A Green’s function decoupling scheme for the Edwards fermion–boson model

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Abstract

Holes in a Mott insulator are represented by spinless fermions in the fermion–boson model introduced by Edwards. Although the physically interesting regime is for low to moderate fermion density, the model has interesting properties over the whole density range. It has previously been studied at half-filling in the one-dimensional (1D) case by numerical methods, in particular using exact diagonalization and the density matrix renormalization group (DMRG). In the present study the one-particle Green’s function is calculated analytically by means of a decoupling scheme for the equations of motion, valid for arbitrary density in 1D, 2D and 3D with fairly large boson energy and zero boson relaxation parameter. The Green’s function is used to compute some ground state properties, and the one-fermion spectral function, for fermion densities \( n = 0.1, 0.5 \) and 0.9 in the 1D case. The results are generally in good agreement with numerical results obtained using the DMRG and dynamical DMRG, and new light is shed on the nature of the ground state at different fillings. The Green’s function approximation is sufficiently successful in 1D to justify future application to the 2D and 3D cases.

1. Introduction

Hubbard-like models provide a paradigm for a large class of strongly correlated systems. A general form for the Hubbard Hamiltonian is

\[
H_{\text{Hu}} = T + U = - \sum_{r,\rho, a, b} t_{\rho a b} c_{r+\rho a}^\dagger c_{ra} + U \sum_r n_{r+} n_{r-}
\]

(1)

and the system is strongly correlated when the repulsive on-site interaction \( U \) is considerably larger than the hopping parameters \( t \). Here \( c_{ra} \) destroys an electron in state \( a \) on lattice site \( r \) and \( c_{r+\rho a}^\dagger \) creates an electron in state \( b \) on a nearest-neighbour site \( r + \rho \). The state indices \( a, b \) are summed over two states denoted by + and −, and the occupation numbers \( n_{r+} = c_{r+}^\dagger c_{r+} \). In general we consider bipartite lattices in one, two and three dimensions. The standard Hubbard model [1] corresponds to the case of a single orbital on each site with states ± corresponding to spin ±\( \frac{1}{2} \) and with hopping parameter

\[
t_{\rho a b} = t_0 \delta_{\rho 0}.
\]

(2)

This model with a 2D square lattice is frequently used to describe the copper–oxygen plane of high-\( T_c \) systems such as doped \( \text{La}_2\text{CuO}_4 \), where the orbital corresponds to a \( \text{Cu} d_{x^2−y^2} \) orbital. A related model on a 2D square lattice in the \( xy \) plane describes a ferromagnetic system in which the on-site states all have the same spin and the states + and − correspond to \( d_{z^2−r^2} \) and \( d_{x^2−y^2} \) orbitals respectively. This ‘\( t_{2g} \) model’ describes ferromagnetic planes in \( \text{Sr}_2\text{VO}_4 \) and also in fluorides such as \( \text{K}_3\text{CuF}_4 \) and \( \text{Cs}_2\text{AgF}_4 \), where in this case a crystal field converts \( e_g \) orbitals into an effective \( t_{2g} \) system [2–6]. In the \( t_{2g} \) model one considers only the dominant hopping processes in which hopping between + states occurs along the \( x \) axis and hopping between − states occurs along the \( y \) axis. Thus in the \( t_{2g} \) model

\[
t_{\rho a b} = t_0 \delta_{\rho 0} |\hat{\rho} \cdot \mathbf{e}_x|,
\]

(3)

where \( \mathbf{e}_x \) and \( \mathbf{e}_y \) are unit vectors along the \( x \) and \( y \) directions respectively and \( \hat{\rho} \) is a unit vector in the nearest-neighbour direction \( \rho \).

In the large-\( U \) limit we can approximate the Hamiltonian \( H_{\text{Hu}} \) by an effective Hamiltonian \( H_{\text{eff}} \) which is defined to act only within a subspace where there is no double occupation of any site. \( H_{\text{eff}} \) takes the form

\[
H_{\text{eff}} = T - \frac{T^2}{U},
\]

(4)
where \( T_1 \) is that part of \( T \) which has matrix elements linking states of no double occupation with states having a single doubly occupied site. If three-site terms are neglected it is straightforward to write the second term of equation (4) in terms of spin (pseudo-spin for the t2g model) operators \( S^i_r = (n_{r+} - n_{r-})/2 \), \( S^x_r = S^z_r \pm i S^y_r = c^\dagger_{r\uparrow} c^\dagger_{r\downarrow} \) and number operator \( n_r = n_{r+} + n_{r-} \). Then, for the Hubbard model, \( \mathcal{H}_{\text{eff}} \) becomes

\[
\mathcal{H}_{\text{eff}} = T_{\text{1-j}} + \frac{1}{2} J \sum_{\langle r, \rho \rangle} (S^i_{r\uparrow} S^i_{r\rho} + \frac{1}{2} (S^z_{r\uparrow} S^z_{r\rho} + S^y_{r\uparrow} S^y_{r\rho}))
\]

where \( J = 4t_0^2/U \) and \( T_{\text{1-j}} \) contains the hopping parameters of equation (2) [7]. For the t2g model the corresponding Hamiltonian is

\[
\mathcal{H}_{\text{t2g}} = T_{\text{1-g}} + \frac{1}{2} J \sum_{\langle r, \rho \rangle} (S^i_{r\uparrow} S^i_{r\rho} - \frac{1}{2} n_r n_{r\rho}),
\]

where \( T_{\text{1-g}} \) contains the hopping parameters of equation (3) [2]. \( \mathcal{H}_{\text{1-j}} \) is known as the t-J model and if the transverse exchange term in equation (5) is omitted we have the t-J model. This differs from \( \mathcal{H}_{\text{t2g}} \) where \( + \) and \( - \) spins only hop along the \( x \) and \( y \) axes respectively. For the case of one electron per site, the undoped case for the oxides being modeled, \( T \) drops out and the system is a Mott insulator. In the Hubbard case the insulator is a Heisenberg antiferromagnet and in the t2g case it is an Ising antiferromagnet with alternating orbital order in the ground state. The transverse exchange terms in equation (5) are not present in equation (6) owing to the directional hopping of the orbitals in the t2g case.

When a hole is introduced into the lattice its motion disturbs the spin or orbital order of the ground state. At each hop through the ordered lattice the hole leaves a spin, or orbital, deviation at the site it vacates. Thus as the hole hops through the lattice the energy of the system increases linearly and the hole is bound to its starting point. This is known as the string effect and it exists in dimensions higher than one [8]. In 1D the spin deviation created by a hole at its first hop, from some initial position, increases the energy but subsequent hops in the same direction merely shift the ordered spin configuration by a lattice spacing without further increase in energy. There is therefore no string effect in the 1D t-J or t-J models. Furthermore in the 2D or 3D t-J models the string effect is relaxed by the exchange terms between lattice sites and can lead to a healing of the spin deviations created by the hole. Thus the hole can move at a speed determined by the healing rate, which leads to a quasiparticle band of width proportional to the exchange parameter \( J \) [9]. This way of relaxing the string effect is not possible in the t-J or the \( \mathcal{H}_{\text{t2g}} \) model owing to the absence of the \( S^i_{r\uparrow} S^i_{r\rho} \) terms. In these models a relaxation mechanism can be introduced by including in the Hamiltonian a term of the form

\[
\sum_{\langle r, \rho \rangle} (S^i_{r\uparrow} + S^i_{r\rho}) = 2 \sum_{\langle r, \rho \rangle} S^x_r
\]

so that the spin part of equation (6) becomes a transverse-field Ising antiferromagnet. In the physical case of the t2g model this transverse field corresponds to an on-site crystal field which mixes the two t2g orbitals just as the transverse magnetic field mixses + and − spins. In both the t-J and the \( \mathcal{H}_{\text{t2g}} \) model inclusion of three-site terms mentioned earlier also relaxes the string effect [2] but we shall not include them here. A hole in the 2D t-J model can also propagate by means of a Trugman path [10] which consists of six hops around a four-site square plaquette. The hole moves to a next-nearest-neighbour site and leaves the antiferromagnetic spin arrangement undisturbed. This cannot occur in the \( \mathcal{H}_{\text{t2g}} \) model owing to the directional hopping [2]. We shall here concentrate on the t-J model and the t-J model in a transverse field although there should be no difficulty in treating the directional hopping of the \( \mathcal{H}_{\text{t2g}} \) model.

Our reference state has one electron on each site with spins (or pseudospins) ordered as in an antiferromagnet. We follow Martinez and Horsch [11] in introducing a spinless fermion operator \( f^\dagger_r \), which creates a hole in the reference state at site \( r \) and a boson operator \( b^\dagger_r \) which creates a spin reversal on site \( r \). Thus \( b^\dagger_r = S^z_r \) for site \( r \) on the + spin sublattice and \( b^\dagger_r = S^z_r \) for \( r \) on the − sublattice. Clearly each nearest-neighbour hop of the hole through the ordered lattice reverses the spin on the site vacated by the hole. Thus the hopping operator \( T_{\text{1-j}} \) may be written as

\[
T_{\text{1-j}} = -t_0 \sum_{\langle r, \rho \rangle} (f^\dagger_{r\rho} f_r + \text{H.c.}).
\]

For the t2g model the corresponding hopping operator \( T_{\text{1-g}} \) is

\[
T_{\text{1-g}} = -t_0 \sum_{\langle r, \rho \rangle} (f^\dagger_{r\rho} f_r + \text{H.c.}) \langle \hat{\rho} \cdot \hat{e}_a \rangle.
\]

Since \( S^z_r = \pm (\frac{1}{2} - b^\dagger_r b_r) \) for \( r \) on the ± spin sublattice we may write the Ising part of the exchange term, in the presence of the hole, as

\[
-\frac{1}{2} J \sum_{\langle r, \rho \rangle} (1 - h_r)(1 - 2b^\dagger_r b_r)(1 - 2b^\dagger_{r\rho} b_{r\rho})(1 - h_{r\rho})
= \frac{1}{2} J \sum_{\langle r, \rho \rangle} b^\dagger_r b_r (1 - h_{r\rho} - b^\dagger_{r\rho} b_{r\rho}).
\]

where \( h_r = f^\dagger_r f_r \) and an irrelevant constant term has been dropped. Here we have used \( h_r b^\dagger_r b_r = 0 \) since the hole and a spin deviation cannot occupy the same site. A spin deviation created on site \( r \) when the hole vacates that site will in general have one neighbouring site \( r + \rho \) occupied by the hole and another by a spin deviation created when the hole arrived on site \( r \). Thus the expression in equation (10) may be written as

\[
\frac{1}{2} J (z - 2) \sum_r b^\dagger_r b_r,
\]

where \( z \) is the number of nearest-neighbours. \( z = 2 \) in 1D so that the above term is zero, which is consistent with the absence of the string effect in the 1D t-J model. Apart from an additive constant the t-J Hamiltonian equation (5) becomes

\[
\mathcal{H}_{\text{t-J}} = -t_0 \sum_{\langle r, \rho \rangle} (f^\dagger_{r\rho} f_r + \text{H.c.})
+ \frac{1}{2} J \sum_{\langle r, \rho \rangle} (b^\dagger_r b^\dagger_{r\rho} + b_r b_{r\rho})(1 - 2\frac{1}{2} J (z - 2) \sum_r b^\dagger_r b_r.
\]
The derivation of this Hamiltonian in [11] gives additional constraints which we will not discuss here.

The relaxation of the string effect occurs when bosons, created by fermion hopping, are spontaneously destroyed in nearest-neighbour pairs by the terms $b_y b_{y+\rho}$. Edwards [12] introduced a simplified model Hamiltonian of the form

$$\mathcal{H}_{\text{Ed}} = -t_0 \sum_{\mathbf{r}\mathbf{\rho}} (f_{\mathbf{r}+\mathbf{\rho}}^\dagger f_{\mathbf{r}} + \text{H.c.}) - \lambda \sum_{\mathbf{r}} (b_{\mathbf{r}}^\dagger b_{\mathbf{r}}) + \omega_0 \sum_{\mathbf{r}} b_{\mathbf{r}}^\dagger b_{\mathbf{r}} + N\lambda^2/\omega_0$$

(13)

in which boson relaxation terms $b_y b_{y+\rho}$ are replaced by the simpler linear ones $b_{\mathbf{r}}$. A unitary transformation $\mathcal{H}_{\text{Ed}} \rightarrow e^{\lambda/\omega_0} \mathcal{S} \mathcal{H}_{\text{Ed}} e^{-\lambda/\omega_0}$, in the absence of fermions, results in $b_{\mathbf{r}} \rightarrow b_{\mathbf{r}} + \lambda/\omega_0$, yields

$$\mathcal{H}_{\text{Ed}} = -2t_0\lambda \sum_{\mathbf{r}\mathbf{\rho}} f_{\mathbf{r}+\mathbf{\rho}}^\dagger f_{\mathbf{r}} - t_0 \sum_{\mathbf{r}} (f_{\mathbf{r}+\mathbf{\rho}}^\dagger f_{\mathbf{r}} b_{\mathbf{r}}^\dagger + \text{H.c.}) + \omega_0 \sum_{\mathbf{r}} b_{\mathbf{r}}^\dagger b_{\mathbf{r}}.$$  

(14)

Thus the second term in $\mathcal{H}_{\text{Ed}}$ is eliminated in favour of the first term in $\mathcal{H}_{\text{Ed}}$ which introduces a coherent hopping channel in addition to the original incoherent one. In the ground state $|0\rangle$ of the Hamiltonian $\mathcal{H}_{\text{Ed}}$, in the absence of fermions, there are no bosons, so that $b_{\mathbf{r}}|0\rangle = 0$. Thus for the ground state $|0\rangle$ of $\mathcal{H}_{\text{Ed}}$ we have $(b_{\mathbf{r}} - \lambda/\omega_0)|0\rangle = 0$ and hence $0|b_{\mathbf{r}}^\dagger b_{\mathbf{r}}|0\rangle = \lambda^2/\omega_0^2$.

From the above discussion it is clear that in 2D the Edwards model corresponds, within a certain range of parameters, to an underlying t-J' type of model with Hamiltonian

$$-t_0 \sum_{\mathbf{r}\mathbf{\rho}} c_{\mathbf{r}+\mathbf{\rho}}^\dagger c_{\mathbf{r}} + \frac{1}{2} \omega_0 \sum_{\mathbf{r}} S_{\mathbf{r}}^x S_{\mathbf{r}+\mathbf{\rho}} - 2\lambda \sum_{\mathbf{r}} S_{\mathbf{r}}^z,$$

(15)

since $2S_{\mathbf{r}}^x = S_{\mathbf{r}}^+ S_{\mathbf{r}}^- = b_{\mathbf{r}}^\dagger b_{\mathbf{r}} + b_{\mathbf{r}}$. This corresponds to a doped antiferromagnetic Ising model in a transverse magnetic field, but this correspondence is only valid for low hole density (low fermion density in the $\mathcal{H}_{\text{Ed}}$ model) and $\lambda \ll \omega_0$. In this case, since $|0(S_{\mathbf{r}}^z)|0\rangle = \frac{1}{2} - |0|b_{\mathbf{r}}^\dagger b_{\mathbf{r}}|0\rangle = \frac{1}{2} - \lambda^2/\omega_0^2$, the holes move in a background medium which is close to a saturated antiferromagnet. For $\lambda \gg \omega_0$ the transverse-field Ising model no longer exhibits antiferromagnetic order [13].

In the 1D correspondence breaks completely since if $\lambda = 0$ and $\omega_0 \neq 0$ the $\mathcal{H}_{\text{Ed}}$ model exhibits the string effect which is absent in the 1D t-J' model. In fact the 1D $\mathcal{H}_{\text{Ed}}$ model has an interesting similarity to the 2D t-J' model. Even for $\lambda = 0$ the string effect is relaxed by an analogue of the Trugman path [14]. This is again a six-step process in which the fermion propagates to a next-nearest-neighbour site with the background medium left undisturbed, i.e. no bosons excited. During the process the fermion excites three bosons which are subsequently destroyed. Unfortunately the model $\mathcal{H}_{\text{Ed}}$ does not represent precisely any physical system that we know of. The underlying Ising exchange is characteristic of the $t_{2g}$ model for vanadates and fluorides but the directional hopping of that model is not included, although this could be remedied in the 2D model. However if the model is considered over the whole $\lambda/t_0$, $\omega_0/t_0$-space it is found, even in the 1D case, to exhibit a surprising number of different physical regimes reminiscent of some found in realistic strongly correlated electron systems and strongly coupled electron–phonon systems. This was first demonstrated for the case of a single fermion ($N_f = 1$) at temperature $T = 0$ by the method of variational Lanczos diagonalization [14]. A great advantage of the model is that the simple treatment of the background medium, in terms of local bosons, makes it possible to obtain essentially exact results in the thermodynamic limit, at least in the 1D case. More recently the half-filled case, with $N_f = N/2$ where $N$ is the number of lattice sites, has been investigated [15–17]. The ground state phase diagram has been mapped out in the whole $\lambda/\omega_0$ plane, using a density matrix renormalization group (DMRG) technique [18, 19]. A quantum phase transition between a metallic Tomonaga–Luttinger liquid and an insulating charge density wave (CDW) was shown to exist.

It is desirable to complement these numerical results with some more analytical approaches. Nearly all analytical work on t-J-like or polaronic models is confined to the case of a single fermion. However very recently the projective renormalization method [20] has been applied to the half-filled 1D $\mathcal{H}_{\text{Ed}}$ model in a study of CDW formation at $T = 0$ [21]. Our initial analytical work on the $\mathcal{H}_{\text{Ed}}$ model was confined to the one-fermion case in 1D at $T = 0$ with the additional restriction $\lambda = 0$ [22]. The latter restriction means that the fermion can only propagate as a coherent quasiparticle by the Trugman-like process discussed above. This requires the coexistence of at least three bosons during the particle’s motion and in [22] the one-fermion Green’s function was calculated within a three-boson approximation. The spectral functions agree well with exact numerical results for $\omega_0/t_0 \gtrsim 1$. In this paper we extend the analytical approach for $\lambda = 0$ to finite fermion density and to higher dimension, 2D and 3D. This is achieved within a two-boson approximation and comparison with exact numerical results in 1D shows that the range of validity is thereby reduced to $\omega_0/t_0 \gtrsim 3$. No Trugman-like processes exist in the two-boson approximation but they are unimportant to understand CDW formation at finite density. Relaxation of the string effect does not depend on them because bosons created by one fermion can be destroyed by other fermions. Near half-filling a CDW state is found, in good agreement with 1D numerical results, and the CDW transition temperature can be calculated formally. However, within the two-boson approximation, the transition is mean-field-like, with no short-range order in the disordered state. This is clearly not realistic in 1D but finite-temperature calculations should certainly be relevant in 3D. In this paper we concentrate on the calculation of spectral functions in 1D at $T = 0$ with various fermion densities and compare with some new DMRG results.

In section 2 we determine the one-fermion Green’s function for $\lambda = 0$ within the two-boson approximation in the cases of one particle ($N_f = 1$) and one hole ($N_f = N - 1$). The method used is different from that previously employed for $N_f = 1$ [22] and provides a wavefunction as a byproduct. In section 3 we study the hierarchy of equations of motion for the Green’s function at arbitrary density and devise a decoupling which leads to the correct Green’s function in the two-boson approximation for $N_f = 1$ and $N - 1$. This two-boson Green’s
function is valid for any bipartite lattice in 1D, 2D or 3D and allows for two distinct sublattices so that the CDW state can be investigated. In section 4 spectral functions and ground state properties calculated from the two-boson Green’s function are compared with those calculated numerically by the DMRG method. In section 5 we draw conclusions and consider the outlook for future work.

2. Limiting cases of the Green’s function within the two-boson approximation

We now derive expressions for the one-fermion Green’s function in the special cases of a single fermion \((N_f = 1)\) and a single hole \((N_f = N - 1)\). For these single-particle cases the Green’s function may be calculated by direct solution of the Schrödinger equation, which also yields the wavefunction.

2.1. The case \(N_f = 1\)

For \(\lambda = 0\) the Hamiltonian equation (13), written in \(k\)-space, takes the form

\[
\mathcal{H}_{\text{Fid}} = \frac{1}{\sqrt{N}} \sum_{kk'} f_k^* f_{k'} \left[ \gamma(k) b^+_k b_{k'} + \gamma(k') b^+_k b_{k'} \right] + \omega_0 \sum_q b^+_q b_q,
\]

where

\[
f_k = \frac{1}{\sqrt{N}} \sum_{r} e^{i k r} f_r, \quad b_q = \frac{1}{\sqrt{N}} \sum_{r} e^{i q r} b_r, \quad \gamma(k) = -t_0 \sum_{\rho} e^{i k \rho}.
\]

The \(\rho\)-summation is over \(z\) nearest-neighbours in a 1D, 2D or 3D bipartite lattice. In the two-boson approximation the wavefunction for a single fermion is of the form

\[
\Psi_k = \left\{ f_k + \sum_{q_1} a(q_1) f_{k-q_1} b^+_{q_1} \right. + \left. \sum_{q_1 q_2} \sum_{q_1 q_2} a(q_1, q_2) f_{k-q_1-q_2} b^+_{q_1} b^+_{q_2} \right\} \langle \text{vac} \rangle,
\]

where \(\langle \text{vac} \rangle\) is the vacuum state and \(a(q_1, q_2) = a(q_2, q_1)\). On substituting this in the Schrödinger equation \(\mathcal{H} \Psi_k = \mathcal{E} \Psi_k\), and multiplying on the left by \(\langle \text{vac} \rangle, \langle \text{vac} | b_q f_{k-q} \rangle\) and \(\langle \text{vac} | b_{q_1} b_{q_2} f_{k-q_1-q_2} \rangle\), we obtain equations of the form

\[
-E + \frac{1}{\sqrt{N}} \sum_{q_1} \gamma(k - q_1) a(q_1) = 0,
\]

\[
\frac{1}{\sqrt{N}} \gamma(k - q_1) a(q_1) + \omega_0 - E
\]

\[
+ \frac{2}{\sqrt{N}} \sum_{q_1} a(q_1, q_2) \gamma(k - q_1 - q_2) = 0,
\]

\[
\frac{1}{\sqrt{N}} \gamma(k - q_1 - q_2) [a(q_1) + a(q_2)]
\]

\[
+ 2a(q_1, q_2) (2 \omega_0 - E) = 0.
\]

On solving equation (21) for \(a(q_1, q_2)\), and substituting in equation (20), we find

\[
a(q_1) \left[ \omega_0 - E - \frac{z t_0^2}{2 \omega_0} \right] = - \frac{1}{\sqrt{N}} \gamma(k - q_1)
\]

\[
+ \frac{1}{\sqrt{N}} \sum_{q_2} \frac{[\gamma(k - q_1 - q_2)]^2 a(q_2)}{2 \omega_0}.
\]

It is easily shown that for a bipartite lattice the last term in this equation vanishes when \(a(q) \propto \gamma(k - q)\). Hence a solution for \(a(q_1)\) is obtained by omitting the last term and, on substituting this solution in equation (19), we find

\[
E + \frac{z t_0^2}{\omega_0} = 0.
\]

The solutions of this equation are the energies of single fermion eigenstates, which are the poles of the one-fermion Green’s function \(G_k(E)\). The left-hand side of equation (23) is \(G_k^{-1}(E)\). The absence of \(k\)-dependence in this expression shows that the single fermion eigenstates are localized. This is due to the string effect which is not relaxed in the two-boson approximation.

2.2. The case \(N_f = N - 1\)

The motion of a single hole in the present model is quite different from that of a single particle. To avoid confusion it should be stressed that the hole discussed here does not correspond to a hole in a t-J-like model, the latter being represented by a fermion in the present model. Clearly when the hole considered here hops to a neighbouring site a boson is created on the arrival site, not on the departure site as in the motion of a single particle. This boson can be destroyed immediately when the hole makes a further hop. There is therefore no string effect and the hole propagates easily. To find the Green’s function in this case we use the Schrödinger equation as in the case of a single particle. The wavefunction for the hole is of the form

\[
\Phi_k = \left\{ f_k + \sum_{q_1} c(q_1) f_{k+q_1} b^+_{q_1} \right. + \left. \sum_{q_1 q_2} \sum_{q_1 q_2} c(q_1, q_2) f_{k+q_1+q_2} b^+_{q_1} b^+_{q_2} \right\} |F\rangle,
\]

where \(|F\rangle\) is the state with every site occupied by a fermion and \(c(q_1, q_2) = c(q_2, q_1)\). The equations corresponding to equations (19)–(21) in the previous case are

\[
-E - \frac{1}{\sqrt{N}} \gamma(k) \sum_{q_1} c(q_1) = 0,
\]

\[
- \frac{1}{\sqrt{N}} \gamma(k) + c(q_1) (\omega_0 - E)
\]

\[
- \frac{2}{\sqrt{N}} \sum_{q_1} c(q_1, q_2) \gamma(k + q_1 - q_2) = 0,
\]

\[
- \frac{1}{\sqrt{N}} [c(q_1) \gamma(k + q_1) + c(q_2) \gamma(k + q_2)]
\]

\[
+ 2c(q_1, q_2) (2 \omega_0 - E) = 0.
\]
From equations (26) and (27) we find that for a bipartite lattice
\[ e(q) = \frac{1}{\sqrt{N}} \left( \frac{\gamma(k)}{\omega_0 - E - \frac{\gamma(q) q^2}{2m_0 - E}} \right). \] (28)
Hence, from equation (25),
\[ -E - [\gamma(k)]^2 \frac{1}{N} \sum_q \omega_0 - E - \frac{\gamma(q) q^2}{2m_0 - E} = 0. \] (29)
The solutions of this equation are the energies of single hole eigenstates and, as expected, they depend on the wavevector \( k \), being functions of \( [\gamma(k)]^2 \). This type of \( k \) dependence arises from the fact that the hole propagates through the lattice, leaving behind no excited bosons, by means of double hops, as discussed at the beginning of this section. As in the one-particle case we deduce that for one hole (\( N = 1 \)) the one-fermion Green’s function is given by
\[ [G_k(E)]^{-1} = -E - [\gamma(k)]^2 \frac{1}{N} \sum_q \omega_0 + E - \frac{\gamma(q) q^2}{2m_0 + E}. \] (30)
The sign of \( E \) has been changed since in equation (29) the energy refers to a hole state.

3. The Green’s function at finite fermion density
In this section we study the hierarchy of equations of motion of the one-fermion Green’s function and find a decoupling which is consistent with the results derived in section 2 for the limiting cases of low (\( N = 1 \)) and high (\( N = N - 1 \) ) fermion density. We allow for two distinct sublattices with different occupation so that the CDW state can be investigated.

The Fourier transform of the one-fermion retarded Green’s function is defined by [23]
\[ G_k(E) = \langle \langle f_k; f_k^\dagger \rangle \rangle = -i \int_{-\infty}^{\infty} \theta(t) \langle \langle f_k(t), f_k^+ \rangle \rangle e^{iEt}, \] (31)
where \( f_k(t) = e^{iHt} f_k e^{-iHt} \) and \( \theta(t) \) is the unit step function. We may write \( f_k \), defined by equation (17), as a sum of two-sublattice components. Thus
\[ f_k = \frac{1}{\sqrt{2}} (f_{k_1} + f_{k_2}), \] (32)
where
\[ f_{k_l} = \sqrt{\frac{2}{N}} \sum_r e^{ik_l r} f_r. \] (33)
Here the summation is over sites \( r \) which belong to sublattice \( l \) (\( l = 1, 2 \)). It follows from equations (31) and (32) that
\[ G_k(E) = \frac{1}{2} \sum_{l=1}^{2} \sum_{m=1}^{2} G_{k_l}^{lm}(E), \] (34)
where
\[ G_{k_l}^{lm} = \langle \langle f_{k_l}; f_{k_m}^\dagger \rangle \rangle = \frac{2}{N} \sum_{r,s} e^{ik_l (r - s)} G_{r,s}, \] (35)
and
\[ G_{r,s} = \langle \langle f_r; f_s^\dagger \rangle \rangle. \] (36)
The equation of motion for the Green’s function \( \langle \langle A; B \rangle \rangle \) is [23]
\[ E \langle \langle A; B \rangle \rangle = \langle \langle [A, B]; 1 \rangle \rangle + \langle \langle [A, \mathcal{H}]; B \rangle \rangle. \] (37)
Hence, noting that \( \mathcal{H} \) is given by equation (13) with \( \lambda = 0 \), we find
\[ E G_{r,s} = \delta_{r,s} - \frac{1}{2} \sum_{\rho} H_{\rho, r,s}^{r, +} - \frac{1}{2} \sum_{\rho} f_{\rho, r,s}^{r, +}, \] (38)
where
\[ H_{\rho, r,s}^{r, +} = \langle \langle f_{\rho, r, s}^{r, +} b_r^\dagger; f_{\rho, s}^\dagger \rangle \rangle, \] (39)
\[ f_{\rho, r,s}^{r, +} = \langle \langle f_{\rho, r, s}^{r, +} b_r^\dagger; f_{\rho, s}^\dagger \rangle \rangle. \] (40)
Here \( r, s \) is a general site on the sublattice \( l \) which complements the sublattice \( \bar{l} \) \((\bar{l} = 2, 1 \) for \( l = 1, 2 \) respectively). \( r_1 + \rho \) is a particular site on the sublattice \( l \) but it is necessary to define \( l \) more generally in order to close the equations of motion. However we first calculate the second term on the right of equation (38).

The equation of motion for \( H \) is
\[ (E - \omega_0) H_{r_1, r_2, \rho}^{r_1, +} = \delta_{r_1, r_2} \langle \langle \rho, r_2 \rangle \rangle, \] (41)
where
\[ H_{r_1, r_2, \rho}^{r_1, +} = \langle \langle f_{r_1, r_2, \rho}^{r_1, +} b_{r_2}^\dagger; f_{r_2}^\dagger \rangle \rangle, \] (42)
\[ f_{r_1, r_2, \rho}^{r_1, +} = \langle \langle f_{r_1, r_2, \rho}^{r_1, +} b_{r_2}^\dagger; f_{r_2}^\dagger \rangle \rangle. \] (43)
The last term of this equation is produced by a process in which a fermion hops from \( r_1 + \rho \) to \( r_1 + \rho + \rho' \) with either the creation of a boson on the vacated site \( r_1 + \rho \) or destruction of a boson on the arrival site \( r_1 + \rho + \rho' \). The latter boson must have been created by another fermion so that the term involving \( b_{r_2}^\dagger \rho, \rho' b_{r_1} \) corresponds to a dynamical interaction between fermions. This should only be included if we consider two-fermion interactions consistently which goes beyond the effective Hartree–Fock treatment introduced below. We therefore neglect this term. In the second term on the right of equation (41) we retain only the part involving fermions on two sites, thus taking \( \rho' = \rho \), and then make a Hartree–Fock type of approximation. Thus the second term becomes
\[ -\omega_0 (\langle \langle (1 - n_{r_1, \rho}) f_{r_1}; f_{r_1}^\dagger \rangle \rangle \sim -\omega_0 (1 - (n_{r_1, \rho}) G_{r_1, s}). \] (44)
In the equation of motion for \( J \) the terms arising from \( [f_{r_1, r_2, \rho}, \mathcal{H}] \) involve three boson operators and we neglect
them. The terms arising from \( [b_{l+\rho} b_r, H] \) may be treated in the Hartree–Fock-like way used to obtain equation (42). Hence

\[
(E - 2\omega_0) \rho^\gamma_{r_1+\rho, r_0} = \delta_{r_1+\rho, r_0} \rho^\gamma_{r_1+\rho, r_0} (b_{r_1+\rho} b_{r_0})
\]

\[
- t_0 (1 - n_l) H^\gamma_{r_1+\rho, r_0}.
\]

(45)

We approximate \( \langle b_{r_1+\rho} b_{r_0} \rangle \) by \( b_l b_l \), where \( b_l = \langle b_r \rangle \). Furthermore it is shown below that \( b_l = 0 \) for the present model, with \( \lambda = 0 \). Hence \( \langle b_{r_1+\rho} b_{r_0} \rangle \) may be neglected. On substituting for \( J \) in equation (43), using equation (45), and summing over \( \rho \), we find

\[
\sum_{\rho} H^\gamma_{r_1+\rho, r_0} = C(E, n_l) \left[ b_l \sum_{\rho} \delta_{l\rho, r_0} - z t_0 (1 - n_l) G_{l, r_0} \right],
\]

where

\[
C(E, n_l) = \frac{1}{E - \omega_0 - \frac{z t_0 (1 - n_l)}{2 \omega_0}}.
\]

(46)

The second term on the right of equation (38) has therefore been determined and to find the third term we must consider the equation of motion of the Green’s function \( I \) defined by equation (40). This takes the form

\[
(E + \omega_0) \rho^\gamma_{r_1+\rho, r_0} = \delta_{r_1+\rho, r_0} \rho^\gamma_{r_1+\rho, r_0} (b_{r_1+\rho} b_{r_0}) + t_0 \sum_{\rho} \langle \{ b_{r_1+\rho} f_{r_1+\rho}^\dagger b_{r_1+\rho} f_{r_1+\rho}^\gamma \} \rangle - t_0 \sum_{\rho} \langle \{ b_{r_1+\rho} (\delta_{r_1+\rho, r_0} + b_{r_1+\rho} b_{r_1+\rho}^\dagger b_{r_1+\rho} b_{r_1+\rho}^\gamma) \} \rangle.
\]

(47)

We now make similar approximations to those used to obtain equation (43). Thus we make an effective Hartree–Fock approximation to the second term on the right of equation (48), retaining it only when \( r_l = r_0 + \rho \), and we neglect the \( b_{r_1+\rho} b_{r_1+\rho} \) term in the third term. Hence

\[
(E + \omega_0) \rho^\gamma_{r_1+\rho, r_0} = \delta_{r_1+\rho, r_0} \rho^\gamma_{r_1+\rho, r_0} (b_{r_1+\rho} b_{r_0}) + t_0 \delta_{r_1+\rho, r_0} \sum_{\rho} G_{r_1+\rho, r_0} = - t_0 \sum_{\rho} \rho^\gamma_{r_1+\rho, r_0} (b_{r_1+\rho} b_{r_1+\rho}^\dagger b_{r_1+\rho} b_{r_1+\rho}^\gamma) \langle f_{r_1+\rho}^\dagger f_{r_1+\rho} \rangle.
\]

(49)

where

\[
K^\gamma_{r_1+\rho, r_0} = \langle \{ b_{r_1+\rho} b_{r_1+\rho}^\dagger b_{r_1+\rho} b_{r_1+\rho}^\gamma \} \rangle.
\]

(50)

We treat the equation of motion of \( K \) in a similar way to that of \( J \). A slight difference is that instead of neglecting all the terms arising from \( [f_{r_1+\rho}, H] \) we retain one which leads to a factor \( b_{r_1+\rho}^\dagger b_{r_1+\rho}^\gamma \). This equals \( \delta_{r_1+\rho, r_0} \sum_{\rho} G_{r_1+\rho, r_0} \) approximately by \( 1 \), neglecting the boson occupation number. Hence, neglecting a \( (b^\dagger b^\gamma) \) correlation function as before, we find

\[
(E + 2\omega_0) K^\gamma_{r_1+\rho, r_0} = - t_0 \delta_{r_1+\rho, r_0} \sum_{\rho} G_{r_1+\rho, r_0}.
\]

(51)

Thus, from equations (49) and (51),

\[
(E + \omega_0) \rho^\gamma_{r_1+\rho, r_0} = \delta_{r_1+\rho, r_0} \rho^\gamma_{r_1+\rho, r_0} (b_{r_1+\rho} b_{r_0}) + \frac{t_0^2}{E + 2\omega_0} \sum_{\rho} \sum_{\rho'} G_{r_1+\rho, r_0} \rho^\gamma_{r_1+\rho', r_0} - \frac{t_0}{E + 2\omega_0} \sum_{\rho} G_{r_1+\rho, r_0}.
\]

(52)

To solve this equation for \( I \) we introduce the Fourier transform

\[
I^m_{kq} = \left( \frac{2}{N} \right)^2 \sum_{r_1 \rho s_0} \rho^\gamma_{r_1+\rho, r_0} e^{i(k+q)\rho - k s_0 - q(n_l + \rho)}
\]

(53)

so that

\[
\left( E + \omega_0 - \frac{n_l [\gamma(k+q)]^2}{E + 2\omega_0} \right) I^m_{kq} = \sqrt{\frac{N}{2}} \delta_{k0} \delta_{l m} b^*_l + \sqrt{\frac{N}{2}} n_l \gamma(k) G^m_{kq}.
\]

(54)

By taking the Fourier transform of equation (38), and using equation (46), we find

\[
E G^m_{kq} = \delta_{l m} - C(E, n_l) [b_l \gamma(k) \delta_{l m} - z t_0^2 (1 - n_l) G^m_{kq}]
\]

(55)

where

\[
I^m_{kq} = \sqrt{\frac{N}{2}} \sum_{r_1 \rho s_0} e^{i(k+q)\rho} \rho^\gamma_{r_1+\rho, r_0} = \sqrt{\frac{2}{N}} \sum_{q} I^m_{kq}.
\]

(56)

By combining the last three equations we obtain

\[
G^m_{kq} [E - z t_0^2 (1 - n_l) C(E, n_l)] = \delta_{l m} - [C(E, n_l) b_l + D(E, k, n_l) b^*_l] \delta_{l m} \gamma(k),
\]

(57)

where

\[
D(E, k, n_l) = \frac{1}{E + \omega_0 - \frac{n_l \gamma(k)}{E + 2\omega_0}}.
\]

(58)

and

\[
\tilde{D}(E, n_l) = \frac{2}{N} \sum_{q} D(E, q, n_l).
\]

(59)

It remains to explain why, as indicated following equation (45), \( b_l = 0 \). This may be shown using

\[
b_l = \langle b_l \rangle = \frac{1}{z} \sum_{\rho} \langle b_{r_1+\rho} f_{r_1+\rho}^\dagger f_{r_1+\rho}^\gamma \rangle = \frac{1}{2 \pi z} \int_{-\infty}^{\infty} \left[H^\gamma_{r_1+\rho, r_1+\rho} (E + i\eta) - H^\gamma_{r_1+\rho, r_1+\rho} (E - i\eta) \right] dE.
\]

(60)

Equations (46) and (57) may be used to show that the above expression is a linear combination of \( b_l \) and \( b^*_l \). Thus \( b_l, b^*_l \) \((l = 2, 1)\) satisfy a set of linear homogeneous equations so that in general \( b_l = 0 \). This result is consistent with the symmetry of the transverse-field \( t-J \) model (equation (15)) which underlies the present model for low fermion density. When \( \lambda = 0 \), as assumed here, the transverse magnetic field vanishes so that by symmetry the expected value of the transverse spin moment \( \langle S^y_0 \rangle = \langle b^*_l + b_l \rangle / 2 = 0 \). The properties \( b_l = 0 \) and the characteristic periodicity in \( k \) are shown in the appendix to be generally true for the model with \( \lambda = 0 \) for all values of \( \omega_0 / t_0 \). As soon as \( \lambda \neq 0 \) we shall in general have \( b_l \neq 0 \) so that, from equation (57), the \( k \) dependence of the Green’s function will involve \( \gamma(k) \) and not only \( [\gamma(k)]^2 \) as is the case for \( \lambda = 0 \).
The final result for the Green’s function in the case \( \lambda = 0 \) is

\[
G_k^{\text{Int}}(E) = \delta_{\text{Im}}[E - \varepsilon_0^2(1 - n)C(E, n_i) - \gamma(k)^2 n_i \bar{D}(E, n_i)]^{-1}
\]

(61)

where \( C \) and \( \bar{D} \) are given by equations (47) and (59) respectively. Also, from equation (34),

\[
G_k(E) = \frac{1}{2}[G_{k1}(E) + G_{k2}(E)],
\]

(62)

where \( G_{k1} = G_{k1}^0 \). In the homogeneous case, where there is no CDW, \( n_1 = n_2 = n \), where \( n = N_i/N \) is the fermion density, and \( G_{k1} = G_{k2} = G_k \). For \( n = 0 \) we recover the \( N_i = 1 \) result for \( G_k^{-1} \), given by the left-hand side of equation (23), and for \( n = 1 \) we recover the \( N_i = N - 1 \) result given by equation (30). Thus \( G_k \) is correct in these two limits within the two-boson approximation. The factor \( \delta_{\text{Im}} \) in equation (61) shows that within the present approximation fermions propagate, by means of double hops, within a single sublattice. The dimension of the bipartite lattice (1, 2 or 3) enters only through the band energy \( \gamma(k) \) (equation (17)) and the function \( \bar{D} \), which may be written as

\[
\bar{D}(E, n_i) = \int \frac{N_0(\gamma)}{E + \omega_0 - \gamma^2 n_i/(E + 2\omega_0)} \, d\gamma,
\]

(63)

where \( N_0(\gamma) \) is the density of states per site for the band energy \( \gamma(k) \).

To complete the present formulation we require the equations which determine the chemical potential \( \mu \) and the self-consistent sublattice densities \( n_1, n_2 \) for a given fermion density \( n \). The density of states per site on sublattice \( l \) is given by

\[
N_l(E) = -\frac{1}{\pi} \text{Im} G_k(E + i\eta)
\]

(64)

and the total spectral function is

\[
S(k, E) = -\frac{1}{\pi} \text{Im} G_k(E + i\eta) = \frac{1}{2}[S_1(k, E) + S_2(k, E)].
\]

(68)

These spectral functions may be used to calculate the occupation number \( n(k) = \langle f_k^\dagger f_k \rangle \) and the related quantity \( d(k) = \langle (f_k^\dagger f_{k+Q})^2 \rangle \), where \( Q \) is a basis vector of the reciprocal lattice. \( f_k \) is given by equation (32) and it follows from equation (33) that \( f_{k+Q} = (f_k - f_{k+Q})/\sqrt{2} \). Hence we find

\[
n(k) = \frac{1}{\pi} \int_{-\infty}^{\mu} [S_1(k, E) + S_2(k, E)] \, dE
\]

(69)

and

\[
d(k) = \frac{1}{\pi} \int_{-\infty}^{\mu} [S_1(k, E) - S_2(k, E)] \, dE.
\]

(70)

These quantities satisfy the sum rules

\[
\frac{1}{N} \sum_k n(k) = n, \quad \frac{1}{N} \sum_k d(k) = nP.
\]

(71)

In section 4 we report numerical results for all the above quantities, based on the approximate Green’s function of equation (61), in the 1D case. Many of the results are compared with those of the DMRG method, both to assess the validity of the present approximation and sometimes to throw new light on the DMRG results. The application of the DMRG method to the present model has been described previously \[16, 17\].

4. Numerical results for ground state and spectral properties in 1D

In 1D the band energy \( \gamma(k) = -2 \cos k \), with the lattice constant taken as 1 and the unit of energy taken as \( \hbar \). The quantity \( \bar{D} \) which appears in the Green’s function (equation (61)) can then be evaluated analytically using equations (58) and (59). The q-summation is most conveniently performed as a contour integral around the unit circle. The result is

\[
\bar{D}(E, n_i) = D_{\omega_0} \theta(a_1^2 - b_1^2) - i D_{\omega_0} \theta(b_1^2 - a_1^2),
\]

(72)

where

\[
D_{\omega_0} = \frac{1}{a_i \sqrt{1 - b_i^2/a_i^2}}, \quad \quad D_{\omega_0} = \frac{1}{\sqrt{b_1^2 - a_1^2}}
\]

(73)

with

\[
a_i = E + \omega_0 - \frac{2n_i}{E + 2\omega_0}, \quad b_i = -\frac{2n_i}{E + 2\omega_0}.
\]

(74)

The density of states, given by equation (64), can be evaluated similarly with the result

\[
N_l(E) = N_{l1}(E) + N_{l2}(E),
\]

(75)
where

\[
N_{11}(E) = \frac{1}{\pi} \left| \theta(a_i^2 - b_i^2) \right| \left[ \frac{\theta(v_i^2 - u_i^2)}{\sqrt{v_i^2 - u_i^2}} + \theta(u_i^2 - v_i^2) \right]
\]

(76)

\[
N_{12}(E) = \frac{\theta(b_i^2 - a_i^2)}{2\pi n_i D_{ii}} \frac{\sqrt{V_i^2 + 4 - |V_i|}}{2|V_i|(\sqrt{V_i^2 + 4})}
\]

(77)

with

\[
u_i(E) = -2n_i D_{ii}(E),
\]

(78)

\[
u_i(E) = \frac{E - 2(1 - n_i)C(E, n_i) - 2n_i D_{ii}(E)}{2n_i D_{ii}(E)}.
\]

(79)

(80)

The results presented below are for the case \(\omega_0 = 3\). By comparison with the DMRG results for the ground state properties \(n(k)\) and \(d(k)\), and with the dynamical DMRG [24] (DDMRG) results for \(S(k, E)\), it is found that this boson energy is large enough for many results of the present analytic approximation to be quite accurate.

4.1. High and low fermion density

The existence of a two-sublattice CDW state in the present model is well established at half-filling \((n = 0.5)\) for sufficiently large \(\omega_0/t_0\) [15–17, 21]. However the general results of the appendix imply that a two-sublattice CDW state is always a possibility, whatever the density \(n\). We therefore used equations (65), (66) and (75) to search for such states even in the high and low density cases of \(n = 0.9\) and \(0.1\). In the case \(n = 0.9\) with \(\omega_0/t_0 = 3\) we find a self-consistent CDW state with order parameter \(P = 0.063\). The flow of the iterative procedure to determine \(P\) indicates that this, not the uniform density \(P = 0\) state, is the stable ground state. In the DMRG calculations convergence to the CDW solution is improved by use of suitable external fields at the boundaries with open boundary conditions. The spectral function \(S(k, E)\) calculated from the Green’s function using equation (68) is compared with the DDMRG results in figure 1. The quasiparticle peaks for the Green’s function decoupling scheme results (left panels) are delta-functions in the limit \(\eta \to 0\), but to make them visible we have taken \(\eta = 0.05\); the same value has been taken in the DDMRG data (right panel). The following main features are in good agreement: the general shape and width of the quasiparticle band crossing the Fermi level, the dispersive peaks just below \(E - E_F = -4\) which vanish for \(k = \pi/2\), the weak flat band at \(E - E_F = 2\) in the left panel and 3 in the right one. The splitting of the quasiparticle peaks due to CDW order is clearly visible in the left panel. This splitting is not clearly resolved in the main right panel but the inset for \(k = \pi/21\) shows a splitting between a peak below \(E_F\) in the photoemission spectrum (PES) and one above \(E_F\) in the inverse photoemission spectrum (IPES). The absence of the splitting for states further from \(E_F\) in the DDMRG data is presumably due to finite lifetime broadening processes which are not included in the Green’s function approximation. It should be noticed that the quasiparticle states at the Fermi level are from only one of the split subbands near \(k = 0\) and \(\pi\), the other subband being fully occupied. This means that the Fermi wavevectors are at \(\pi/10, \pi - \pi/10\) so that the hole pockets near \(k = 0\) and \(\pi\) have the correct Fermi surface ‘volume’, this being twice what it would have been in the absence of the CDW (\(P = 0\)). Discontinuities at these wavevectors are clearly seen in figure 3 where \(n(k)\) and \(d(k)\), calculated from equations (69) and (70), are plotted. The agreement with the DMRG data, also plotted, is excellent. In the Green’s function calculations the discontinuities are not perfectly sharp owing to the use of a finite \(\eta =0.0005\) near \(E_F\). There are two contributions to \(n(k)\) and \(d(k)\), one arising from the coherent quasiparticle bands and the other from spectral weight further below the Fermi level. In the present case \(d(k)\) arises almost entirely from the coherent contribution and the quasiparticle states have slightly more weight on the minority sublattice, hence the negative value of \(d(k)\) over most of the zone. Near \(k = 0\) and \(\pi\), however, only the subband associated with the majority sublattice is occupied,
hence the strong positive contribution. Correlations giving rise to Luttinger liquid behaviour in 1D are beyond the present Green’s function approach. It should be noted that at $n = 0.9$ the CDW state is metallic, whereas at $n = 0.5$ it is an insulator, as discussed in section 4.2.

In the case $n = 0.1$ with $\omega_0/t_0 = 3$ we find a self-consistent CDW state with order parameter $P = 0.61$. The discrepancy in order of magnitude between this value and the much smaller one for $n = 0.9$ is due to the consistent use of fermion density in the definition of $P$ (equation (66)), rather than changing to hole density for the $n = 0.9$ case. For $n = 0.1$ it has proved difficult to converge to a CDW solution in DMRG. This may indicate that the Green’s function approximation is failing in this low density case. If so, the nature of the ground state is unclear. Nevertheless in figure 2 we show results for the spectral function and in figure 3 we plot $n(k)$ and $d(k)$. The bottom of the narrow quasiparticle band near $E_F$ is at $k = \pi/2$ and only the majority subband is occupied. Hence $d(k)$ is strongly positive near $k = \pi/2$ and almost equal to $n(k)$. Over most of the zone the only contribution is from spectral weight further below the Fermi level; this vanishes at $k = \pi/2$ and makes a negative contribution to $d(k)$ owing to more weight residing on the minority sublattice. The discontinuities in $n(k)$ and $d(k)$ at the expected values $\pi/2 \pm \pi/10$ are clearly seen.

4.2. The CDW state at half-filling

We now consider a self-consistent CDW state for the half-filled band with $n = 0.5$ and $\lambda = 0$, $\omega_0 = 3$. As before the Fermi energy $E_F$ (or chemical potential $\mu$ at $T > 0$) and order parameter $P$ are determined by equations (66) and (65), with the sublattice densities of states given by equation (75). For the above parameters we find $\mu = -0.211$, $P = 0.765$. $E_F$ lies in a gap as is appropriate for an insulator.

The spectral functions calculated from the Green’s function method and the DDMRG are compared in figure 4. The agreement is generally good. The Fermi level lies in a gap between two quasiparticle bands, a broad upper unoccupied one and a narrower lower occupied one. The main discrepancy is that in the DDMRG case the lower quasiparticle band is extremely narrow whereas in the Green’s function method it has a significant width. This shortcoming of the Green’s function approximation is discussed thoroughly.
**Figure 4.** Single-particle spectral function $S(k, E)$ at $n = 0.5$ for $\lambda = 0$, $\omega_0 = 3$ from equation (68) (left panel) compared with the DDMRG result (right panel).

**Figure 5.** Density of states $D(E)$ near the Fermi level for $\lambda = 0$ at half-filling from equation (75) (left panel) and corresponding quasiparticle band dispersion $E(k)$ (right panel).

**Figure 6.** Bloch state occupation number $n(k)$ (left panel) and related correlation function $d(k)$ (right panel) for the half-filled band case with $\lambda = 0$, $\omega_0 = 3$.

in section 4.3. Figure 5 shows the density of states near the Fermi level, projected onto the majority and minority sublattices, calculated from equation (64). It is remarkable that states in the occupied quasiparticle band are entirely confined to the majority sublattice, whereas those in the unoccupied quasiparticle band reside entirely on the minority sublattice. The dispersion curves $E(k)$ of the quasiparticle bands, obtained by plotting the loci of quasiparticle peaks in the spectral function, are shown in figure 5. The approximate Green’s function therefore predicts an indirect gap, with the top of the occupied quasiparticle band at $k = 0$, $\pi$ and the bottom of the unoccupied band at $k = \pi/2$.

In figure 6 left and right panels we plot curves for $n(k)$ and $d(k)$, respectively, calculated from equations (69) and (70), together with points obtained by the DMRG method for finite systems of different sizes. The agreement is very
striking, particularly since in DMRG these quantities are calculated directly from the ground state whereas in the Green’s function method they are obtained as integrals over the spectral function. It should be noted that $n(\pi/2) = d(\pi/2)$ because for $k = \pi/2$ all the spectral weight of the occupied states resides in the quasiparticle peak which is entirely based on the majority sublattice. From equations (69) and (70) it is clear that the weight in this peak is $2n(\pi/2) \simeq 0.88$.

Finally in figure 7 we plot the spectral function (left panel) and quasiparticle energy (right panel) in the high-temperature limit, where all short-range order has disappeared ($P = 0$). The Fermi wavevectors are close to $\pi/4$ and $3\pi/4$ with the correct Fermi surface ‘volume’. This is the state from which the CDW evolves as the temperature is lowered. Clearly the situation is quite different from the usual one in which a CDW evolves because of nesting between Fermi wavevectors at $k = \pm \pi/2$. The origin of this difference is that for $\lambda = 0$ only next-nearest-neighbour hopping, resulting in $\pi$-periodicity in $k$-space, occurs even in the disordered state.

4.3. Discussion

The most notable difference between the Green’s function and DDMRG results lies in the width of the occupied quasiparticle band for $n = 0.5$. A very narrow band is also found in earlier calculations for $\omega_0 = 2$ using exact diagonalization where there is also no tendency for the top of the band to be at $k = 0, \pi$ rather than $\pi/2$ [15]. The band seems to be almost as narrow as one would have from the Trugman-like six-step process in a perfect CDW ($P = 1$). This is quite surprising since for $\omega_0 = 2$ the densities on the two sublattices are about 0.8 and 0.2 [16] which is far from a perfect CDW. Satellites below the main quasiparticle peak are suggestive of finite-size effects in the systems considered with 12 and 16 sites.

To gain more insight into the Green’s function approximation we may consider the limit of large $\omega_0$ where all the weight is concentrated in the two quasiparticle bands. From equation (61) it follows that these bands are given by

$$E(k) = \frac{\pm 2P + (1 \pm P) \cos(2k)}{\omega_0}$$

where the upper signs (+) correspond to the upper band and the lower signs (−) to the lower band. The widths $2(1 \pm P)/\omega_0$ of these bands can be understood by inspection of the following processes, where a bullet represents a vacant site, a star represents a boson and a circle represents a fermion:

$$| \bullet \circ \circ \rangle \rightarrow | \circ \bigcirc \circ \rangle \rightarrow | \circ \circ \bullet \bigcirc \rangle$$

$$| \circ \circ \bullet \bigcirc \rangle \rightarrow | \circ \bigcirc \circ \bullet \rangle \rightarrow | \bullet \circ \circ \rangle.$$
equal in number, to form bound pairs. We conclude that for \( n = 0.5 \) important correlations exist in the ground state which are not included in the Hartree–Fock-like approximation of the Green’s function decoupling.

5. Conclusions and outlook

Doped Mott insulators remain at the forefront of physics research largely because of their relevance to high-temperature superconductivity. In the cuprate superconductors holes in the copper oxide planes move in a background of antiferromagnetic order. In other systems of interest the background is one of alternating orbital order. The progress of a hole through such a background is hindered by the string effect. This effect has been known for a long time [8] but the string picture is central to much recent work (e.g. [25–28]).

The Edwards fermion–boson model considered in this paper was introduced to describe this effect in the simplest possible way [12]. The spinless fermions correspond physically to the holes in the Mott insulator. The ordered background does not appear explicitly in the model; the essence of the string effect actually relies only on the existence of substantial short-range order without the necessity of true long-range order. Clearly the physical interest lies mainly in 2D [29], but so far most calculations for the Edwards model have been made in 1D [14–17, 21, 22]. In this paper we describe an analytical approximation to the one-fermion Green’s function which is valid in 1D, 2D and 3D. Its main limitations are that the boson approximation to the one-fermion Green’s function which is associated with the CDW. There is also generally good agreement for the one-fermion spectral function although relaxation is neglected (\( \omega_0^2 \approx 0 \)). The principal objective of this paper is to test the accuracy of the Green’s function method, within its expected domain of validity, by comparing with numerical results obtained in 1D by the DMRG and DDRMG methods.

This comparison has been made in detail for \( \omega_0 = 3 \) and for various fermion densities. For the half-filled band case \( (n = 0.5) \) excellent agreement is obtained for ground state properties. These include the CDW order parameter, the Bloch state occupation number \( n(k) \) and a related quantity \( d(k) \) associated with the CDW. There is also generally good agreement for the one-fermion spectral function although the Green’s function method predicts much too wide an occupied quasiparticle band in section 4.3 the reason for this discrepancy is traced to missing correlations in the Green’s function approximation. The Green’s function method predicts that the CDW state evolves from a high-temperature disordered state with Fermi wavevectors close to \( \pi/4 \) and \( 3\pi/4 \), which is quite different from the usual case where a CDW evolves because of nesting between Fermi wavevectors at \( k = \pm \pi/2 \). The origin of the difference is that only next-nearest-neighbour hopping, resulting in \( \pi \)-periodicity in \( k \)-space, occurs even in the disordered state.

There is excellent agreement between the two methods for the dilute hole \( (n = 0.9) \) case with the usual parameters \( \lambda = 0, \omega_0^2 / t_0 = 3 \). Somewhat surprisingly, there is also a two-sublattice CDW state which in this case is metallic. In the appendix it is shown how the appearance of two-sublattice CDW states is related to a symmetry property of the model with \( \lambda = 0 \). The comparison between the two methods proved to be more difficult in the dilute fermion \( (n = 0.1) \) case. The Green’s function method again predicts a metallic CDW state but the DMRG fails to confirm this. The true nature of the ground state in this case remains unclear.

It may be concluded that the rather simple Green’s function approximation derived here is sufficiently successful in 1D, which is probably the least favourable case, to envisage future applications to the 2D t-J model and, with a slight extension of the model, to the \( t_2g \) model of alternating orbital order. The situation of physical interest will be low to moderate fermion density, the fermions corresponding to holes in the relevant Mott insulator.

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Appendix

Through the particular form of the fermion–boson hopping term at \( \lambda = 0 \) (cf equation (14)) the transfer of a fermion between neighbouring lattice sites coincides with a change of the number of bosons by one. As a consequence we may think of the bosons as tracking the motion of the fermions. This picture becomes exact through the identification of a conserved quantity. Let us define operators \( N_{A}^0 \) and \( N_{B}^0 \) which count the number of fermions on the \( A \)- or \( B \)-sites of a bipartite lattice, and similar operators \( N_{A}^1 \) and \( N_{B}^1 \) for bosons. Then \( N_{A} = N_{A}^0 - N_{B}^0 + 2(N_{A}^1 - N_{B}^1) \) commutes with the Hamiltonian \( H_{\text{Ed}} \), when \( \lambda = 0 \), and is therefore a conserved quantity. The eigenvalues of \( N_{A} \) can be used to classify the eigenstates of the Hamiltonian. We note that \( N_{B} \) is not conserved for \( \lambda \neq 0 \).

The existence of \( N_{B} \) has two major consequences. First, fermion operators such as \( f_{\alpha}^\dagger f_{\beta} \) change \( N_{B} \) by 2, so that in any eigenstate of the Hamiltonian the expectation value \( \langle f_{\alpha}^\dagger f_{\beta} \rangle = 0 \) for arbitrary sites \( nA \) and \( mB \) on the respective sublattices. Similarly \( \langle b_{nA}^\dagger \rangle = 0 \). Hence quantities such as the spectral function \( S(k, E) \) or the momentum distribution \( n(k) \) have the periodicity in \( k \) of the reciprocal lattice of the real-space A or B sublattice, e.g. \( \pi \)-periodicity in the 1D case. Second, it implies that those eigenvalues of the Hamiltonian corresponding to \( N_{B} \neq 0 \) are degenerate. This follows because the translation operator \( T \) commutes with \( H \), but changes the sign of \( N_{B} \). The energy eigenvalue of an eigenstate \( |\psi\rangle \) with \( \langle \psi | N_{B} | \psi \rangle \neq 0 \) must therefore be (at least) two-fold degenerate, since the state \( T|\psi\rangle \) belongs to the same energy but differs from \( |\psi\rangle \) due to the change of \( N_{B} \). This degeneracy corresponds to a breaking of translational symmetry, as for a two-sublattice CDW state. Clearly if the ground state at \( \lambda = 0 \) is not such a CDW state it must have \( N_{B} = 0 \).

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