Effective interactions and superconductivity in the t-J model in the large-N limit

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(March 28, 2021)

Abstract

The feasibility of a perturbation expansion for Green’s functions of the $t-J$ model directly in terms of X-operators is demonstrated using the Baym-Kadanoff functional method. As an application we derive explicit expressions for the kernel $\Theta$ of the linearized equation for the superconducting order parameter in leading order of a $1/N$ expansion. The linearized equation is solved numerically on a square lattice taking instantaneous and retarded contributions into account.

Classifying the order parameter according to irreducible representations $\Gamma_i$, $i = 1, \ldots, 5$, of the point group $C_{4v}$ of the square lattice and according to even or odd parity in frequency we find that a reasonably strong instability occurs only for even frequency pairing with d-wavelike $\Gamma_3$ symmetry. The corresponding transition temperature $T_c$ is $\sim 0.01|t|$ where $t$ is the nearest-neighbor hopping integral. The underlying effective interaction consists of an attractive, instantaneous term and a retarded term due to charge and spin fluctuations. The latter is weakly attractive at low frequencies below $\sim J/2$, strongly repulsive up to $\sim |t|$ and attractive towards even higher energies. $T_c$ increases with decreasing doping $\delta$ until a d-wavelike bond-order wave instability is encountered near optimal doping at $\delta_{BO} \sim 0.14$ for $J = 0.3$. $T_c$ is essentially linear in $J$ and rather insensitive to an additional second-nearest neighbor hopping integral $t'$. A rather striking property of $T_c$ is that it is hardly affected by the soft mode associated with the bond-order wave instability or by the Van Hove singularity in the case with second-nearest neighbor hopping. This unique feature reflects the fact that the solution of the gap equation involves momenta far away from the Fermi surface (due to the instantaneous term) and many frequencies (due to the retarded term) so that singular properties in momentum or frequency are averaged out very effectively.

PACS numbers: 74.20-z, 74.20Mn
I. INTRODUCTION

Many studies of the $t - J$ model suggest that this model is able to describe a large body of the low-energy physics of real high-$T_c$ superconductors\cite{1,2}. This is true for many normal state properties of high-$T_c$ oxides where accurate numerical predictions of the $t - J$ model are available for the comparison with experiment. Whether the phenomenon of high-$T_c$ superconductivity itself can be explained within this model is presently not so clear. There are several calculations yielding instabilities of the normal state with respect to $d$-wave superconductivity\cite{2,3,4} and also reasonably high values for the transition temperature $T_c$\cite{6,7,8,9}. These calculations, however, use often somewhat uncontrolled assumptions, making definite conclusions difficult. There is also the view\cite{1} that the large observed values for $T_c$ are not directly related to large mean field $T_c$’s in isolated $CuO_2$ planes described by the $t - J$ model. Instead it is argued that pair tunneling between planes enhances strongly weak, plane-related superconducting instabilities producing in this way the phenomenon of high-$T_c$ superconductivity.

In this paper we present a new attempt to calculate $T_c$ for the $t - J$ model in a well controlled way. Similar as in Refs.\cite{3,4} we do not assume the validity of Migdal’s theorem or approximate the self-energy by the lowest skeleton graphs. Instead we assume that $1/N$ can be considered as a small parameter where $N$ is the number of electronic degrees of freedom per site. $N$ consists of two spin directions times $N/2$ copies of the local electronic orbital counted by a flavor index. Similar like in many slave boson calculations\cite{3,4,10} the flavor index is introduced in a somewhat artificial way just to make $N$ a large integer. The $SU(2)$ spin symmetry of the original model is thus enlarged to the symplectic group $Sp(N/2)$. The original constraint of having no double occupancies of sites is modified to the condition that at most $N/2$ electrons can occupy the $N$ states at each site. Compared to Refs.\cite{3,4,10} our treatment exhibits two novel features. First, the constraint is implemented in a different and more rigorous way yielding differences in the equation for the superconducting gap already in the leading order $O(1/N)$. Secondly, we have solved the linearized gap equation numerically obtaining also values for $T_c$. Our conclusions about the occurrence of superconductivity in the $t - J$ model are thus no longer based only on Fermi surface averaged, static coupling strengths as in Refs.\cite{3,4,10,11}.

Regarding constraints their implementation in the X-operator approach is trivial.\cite{12} Mathematically, the constraint means that the sum over diagonal elements of X-operators at a given site has to be equal to $N/2$. This sum commutes with every X-operator and thus is a multiple of the identity operator in any irreducible representation of X-operators. Enforcing the constraint means therefore just to select the correct subspace of Hilberts space where the eigenvalue of this sum is equal to $N/2$. In slave boson theory the X-operators are represented by product of slave operators and the Hilbert space is enlarged to the Hilbert space of slave particles. The constraint means now that at each site the number operator of slave particles has to be equal $N/2$. This condition can obviously not hold as an operator identity, i.e., cannot be enforced at any position of an expectation value of slave operators. However, it should be enforced at all positions which originally separated X-operators. Such an enforcement for a fixed $N$ seems to be in conflict with the Bose condensation of the bosonic slave particles as well as the independence of fermion and bosonic slaves in the limit $N \rightarrow \infty$. Using the Dirac method to enforce constraints on the operator level it has indeed
been shown that the commutator relations for the slave particles have to be changed and that, for instance, the fermionic and the bosonic slave operators no longer commute with each other. In view of this open problems we use in sections II and III a perturbation expansion directly in X-operators following the work of Ref. This procedure will give us also the opportunity to compare the O(1/N) expression for the gap equation with that of the slave boson approach to see whether the 1/N expansions are really the same in the two cases. The final analytic results for the gap equation have already been presented in Ref., however, without derivations. These derivations can be found in sections II and III.

The second new feature of our work deals with the solution of the gap equation. Previous work using 1/N expansions concluded from Fermi surface averaged, static coupling constants on the occurrence of superconductivity. Our previous work has shown that superconducting instabilities occur for any symmetry channel, doping and both for J = 0 and J ≠ 0. The point is more whether the corresponding Tc is very small or large enough to be relevant. Tc here is understood as the transition temperature within mean-field theory. This means that the lowering of Tc due to fluctuations in the superconducting order parameter is not taken into account. This assumption is justified if one compares with real, three-dimensional superconductors but does not apply, of course, to strictly two-dimensional models where fluctuations drive the order parameter to zero at any finite temperature. In order to determine Tc one has to solve the gap equation which has several non-BCS features: The kernel of the linearized gap equation consists of an instantaneous and a retarded term and both are characterized by different cutoffs. Moreover, the presence of the instantaneous term does not allow to restrict the momenta to the Fermi surface. Consequently, we solved the gap equation by numerical means. Our method is also suitable to investigate the question of odd frequency pairing in our model. In this case the instantaneous term drops out but the full frequency dependence of the kernel must be kept in addition to the momentum dependence along the Fermi line. Results for odd frequency pairing will also be presented in section IV together with the conclusions.

II. MODEL AND GENERAL EQUATIONS FOR THE ELECTRON GREEN’S FUNCTION

The Hamiltonian of the t – J model can be written as

\[
H = \sum_{p=1}^{N} \frac{t_{ij}}{N} X_i^{p0} X_j^{0p} + \sum_{ij} \sum_{p,q=1}^{N} \frac{J_{ij}}{4N} X_i^{pq} X_j^{qp} - \sum_{ij} \sum_{p,q=1}^{N} \frac{J_{ij}}{4N} X_i^{pp} X_j^{qq}.
\]

(1)

Let us consider first the case N = 2. Assuming one orbital per site and excluding double occupancies of sites there are three states \( |p_i> \) at each atom i. p = 0 denotes the empty state, and p = 1, 2 singly occupied states with spin up and down. The Hubbard operators \( X_i^{pq} \) can be represented as projection operators \( X_i^{pq} = \phi^p_i \phi^q_i \) and obey the following commutator and anticommutator relations

\[
[X_i^{pq}, X_j^{rs}]_{\pm} = \delta_{ij}(\delta_{qr} X_i^{ps} \pm \delta_{sr} X_i^{rq}),
\]

(2)

and the completeness relation
\[ \sum_{p=0}^{2} X_{i}^{pp} = 1. \quad (3) \]

The upper (lower) sign in Eq.(2) holds for bosonlike or mixed (fermionlike) Hubbard operators defined by \( p, q > 0 \) or \( p = q = 0 \) (\( p = 0, q > 0 \) or \( p > 0, q = 0 \)). For \( i = j \) both the upper and lower signs hold in each case. The first term in Eq.(1) describes the hopping of particles between the sites \( i \) and \( j \) with matrix elements \( t_{ij} \). The second term in Eq.(1) denotes the Heisenberg interaction between the spin densities at site \( i \) and \( j \) with the exchange constants \( J_{ij} \). The third term in Eq.(1) represents the charge-charge interaction of the \( t - J \) model.

In the following we consider \( J_{ij} \) only between nearest neighbors (\( J_{ij} = J \)) and \( t_{ij} \) between nearest (\( t_{ij} = t \)) and next nearest (\( t_{ij} = t' \)) neighbors. We also use always \(|t|\) as energy unit.

The Hamiltonian in Eq.(1) is a generalization from \( N = 2 \) to an (even) arbitrary integer \( N \). The orbital index \( p \) consists now of the spin and a flavor index, the latter enumerating \( N/2 \) identical orbitals at a site. The symmetry group of \( H \) is the symplectic group \( \text{Sp}(N/2) \).

For \( N > 2 \) the operators \( X \) can no longer be written as projection operators in some basis. Instead, fermionlike (bosonlike or mixed ones) operators are assumed to satisfy the anticommutator (commutator) relations of Eq.(2). Only some of the diagonal operators can assumed to retain their projection properties, namely, \( (X_{i}^{pp})^2 = X_{i}^{pp} \) for \( p > 0 \). Most other relations characteristic of projection operators such as \( X_{i}^{10}X_{i}^{01} = X_{i}^{11} \) are lost for \( N > 2 \). The completeness relation Eq.(3) is replaced by the constraint

\[ Q_{i} = \sum_{p=0}^{N} X_{i}^{pp} = N/2. \quad (4) \]

By explicit construction of the Hilbert space and the action of the \( X \)'s on its vectors one can show that the above properties, together with \( H \), specify completely the problem. In particular, \( X_{i}^{p0} \) is a non-negative operator. As a result, Eq.(4) means that at most \( N/2 \) particles can occupy the \( N \) available states at a site. In this way one may expect that expectation values of observables approach smoothly the physical case \( N = 2 \) from large \( N \)'s yielding a basis for \( 1/N \) expansions. We also would like to point out that slave boson treatments of \( H \) have many features in common with our approach. However, the Hilbert spaces are different in the two cases as well as the enforcement of the constraint Eq.(4). \( Q_{i} \) commutes with all Hubbard operators and thus is proportional to the identity in any irreducible representation of the \( X \)-operators. Enforcement of the constraint means in our case just the selection of the correct subspace of the Hilbert space where Eq.(4) is satisfied as an operator identity.

Following the Baym-Kadanoff procedure we define a non-equilibrium Matsubara Green’s function of two fermionic operators \( X(1) \) and \( X(2) \) by

\[ G(12) = - \frac{< T S X(1) X(2) >}{< S >}, \quad (5) \]

\[ S = T e^{\int d\tau X(1) K(1)}. \quad (6) \]

The number 1 stands for an internal pair index \( p, q \) as well as a site and a (imaginary) time index \( i, \tau \). \( \int d\tau \) means \( \sum_{p,q,i} \int_{0}^{\beta} d\tau \), where \( \beta \) is the inverse temperature. In Eqs.(5) and (6)
$T$ is the time ordering operator and $K$ an external source term which is assumed to couple only to bosonic X-operators. We also introduce the non-equilibrium expectation value of bosonic X-operators by

$$L(1) = \langle T S X(1) \rangle / \langle S \rangle .$$  \hspace{1cm} (7)

Using the Hamiltonian Eq.(1) the Heisenberg equation of motion for fermionic operators becomes

$$\frac{\partial}{\partial \tau_1} X(1) = \int d2d3t(123) X(2) X(3) ,$$  \hspace{1cm} (8)

with

$$t(123) = \delta(\tau_2 - \tau_1) \delta(\tau_3 - \tau_1) \left[ \frac{t_{i_1 i_2}}{N} \left( \delta_{q_1 0} \delta_{q_3 0} (1 - \delta_{p_3 0}) \delta_{p_2 0} \delta_{q_2 q_1} \delta_{p_3 p_1} + \delta_{p_2 p_1} \delta_{q_2 q_3} \right) ight] ,$$  \hspace{1cm} (9)

Eq.(9) agrees with Eq.(5) of Ref.\cite{19} if the different sign convention for the hopping term is taken into account. Using Eq.(8) and rewriting higher-order correlation functions in terms of functional derivatives with respect to $K$ the equation of motion for $G$ can be written as

$$\int d2 (G^{-1}_0(12) - \Sigma'(12)) G(21') = Q(11'),$$  \hspace{1cm} (10)

$$\Sigma'(12) = - \int d3 t(132) L(3) + \int d3 d4 d5 t(134) G(45) \Gamma(52, 3),$$  \hspace{1cm} (11)

$$\Gamma(12, 3) = \delta G^{-1}(12) / \delta K(3).$$  \hspace{1cm} (12)

$G_0$ is the unperturbed Green’s function

$$G^{-1}_0(12) = - \delta(1 - 2) \frac{\partial}{\partial \tau_2} - \delta(\bar{1} - 2)(K_{00}(\bar{1}) \delta_{q_1 q_2} - K^{q_1 q_2}(\bar{1})).$$  \hspace{1cm} (13)

$Q$ is given by

$$Q(11') = \delta(1 - 1')(L^{pq'}(\bar{1}) \delta_{q' p'} + L^{q' q}(\bar{1}) \delta_{q' p'}).$$  \hspace{1cm} (14)

where the index $1 = (pq, i, \tau)$ has been split into $1 = (pq, \bar{1})$ with $\bar{1} = (i\tau)$. $L$ can be expressed by $G$ so the above system of equations for $G$, the self-energy $\Sigma'$, and the vertex $\Gamma$ is closed.

For our purposes it is more convenient to use a normalized Green’s function $g$ which obeys a Dyson equation with the usual $\delta$-function on its right-hand side. Writing

$$g(11') = \int d2 G(12) Z^{-1}(21'),$$  \hspace{1cm} (15)

and requesting that $Z$ satisfies the following equation...
\[ Z(11') = Q(11') - \int d3d4d5t(134)g(45)\frac{\delta Z(51')}{\delta K(3)} , \]  

(16)

Eqs(10)-(12) assume the form
\[
\int d2(G_0^{-1}(12) - \Sigma(12))g(21') = \delta(1 - 1'), \tag{17}
\]

\[
\Sigma(12) = - \int d3t(132)L(3) + \int d3d4d5t(134)g(45)\gamma(52, 3), \tag{18}
\]

\[
\gamma(12, 3) = \delta g^{-1}(12)/\delta K(3). \tag{19}
\]

From Eqs.(17) and (19) follows, moreover, the equation for the vertex
\[
\gamma(11', 3) = \alpha(11'; 3) + \int d4d5\Theta(11', 45)\gamma(45, 3), \tag{20}
\]

with
\[
\alpha(11', 3) = \frac{\delta G_0^{-1}(11')}{\delta K(3)} - \int d4\frac{\delta \Sigma(11')}{\delta L(4)} \cdot \frac{\delta L(4)}{\delta K(3)}. \tag{21}
\]

\[
\Theta(11', 45) = \int d6d7\frac{\delta \Sigma(11')}{\delta g(67)}g(64)g(57). \tag{22}
\]

In Eq.(22) we have replaced the functional derivative of \( g \) with respect to \( K \) by the vertex \( \gamma \) using Dyson’s equation which also leads to a sign change. The above equations are exact and hold both in the normal and the superconducting state. In the normal state the two internal index pairs in \( G(12) \) must obey either \( p_1 = q_2 = 0 \) or \( p_2 = q_1 = 0 \). From Dyson’s equation follows then that the same holds for \( Q(12) \) whereas the possible indices in \( G_0(12), \Sigma(12), \) and \( g(12) \) obey either \( p_1 = p_2 = 0 \) or \( q_1 = q_2 = 0 \). In the superconducting state there are no such restrictions; the only general requirement is that all these indices are associated with fermionic operators.

The self-energy is a functional of \( g \) and \( L \), i.e.,
\[
\Sigma = \Sigma[g, L]. \tag{23}
\]

In order to derive the linearized equation for the anomalous self-energy we split \( \Sigma \) into the normal part \( \Sigma_N \) and a small anomalous part \( \Sigma_{an} \). Expanding Eq.(23) up to linear terms we obtain
\[
\Sigma_{an}(11') = \int d2d3(\frac{\delta \Sigma(11')}{\delta g(23)})_N \delta g(23) + \int d2(\frac{\delta \Sigma(11')}{\delta L(2)})_N \delta L(2), \tag{24}
\]

where \( \delta g \) and \( \delta L \) are of first order in the anomalous part and the subscript \( N \) means that the functional derivatives are to be taken in the normal state. Now \( \delta L \) has no linear contribution.
due to gauge invariance and thus drops out in Eq.(24). Calculating \( \delta g \) from Dyson’s equation Eq.(17) in linear approximation we obtain

\[
\Sigma_{an}(11') = \int d2d3\Theta(11', 23)\Sigma_{an}(23),
\]

(25)

where \( \Theta \) is given by Eq.(22) with all quantities taken in the normal state. Eq.(25) represents the linearized equation for a general order parameter for superconductivity. In order to have superconductivity the homogenous equation Eq.(25) must have a nonvanishing solution for \( \Sigma_{an} \). The highest temperature where this occurs is the transition temperature \( T_c \).

The basic quantity to be calculated is according to Eq.(25) the kernel \( \Theta \). In order to find a functional equation for \( \Theta \) we get from Eq.(18)

\[
\frac{\delta \Sigma(11')}{\delta g(78)} = \int d2t(127)g(81', 2) + \int d2d3d6d7d8t(123)g(36)\frac{\delta g(78)}{\delta g(78)},
\]

(26)

and from Eq.(20)

\[
\frac{\delta \gamma(61', 2)}{\delta g(78)} = \frac{\delta \alpha(61', 2)}{\delta g(78)} + \int d9d10d9'd10'\Theta(61', 9'10')\frac{\delta \gamma(9', 10')}{\delta g(78)} + \int d9d10'\frac{\delta \Theta(61', 9'10')}{\delta g(78)}g(9', 10', 2).
\]

(27)

Solving Eq.(27) for \( \delta \gamma/\delta g \), inserting the result into Eq.(26) and inserting Eq.(26) into Eq.(22) yields an exact functional equation for \( \Theta \).

III. 1/N EXPANSION FOR THE KERNEL \( \Theta \)

The equation for \( \Theta \) obtained in the previous section is too difficult to be solved directly. On the other hand it is exact and holds for any \( N \) so it may serve as a starting point for approximate treatments. In the following we assume that \( 1/N \) may be used as a small parameter and calculate \( \Theta \) up to order \( 1/N \). This will allow us to obtain the gap equation Eq.(25) in leading order of the \( 1/N \) expansion.

The \( N \)-dependence of an equilibrium quantity is determined by the number of coupling constants it contains and the number of free internal summations. In equilibrium, i.e., without the source term \( K \), we have, for instance,

\[
g(0q_1 0q_2) = \delta_{q_1q_2}g(\bar{1} - \bar{2}),
\]

(28)

\[
\Sigma(0q_1 0q_2) = \delta_{q_1q_2}\Sigma(\bar{1} - \bar{2}),
\]

(29)

\[
\gamma^{0q_1 0q_2, p_3q_3}(\bar{1}, 2; \bar{3}, 3) = -\frac{1}{N}\delta_{p_3q_3}\delta_{q_1q_2}\gamma_c(\bar{1}, \bar{2}, 3) + \delta_{q_1p_3}\delta_{q_2q_3}\gamma_s(\bar{1}, \bar{2}, 3),
\]

(30)

and similar expressions for \( \alpha \) and \( \Theta \). In Eqs.(28)-(30) all the indices \( q_1, q_2, \ldots \) are assumed to be larger than zero. The equilibrium Green’s function \( g \), the self-energy \( \Sigma \), the charge
vertex $\gamma_c$ and the spin vertex $\gamma_s$ are free of internal indices. In the following we need only the leading $O(1)$ contributions for these quantities. From Eqs.(13), (17) and (19) follows that the spin vertex becomes then simply equal to $\delta(\bar{1} - 2)\delta(\bar{1} - 3)$. Using the constraint Eq.(4) one also recognizes that the charge vertex $\gamma_c$ defined in Eq.(30) is equal to the element $q_1 = q_2 > 0, p_3 = q_3 = 0$ of the general vertex $\gamma$ on the left-hand side of Eq.(30) which is the motivation to use prefactors in Eq.(30) in defining $\gamma_c$.

Decomposing the labels $1,1', \cdots$ into their internal and external parts the anomalous self-energy or order parameter has the form $\Sigma_{an}(\gamma_1 \gamma_1', \gamma_1' \gamma_1)$, or in more detail, $\Sigma_{an}(\gamma_1 \gamma_1', \gamma_1' \gamma_1)$. In the following we are interested in order parameters which are structureless in the flavor indices, i.e., we assume always $m_1 = m_1'$. With respect to spin indices we consider either singlet or triplet pairing. Both cases are included if we put $\sigma_1 = -\sigma_1'$. Correspondingly we will write for the internal indices $p_1, \bar{p}_1$ thus indicating the special relationship between these two indices. Exposing explicitly the internal indices the gap equation Eq.(25) becomes

$$\Sigma_{an}(p_1 \gamma_1 0 \bar{p}_1) = \sum_{p_2} \int d2d3\Theta(p_1 \gamma_1 0 \bar{p}_1, p_2 \gamma_2 0 \bar{p}_2) \Sigma_{an}(p_2 \gamma_2 0 \bar{p}_2),$$  \hspace{1cm} (31)$$

where $\Theta$ is to be taken in the normal state. Eq.(31) also makes visible which indices for $\Theta$ are actually needed in the gap equation. A closer examination shows that the first two contributions on the right-hand side of Eq.(27) are of higher orders in $1/N$ than the third term in this equation for the needed combination of indices. Dropping these terms and inserting Eqs.(26) and (27) into Eq.(22) yields the following equation for $\Theta$ valid up to $O(1/N)$ and for the above combinations of internal indices:

$$\Theta(11', 910) = \int d2d7d8t(127)\gamma(81', 2)g(79)g(108) +$$

$$\int d2d3d6d7d9d10't(123)g(36)\frac{\delta\Theta(61', 9'10')}{\delta g(78)}\gamma(9'10', 2)g(79)g(108).$$ \hspace{1cm} (32)$$

The first term on the right-hand side of Eq.(32) could have been also obtained directly from Eq.(18). Considering only anomalous contributions to $\Sigma$ the first term on the right-hand side of Eq.(18) cannot contribute, whereas the second one can contribute in two ways: a) via an anomalous $g$ and a normal vertex $\gamma$, b) via a normal $g$ and an anomalous vertex $\gamma$. Case a) corresponds to the usual situation of Eliashberg theory and also of the slave boson treatment of superconductivity in the $t - J$ model in $O(1/N)$: $\Sigma_{an}$ consists then of a Fock diagram containing an anomalous Green’s function and an effective interaction taken in the normal state. Linearizing $g$ in $\Sigma_{an}$ case a) immediately yields the first term on the right-hand side of Eq.(32). Its evaluation is straightforward and shows that it is of $O(1/N)$ for the relevant combination of internal indices. This indicates that $T_c \rightarrow 0$ for $N \rightarrow \infty$ which agrees with the fact that at $N = \infty$ the system consists of renormalized, but non-interacting fermions.

The leading contribution of the second term on the right-hand side of Eq.(32) is obtained by taking the spin-flip part in the hopping matrix element $t$ and the spin vertex $\gamma_s$ in $O(1)$ for the vertex $\gamma$. As a result only one sum over internal indices survives. The second term in Eq.(32) is of $O(1/N)$ if the two functions
\begin{align}
  g^{(1)}(\bar{6}\bar{1}', 2|78) &= \sum_{p_3} \frac{\delta \Theta \left( p_{3}^{0,0} p_{1}^{0}, p_{3}^{0} p_{1}^{0} \right)}{\delta g \left( p_{3}^{7} p_{1}^{8} \right)} , \tag{33}
  \\
  g^{(2)}(\bar{6}\bar{1}', 2|78) &= \sum_{p_3} \frac{\delta \Theta \left( p_{3}^{0,0} p_{1}^{0}, p_{3}^{0} p_{1}^{0} \right)}{\delta g \left( p_{3}^{7} p_{1}^{8} \right)} , \tag{34}
\end{align}

are of \( O(1) \). Equations for \( g^{(i)}, i = 1, 2 \) can be obtained from Eq.(32) by taking appropriate derivatives. An examination of the various terms shows that terms with a second functional derivative of \( \Theta \) are smaller by a factor \( 1/N \) compared to the leading ones and thus can be neglected. One then finds that the functions \( g^{(i)} \) have the form

\begin{align}
  g^{(1)}(\bar{6}\bar{1}', 2|78) &= \delta(2 - 8) g^{(1)}(\bar{6}\bar{1}', 2|7) , \tag{35}
  \\
  g^{(2)}(\bar{6}\bar{1}', 2|78) &= \delta(2 - 7) g^{(1)}(\bar{6}\bar{1}', 2|8) . \tag{36}
\end{align}

The reduced functions \( g^{(i)}(\bar{6}\bar{1}', 2|\bar{8}) \) (the use of the same name for corresponding functions with different number of arguments should not cause any confusion) satisfy the integral equation

\begin{align}
  g^{(i)}(\bar{1}\bar{1}', 2|7) &= h^{(i)}(\bar{1}\bar{1}', 2|7) - \int d\bar{3}d\bar{4}d\bar{6} N t(\bar{1}\bar{3}4) g(\bar{6}\bar{4}) g^{(i)}(\bar{6}\bar{1}', 3|7) g(23) , \tag{37}
\end{align}

with

\begin{align}
  h^{(1)}(\bar{1}\bar{1}', 2|7) &= N t(\bar{1}\bar{1}7) g(21') , \tag{38}
  \\
  h^{(2)}(\bar{1}\bar{1}', 2|7) &= - \int d\bar{3}d\bar{4} N t(\bar{1}\bar{3}4) \gamma_{c} (\bar{7}\bar{1}', 4) g(23) . \tag{39}
\end{align}

Here \( t(123) \) denotes the external part of \( t(123), \text{i.e.,} \) the first three factors outside of the bracket in Eq.(9). Inserting the explicit expression for \( t \) into Eq.(37) one finds that Eq.(37) represents an integral equation for \( g^{(i)} \) with a kernel consisting of not more than 6 separable contributions. The solution of Eq.(37) thus reduces to the inversion of at most 6x6 matrices similar as in the case of the charge vertex \( \gamma_{c} \) dicussed in Ref. Inserting the resulting expressions back into Eq.(32) and performing Fourier transforms one obtains after some elementary, but tedious algebra the following results. The linearized gap equation Eq.(25) becomes

\begin{align}
  \Sigma_{an}(k) &= - \frac{T}{N N_{c}} \sum_{k'} \Theta(k, k') \frac{1}{\omega_{n'}^{2} + \epsilon^{2}(k')} \Sigma_{an}(k') . \tag{40}
\end{align}

\( N_{c} \) is the number of cells and \( k \) the supervector \( k = (n, k) \), where \( n \) denotes a fermionic Matsubara frequency \( \omega_{n} = (2n + 1) \pi T \) with \( T \) being the temperature. \( \epsilon(k) \) is the one-particle energy with momentum \( k \) in the limit \( N \rightarrow \infty \) given by the implicit equation

\begin{align}
  \epsilon(k) &= \frac{\delta}{2} t(k) - J(k) \cdot \frac{1}{N_{c}} \sum_{P} \cos(p_{z}) f(\epsilon(p - \mu)) \tag{41}
\end{align}
where $\delta$ is the doping, $t(k)$ and $J(k)$ the Fourier transforms of the coupling constants $t_{ij}$ and $J_{ij}$, respectively, $\mu$ is the chemical potential and $f$ the Fermi function.

The kernel $\Theta$ in Eq.(40) consists of four different terms

$$\Theta(k, k') = \Theta^{(1)}(k, k') + \Theta^{(2)}(k, k') + \Theta^{(3)}(k, k') + \Theta^{(4)}(k, k').$$

(42)

The first two terms are given by

$$\Theta^{(1)}(k, k') = \mp t(k') \mp J(k - k') - t(k) - J(k - k'),$$

(43)

$$\Theta^{(2)}(k, k') = (t(k') + J(k - k'))(\gamma_c(k, k' - k) + 1).$$

(44)

Here, $\gamma_c$ is the charge vertex in $O(1)$ determined by

$$\gamma_c(k, q) = -1 + \sum_{\alpha, \beta = 1}^{6} F_{\alpha}(k)(1 + \chi(q))^{-1}_{\alpha \beta} \chi_{\beta 2}(q),$$

(45)

with the susceptibility matrix

$$\chi_{\alpha \beta}(q) = \frac{1}{Nc} \sum_k E_{\alpha}(k, q) F_{\beta}(k) \frac{f(\epsilon(k + q)) - f(\epsilon(k))}{\epsilon(k + q) - \epsilon(k) - i\nu_n}.$$

(46)

$q$ is the supervector $q = (q, i\nu_n)$ where $\nu_n$ denotes the bosonic Matsubara frequency $2\pi n T$.

The two vectors $E$ and $F$ are given by

$$E_{\alpha}(k, q) = (1, t(k + q) + J(q), \cos k_x, \sin k_x, \cos k_y, \sin k_y),$$

(47)

$$F_{\beta}(k) = (t(k), 1, J\cos k_x, J\sin k_x, J\cos k_y, J\sin k_y).$$

(48)

The explicit expressions for the third and fourth contributions to $\Theta$ are

$$\Theta^{(3)}(k, k') = -\sum_{r=1}^{5} \tilde{E}_r(-k) \tilde{\chi}_{2r}(k - k') \gamma(k, k' - k) + \sum_{r,s=1}^{5} \tilde{E}_s(-k) \tilde{\chi}_{rs}(k - k') \tilde{\gamma}_r(k - k'),$$

(49)

$$\Theta^{(4)}(k, k') = \pm \sum_{r,s=1}^{5} (\tilde{E}_r(-k) \tilde{\chi}_{rs}(k + k'))(\tilde{\gamma}_r(-k') + \tilde{\gamma}_r(k + k')),$$

(50)

with the vectors $\tilde{V}$ and $\tilde{W}$

$$\tilde{V}_r(k - k') = \sum_{s=1}^{5} (1 + \tilde{\chi}(k - k'))^{-1}_{rs} \tilde{\chi}_{2s}(k - k') \gamma_c(k, k' - k),$$

(51)

$$\tilde{W}_r(k + k') = -\sum_{s,t=1}^{5} (1 + \tilde{\chi}(k + k'))^{-1}_{rs} \tilde{\chi}_{ts}(k + k').$$

(52)
The vectors $\tilde{E}$ and $\tilde{F}$ have five components and are obtained from the six-component vectors $E$ and $F$ by omitting the second component. In an analogous way, $\tilde{\chi}_{2,s}$ is obtained from the second row of the matrix $\chi$ by omitting the second element. Similarly, the 5x5 matrix $\tilde{\chi}$ is obtained from the 6x6 matrix $\chi$ by dropping the second row and second column. The upper (lower) signs in the contributions $\Theta^{(i)}$ refer to singlet (triplet) pairings.

The kernel $\Theta(k, k')$ is invariant under the transformations of the point group $C_{4v}$ of the square lattice. As a result $\Sigma_{an}(k)$ transforms corresponding to one of the five irreducible representations $\Gamma_i$ of $C_{4v}$, and it can be chosen to be either even or odd in Matsubara frequencies. $\Gamma_1$ corresponds to singlet s- or extended s-wave, $\Gamma_3$ to singlet d-wave- and $\Gamma_5$ to triplet p-wave pairing for even frequency pairing. The linearized gap equation Eq.(40) splits completely into its irreducible parts. This also means that the momenta $k,k'$ can be restricted to 1/8 of the Brillouin zone which greatly simplifies the numerical solution of Eq.(40). In the following we will deal with even frequency pairing unless the opposite is explicitly stated and drop for simplicity the index which differentiates between even and odd frequency-pairing. The irreducible kernels and order parameters will thus be denoted by $\Theta_i(k, k')$ and $\Sigma_{an,i}(k)$, respectively.

IV. DISCUSSION OF THE GAP EQUATION AND NUMERICAL RESULTS

According to Eq.(42) the total kernel $\Theta$ is the sum of four contributions. The first one, $\Theta^{(1)}$, represents the instantaneous part of $\Theta$. From Eq.(40) follows that positive values for $\Theta^{(1)}$ correspond to repulsion, negative values to attraction between electrons. Keeping only $\Theta^{(1)}$ and decomposing it into its irreducible symmetry components the gap equation Eq.(40) becomes for the representation $i=1...5$ of the point group $C_{4v}$ of a square lattice

$$\Sigma_{i,an}(k) = \frac{-1}{NN_c} \sum_{k'} \Theta^{(1)}_i(k, k') \chi(k') \Sigma_{i,an}(k'),$$

with

$$\chi(k') = \frac{\tanh((\epsilon(k') - \mu)/T)}{2(\epsilon(k') - \mu)},$$

$$\Theta^{(1)}_2 = \Theta^{(1)}_4 = \Theta^{(1)}_5 = 0,$$

and

$$\Theta^{(1)}_1(k, k') = 8|t|\eta_1(k') - 8J\eta_1(k)\eta_1(k'),$$

$$\Theta^{(1)}_3(k, k') = -8J\eta_3(k)\eta_3(k').$$

In Eqs.(55)and (56) we assumed nearest-neighbor hopping with $t_{ij} = -|t|$ and used the basis functions $\eta_1(k) = (\cos(k_x) + \cos(k_y))/2$, $\eta_3(k) = (\cos(k_x) - \cos(k_y))/2$. Inserting the kernels Eqs.(55) and (56) into the gap equation yields the following conditions for a finite $T_c$:

$$s - \text{wave pairing (} \Sigma_{an} \sim \text{constant) : } 1 + a|t| + \frac{ab_1|t|J}{1 - Jb_1} = 0,$$
extended s-wave pairing \((\Sigma_{an} \sim \eta_1(k))\): 
\[1 - b_1 + \frac{a b_1 |t| J}{1 + a |t|} = 0,\] \(58\)

d-wave pairing \((\Sigma_{an} \sim \eta_3(k))\): 
\[1 - b_3 J = 0.\] \(59\)

Here we used the following abbreviations
\[a = \frac{8}{NN_c} \sum_{k'} \chi(k') \eta_1(k'),\] \(60\)
\[b_l = \frac{8}{NN_c} \sum_{k'} \chi(k') \eta_l^2(k'),\] \(61\)

for \(l = 1, 3\). A numerical evaluation shows that in the interesting parameter region \(0 \leq J \leq 0.3, 0 \leq \delta \leq 0.8\) (\(\delta\) is the doping) and at low temperatures \(a, b_1, b_3\) are positive and \(b_1 < 3\). This means that there is never an instability with respect to constant s-wave pairing. \(b_1\) is an increasing, \(b_3\) a strongly decreasing function with increasing \(\delta\) and these two functions cross around \(\delta \sim 0.6\). Thus d-wave pairing is stable in any case for \(\delta < 0.5\). For \(0.5 \leq \delta \leq 0.8\) \(a |t|\) is much larger than 1 so that the third term in Eq.(58) cancels the second term to a large extent making extended s-wave pairing also in this region unfavourable. Thus we find that the instantaneous contribution to the kernel strongly favours d-wave pairing in the interesting parameter regime. The reason for this is to a large extent a band structure effect: \(b_3\) is much larger than \(b_1\) at small dopings because of large contributions around the X-point. At larger doping the competing extended s-wave pairing is strongly suppressed by its coupling to the hopping term, which ultimately is caused by the constraint.

\(\Theta^{(2)}, \Theta^{(3)},\) and \(\Theta^{(4)}\) are retarded contributions to \(\Theta\). \(\Theta^{(2)}\) is mainly determined by collective charge fluctuations due to the poles of \(\gamma_c\). \(\Theta^{(3)}\) and \(\Theta^{(4)}\) originate from the anomalous part of the vertex and involve both charge and spin fluctuations. The latter dominate in \(\Theta^{(4)}\) in agreement with the sign change between its singlet and triplet contribution. Quantitatively, \(\Theta^{(4)}\) is by far the largest of the retarded contributions. Our \(\Theta\) does not contain terms which are related to the magnons of the undoped case. Such contributions are of higher order in \(1/N\) than those considered above and are thus neglected.

Our expression for \(\Theta\) is different from that of the slave boson approach. \(\Sigma_{an}\) in the slave boson approach essentially consists of a spinon Fock term with an effective interaction taken in the normal state times an anomalous spinon Green’s function. Such an contribution clearly corresponds to the case a) discussed after Eq.(32), i.e., to the first term on the right-hand side of Eq.(32), or, equivalently, to the sum of \(\Theta^{(1)}\) and \(\Theta^{(2)}\). Our contributions \(\Theta^{(3)}\) and \(\Theta^{(4)}\) have no analogue in the slave boson approach though they are clearly also of \(O(1/N)\). The differences between the two approaches can be made more explicitly by considering two limiting cases. For \(J \to 0\) \(\Theta\) reduces to Eq.(9) of Ref.\[11\] The underlying interactions are still spin-dependent as can be seen from the singlet versus triplet case. Such a dependence on spin is not present in the corresponding slave boson expression\[12\]. Furthermore, the high-frequency limit of \(\Theta\), i.e., \(\Theta^{(1)}\) should be identical to that of the slave boson expression if the two approaches are equivalent. The slave boson result for \(\Theta^{(1)}\) differs, however, from Eq.(43): the argument in one of the two hopping terms is replaced by \(k\) and there is an additional
term proportional to $1/\delta$. Eq.(5) agrees, however, with previous X-operator results based on the diagrammatic methods for X-operators, for instance, Eq.(35) of Ref.20. The expression for $\Theta^{(1)}$ of Ref.7 contains only one of the two hopping terms $t(k')$ of Eq.(5) which, we think, is incorrect. The general question then arises why it is possible to obtain different $O(1/N)$ expressions for the same quantity in the two approaches using the same Hamiltonian. The differences arise because different Hilbert spaces are used in the two cases. This difference is already indicated in the definition of the order parameters for superconductivity. In the X-operator approach all vectors of the Hilbert space are eigenvectors of the constraints $Q_i$ with eigenvalue $N/2$. Expectation values of operators are only nonzero if, in the slave boson language, the number of created slave particles is equal to the number of annihilated slave particles in these operators. The superconducting order parameter of slave boson theory, on the other hand, consists of the expectation values $\langle b \rangle$ and $\langle c^{\dagger}c^{\dagger} \rangle$. Clearly, these expectation values have no analogues in the X-operator approach.

A. Eigenvalues of $\Theta$ in the static limit

In the weak-coupling case it is often assumed that the solution of the gap equation Eq.(40) is of BCS-type for each symmetry $\Gamma_i$, i.e.,

$$T_c = 1.13 \omega_c e^{1/\lambda_i}. \quad (62)$$

$\omega_c$ is a suitable cut-off and $\lambda_i$ the smallest eigenvalue of a matrix which is essentially equal to the static limit of the kernel $\Theta$ and given by Eq.(10) of Ref.11. This matrix includes also the prefactor $1/N$ on the right-hand side of Eq.(40) and we put from now on $N$ always equal to 2. Actually we will find that Eq.(62) is a rather poor approximation for $T_c$ because $\Theta$ contains instantaneous and retarded contributions which have different cut-offs. Nevertheless the study of the static kernel $\Theta$ and its lowest eigenvalues has traditionally played a great role in discussing superconductivity in $t$-$J$ models.

Fig. 1a) shows the terms $\Theta^{(1)}$ and $\Theta^{(2)}$ and Fig. 1b) the terms $\Theta^{(3)}, \Theta^{(4)}$ and $\Theta$ for singlet pairing at zero frequencies for a fixed first momentum $k = (2.465, 0.309)$ as a function of the second argument $k'$. $k'$ moves counterclockwise around the Fermi line passing through the points $X, Y (\bar{X}, \bar{Y})$ along the positive (negative) x- and y-axis, respectively. Note that the letters $X, Y$ denote not the $k$-points $(\pi, 0), (0, \pi)$ but the points on the Fermi line between the $\Gamma$-point and the points $(\pi, 0), (0, \pi)$, respectively. The doping is $\delta = 0.17$ and $J = 0.3$. We used 22 $k$-points along 1/8 of the Fermi line and a net of 300x300 $k$-points in the Brillouin zone. Positive values of $\Theta$ mean repulsion, negative ones attraction between electrons in the s-wave channel. $\Theta^{(1)}$ and $\Theta^{(2)}$ originate from the normal, $\Theta^{(3)}$ and $\Theta^{(4)}$ from the anomalous vertex. $\Theta^{(1)}$ is due to instantaneous, the other terms due to retarded interactions. The Figures show that $\Theta^{(4)}$ and $\Theta^{(1)}$ are by far the largest contributions. Both are dominated by the d-wave component and add up in a coherent way. $\Theta^{(4)}$ has different signs for singlet and triplet pairings. This as well as the explicit calculation shows that important contributions to $\Theta^{(4)}$ come from spin fluctuations within the partially filled band. There are no spin contributions related to the Heisenberg term and the magnon spectrum at zero doping because these terms are at least one order in $1/N$ smaller than the considered ones. $\Theta^{(3)}$ is due to charge (and spin) excitations and it is very small. Finally, $\Theta^{(2)}$ is due to charge
fluctuations and consists of an attractive s-wave component (which, however, is cancelled by a repulsive s-wave term in $\Theta^{(1)}$) and a d-wave part which cancels partially that of the $\Theta^{(1)}$ term. Figs.1 demonstrate the importance of anomalous vertex contributions to $\Theta$, the d-wave character of the leading contributions, and the competition between instantaneous and retarded interactions.

Fig. 2 shows the lowest eigenvalues $\lambda_i$ for each of the five representations $\Gamma_i$. In the calculation we used 5 $k$-points along 1/8 of the Fermi line and a net of 300x300 $k$-points in the Brillouin zone. The eigenvalues $\lambda_i$ decrease with decreasing doping $\delta$ and diverge at $\delta \sim \delta_{BO} \sim 0.13$. At this doping value one of the six eigenvalues of the 6x6 matrix $1 + \chi$ in Eq.(45) goes through zero. As a result there is a soft mode which freezes into a static incommensurate bond-order wave of d-wave symmetry for $\delta < \delta_{BO}$. This instability causes a divergence in some of the contributions to $\Theta$ at $\delta_{BO}$ and, as a precursor, the large negative values seen in Fig. 2 for $\delta \leq 0.15$. The solid line in Fig. 2 exhibits the lowest eigenvalue $\lambda_3$ associated with d-wave pairing. Its absolute magnitude is much larger than all the other eigenvalues and this is true for all dopings. Inserting $\lambda_i$ into Eq.(40) it is clear that superconducting instabilities for non-d-wave symmetries are more of academic interest because the corresponding $T'_c$s would be extremely small. One concludes that the normal state is generally unstable against superconductivity in every symmetry channel but that only the d-wave ($\Gamma_3$) instability is strong enough to account for high transition temperatures.

B. Frequency dependence of $\Theta$

For either even or odd frequency pairing the frequency dependence of $\Theta$ can be rewritten in terms of one frequency argument $\omega_n$ which is equal to the difference of the two original frequency variables. In order to illustrate the frequency dependence of the retarded part of $\Theta$, $\Theta^{ret}$, in the case of d-wave scattering, we average the momenta in $\Theta^{ret}$ over the Fermi line in the irreducible Brillouin zone using the eigenvector of the lowest eigenvalue $\lambda_3$. Finally we perform the analytic continuation $i\omega_n \rightarrow \omega + i\eta$.

Fig.3 shows the negative imaginary part of $\Theta_3^{ret}(\omega + i\eta)$ for $\eta = 0.005$. It is the analogue of the familiar function $\alpha^2 F(\omega)$ of Eliashberg theory for phonon-induced s-wave superconductivity. In our case this function is no longer positive-definite. This is a result of the projection of the total kernel on the $\Gamma_3$ symmetry. This projection involves a sum of $k'$ over the 8 symmetry-related points along the Fermi line with factors determined by the representation $i$. In the case of d-wave pairing these factors are positive for small and large momentum transfers (of the order or a reciprocal lattice vector) and negative for intermediate momentum transfers of about half of a reciprocal lattice vector. For collective density fluctuations described by $\Theta^{(2)}$ this means that high-frequency contributions appear with a negative and low-frequency contributions with a positive sign in the d-wave kernel. A similar behavior is found for the other contributions. As a result $-Im\Theta_3^{ret}$ is positive for $\omega \leq \omega_2 \sim |t|$ and negative for $\omega \geq \omega_2$. Using a momentum average with a constant weight of $\Theta^{ret}$ corresponding to constant s-wave pairing would yield a curve which is similar to that in Fig.3 for $\omega \leq \omega_2$ but opposite in sign for $\omega \geq \omega_2$. $-Im\Theta_3^{ret}$ describes the spectral distribution and spectral weight of spin and charge excitations involved in d-wave scattering. This function extends over a wide frequency region of about $2|t|$. Its high-frequency part is typical for collective charge fluctuations. It also contains substantial spectral weight at
lower frequencies exhibiting a rather linear increase in frequency at the low-frequency end of the spectrum. Fig. 4 shows the corresponding real part of $\Theta_3^{ret}$. It is weakly negative at low frequencies up to about $\omega_1 \sim J/2$, changes then from negative to positive values until about $\omega \sim \omega_2$. The retarded interaction between electrons is thus attractive for $0 \leq \omega \leq \omega_1$ and strongly repulsive for $\omega_1 \leq \omega \leq \omega_2$. Somewhat above $\omega_2$ the real part of $\Theta_3^{ret}$ jumps to large negative values and approaches zero from below in the high-frequency limit. The inset of Fig. 4 shows the real part of $\Theta_3$ at small frequencies with a larger resolution using $\eta = 0.002$.

C. Transition temperatures

Figs. 1a) and b) show that the instantaneous and the retarded contributions to $\Theta$ are of similar magnitude and often compete with each other. In the calculation of $T_c$ they are associated with quite different cut-offs. The instantaneous part is characterized by a cut-off determined by the width of the effective band whereas the cut-off relevant for the retarded part is set by the frequency range where attraction dominates characterized roughly by $J$. In view of these complications we developed a method to solve Eq.(40) directly, avoiding the use of pseudopotentials. The only simplification we use is to put the momenta of the retarded kernel on the Fermi line. The validity of this approximation has been checked numerically and holds very well in our case. Using the fact that the instantaneous kernel consists only of a few separable contributions Eq.(40) can be reduced to a linear matrix problem. The number of rows and columns are given by the number of considered Matsubara frequencies times the number of $k$-points on the Fermi line in the irreducible Brillouin zone. Numerical tests showed that well converged results can be obtained with about 300 Matsubara frequencies and 5 $k$-points along 1/8 of the Fermi line down to values for $T_c$ of $\sim 0.002$. Details of our method will be given elsewhere.

The squares in Fig. 5 (joined by a solid line) show $T_c$ for $\Gamma_3$-symmetry for doping $\delta > \delta_{BO}$. The broken line exhibits $T_c$ if only the instantaneous part of the kernel is used. The dotted line corresponds to $T_c$ if the charge-charge contribution (last term in Eq.(1)) is dropped. Finally the dashed-dotted line describes $T_c$ if only the retarded kernel is taken into account.

In order to understand the curves in Fig. 5 we consider the real part of the retarded kernel as a function of frequency, as shown in Fig. 4. Taking only the part between zero and $\omega_2$ into account would yield rather large values for $T_c$. To realize this one can decompose the real part into a large, constant repulsive part between $\omega = 0$ and $\omega = \omega_2$ plus the difference which is non-zero and attractive at low frequencies. Changing the cut-off from $\omega_2$ to $\omega_1$ using a pseudopotential description decreases strongly the effective repulsion yielding a large net attraction between 0 and $\omega_1$ and thus a high value for $T_c$. Model calculation show that the large negative part in the real part of $\Theta_3$ above $\omega_2$ is very harmful to superconductivity: Lowering first the cut-off from $\sim 3|t|$ to $\omega_2$ increases the effective potential, reducing further the cut-off to $\omega_1$ decreases again the effective potential. The overall result is a rather modest attraction between $\omega = 0$ and $\omega = \omega_1$. This explains the rather low values for $T_c$ calculated from the retarded part of $\Theta$ alone, as shown by the dash-dotted line in Fig. 5. An alternative consideration would start from the instantaneous part in $\Theta$ leading to the transition temperatures shown by the broken line in Fig. 5. Using a realistic value for the energy unit of about 8000K (note that the usual effective hopping $t_{\text{exp}}$ corresponds
according to Eq.(1) to \(|t|/2\) these values are already of the order of 100\(K\) and comparable to the experimental ones. Adding the retarded part of the kernel does not lead to a further increase in \(T_c\) but rather to a slight decrease as illustrated by the squares in Fig. 5. Model calculations show that a purely attractive \(\Theta^{ret}\) would always increase \(T_c\). The observed lowering of \(T_c\) must therefore be due to the repulsive part in \(\Theta^{ret}\) above \(\omega_1\). This part plays a role because the \(T_c\) due to the instantaneous part alone is large enough to couple strongly the frequencies above and below \(\omega_1\) in the gap equation. The doping dependence of \(T_c\) is determined by various processes which compete with each other. For instance, the density of states decreases with doping which decreases the effective coupling. On the other hand, the energy scale for \(T_c\) is in the case of the instantaneous term set by the effective band width which increases with doping. Fig. 5 shows that the net effect of these and other processes is to lower \(T_c\) substantially with increasing doping for \(\delta > \delta_{BO}\). The eigenvalue \(\lambda_3\) of the static kernel assumes according to Fig. 2 large negative values near \(\delta_{BO}\) anticipating the incipient bond-order wave instability. The associated soft mode does not affect the instantaneous but the retarded part. Fig. 5 shows that the dashed line indeed increases steeply approaching \(\delta_{BO}\) from above. On the other hand little effects are seen in the squares representing the full calculation. The reason for this is that at the low \(T_c\)'s of the dashed curve the relevant frequencies lie in the attractive region of \(\Theta^{ret}_3\) whereas for the high \(T_c\) values of the squares these frequencies lie already in the repulsive part. From this one may conclude that neither soft modes nor large negative eigenvalues of the static kernel guarantee large \(T_c\) values. In particular, there is no simple connection between \(T_c\) and \(\lambda_3\).

The squares (connected by a dashed line) in Fig. 6 show \(T_c\) as a function of the exchange constant \(J\) for a doping \(\delta = 0.17\). \(T_c\) approaches 0 for \(J \to 0\) in agreement with previous findings\(\textsuperscript{11}\). Except at small values of \(J\) \(T_c\) depends linearly on \(J\). This is quite in contrast to the BCS-formula with an exponential dependence on \(J\). The main reason for this is that the instantaneous term plays a major role and the momenta in it cannot confined to the Fermi surface in calculating \(T_c\). Eq.(53) is then a more appropriate formula for \(T_c\). It is, however, too simple to argue that \(b_3\) in that formula is rather independent on \(J\) and \(\sim 1/T\) for \(T \geq 0.01\). The rather perfect linear behavior is the result of more subtle dependencies such as the \(J\)-dependence of the one-particle energies and the non-rigidity of the bands as a function of \(J\).

It has been argued\(\textsuperscript{22}\) that the nearest-neighbor Coulomb potential \(V\) is not neglegible in the cuprates. Adding this term

\[
H' = \frac{V}{2N} \sum_{p,q=1}^{N} X_{pq}^I X_{pq}^q, \tag{63}
\]

to the Hamiltonian Eq.(1) we have calculated \(T_c\) as a function of \(V\). The circles (connected by a solid line) in Fig. 6 present the result for \(J = 0.3\) and \(\delta = 0.17\). \(T_c\) decreases strongly with increasing \(V\) and is extremely small for \(V \geq 3J/4\). This may be understood by looking just at the instantaneous term: Eq.(63) yields an additional term \(2V(k-k')\) on the right-hand side of Eq.(43), canceling the \(J\)-terms exactly for \(V = J\). \(T_c\) would thus vanish if only the instantaneous term would be present. For \(V = J/2\) \(H'\) cancels the charge term in Eq.(1). According to Fig. 6 this means a drop of \(T_c\) by about a factor 5 compared to the value at \(V = 0\) which also agrees with Fig. 5. Such a big drop of \(T_c\) due to the charge term of the \(t-J\) model seems very surprising because the latter can only produce effects \(\sim \delta^2\)
and is therefore often omitted. Our calculation shows that this term cannot be neglected in a calculation of \( T_c \) and increases \( T_c \) substantially.

Fig. 7 shows \( T_c \) as a function of \( \delta \) for \( J = 0.3 \) and \( V = 0.15 \). Due to the Coulomb repulsion the \( T_c \) values are rather low. The important frequencies in the solution of the gap equation are also low and mainly located in the attractive region of the retarded kernel, Adding the retarded to the instantaneous part thus increases \( T_c \). The two curves in Fig. 7 demonstrate this effect.

Figs. 8 and 9 show the influence of a second-nearest neighbor hopping term \( t' \) on \( T_c \). In these figures a lower value of 0.2 has been chosen for \( J \) similar as in Ref. 9 in order to have a reasonably low \( \delta_{BO} \) and a Van Hove singularity not too far away from optimal doping. Assuming a linear dependence of \( T_c \) on \( J \) the absolute values for \( T_c \) are similar in the corresponding Figs. 5 and 8. \( T_c \) is thus rather robust to changes in \( t' \). \( T_c \) decreases in Fig. 8 somewhat slower than in Fig. 5 due to the larger density of states in the surroundings of the Van Hove singularity. The difference between squares and circles is also larger in Fig. 8 and decreases much slower with increasing doping. These effects are caused by the retarded part of the kernel which in Fig. 8 is less attractive below \( \omega_1 \) and more repulsive above \( \omega_1 \) as shown in Fig. 2 of Ref. 9 (Note that the curve in that Figure includes a constant instantaneous contribution of -0.29). The Van Hove singularity is located near \( \delta = 0.28 \). There is nearly no effect of the Van Hove singularity on \( T_c \) for the same reasons as in the above case of the incipient bond-order wave instability. (The curves in Fig. 3 of Ref. 9 were calculated without the charge-charge term and, erroneously, without the factor 1/2 in the instantaneous part. Correcting this error amounts essentially to lower the curves in that Figure by about a factor 4). Fig. 9 should be compared with Fig. 6. \( T_c \) decreases in both cases strongly with increasing \( V \) and practically vanishes for \( V \geq J \).

We have also searched for superconducting instabilities with order parameters which are odd in frequency. The sum over Matsubara frequencies is always zero in this case so that the constraint of having no double occupancies of sites at the same time is automatically fulfilled for the two particles of the Cooper pair. There exists no static approximation for the kernel in this case. Possible instabilities are again determined by the linearized gap equation Eq.(40) where \( \Sigma_{an} \) and \( \Theta \) have to be projected on the odd frequency parts. The transition temperature is determined by the condition that the determinant of a matrix consisting essentially of the kernel and the unity matrix is zero. In Fig. 10 we have plotted this determinant as function of the temperature for each of the five irreducible representations using \( J = 0.3, t' = 0.1, \) and \( \delta = 0.17 \). The Figure clearly shows that none of the curves tends to zero in the investigated temperature interval ruling out any odd frequency pairing instability with a \( T_c \) larger than \( \sim 0.002 \). This also can be seen directly from Fig. 11 where the real part of \( \Theta_3(\omega + i\eta) \) is shown for odd frequency pairing with \( \Gamma_3 \)-symmetry corresponding also to triplet pairing. The effective interaction is repulsive up to energies \( \sim t \) ruling out an instability towards superconductivity in this channel.

**D. Conclusions**

Sections II and III demonstrate the feasibility of developing a perturbation expansion for the \( t - J \) model in terms of X-operators and obtaining explicit expressions for the leading contributions to the anomalous self-energy of physical electrons using an \( 1/N \) expansion. The
nonapplicability of Wick’s theorem to X-operators does not cause any problem in such an approach. One also should note that the more involved and sophisticated parts of sections II and III deal with contributions which are far beyond those considered in previous treatments. For instance, the slave boson 1/N result for the anomalous self-energy corresponds in our approach just to the inhomogenous term in the integral equation Eq.(32). Similarly, only part of the first contribution to the kernel, \( \Theta^{(1)} \), was calculated in Ref.\[7\] using a diagram technique for X-operators. This shows in our opinion that the employed functional approach is at least as suitable as other approaches to treat highly correlated Fermi systems. Taking also the numerical results of section 3 into account our main conclusions can be summarized as follows:

a) Our explicit expression for the O(1/N) anomalous self-energy is clearly different from the corresponding expression of the slave boson theory. In particular, the largest contribution to the retarded kernel comes in the present approach from the anomalous part of the vertex and has no analogue in the slave boson approach. The presented expressions show for the first time in an explicit way that the 1/N expansions are really different in the two approaches. This difference can be traced back to different Hilbert spaces and different enforcement of the constraint. Even the order parameters for superconductivity are not related in a simple way: The leading slave boson order parameter involves necessarily (small) violations of local constraints in order to be nonzero whereas such violations are ruled out in our approach.

b) The kernel of the linearized gap equation consists of an instantaneous and an retarded part and both are similar in magnitude at low frequencies and for momenta near the Fermi surface. We found that there are superconducting instabilities in each symmetry channel and for all dopings. The true ground state thus never describes a Fermi liquid but a superconductor similar as in the weak coupling case\[23\]. However, these instabilities are in general extremely weak leading to academically low transition temperatures. The only clear and robust exception is the d-wave-like \( \Gamma_3 \) symmetry where a strong instability towards superconductivity occurs. Odd symmetry pairing mechanisms turned out to be very weak and can be ruled out as a mechanism for high-\( T_c \) superconductivity in our model.

c) In the case of \( \Gamma_3 \) symmetry the real part of the retarded kernel is weakly attractive at low frequencies on an energy scale of \( J \) or a fraction thereof and strongly repulsive at higher frequencies whereas the instantaneous part is attractive. Solving numerically the linearized gap equation the obtained transition temperatures \( T_c \) are for \( N = 2 \) of the order of 0.01|\( t \)| and thus in principle large enough to account for the phenomena of high-\( T_c \) superconductivity. It is interesting to note that the Hubbard model at small or intermediate couplings also shows d-wave superconductivity with similar values for \( T_c \).\[24\] The instantaneous term is instrumental in getting these large values for \( T_c \): First, its cutoff is given by the effective band width and thus in general larger than \( J \). Secondly, due to the large \( T_c \), the solution of the gap equation involves large frequencies where the retarded term is strongly repulsive. As a result the retarded term is of less importance because attractive and repulsive parts of it cancel each other to a large extent. The dominance of the instantaneous part and the presence of two cutoffs lead to strong deviations from BCS-behavior. For instance, \( T_c \) depends linearly and not exponentially on \( J \) except at very small values for \( J \).

d) \( T_c \) is rather insensitive to the addition of a second-nearest neighbor hopping term \( t' \) and to a Van Hove singularity. The latter can be understood by noting that the solution of the gap equation involves momentum and frequency averages in the instantaneous and
retarded part, respectively, so that singularities in the density of states are washed out. \( T_c \) depends, however, sensitively on a nearest-neighbor Coulomb repulsion \( V \) and becomes very small if \( V \) is substantially larger than \( J \).

e) Our calculations are limited to dopings larger than \( \delta_{BO} \sim 0.14 \) for \( J = 0.3 \) where an instability towards an incommensurate bond-order wave of d-symmetry occurs. The associated soft mode causes \( \lambda_3 \rightarrow -\infty \) for \( \delta \rightarrow \delta_{BO} \) whereas \( T_c \) is nearly unaffected. In the underdoped regime \( \delta < \delta_{BO} \) we expect a competition between bond-order and antiferromagnetic fluctuations which is beyond the leading order of the \( 1/N \) expansion considered in this investigation.

**Acknowledgement:** The second author (A. Greco) thanks the International Bureau of the Federal Ministry for Education, Science, Research and Technology for financial support (Scientific-technological cooperation between Argentina and Germany, Project No. ARG AD 3P) and the MPI-FKF for hospitality. Both authors acknowledge useful discussions with P. Horsch.
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FIGURES

FIG. 1. Dependence of various contributions to the total kernel $\Theta$ on the second momentum $k'$ along the Fermi line for a fixed first momentum $k = (2.465, 0.309)$; a) contributions $\Theta^{(1)}, \Theta^{(2)}$; b) contributions $\Theta^{(3)}, \Theta^{(4)}$, and the total $\Theta$.

FIG. 2. Lowest eigenvalues $\lambda_i$ of the static kernel $\Theta$ for the five representations $\Gamma_i$ of $C_{4v}$ as a function of the doping $\delta$.

FIG. 3. Negative imaginary part of the d-wave projected kernel $\Theta_3(\omega + i\eta)$ as a function of the frequency $\omega$ using $\eta = 0.005$.

FIG. 4. Real part of the d-wave projected kernel $\Theta_3(\omega + i\eta)$ as a function of the frequency $\omega$ for $\eta = 0.005$. Inset: The same for small frequencies using $\eta = 0.002$.

FIG. 5. Transition temperature $T_c$ (with $|t|$ as energy unit) for $\Gamma_3$ pairing using the total kernel (squares), the instantaneous part (circles), the retarded part (dimaonds) of the kernel, and the total kernel without charge-charge term (triangles).

FIG. 6. Transition temperature $T_c$ for $\Gamma_3$ pairing versus $V$ (circles) and versus $J$ (squares).

FIG. 7. Transition temperature $T_c$ for $\Gamma_3$ pairing for $V = 0.15$ using the total kernel (squares) and the instantaneous part of the kernel (circles).

FIG. 8. Transition temperature $T_c$ for $\Gamma_3$ pairing for $J = 0.2$ and $t' = -0.35$ using the total kernel (squares) and the instantaneous part of the kernel (circles).

FIG. 9. Transition temperature $T_c$ for $J = 0.2$, $t' = -0.35$, $\delta = 0.23$ as a function of $V$ using the total kernel (squares) and the instantaneous part of the kernel (circles).

FIG. 10. Determinant associated with the gap equation for odd frequency pairing with symmetry $\Gamma_i$ as a function of temperature $T$.

FIG. 11. Real part of the odd frequency, $\Gamma_3$ kernel $\Theta_3(\omega + i\eta)$ as a function of the frequency $\omega$ for $\eta = 0.005$. 

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a)

$J = 0.3$

$\delta = 0.17$

momentum $k'$
b) $J = 0.3$

$\delta = 0.17$
The graph shows the lowest eigenvector $\lambda_i$ as a function of doping $\delta$ for different values of $J=0.3$. The eigenvectors are labeled $\Gamma_1$, $\Gamma_2$, $\Gamma_3$, $\Gamma_4$, and $\Gamma_5$. The graph indicates a progressive decrease in the eigenvector value as the doping increases.
$J = 0.3$

$\delta = 0.17$
J = 0.3
δ = 0.17

real part of \( \Theta_3(\omega + i \eta) \)

frequency \( \omega \)
$J=0.3$

d–wave ($\Gamma_3$) symmetry
The transition temperature $T_c$ is plotted against the coupling constants $V$ and $J$. The figure shows two distinct lines:

- A solid line representing $T_c$ vs $V$.
- A dashed line representing $T_c$ vs $J$.

Key values:

- $J = 0.3$
- $\delta = 0.17$
$J = 0.3$

$V = 0.15$

d–wave ($\Gamma_3$) symmetry
$J = 0.2$
$t' = -0.35$

d-wave ($\Gamma_3$) symmetry

transition temp. $T_c$ vs. doping $\delta$
\[ J = 0.2 \]
\[ t' = -0.35 \]
\[ \delta = 0.23 \]
\[ d\text{-wave } (\Gamma_3)\text{symmetry} \]
The diagram shows the determinant for odd frequency pairing as a function of temperature $T$. The parameters $J=0.3$ and $\delta=0.17$ are indicated. The lines represent different odd frequency parings, labeled as $\Gamma_1$, $\Gamma_2$, $\Gamma_3$, $\Gamma_4$, and $\Gamma_5$. The x-axis represents temperature $T$ ranging from 0 to 0.02, and the y-axis represents the determinant value.
$\Gamma_3$–odd frequency pairing

real part of $\Theta_3(\omega+i\eta)$

$J=0.3$

$\delta=0.17$