (N_{\uparrow} + N_{\downarrow})/N \equiv N_{\text{hole}}/N,$$

(2)

where we have used $N_{\text{hole}}$ to denote the number of holes in the system. Due to particle–hole symmetry, we confine ourselves to the region where $N_{\uparrow} + N_{\downarrow} \leq N$. Therefore the total density is given by $(n) = 1 - h$.

Standard linearization of equation (1) leads to the mean-field HF Hamiltonian:

$$H_{\text{HF}} = H_{\text{HF}}^\sigma + H_{\text{HF}}^L,$$

(3)

with

$$H_{\text{HF}}^\sigma = -t \sum_{\{rr\}} (c^\dagger_{r\sigma} c_{r'\sigma} + c^\dagger_{r'\sigma} c_{r\sigma}) + U \sum_r n_{r\sigma} \langle n_{r\sigma} \rangle - \frac{1}{2} \sum_r \langle n_{r\uparrow} \rangle \langle n_{r\downarrow} \rangle,$$

(4)

where $\tilde{\sigma}$ is the conjugate of $\sigma$ and $\langle n_{r\sigma} \rangle$ is an average density. The mean-field decoupling employed in equation (3)
assumes the z-axis as the quantization direction, thus breaking the spin rotational symmetry of the Hubbard Hamiltonian (1). After fixing the quantization orientation and requiring no spin polarization, the solution of the HF Hamiltonian is restricted to the $S^z = 0$ sector, i.e. spin textures in the $x$-$y$ plane, for instance spiral SDWs, are excluded. (At low $U$, the solutions turn out to be 1-SDWs. Then a single spiral cannot be the ground state, since a left-handed spiral can always be combined with a right-handed one, or vice versa, to make an 1-SDW which has lower energy [16].)

For a given set of parameters $(U, N, N_1, N_1^*)$, the HF Hamiltonian (3) is numerically solved using a self-consistent scheme. We begin the procedure by selecting a trial solution in the form of a single Slater determinant for each spin component:

$$\Phi_\sigma^{(0)} = \begin{pmatrix} \phi_1^{1\sigma} & \phi_2^{1\sigma} & \cdots & \phi_{N_\sigma}^{1\sigma} \\ \phi_1^{2\sigma} & \phi_2^{2\sigma} & \cdots & \phi_{N_\sigma}^{2\sigma} \\ \vdots & \vdots & \ddots & \vdots \\ \phi_1^{N_\sigma} & \phi_2^{N_\sigma} & \cdots & \phi_{N_\sigma}^{N_\sigma} \end{pmatrix}, \quad \text{(5)}$$

where each column is normalized to 1.

In the restricted HF (RHF) method, spin-$\uparrow$ and spin-$\downarrow$ parts of the total wavefunction are the same: $\Phi_\uparrow^{(0)} = \Phi_\downarrow^{(0)}$. The RHF method always gives the non-interacting solution in the system studied in this paper. In the UHF method, which is adopted here, $\Phi_\uparrow^{(0)}$ and $\Phi_\downarrow^{(0)}$ are allowed to differ and they converge via the coupled equation (4). The trial densities at site $r$ can be expressed as

$$\langle \rho_\sigma^{(0)} \rangle = \left[ \Phi_\sigma^{(0)} \left( \Phi_\sigma^{(0)} \right)^H \right]_{rr}, \quad \text{(6)}$$

where $'H'$ indicates conjugate transpose of the matrix and we have assumed that the orbitals in $\Phi_\sigma^{(0)}$ are orthonormal. An $N \times N$ matrix $\Sigma$ ($M^*$) for $\mathcal{H}_\text{HF}$ ($\mathcal{H}_\text{HF}$) is then constructed from the densities. By exactly diagonalizing $\Sigma^2$, we obtain the energy

$$E^{(1)}_\sigma = \sum_{i=1}^{N_\sigma} \lambda^{(1)}_{\sigma i}, \quad \text{(7)}$$

where $\lambda^{(1)}_{\sigma 1} < \lambda^{(1)}_{\sigma 2} < \cdots < \lambda^{(1)}_{\sigma N_\sigma}$ are the lowest $N_\sigma$ eigenvalues of $\Sigma^2$. The wavefunction $\Phi^{(1)}_\sigma$ is obtained by filling up $N_\sigma$ corresponding orbitals of $\lambda^{(1)}_{\sigma i}$. The new density $\langle \rho^{(1)}_\sigma \rangle$ is then calculated from $\Phi^{(1)}_\sigma$, which is used to update $M^* (M^*)$. We iterate this process until the total energy $E^{(t)} = E^{(t)}_\uparrow + E^{(t)}_\downarrow$ and the density $\langle \rho^{(t)}_\sigma \rangle$ is converged.

Care must be taken when updating the density during the iteration. As is typical in self-consistent algorithms, convergence to a fixed point is not guaranteed if $\langle \rho^{(t-1)}_\sigma \rangle$ is taken directly as an input for the $t$th step. To improve convergence, we adopt a mixing scheme: The $t$th input density is constructed as a linear combination of previous input and output densities as:

$$\langle \rho^{(t)}_{\sigma \text{in}} \rangle = (1 - \alpha) \langle \rho^{(t-1)}_{\sigma \text{in}} \rangle + \alpha \langle \rho^{(t-1)}_{\sigma \text{out}} \rangle, \quad \text{(8)}$$

where ‘in’ indicates the input density to construct $M^*$, and ‘out’ denotes the output density calculated by diagonalizing $M^*$. The mixing parameter $\alpha$ is typically chosen to be between $\sim 0.5$ and 0.75.

Due to non-linearity of the coupled equation (4), we implement two additional procedures to help the system reach the global minimum. Firstly, different initial wavefunctions are used and the consistency between the results is checked. Secondly, we perform multiple annealing cycles: in each cycle a random perturbation (whose strength can be controlled) is applied to the converged solution and the self-consistent process is repeated.

To reduce shell and one-body finite-size effects, we use twist-averaged boundary conditions (TABC) [25–27], under which the wavefunction $\Psi(r_1, r_2, \ldots)$ gains a phase when electrons hop around lattice boundaries

$$\Psi(\ldots, r_j + \mathbf{L}, \ldots) = e^{i\hat{\mathbf{L}} \cdot \hat{\Theta}} \Psi(\ldots, r_j, \ldots), \quad \text{(9)}$$

where $\mathbf{L}$ is the unit vector along $\mathbf{L}$, and the twist angle $\Theta = (\theta_x, \theta_y)$ is an input parameter which is randomly chosen in this work. For a given $\Theta$, the TABC is the same as a random shift of the momentum space grid. This reduces the discretization error in the integration. In the HF solution, the TABC is applied to each orbital, i.e. each column in equation (5). With a generic $\Theta$, there will be no degeneracy in the one-electron energy levels. We often average the results over many random twist angles [27] in each system to improve convergence to the thermodynamic limit. As can be seen from the energy results in section 3, this procedure produces a smooth curve versus doping, where the one-body finite-size effect is minimized. Additional finite-size errors, which result from the interaction and the formation of long wavelength collective modes [28], are not removed from this approach. We use rectangular lattices in our simulations to help detect the 1-SDWs with long modulating wavelengths, as discussed in section 3.

### 3. Numerical results

Various observables are computed with the converged UHF wavefunction. Two quantities examined throughout the paper are the charge-density (CD) $\rho(r)$ and the spin-density (SD) $s(r)$ defined as

$$\rho(r) \equiv \langle n_{\uparrow r} \rangle + \langle n_{\downarrow r} \rangle, \quad \text{(10)}$$

$$s(r) \equiv \langle \sigma_{\uparrow r} \rangle - \langle \sigma_{\downarrow r} \rangle. \quad \text{(11)}$$

We will also study the converged UHF eigenvalues $\lambda_{\sigma \mathbf{p}}$ and momentum distribution $n_{\mathbf{p} \sigma} = \langle c_{\mathbf{p} \sigma}^\dagger c_{\mathbf{p} \sigma} \rangle$, where $c_{\mathbf{p} \sigma}$ is, as usual, defined by the Fourier transform of $c_{\sigma r}$. Figure 1 illustrates the behavior of the reference system at $h = 0$, which is an AFM state with constant $\rho(r) = 1$.

In figure 2, CD in the 16 × 16 reference system is plotted as holes are doped into the lattice. As doping $h$ is varied, holes tend to cluster and form different patterns. These patterns have a strong $h$-dependence, which is a result of strong finite-size effects. Here the system is at an intermediate interaction strength of $U = 4.0$. As the interaction becomes weaker, we find that the variations in the patterns become larger and
depend sensitively on $\Theta$ (not shown). This is similar to what is seen in the UHF solution of an electron gas [16].

3.1. Linear spin-density wave (l-SDW) state

We first focus on low to moderate interacting strengths ($U < \text{half of the bandwidth}$) and small doping ($h \lesssim 0.1$), and examine the properties of the UHF solution as a function of doping $h$, i.e. as the system moves away from half-filling ($h = 0$). We will show that the UHF ground state at low and moderate $U$ is a linear spin-density wave (l-SDW) along the [01] direction. Figure 3 shows the results from a $32 \times 32$ supercell. An l-SDW is seen whenever the density is such that an l-SDW can be accommodated in the supercell. (The choice between $x$- and $y$-directions in the broken-symmetry UHF state is of course random. To help visualization in the figures, we have selected the same direction, either by an initial bias or by rotating the final result.) At incommensurate densities, strong finite-size effects are present, where the pattern of the cluster is not scalable to the thermodynamic limit. An example is seen by comparing $N_{\text{hole}} = 16$ in figure 2 (not long enough for one period of SDW) and $N_{\text{hole}} = 64$ in figure 3: in both cases $h = 1/16$. $N_{\text{hole}} = 24$ in figure 2 versus $N_{\text{hole}} = 96$ in figure 3 is another (both have $h = 3/32$). The finite-size effects will be further discussed below.

Although significantly larger lattice sizes are reached, the pattern variation clearly indicates that care must be taken in a numerical calculation, and additional ingredients are needed, in order to better approach the thermodynamic limit. We use two additional ingredients in our numerical simulations: TABC and rectangular supercells. To reduce the one-body finite-size effects, most of our results are averaged over $\sim 20$ random $\Theta$ values. In plots showing $\Theta$-averaged results, the statistical uncertainties from the twist angles are indicated by the error bars. The residual (two-body) finite-size effects are reduced by the use of rectangular supercells. This allows us to study longer wavelength modes without increasing the computational cost (compared to a square lattice of the same number of lattice sites, $N$). Obviously rectangular lattices break the symmetry between $x$- and $y$-directions, and can introduce an additional bias. To minimize the effect, we carry out calculations with different supercells with varying aspect ratios to check consistency in the results.

An illustrative set of results is shown in figure 4. We adjust $L_x$ and $L_y$ while keeping the size $N = L_x \times L_y = 32 \times 32$ fixed. An l-SDW solution is seen in a rectangular supercell whenever $L_y$ is sufficiently large to accommodate a wave.
Figure 3. Contour plots of SD for a supercell of $32 \times 32$ at $U = 3.0$ as $N_{\text{hole}}$ is increased. An l-SDW exists when the density is such that the supercell size is sufficient to accommodate the l-SDW, as shown in the left top or the right bottom plot.

Figure 4. Contour plots of CD for systems of fixed $N = L_x \times L_y = 32 \times 32 = 16 \times 64 = 8 \times 128 = 1024$ (from top to bottom) and fixed doping $h = 3/32$ ($N_{\text{hole}} = 96$) at $U = 3.0$. A stable l-SDW solution emerges when the supercell is commensurate. Note that only the accompanying CDW is shown here.

Note that the rectangular supercell does not bias the SDW in the $y$-direction (when $L_y > L_x$). An l-SDW is observed along the $x$-direction if $L_x$ is commensurate with the SDW wavelength. (An example of this is in figure 6 below, where the solution in the $20 \times 36$ lattice is two waves propagating along the $x$-direction.)
Figure 5. Contour plots of CD (left) and SD (right) versus doping. The system is an \(8 \times 64\) supercell at \(U = 2.0\) with doping of \(h = 1/32, 2/32\) and \(3/32\) (from left to right). The wavelength of the l-CDW is \(\lambda_{l-CDW} = 1/h\) and that of the l-SDW is \(\lambda_{l-SDW} = 2/h\).

From the results in figures 2 and 3, it is clear that the wavelengths of the l-SDW and l-CDW vary with doping \(h\). The results of an \(8 \times 64\) lattice with various values of \(h\) are shown in figure 5. As can be seen, the wavelength of the l-CDW/SDW decreases with \(h\). Unlike in figure 3, the lattice size in this case has been chosen so that \(L_y\) is commensurate with the wavelength in each figure. For example, there are exactly two CD waves at \(h = 1/32\), giving a wavelength of \(L_y/2 = 32(= 1/h)\). The wavelengths of SDW (right panel) are twice those of CDW. When the doping is doubled or tripled, the number of waves being accommodated changes accordingly, i.e. the wavelength shortens by 1/2 or 1/3, respectively. The modulating wavelengths of the l-CDW and l-SDW are thus given by

\[
\lambda_{l-CDW}(h) = \frac{1}{h}, \quad (12)
\]

\[
\lambda_{l-SDW}(h) = \frac{2}{h}. \quad (13)
\]

The wavelength relations are verified with many different choices of the aspect ratio.

The variational energy of the UHF ground state is examined in figure 6. A series of supercells are studied with a fixed \(N = L_x \times L_y = 720\) and \(h = 0.1\), while varying \(L_x\) and \(L_y\). It is seen that, for all supercell choices commensurate with the predicted wavelength, the energies are consistent and are lower. In systems which are incommensurate and cannot accommodate the l-SDW/CDW, the resulting ground state energies from the UHF solution are higher, indicating the frustration effect in the variational solution because of the finite size of the supercells. In section 4, we will present an analysis showing why in general the l-SDW is favored at low \(U\).

More lattice sizes at various dopings and interacting strengths are studied. The amplitudes of the l-SDW in the obtained solution are summarized in figure 7. It can be seen that at each fixed density, the l-SDW amplitude decreases as \(U\) is decreased and eventually vanishes, indicating the disappearance of the broken-symmetry UHF solution at a critical interaction strength \(U_c\). Below \(U_c\) only a RHF solution exists. The critical value \(U_c\) appears to decrease with \(h\) and approaches 0 at zero doping. This is consistent with the situation at half-filling (\(h = 0\)), where the Fermi surface (FS) is an open shell and a UHF state can be formed by ‘pairing’ [16] across it with no cost to the kinetic energy. For
As the interaction strength $U$ is further increased, the UHF ground state changes character. Figure 9 shows the CD and SD along the $y$-direction, along which the linear wave propagates. Above $U_c$, the amplitude of the 1-SDW (CDW) grows with $U$. As $U$ is further increased, the CD reaches 1 and starts saturating, creating deeper density valleys at the nodes of the 1-SDW. The maximum and minimum of CD and the 1-SDW amplitude as a function of $U$ are plotted in figure 10 to further illustrate this. As discussed in section 3.1, CD/SD orders are developed beyond $U \sim 1.5$. The 1-SDW amplitude is much greater than that of the 1-CDW. The CD maximum saturates at 1 above $U \sim 2.5$, indicating the formation of a linear stripe (l-stripes) state. The stripe or domain wall states differ from the SDW state because of CD saturation, forming hole-free domains that separate regions in which the holes are localized. The SDW state, in contrast, is a wave state in which the CD spatially oscillates but does not reach 1, and the holes are delocalized.

Thus at low dopings (high densities, $h \lesssim 0.1$), the 1-SDW state turns into an l-stripes state as $U$ is increased, with the l-stripes along the same direction ($x$- or $y$-) and having the same characteristic wavelength. When $U$ is further increased...
Figure 9. CD (top) and SD (bottom) along the y-direction versus $U$. The system being studied is an $8 \times 64$ supercell with doping of $1/32$ at $U = 1.0, 1.3, 1.5, 2.0, 4.0$. Each curve is a 1D cut in which the linear wave propagates. Beyond $U_c$, the l-CDW and l-SDW amplitudes increase with $U$ and the ground state ends up in an l-stripes state. The CDW amplitude is much weaker than that of the SDW.

Figure 10. Maximum and minimum of the CD and SDW amplitude for an $8 \times 64$ supercell with doping of 1/16.

the solution changes orientation, turning into a stripes state with modulation along the [11]-direction, a diagonal stripes (d-stripes) state.

At somewhat larger doping (0.1 $\lesssim h \lesssim 0.3$), the evolution of the l-SDW state with $U$ is different. The SDW state changes its modulation direction from the [10]-direction to diagonal. (d-SDW has been discussed in [22], for example.) Figure 11 shows an example for doping of $h = 1/6$. We see that the modulating wave changes direction from [10] at $U = 4.0$ to [11] at $U = 5.0$, and the d-SDW saturates to become a d-stripes state at $U = 9.0$.

We have scanned different parameter combinations to map out the sequence of the evolution of the UHF ground state. In section 5, a phase diagram is sketched to summarize the properties of the UHF ground state in the part of the phase space on which we have focused. The difference in the pairing mechanism of the d-SDW state from that of the l-SDW state is briefly discussed in section 4.4.

4. Analytic calculations

In this section we present a phenomenological model of the l-SDW state at low $U$ and small $h$. The model will help explain the numerical findings and provide a simple physical picture that captures the basic features of the exact UHF solutions in this parameter regime. The numerical studies are independent of the analysis here, but together they will give a more complete description of the UHF states. Below we first discuss the basic pairing model [29, 16], then carry out calculations in detail in the limit of small $U$ and $h$ for the l-SDW state, which is the focus of the present work. Some quantitative comparisons and validations of the pairing analysis are then presented, using the numerical data from calculations presented in section 3.1. We then briefly discuss the mechanism for d-SDW and d-stripes orders at higher $U$.

4.1. Pairing model

At low $U$, the region of interest in momentum space is the immediate vicinity of the FS, where pairing effects of electrons determine the nature of the UHF solution. (Often the effect has been discussed in the context of nesting. We refer to the mechanism as pairing since, although nesting greatly facilitates pairing in the Hubbard model, it is not required for the pairing mechanism to be realized, as seen in the electron gas [16].) In the fully filled region inside the FS, the electron density is uniform:

$$n_{p\sigma}(p) = \tilde{n}_\sigma = \frac{N}{4\pi^2}. \quad (14)$$

We first specify the pairing mechanism [16] more explicitly. Recall that the non-interacting energy for the state $|p\rangle$ is

$$\epsilon_p = -2(\cos p_x + \cos p_y). \quad (15)$$

The plane-wave state is $|p\rangle = \frac{1}{\sqrt{N}}e^{ip\cdot r}$, with $r = (x, y)$, where $x$ and $y$ are integer coordinates denoting lattice sites. Consider a pair of spin-$\uparrow$ and spin-$\downarrow$ electrons in a $p$ state, where $\epsilon_p \leq \epsilon_F$, with $\epsilon_F$ the Fermi energy. In the pairing model [16], this pair is made to partially occupy a $p'$ state:

$$|\uparrow\rangle = u_p|p\rangle + v_p|p'\rangle, \quad |\downarrow\rangle = u_p|p\rangle - v_p|p'\rangle. \quad (16)$$

where $|u_p|^2 + |v_p|^2 = 1$. This pair gives the following contribution to the local density in real space:

$$n_\uparrow(r, p) = \frac{1}{N}(1 + 2|u_pv_p|\cos(|p' - p| \cdot r)), \quad (17)$$

$$n_\downarrow(r, p) = \frac{1}{N}(1 - 2|u_pv_p|\cos(|p' - p| \cdot r)). \quad (18)$$

And an SDW state will result from the state in equation (16), with local spin

$$s(r, p) = n_\uparrow(r, p) - n_\downarrow(r, p) = \frac{4}{N}|u_pv_p| \cdot \cos(|p' - p| \cdot r). \quad (19)$$
Figure 11. Contour plots of CD (top) and SD (bottom) versus interacting strengths. The system being studied is a 36 × 36 supercell with doping of $h = 1/6$ at $U = 4.0, 5.0$ and $9.0$ (from left to right), representing l-SDW, d-SDW and d-stripes state respectively.

The SDW state lowers the interaction energy contribution of the pair compared to the non-interacting solution (i.e. the solution when $v_p = 0$) by the amount:

$$\Delta \mathcal{E}_V(p) = U \sum_r \left[ n_{\uparrow}(r, p) n_{\downarrow}(r, p) - \frac{1}{N_N} \right]$$

$$= -\frac{U}{4} \sum_r s^2(r, p).$$

(20)

If we have multiple pairs each formed as in equation (16), the change in interaction energy follows the same relation:

$$\Delta \mathcal{E}_V = -\frac{U}{4} \sum_r s^2(r) = -\frac{U}{4} \sum_r \left[ \sum_p s(r, p) \right]^2.$$  (21)

where the sum over $p$ is over all pairing plane waves (one of the four sides is illustrated by striped areas in figure 12).

At half-filling, the shell at the Fermi level, i.e. on the border of the diamond, is open, with the number of degenerate $p$ states equal to twice the number of spin-$\uparrow$ (or spin-$\downarrow$) electrons that need to be accommodated. Pairing can be achieved by choosing $p' - p = q_0 = (\pi, \pi)$, i.e. having electrons occupy two states in the open shell across the FS. This is perfect nesting and the SDW formed has perfect AFM order. Because pairing occurs in the open shell at the FS, the reduction in interaction energy from the SDW has no penalty, i.e. no increase in the kinetic energy.

4.2. The linear spin-density wave state

We next consider the case of low $U$, slightly doped ($U \ll$ bandwidth), to help understand the mechanism of the l-SDW state. As the FS shrinks with small doping, we assume that it remains approximately the shape of a diamond. The distance between the FS at half-filling and the doped FS is determined by

$$d = \frac{\sqrt{2}}{4} h \pi.$$  (22)

Figure 12. Illustration of the pairing model at small $U$ and $h$. The half-filling FS is the large diamond (red dashed). The non-interacting FS at low doping remains approximately the shape of a diamond (blue solid). AFM order arises from $q_0$, the pairing vector across the half-filling FS. The pairing vector is $q$ across the doped FS. The difference between $q_0$ and $q$, $\Delta q$, determines the characteristic modulating wavelength of the l-SDW.

As the interaction is turned on, it can become advantageous for some of the electrons near the FS to be partially excited. Partially occupied states around the FS can then allow pairing across the FS, which causes a correlation between electrons of opposite spins to generate an SDW. The presence of the SDW will lower the interaction energy. However, in this more general case there will also be an increase in the kinetic energy. When the lowering of the interaction energy surpasses the increase in kinetic energy...
an overall lower energy state is found compared to the free-electron (or RHF) solution.

We first determine the kinetic energy change. At low \( U \), pairing occurs near the FS. Electrons from a small region immediately inside the FS are excited. As a crude model [29, 16], we assume that a fraction \( f \) of the electrons within a distance \( \delta \) of the FS are excited, as illustrated by the horizontally striped region in figure 12. The excited electrons occupy the region (vertically striped) immediately above the FS, also of thickness \( \lambda \). We take \( \varepsilon_p \) and \( \varepsilon_p \) in the pairing state in equation (16) to be independent of \( p \); \( \varepsilon_p = u \) and \( \varepsilon_p = v \). Thus the vertically striped area has uniform density, and \( f = |v|^2 \). An upper bound to the kinetic energy increase due to this process is easily estimated. It is, for each excited electron, given by:

\[
\Delta E_K(p) = \nabla v^p \cdot \Delta p,
\]

where \( \Delta p = \delta(1,1)\sqrt{2}/2 \). The total kinetic energy increase is then

\[
\Delta E_K = 8f/\pi \int S \Delta E_K(p) dS = N\delta^2 \pi^2 8f \left[ 1 + \cos \frac{h\pi}{2} \right].
\]

(24)

where \( S \) in the integral is over the horizontally striped area inside the FS, and the factor of 8 accounts for the 4 sides and 2 spin species.

We now determine the interaction energy change, and show that the optimal SDW is along the \( y \)- or \( x \)-direction. From equation (21) we see that the maximum reduction is achieved by maximizing the quantity

\[
I_V = \sum_{\mathbf{q}} \left( \sum_{\mathbf{r}} \cos(\mathbf{q} \cdot \mathbf{r}) \right)^2,
\]

where the sum over \( \mathbf{q} \) is over all pairing states, with \( \mathbf{q} = \mathbf{p'} - \mathbf{p} \). This is realized if all the electron pairs line up their pairing vectors. There are two groups of pairing states, corresponding to the two diagonal directions. Within each group, the optimal choice is for all pairs to have one common pairing wavevector \( \mathbf{q} \). Let us denote the pairing wavevectors along \([11]\) and \([-11]\) by \( \mathbf{q} \) and \( \mathbf{q}' \), respectively, and write: \( \mathbf{q} = (\pi, \pi) - \Delta \mathbf{q} \) and \( \mathbf{q}' = (-\pi, \pi) - \Delta \mathbf{q}' \). We then obtain:

\[
I_V \propto \sum_{\mathbf{q}} \left( \sum_{\mathbf{r}} \cos(\mathbf{q} \cdot \mathbf{r}) \right)^2 N + \sum_{\mathbf{q}} \left( \sum_{\mathbf{r}} \cos(\mathbf{q} + \mathbf{q}') \cdot \mathbf{r}) + \cos(\mathbf{q} - \mathbf{q}') \cdot \mathbf{r} \right).
\]

(26)

The maximum is achieved in equation (26) when \( \Delta \mathbf{q} = \pm \Delta \mathbf{q}' \). This occurs when \( \mathbf{q} \) and \( \mathbf{q}' \) are such that the SDW modulation from the two groups of pairing states are the same, leading to a positive ‘interference’ between them. The direction of the modulating wavevector must be along \([01]\) or \([10]\). The magnitude is given by

\[
|\Delta \mathbf{q}| = 2\sqrt{2} \delta = h\pi.
\]

(27)

as illustrated in figure 12. This leads to the following total reduction in interaction energy:

\[
\Delta E_V = -\frac{N\delta^2}{\pi^2} 4|u|^2 fU \left( 1 - \frac{h}{2} \right)^2.
\]

(28)

Thus the lowest energy state is an \( l \)-SDW with broken \( x-y \) symmetry, with the modulation along either the \( x \)- or the \( y \)-direction. The modulating wavelength is \( \lambda_{l-SDW} = 2/h \), consistent with our numerical result.

To reach an SDW state of lower energy than the non-interacting solution, the condition

\[
|\Delta E_V| \geq |\Delta E_K|
\]

must be satisfied. From equations (24) and (28), we obtain

\[
|u|^2 U \left( 1 - \frac{h^2}{2} \right)^2 \geq 2 \left[ 1 + \cos \frac{h\pi}{2} \right].
\]

(30)

Taking \(|u| \sim 1\) on the left-hand side, we obtain a rough estimate to the critical value which \( U \) must exceed:

\[
U_c = \frac{2[1 + \cos \frac{h\pi}{2}]}{(1 - \frac{h^2}{2})}.
\]

(31)

The absolute value of the kinetic and interaction energy changes in equations (24) and (28) are plotted versus \( h \) in the left panel in figure 13. \( \Delta E_K \) is independent of \( U \), while \( \Delta E_V \) is proportional to \( U \), for which several curves are plotted for various values of \( U \). It is seen that a critical value of \( U \) exists for a doped system \( (h \neq 0) \). Above \( U_c \), the two curves cross at a critical \( h_c \), below which the broken-symmetry l-SDW state exists. As \( U \) increases, the point of crossing, \( h_c \), moves to the right. Equivalently, the critical \( U_c \) decreases as doping is reduced. In the right panel the curve of \( U_c \) versus \( h \) is plotted to illustrate this.

4.3. Comparison with numerical results

The simple model and analysis above capture most of the properties of the exact UHF ground state at low \( U \) and small \( h \). It gives the correct \( l \)-SDW modulating wavelength, and explains the existence of \( U_c \) and how it varies with doping. Because of the crudeness of the model, the values of \( U_c \) and other quantitative features are not very accurate compared to the exact numerical results in section 3.1. Larger discrepancies can be expected further away from its domain of validity, namely small doping and modest interaction (although it also incorrectly predicts \( U_c = 4 \) as \( h \to 0 \)).

The model considers only pairing of two electrons, so CDW is excluded. This is consistent with the numerical result that at low \( U \), CDW is much weaker than SDW order. The exact UHF solution will necessarily involve more electrons in the pairing [16], which will lead to a larger energy lowering \( |\Delta E_V| \) (and thus lower \( U_c \)) and will result in CDW, as observed in the numerical results.

Figure 14 shows the modification to the RHF band structure in the UHF solution as a function of interaction strength. The difference between the UHF eigenvalue \( \lambda_{\mathbf{p}} \) and the RHF spectrum \( \epsilon_p \) in equation (15) is plotted for all momentum values \( \mathbf{p} \). As discussed in figure 8, the eigenvalue \( \lambda_{\mathbf{p}} \) is identified with the momentum \( \mathbf{p} \) with which the corresponding eigenstate has the maximum overlap. We see that, just above \( U_c \), a small fraction of the states on the FS are involved in pairing, which creates a small energy lowering.
Figure 13. Absolute values of kinetic energy gained and interaction energy lost in the pairing model. On the left the energies are plotted as a function of $h$ for several values of $U$. On the right, $U_c$ is plotted versus $h$.

Figure 14. Energy plot of the modification to the RHF band structure in the UHF solution, for a sequence of $U$ values in the $l$-SDW regime. Shown are the values $\lambda_{\sigma p} - \epsilon_p$ versus $p$, where $\epsilon_p$ is given in equation (15). The system is a $16 \times 48$ supercell with doping of $h = 1/24$.

that leads to the UHF solution. The plot is for a single twist angle. In a finite system, the shift in momentum space from the twist creates a small asymmetry between each pair of surfaces diagonally opposite. At small $U > U_c$, this is reflected in the solution as an asymmetry in the gaps on the two surfaces. As $U$ increases, excitation spans a wider region at the FS, and the gap structure from pairing becomes more pronounced.

The momentum distribution from a numerical UHF solution at $h = 1/12$ is shown in figure 15. The left panel plots $n(p)$ minus the non-interacting value $n_0(p)$: $\Delta n(p) = n(p) - n_0(p)$. Electrons are excited from the darker area to the lighter. The right panel shows the two-point correlation function from the left panel: $\Delta n(p)\Delta n(p')$ versus $(p - p')$. Negative peaks are seen at $(\pm \pi, \pm(\pi - \pi/12))$ on the right, which result from the pairing between the negative just inside the FS (where electrons are excited from) and the positive immediately above the FS (where electrons are excited to) in the left panel. The position of the negative peaks indicates a
Figure 15. Density plots of $\Delta n(p)$, the momentum distribution difference from RHF solution (left) and its correlation $\Delta n(p)\Delta n(p')$ (right). The system is a $16 \times 24$ supercell with doping of $1/12$ at $U = 3.0$. Negative peaks at $(\pm \pi, \pm (\pi - \pi/12))$ in the correlation result from the pairing.

Figure 16. Illustration of the pairing scheme for d-SDW order. The left panel shows $n(p)$ and the right panel $\Delta n(p)$, the difference from the non-interacting solution. The momentum distribution is actual numerical data from a system of $36 \times 36$ with doping of $1/6$ at $U = 5.0$. Electrons are excited from the FS across the $(-\pi, \pi)$-direction to the FS across the other direction $(\pi, \pi)$, such that the FS along the latter reaches the half-filling FS. This enables two groups of pairings to maintain interference, with $\Delta q = \Delta q'$, to lower the energy.

The system is a $16 \times 24$ supercell with doping of $1/12$ at $U = 3.0$. Negative peaks at $(\pm \pi, \pm (\pi - \pi/12))$ in the correlation result from the pairing.

4.4. Diagonal spin-density wave states

As mentioned before, diagonal modulations lose the interference between $[11]$ and $[-11]$, so a diagonal (or any orientation other than $[10]$ and $[01]$) SDW is not the solution at small $h$ and moderate $U$. This does not exclude it as a solution as we move away from this parameter regime, when the distortion to the FS becomes more severe.

This situation happens when the doped FS is deformed sufficiently away from the half-filling shape of a diamond and the area of excitation becomes sufficiently large to reach the half-filling FS. The number of pairs that could participate in the ‘interference’ of the l-SDW is decreased, because the FS no longer has the shape of a diamond. Eventually it becomes energetically more favorable to have the FS be longer in one diagonal direction than the other, i.e. to break the four-fold rotational symmetry. As illustrated in figure 16, it is then possible to create two different types of pairing states along the two diagonal directions, such that they share a common modulating wavevector along one diagonal direction: $\Delta q = \Delta q'$. The two groups of pairs will achieve interference, similar to the case of l-SDW. As in section 4, the pairing vector is determined by $h$, giving $\Delta q = (h\pi, h\pi)$, which gives rise to an SDW with modulating wave along $[11]$-direction, and of wavelength $\lambda_{d-SDW} = \sqrt{2}/h$. The corresponding wavelength for d-CDW is $1/\sqrt{2}h$. This is consistent with the numerical results in section 3.2.

5. Discussion

We can now place our 2D results in the context of an HF phase diagram for the Hubbard model. Our numerical calculations have focused on small and intermediate dopings ($h$ from 0 to $\sim 0.3$), and small to moderate interactions ($U$ from 0 to $\sim 10$), because of possible connections with the many-body ground state at moderate interacting strengths. The analytic calculations are for small $h$ and low $U$, where our pairing model captures the physics in the HF framework. Our numerical results are sufficiently detailed such that we could determine some phase boundaries as shown in figure 17. We fitted the numerical locations for the phase transition or crossover using power functions, except for the AFM to FM transition which was fitted by an exponential. Because of the limited number of data points and the finite resolution with which the transition was scanned, there are significant pairing vector of $\Delta q = (0, h\pi)$, consistent with the pairing vector in the analytical model.
uncertainties in the fits, of several line widths or larger. The phase boundaries are thus only meant as rough guides.

At half-filling, the UHF solution is an AFM state. Upon doping, there is a phase boundary \( U_c(h) \), shown as the blue line in figure 17, below which is the PM phase. Above \( U_c(h) \) is an AFM region where a rich set of sub-regions exhibit different characters, including the l-SDW states we have focused on in this work; we describe this region in further detail below. Above the AFM phase is an FM phase. Our numerical UHFs show that the FM state has lower energy above the green solid line. The RHF approach, naturally, predicts an earlier transition to FM. This is the theoretical phase boundary from the Stoner criterion, and is shown as the black dotted line. Recall that we have excluded spiral SDWs. As we discussed, this is not the ground state at low \( U \) (see also [19, 23]). However, at large \( U \), spiral orders can become more favorable deep in the d-stripes region.

Between the PM and FM phases is the AFM phase. In this region, at low and intermediate \( U \), we see an l-SDW state with a long wavelength modulation along the [10]-direction; a weaker CDW accompanies the SDW. Near half-filling, as \( U \) is increased the l-SDW state evolves into a l-stripes state which shares the same characteristic wavevector as the l-SDW, but whose CD saturates to 1 in regions separated by 'stripes' anchored by the nodal positions defined by the SDW. The holes are localized in these stripes. This is consistent with the observation in [22] of SDW deforming into domain walls with increasing \( U \). The transition from delocalized holes (such as the l-SDW state) to localized holes is denoted by the red dashed line in figure 17. As we move further away from half-filling, the l-SDW at lower interaction changes its direction of modulation as \( U \) is increased. This forms a d-SDW state. The transition from a state with modulation along the [10]-direction to one with diagonal modulation is denoted by the cyan dot-dashed line. We see that the two dashed lines cross each other. At low doping (\( h \lesssim 0.1 \)), the system reaches an l-stripes state first before changing the direction of modulation to a d-stripes state. At higher doping, the order is reversed. The l-SDW first changes into a d-SDW state. As \( U \) is further increased, density saturation appears, and holes become localized in a d-stripes state.

It is important to keep in mind that the results we have discussed and the phase diagram above are for HF theory. For strong interactions in particular, the HF results are expected to be severely biased and correlation effects can fundamentally change the nature of the many-body state. For example, the FM phase was shown not to exist at low density (\( h > 0.5 \)) in the 3D Hubbard model [30].

The present work was in part motivated by a recent quantum Monte Carlo (QMC) calculation [28] which indicated that the ground state of the 2D Hubbard model has a long wavelength SDW collective mode. Upon doping, the AFM order at half-filling was found to evolve into an SDW state with a long wavelength modulation which has essentially a constant charge–charge correlation at low to intermediate interaction strengths. Given that the UHF solution is qualitatively correct at half-filling, it was natural to ask to what extent the UHF solution contains any of these features upon doping.

We see from the numerical results in this work that the UHF solution appears to qualitatively capture the basic features of the magnetic correlations in the ground state upon doping, as it does at half-filling. Of course the UHF solution gives a static modulated SDW, while the many-body ground state in the QMC preserves translational invariance and the SDW correlation is only seen in the correlation functions [28]. This is similar to the situation at half-filling.

In the UHF solution, the tendency for the holes to localize is much overestimated. This was part of the reason to focus on low \( U \) in the present study. A CDW correlation almost always accompanies the SDW in the UHF solution, and holes appear to localize (leading to domain walls or stripes) at \( U \sim 4 \). In contrast, holes remain delocalized (wave-like) in the many-body solution [28], with essentially constant charge–charge correlation, until the strong interaction regime (\( U \gtrsim 10 \)). It is an interesting question whether diagonal order, which is present in the HF solution at larger \( U \), is present in the true many-body ground state.

The UHF solution thus provides a useful starting point for understanding the magnetic and charge correlations in the ground state of the Hubbard model at intermediate interactions. In addition, the ability to reliably determine the true UHF ground state numerically could prove valuable in QMC calculations, which often require a trial wavefunction and where the qualitative correctness of the trial wavefunction can make a significant difference. Although the physics in the UHF solution is sensitive to the particular many-body Hamiltonian, the basic approach we have used and the
6. Conclusion

In summary, we have performed exact numerical calculations for the UHF ground state of the Hubbard model systematically for a wide range of lattice sizes, initial conditions, doping and interaction strengths. Special care has been taken to reduce finite-size effects in order to obtain the solution at the thermodynamic limit. These results allow us to map out the magnetic phase diagram for regimes most relevant in modeling condensed matter systems.

A broken-symmetry UHF solution exists above a critical $U_c(h)$, whose value increases with doping. Above $U_c(h)$, the ground state is a static l-SDW/CDW, with a modulation whose wavelength is inversely proportional to doping at small $h$. The amplitude of the SDW/CDW decreases with $h$ and increases with $U$. At low $U$, the SDW amplitude is much stronger than that of the accompanying CDW, and the holes are essentially delocalized. For larger $U$, the SDW and CDW amplitudes become more comparable. At small doping, the solution turns into the l-stripes state with the same characteristic modulating wavevector and holes localized at the nodal positions, before eventually entering the d-stripes state. At larger doping, the l-SDW state first turns into the d-SDW state before eventually entering the d-stripes state at larger interactions.

We have also presented an analytic theory to explain the mechanism for the formation of the SDW state. The model provides a conceptual understanding of the physics of SDW which can be applied in systems beyond the 2D Hubbard model. Comparison with recent QMC results shows that the UHF solution captures the magnetic correlations in the true many-body ground state at intermediate interactions.

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