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Finite-temperature superconducting correlations of the Hubbard model

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We utilize numerical linked-cluster expansions (NLCEs) and the determinantal quantum Monte Carlo algorithm to study pairing correlations in the square lattice Hubbard model. To benchmark the NLCE, we first locate the finite-temperature phase transition of the attractive model to a superconducting state away from half filling. We then explore the superconducting properties of the repulsive model for the d-wave and extended s-wave pairing symmetries. The pairing structure factor shows a strong tendency to d-wave pairing and peaks at an interaction strength comparable to the bandwidth. The extended s-wave structure factor and correlation length are larger at higher temperatures but clearly saturate as temperature is lowered, whereas the d-wave counterparts, which start off lower at high temperatures, continue to rise near half filling. This rise is even more dramatic in the d-wave susceptibility. The convergence of NLCEs breaks down as the susceptibilities and correlation lengths become large, so we are unable to determine the onset of long-range order. However, our results extend the conclusion, previously restricted to only magnetic and charge correlations, that NLCEs offer unique window into pairing in the Hubbard model at strong coupling.

I. INTRODUCTION

Despite several decades of intensive theoretical research, the question of whether a non-local attraction can dominate in a fermionic Hubbard model with local repulsive interaction has remained largely unanswered for parameters relevant to cuprate high-temperature superconductors.1–6 Controlled theoretical approaches confirm this possibility, however, only when the strength of the local repulsion is much smaller than the hopping amplitude of fermions on a square lattice.7

Numerical methods provide important data for strongly-correlated quantum Hamiltonians, and, in particular, for phenomena like superconductivity, magnetism, and Mott metal-insulator transitions. Although many developments have made these approaches increasingly powerful over the last decade, significant limitations remain, especially for fermions. The density matrix renormalization group8,9 and related techniques, function best in one dimension. Diagrammatic quantum Monte Carlo techniques10,11 are restricted to weak-coupling regimes. Determinant quantum Monte Carlo (DQMC)12,13 and cluster extensions of the dynamic mean-field theory14,15 are limited to real space or momentum space clusters of tens to hundreds of sites. Moreover, the “sign problem”16,17 remains an unsolved problem which limits accessible temperatures unless special symmetries prevail.

These limitations emphasize the need for continued algorithm development. Recently developed numerical linked-cluster expansions (NLCEs)18,23 are especially promising as an approach to access strong coupling regimes, which are inaccessible to QMC methods, as a consequence both of the sign problem and of large and even diverging statistical fluctuations. For instance, analysis of magnetic correlations and Mott phases in trapped atoms on optical lattices22,23 where strong coupling is present at the cloud edge, would not have been possible without NLCEs.

A natural next step is the application of NLCEs to superconductivity. In this paper, we show this method can be developed and successfully used to study the pairing correlations in the square lattice Hubbard model,

$$H = -t \sum_{\langle ij \rangle \sigma} c_{i \sigma}^\dagger c_{j \sigma} + U \sum_i n_{i \uparrow} n_{i \downarrow} - \mu \sum_{i \sigma} n_{i \sigma},$$  

(1)

where $c_{i \sigma} \left( c_{i \sigma}^\dagger \right)$ annihilates (creates) a fermion with spin $\sigma$ on site $i$, $n_{i \sigma} = c_{i \sigma}^\dagger c_{i \sigma}$ is the number operator, $U$ is the onsite Coulomb interaction, and $t$ is the nearest neighbor hopping integral. We set $k_B = 1$, and $t = 1$ as the unit of energy throughout the paper.

We complement our NLCE results with those obtained from (numerically unbiased) DQMC simulations on a large lattice. We find excellent agreement between the two in parameter regions accessible to both, and show that the lowest temperatures achievable in the NLCE are similar to, or often lower than, those of the DQMC. For the attractive model ($U < 0$), or in the weak-coupling regime of the repulsive model, where the sign problem is either absent or less severe, DQMC can generally access lower temperatures than the NLCE. On the other hand, the series converges to lower temperatures in the strong-coupling regime, where DQMC runs into sampling difficulties and faces an unforgiving sign problem.

We find that, for an interaction strength $U$ equal to the bandwidth, the s-wave pairing structure factor of the attractive model away from half filling shows divergent behavior at low temperatures, and points to a finite transition temperature that is consistent with findings of previous large-scale DQMC studies24,25. For the repulsive model, we consider several values of $U$ and doping and study pairing in the nonlocal channels of extended s-wave ($s^\ast$-wave) and d-wave. While the structure factor for the former symmetry tends to saturate at increasingly high temperatures as the doping is increased, for the latter symmetry, no such tendency is observed. We examine results at 10% doping more
closely and find that the low-temperature structure factor is maximum around \( U = 8 \). On the other hand, the pair-field susceptibility, while larger for smaller values of \( U \) in the intermediate temperature region, shows a sharp upturn at the lowest accessible temperatures for the largest interactions considered.

**II. NUMERICAL METHODS**

In NLCEs, an extensive property of the lattice model, when normalized to the number of sites, is expressed in the thermodynamic limit in terms of contributions from finite clusters of various sizes and topologies that can be embedded in the lattice. Thus, NLCEs use the same basis as the high-temperature expansions (HTEs). However, the calculation of the extensive quantities at the level of individual clusters is left to an exact numerical method, such as exact diagonalization, as opposed to a perturbative expansion in terms of inverse temperature in the HTEs. A typical expansion involves clusters up to a certain size that are chosen according to a self-consistent criterion. Despite the lack of an explicit small parameter, having a finite number of clusters in the series inevitably leads to the loss of convergence below a certain temperature, where the correlations in the system extend beyond a length of the order of the largest sizes considered. However, the exact treatment of clusters leads to convergence temperatures that are lower than those of HTE with a comparable number of orders.

Similar to the analytic Padé approximations used extensively in HTEs, here we take advantage of two numerical resummation techniques to improve the convergence of our series at low temperatures. We use the Euler algorithm\(^{29}\) to resum the last 4-6 terms of the series or the Wynn algorithm\(^{29}\) with 3 and 4 cycles of improvement (details of these techniques can be found in Ref. [19]). We then take the average of values from the last two orders after the Euler, and the last two orders after the Wynn transformations as our best estimate. To quantify our confidence in the accuracy of the resummed results, we define a “confidence region” around this average where all the values that contribute to the average fall. Thus, the errorbars in our figures simply mark the boundaries of this region and should not be confused with statistical errorbars.

We study the superconducting properties of the model at several values of the interaction strength and on a fine grid of temperature and chemical potential. The latter allows us to study the calculated quantities at constant electronic densities after numerical conversion. As with previous studies of Hubbard models using the NLCE,\(^{31-33}\) we employ the site expansion in which the order to which each cluster belongs is determined by the number of sites it has. In order \( l \), we consider all the open boundary clusters of various shapes and topologies on the square lattice that have \( l \) sites, and use exact diagonalization to solve for their properties.

For the pairing correlations, the Hamiltonian matrices are block-diagonalized in each particle number sector. So, we are able to carry out the expansion to the ninth order. For the pairing susceptibility, on the other hand, we can only carry out the expansion to the seventh order since not only particle number is not conserved during the time-dependent measurements (see Eq. [2]), but also the majority of the computational time is spent on obtaining the off-diagonal expectation values, which, like the diagonalization, scales like \( O(N^3) \).

DQMC simulations are performed on a \( 10 \times 10 \) lattice, which is large enough to have only small finite size effects at the temperatures studied here. Results represent averages of at least 8 independent runs with 10,000 sweeps each. To fix the density, \( n \), away from half filling at each temperature and \( U \) value, the chemical potential needs to be tuned starting from an estimate provided by the NLCE. Therefore, we repeat the calculations for several values of \( \mu \) to achieve an accuracy of about 0.01% for the density. For the structure factor, we extrapolate our results to the continuous imaginary time limit using the outcome of two separate simulations with a discretization of the inverse temperature \( \beta = L \Delta \tau \) corresponding to \( \Delta \tau = 1/16 \) and \( 1/12 \). In the case of the susceptibility, we choose an even smaller \( \Delta \tau = 1/50 \), in order to perform the imaginary time integration accurately. This value leads to Trotter errors that are negligible in comparison to the statistical ones.

One of the quantities we calculate is the equal-time pairing structure factor,

\[
S^\alpha(q) = \sum_r e^{iqr} P^\alpha(r),
\]

where

\[
P^\alpha(r_{ij}) = (\Delta^\alpha_{ij}(0) + \Delta^\alpha_r(0)\Delta^\alpha_{r+}(0))
\]

is the equal-time pair-pair correlation function. Here, the pairing operator for the symmetry \( \alpha \) is defined as

\[
\Delta^\alpha_r(\tau) = \frac{1}{2} \sum_j f^\alpha_{ij} e^{rH}(c_{\alpha i}c_{\alpha j} - c_{\alpha ji}^\dagger c_{\alpha j}^\dagger) e^{-\tau H}.
\]

We consider three pairing symmetries in this study; (local) \( s \)-wave, \( d \)-wave, and \( s^* \)-wave. For the \( s \)-wave symmetry, \( f^\alpha_{ij} = \delta_{ij} \). In the case of \( s^* \)-wave, \( f^\alpha_{ij} \) is +1 if \( i \) and \( j \) are nearest neighbors and \( j > i \) (to avoid double counting) and zero otherwise. \( f^\alpha_{ij} \) for the \( d \)-wave symmetry is the same as \( f^\alpha_{ij} \) except it takes the value \( -1 \) if the bond connecting \( i \) and \( j \) is along the \( y \) axis. Here, we consider only the uniform pairing structure factor, \( S^\alpha(0) \). The correlated structure factors, \( S^\alpha_{\text{corr}} \), is obtained by first subtracting off the uncorrelated parts of the expressions in Eq. [3].

Having the uniform structure factor, the correlation length, \( \xi \), can also be calculated using

\[
(c^\alpha_{\text{corr}})^2 = \frac{1}{2dS^\alpha_{\text{corr}}(q = 0)} \sum_i |r_i|^2 P^\alpha_{\text{corr}}(r_n),
\]
where \( d = 2 \) is the dimension.

The other quantity of interest for superconductivity is the uniform pairing susceptibility, which is defined as

\[
\chi^\alpha = \frac{1}{N} \int_0^\beta d\tau \langle O^\alpha(\tau)O^{\alpha\dagger}(0) \rangle,
\]

where \( O^\alpha = \sum_i \Delta_i^\alpha(\tau) \).

### III. RESULTS

We start with the attractive Hubbard model, for which we know there exists a finite-temperature Kosterlitz-Thouless (KT) phase transition to an \( s \)-wave superconducting state away from half filling.\textsuperscript{24,28} In Fig. 1(a), we show the correlated part of the \( s \)-wave pairing structure factor from the NLCE for \( U = -8 \) and at \( n = 0.85 \), where the superconducting transition temperature is expected to be maximal.\textsuperscript{28} Results are in excellent agreement with the corresponding DQMC results, plotted as empty circles in that figure. As can be inferred from previous DQMC simulations with a smaller \( U \), finite-size effects in DQMC will not play a role here at temperatures as low as \( T = 0.25 \). Whereas the raw NLCE results (before resummations) converge only to \( T \sim 0.11 \), which is in good agreement with results of past DQMC simulations.\textsuperscript{24} The correlation length, which shows an exponential growth, is also plotted in Fig. 1(b). Its behavior is consistent with the trend seen in Fig. 1(c) for the pairing correlations growing faster at longer length scales as the temperature is decreased.

We now turn our focus to the main subject of this study; pairing in the repulsive Hubbard model. We know that if a similar finite-temperature transition to a superconducting phase takes place in the latter model, the pairing symmetry has to be nonlocal because of the onsite Coulomb repulsion. Therefore, in this case, we only explore the \( d \)-wave and the \( s^* \)-wave symmetries. We also expect the corresponding temperature scales to be much smaller than those for the attractive model since we are looking for attraction in a repulsive model.

In Fig. 2 we show the correlated part of the uniform structure factor for the two pairing symmetries when \( U = 8 \) and at various average densities. At half filling, the series converges to a low enough temperature to make clear that \( S_{\text{corr}} \) eventually saturates as we decrease the temperature. In the absence of the ‘sign problem’ at this filling, DQMC can easily access lower temperatures. We see in Fig. 2(a) that, while agreeing excellently with NLCE at high temperatures, results from DQMC simulations confirm the saturation at lower \( T \). As we move away from half filling into the hole-doped region \(( n < 1.0 )\), an interesting trend is observed; the saturation of the \( s^* \)-wave structure factor is seen to take place at higher temperatures whereas the \( d \)-wave structure factor continues to grow at the lowest temperatures accessible to us, although its over all values decrease as we increase the doping. Hence, if there is an instability to pairing away from half filling in this model, it would be in the
shows the inverse of the structure factor $S$ that can explain the higher convergence temperature of the $s$-wave pairing of the corresponding strong-coupling regime, the only relevant energy scale $|t| \ll U = 8$, and then slowly decreases. Beyond $U = 8$, we expect this quantity to scale as $1/U$ as, in the strong-coupling regime, the only relevant energy scale will be the exchange interaction of the corresponding low-energy $t = J$ model with $J = 4t^2/U$. The bottom inset in Fig. 3 shows the inverse of the structure factor at $U = 8$. Unfortunately, we are not close enough to a transition temperature to be able to make any quantitative statement about its value. However, the best estimate from the DCA for a close value of the interaction ($U = 7$), puts $T_c$ around 0.05, which is consistent with a KT fit to our results for $T < 0.6$.

Finally, we turn to the pair-field susceptibility. Figure 4 shows the $\chi^d$ vs temperature at $n = 0.9$ for different interaction strengths. Our results for the susceptibilities match the DQMC ones very well for smaller $U$ values and for larger $U$ values when the temperature is not too low. This includes the susceptibility at $U = 4$ and $n = 0.875$, which is consistent with a KT fit to our results for $T < 0.6$.

In summary, we have employed two unbiased methods, the NLCE and the DQMC to study finite-temperature superconducting properties of the square lattice Fermi-Hubbard model. To benchmark our NLCE approach, we first explore the $s$-wave pairing in the attractive model away from half filling. By fitting our low-temperature pairing structure factor to known forms, we obtain a $T_c$ that is consistent with the best estimate from large-scale QMC simulations. We then investigate the nonlocal $s^*$-wave and $d$-wave pairing instabilities in the repulsive model at various dopings and for several interaction strengths. We find that the $d$-wave symmetry has the tendency to be dominant at low temperatures and that its structure factor has a maximum at $U \sim 8$. We also calculate the pairing susceptibility, which shows a similar divergent behavior in the $d$-wave channel and a sharp upturn at low temperatures for large interactions.

An important potential application of the results

**FIG. 3.** Temperature dependence of the $d$-wave pairing structure factor at $n = 0.90$ for $U = 4$, 6, 8, and 12. Symbols are the DQMC results. Top inset shows the same structure factor vs $U$ at a fixed temperature $T = 0.43$. The bottom inset shows the inverse of the structure factor vs $T$, along with a fit to the function $A \exp(B/\sqrt{T - T_c})$ for $T < 0.6$, which results in $T_c = 0.048$.

**FIG. 4.** The $d$-wave pairing susceptibility at $n = 0.90$ vs temperature for $U = 4$, 6, 8, and 12. Bare NLCE results before resummations for the last two orders, 6th and 7th, are shown as thin dotted and dashed lines, respectively. Symbols are the DQMC results. The inset shows the inverse of the susceptibility vs temperature for the same values of the interaction strength.
described here is to ongoing emulation of model Hamiltonians which describe fermionic atoms in optical lattices. NLCEs allow the rapid evaluation of physical properties on a dense mesh of Hamiltonian parameters, a requirement for accurate modeling of optical lattice experiments, where the confining potential leads to spatially varying chemical potential, interaction strength, and hopping matrix elements. Here we have shown the potential importance of NLCEs as a tool to analyze pairing in these systems.

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34. In the ninth order alone, we need to distinguish all 1285 clusters that may have the same topology (the same Hamiltonian matrix), but are not related by point group symmetry. The largest matrices to be diagonalized have a linear size of $N = 15876$. In the seventh order, there are 108 symmetrically distinct clusters. The largest matrices to be diagonalized have a linear size of $N = 3432$.
35. For example, for the s-wave channel, we have $P_{\alpha \beta}^{\text{corr}}(r_{ij}) = P_{\alpha \beta}(r_{ij}) - 2\langle c_{\alpha \sigma} c_{\beta \sigma} \rangle^2 - \delta_{ij} \left[ 1 - 2\langle c_{\alpha \sigma} c_{\beta \sigma} \rangle \right]$.
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