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The Mott-Anderson transition in the disordered one-dimensional Hubbard model

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Abstract

We use the density matrix renormalization group to study the quantum transitions that occur in the half-filled one-dimensional fermionic Hubbard model with onsite potential disorder. We find a transition from the gapped Mott phase with algebraic spin correlations to a gapless spin-disordered phase beyond a critical strength of the disorder $\Delta_c \approx U/2$. Both the transitions in the charge and spin sectors are shown to be coincident. We also establish the finite-size corrections to the charge gap and the spin-spin correlation length in the presence of disorder and using a finite-size-scaling analysis we obtain the zero temperature phase diagram of the various quantum phase transitions that occur in the disorder-interaction plane.

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behavior as a function of either the interaction strength or the degree of disorder or both [1]. A clean system at certain commensurate fillings may develop a gap in the energy spectrum as the strength of the repulsion is increased and turn into a Mott insulator [2]. On the other hand, a system of noninteracting electrons can undergo a transition from metal to insulator as the degree of randomness is increased and turn into an Anderson localized insulator [3]. The interplay of electron-electron interaction and disorder raises interesting possibilities of a new type of transition, distinct from the clean correlation induced Mott transition or the disorder induced Anderson transition.

The repulsive Hubbard model in one dimension (1D) is probably the simplest model which shows a Mott transition at halffilling for arbitrary values of the interaction strength U > 0 [4]. One of the most attractive features of this model is that the Mott transition is unaccompanied by a spindensity-wave gap as e.q. happens in the 2D netic long-range order (MLRO) [5]. Instead sible fixed points in parameter space. Com-

Electronic systems can undergo quantum the 1D model shows algebraically decaying phase transitions from metallic to insulating spin correlations as the maximal remnant of MLRO in 1D. The properties of the clean 1D Hubbard model are well established: The dependence of the Mott gap on the interaction parameter [4], the asymptotic behavior and the critical exponents of various correlation functions [6] have all been computed using Bethe Ansatz and the finite-size-scaling approach of conformal quantum field theory [7]. All this makes this model particularly attractive to study the effect of disorder on the Mott state.

Recent studies of the half-filled disordered 1D Hubbard model have proceeded numerically using the Quantum Monte Carlo (QMC) method [8] and analytically using bosonization and the renormalization group method [9]. The QMC results give a very accurate description of the finite temperature properties of the system. However, the low temperature properties can only be inferred by extrapolating the finite temperature data and going to larger system sizes. On the other hand, bosonization methods using perturbative renormalization group techniques Hubbard model due to the existence of mag- only give an indication of the various plaucoupling fixed points is then impossible as the coupling constants are driven away from indicates the emergence of a spin-disordered their weak coupling values. Nevertheless, the results of both approaches indicate that at half-filling a *finite* amount of potential disorder is needed to cause a transition from Mott (gapped) to Anderson (gapless) insulating behavior. This is in qualitative agreement with arguments put forward by Ma [10].

produced numerically reliably. We then con-

puting correlation functions at the strong power-law remnant of the MLRO persists, whereas for $\Delta > \Delta_c$, the spin-spin correlation phase.

> The Hubbard Hamiltonian with additional potential disorder on a chain of L sites is given as

$$H = -t \sum_{\substack{x=1\\\sigma=\uparrow,\downarrow}}^{L} (c_{x+1\sigma}^{\dagger} c_{x\sigma} + h.c.) + U \sum_{x=1}^{L} n_{x\uparrow} n_{x\downarrow} + \sum_{x=1}^{L} \mu_{x} n_{x\downarrow}$$
(1)

In the present Letter, we have studied where -t is the hopping amplitude between the ground state properties of the disordered nearest-neighbor, $c^{\dagger}_{x\sigma}(c_{x\sigma})$ the fermion cre-1D Hubbard model at zero temperature with ation (annihilation) operator at site x with the help of the density-matrix renormaliza- spin σ , $n_{x\sigma} = c^{\dagger}_{x\sigma}c_{x\sigma}$ the number operator, tion group (DMRG) [11]. This method has $n_x = n_{x\uparrow} + n_{x\downarrow}$ and U is the onsite repulbeen previously shown to be highly successful sive energy. The onsite chemical potential for 1D quantum systems [12,13] and may be μ_x is a random number which we take to be seen as a numerical variational-wave-function uniformly distributed between $\pm \Delta$ such that approach [14]. After introducing the parame- $\Delta = 0$ corresponds to the clean case. We ters of our DMRG, we show that in the clean work at half-filling, e.g. $N = \langle \sum_{x=1}^{L} n_x \rangle = L$ case, the previously known results can be re- and the energy scale is set by choosing t = 1.

We follow the standard open chain sider finite disorder and show that the charge DMRG algorithm of White [11]. In each itergap G^c remains open for small disorder up to ation, we diagonalize the Hamiltonian matrix a critical disorder strength $\Delta_c \approx U/2$. The of a super-block denoted by $B_{L/2-1}^l \bullet \bullet B_{L/2-1}^r$ transition in the spin sector is seen by study- of L sites and obtain the energy and the ing the behavior of the spin-spin-correlation wave function $|\psi_{0L}\rangle$ of the ground state [13]. function $\langle S^-(r)S^+(0)\rangle$. For $\Delta < \Delta_c$, the Here • represents a single site. Using $|\psi_{0L}\rangle$

the right sub-block $B_{L/2}^r$. The above steps are then repeated for the new super-block tion.

as the target state [11] we compute the re- peat the DMRG steps with $N_{\uparrow} = L/2 + 1$ and duced density matrix ρ^l of the left sub-block $N_{\downarrow} = L/2$ and also with $N_{\uparrow} = L/2 - 1$ and $B_{L/2}^l \equiv B_{L/2-1}^l \bullet$ of size L/2. Diagonalizing $N_{\downarrow} = L/2$. We denote these two ground state ρ^l we obtain its eigenstates and the eigen- energies by $E_1(L)$ and $E_{-1}(L)$, respectively. values. These eigenstates with the highest The charge gap is then defined as the disconeigenvalues are the most probable states of tinuity of the chemical potential at half-filling the left sub-block when the super-block is in [4], *i.e.*, $G_L^c = E_1(L) + E_{-1}(L) - 2E_0(L)$. In the state $|\psi_{0L}\rangle$ and so can be used for trun- the presence of disorder G_L^c is computed for cation. Keeping M eigenstates correspond- at least 10 different disorder realizations and ing to the largest M eigenvalues of ρ^l as the then averaged over all such realizations. We new basis of the left sub-block, we trans- also compute the spin-spin-correlation funcform the Hamiltonian and all further oper- tion $\Gamma_L^s(r) = \langle \psi_{0L} | S^-(r) S^+(0) | \psi_{0L} \rangle$ and the ators into this new basis. Since every single second moment of the staggered antiferro-• has 4 states, we thus truncate the original magnetic (AFM) correlation function $(\xi_L^s)^2 =$ 4M states of $B_{L/2}^l$ to only M states. Usually $\sum_r r^2(-1)^r \Gamma_L^s(r)$. Here, $S^+(r) = c_{r\uparrow}^\dagger c_{r\downarrow}$. In we have used M = 128. Because the disor- the presence of disorder, the correlation funcder destroys translational symmetry, the left tions are first averaged over the disorder realand right sub-blocks are non-identical. Hence – izations and then the correlation length ξ_L^s of a similar procedure needs to be followed for the averaged staggered correlation function is found.

The clean case $\Delta = 0$: It has been shown $B_{L/2}^{l} \bullet \bullet B_{L/2}^{r}$ with L + 2 sites and thus the in Ref. [4] that the half-filled repulsive 1D system increases by two sites at each itera- Hubbard model exhibits a charge gap for all non-zero values of the interaction strength. Let $E_0(L)$ denote the ground state energy We find that the functional dependence of the of the Hubbard chain of length L with N_{\uparrow} = thermodynamic value of the charge gap G_{∞}^{c} $N_{\downarrow} = L/2$. In order to compute the charge on the interaction strength U as obtained by gap G_L^c for a system of finite length L, we re- DMRG (Fig. 1) is in excellent agreement with the exact solution computed in Ref. [4],

$$G^{c} = U - 4 + 8 \int_{0}^{\infty} dw \frac{J_{1}(w)}{w(1 + \exp(wU/2))}$$
(2)

with $J_1(w)$ a Bessel function. We note that the DMRG algorithm using open boundary conditions gives consistent results for the exwhich has been derived using periodic boundcurs at U = 0.

ies have predicted the existence of a critical analysis for U = 3 and 5 where a well devel-

disorder $\Delta_c > 0$ beyond which the Mott gap vanishes. In the limit of large U, this can be motivated [15] by potential energy considerations: any rearrangement of the one particle per site configuration in the half-filled Hubbard model would necessarily cost an energy U due to double occupancy and would gain trapolated charge gap G_{∞}^{c} with that of Eq. 2 in the local site potential energy a maximum of 2Δ . Hence, in order for it to be feasible for ary conditions. Nevertheless, we now em- the electrons to take advantage of the random ploy a finite-size-scaling (FSS) analysis of the site energies we should have $\Delta_c \ge U/2$ in the charge gap G_L^c in order to remove any finite- case of bounded disorder. In Fig. 3 we show size effects that could arise in the extrapola- G^c_{∞} as a function of the disorder strength Δ tion. In Fig. 1 (inset) we show that the lead- for U = 2. We see that the system undergoes ing order finite-size corrections to G^c_{∞} fall off a transition from a gapped Mott insulator as 1/L, *i.e.* $G_L^c(U) = G_\infty^c(U) + g(U)/L$. Hav- phase to a gapless phase for $\Delta_c \approx 1 = U/2$. ing determined the explicit scale dependence Fig. 3 (inset) shows the finite-size corrections of G_L^c , a plot of LG_L^c versus U as in Fig. 2 to G_∞^c in the presence of two representative shows curves for different L coalescing as the weak ($\Delta = 0.1$) and strong ($\Delta = 1.0$) disorcharge gap vanishes. This finite-size behav- der strengths for U = 2. We note that the ior allows a numerically accurate determina- corrections continue to fall off as 1/L even tion of the critical value of the interaction in the presence of disorder. The FSS plot of strength and we find in accordance with the LG_L^c versus Δ (Fig. 4) shows that curves for result of Ref. [4] that the Mott transition oc- different L coalesce at $\Delta_c = 1.0 \pm 0.1$ again indicating the transition from a gapped to a The disordered case $\Delta \neq 0$: Quantum gapless phase as the strength of the disorder Monte Carlo [8] and bosonization [9] stud- is increased. We have also done the same

is more stable [11]. Our results indicate that the transition into the gapless phase takes place at $\Delta_c \approx U/2$.

Spin-spin-correlation function: For $\Delta = 0$, the half-filled 1D Hubbard model shows algebraically decaying antiferbosonization [6],

$$\Gamma^{s}(r) \approx A_{1} \frac{\cos(\pi r + \phi_{1})}{r} + \mathcal{O}(1/r^{2}), \quad (3)$$

off as 1/L such that $1/\xi_L^s(\Delta) = 1/\xi_\infty^s(\Delta) + \text{ dered half-filled Hubbard model.}$ $\zeta(\Delta)/L$ and we can also apply the previous larger Δ beyond Δ_s , the values of L/ξ_L^s do tional dependence of $G_\infty^c(U)$ on the interac-

oped Mott gap exists and where the DMRG not coalesce any more. This indicates the transition from power-law correlations into a short-ranged spin-disordered phase with a finite correlation length. Again, the same analysis of the spin-spin-correlation function for U = 3 and 5 confirms that $\Delta_s \approx U/2$.

Thus the two transitions seen above viz. romagnetic (AFM) correlations as computed (I) gapped Mott insulator phase to a gapless by the methods of conformal field theory and phase at Δ_c and (II) long-ranged AFM phase to a short-ranged spin-disordered phase at Δ_s , are coincident, and $\Delta_c = \Delta_s$ within the numerical accuracy. Beyond the crit-When the spin-spin-correlation function de- ical disorder $\Delta_c \approx U/2$ the gapped Mott cays as a power-law, as in Eq. 3, the AFM insulator with power-law AFM correlations correlation length ξ_{∞}^{s} diverges in the thermo- goes over to a gapless short-ranged spindynamic limit. However, in the absence of disordered phase. This is the main result of long-ranged correlations, ξ_{∞}^{s} remains finite. our work and allows us to show in Fig. 6 the Fig. 5 (inset) shows that for $\Delta = 0.1$ the phase diagram of the zero temperature quanfinite-size correction to $1/\xi_{\infty}^{s}$ continues to fall turn transitions that occur in the 1D disor-

In summary, the DMRG allows us to an-FSS analysis to the correlation length. In alyze the various quantum transitions that Fig. 5 we plot L/ξ_L^s as a function of Δ . The occur in the half-filled 1D Hubbard model in data for different L coalesce until the disor- the presence of onsite disorder. For $\Delta = 0$ der $\Delta_s \approx 0.9 \pm 0.1$. Thus for U = 2 the we find that the finite-size corrections to the staggered $\Gamma^s(r)$ continues to fall off with a thermodynamic value of the charge gap G^c_{∞} power-law up to this critical disorder Δ_s . For scale as 1/L (Fig. 1 (inset)). The func-

tion strength (Fig. 1) is in excellent agreement with the exact solution computed in Ref. [4] in support of our DMRG approach. We have further shown that for $\Delta \neq 0$, the finite size corrections to $G^c_{\infty}(\Delta)$ continue to fall off as 1/L. For small $\Delta < U/2$, the Mott gap is shown to survive. Curves of LG_L^c versus Δ for different L (Fig. 4) come together at $\Delta_c ~(\approx 1.0)$ and coalesce after that indicating a transition from the gapped Mott insulator phase to a gapless phase. A finitesize analysis of the staggered AFM correlation length ξ_L^s shows that the values of L/ξ_L^s for various disorders Δ (Fig. 5) also coalesce at $\Delta_s \approx 1.0 \pm 0.1$. These transitions in the charge and the spin sector are shown to be coincident $\Delta_c = \Delta_s$. Thus in the gapped Mott phase $\Gamma^{s}(r)$ decays algebraically and is short ranged in the disordered gapless phase. We obtain the (U, Δ) phase diagram (Fig. 6) showing the phase boundary which separates the gapped Mott insulator phase with algebraic spin correlations and the spindisordered gapless phase.

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FIGURES



FIG. 1. Filled squares (**n**) show the charge gap G_{∞}^c as a function of the interaction strength U. The exact result [4] is shown by the solid line. Inset: G_L^c vs 1/L for U = 0.0, 2.5, 3.0 and 3.5. The value of the intercept gives G_{∞}^c .



FIG. 2. LG_L^c as a function of the interaction strength U showing the coalescence of curves for different L at U = 0. This indicates that the critical value of the interaction strength at which the Mott transition occurs is U = 0.



FIG. 3. Filled squares (**n**) show the charge gap G_{∞}^c as a function of the disorder Δ for U = 2. Inset: G_L^c vs 1/L for $\Delta = 0.1$ and 1.0. The value of the intercept gives G_{∞}^c .



FIG. 4. LG_L^c as a function of the disorder Δ for U = 2 showing the coalescence of the curves for different L at $\Delta = 1.0$. This indicates the transition from a gapped Mott insulator phase to a gapless phase.



FIG. 5. L/ξ_L^s as a function of the disorder Δ for U = 2 showing the coalescence of the curves for different L at $\Delta \approx 0.9 - 1.0$. This indicates the transition from a gapped Mott insulator phase to a spin-disordered gapless phase. Inset: $1/\xi_L^s$ vs 1/L for $\Delta = 0.1$.



FIG. 6. The phase diagram of the disordered 1D Hubbard Hamiltonian (1) at half-filling showing the Mott insulator with algebraic spin correlations (I) and the gapless spin disordered phase (II). The phase boundary has been drawn through the computed points (filled squares (\bullet) with error bars) and is close to $\Delta = U/2$.