A numerical finite size scaling approach to many-body localization

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We develop a numerical technique to study Anderson localization in interacting electronic systems. The ground state of the disordered system is calculated with quantum Monte-Carlo simulations while the localization properties are extracted from the “Thouless conductance” g, i.e. the curvature of the energy with respect to an Aharonov-Bohm flux. We apply our method to polarized electrons in a two dimensional system of size $L$. We recover the well known universal $\beta(g) = d \log g / d \log L$ one parameter scaling function without interaction. Upon switching on the interaction, we find that $\beta(g)$ is unchanged while the system flows toward the insulating limit. We conclude that polarized electrons in two dimensions stay in an insulating state in the presence of weak to moderate electron-electron correlations.

Since the early days of Anderson localization [1], it is believed that in the thermodynamic limit, an arbitrary small disorder is enough to drive a two dimensional electron gas toward an insulator [2]. At the origin of this prediction is the scaling theory of localization [2] which conjectured that the evolution of the conductance with the system size obeyed a simple one parameter scaling function. An important numerical effort has since been devoted to establish the presence of this scaling [3, 4] and calculate the scaling function. While this one electron localization picture is now reasonably well understood, the corresponding many-body problem, where not only disorder but also electron-electron interactions are considered, is yet unsolved. An important literature has been devoted to the very strong [5] and weak disorder limit [6, 7] but very little on the interplay between interaction and localization itself [8]. It was generally assumed that electron-electron interactions did not modify drastically the one electron physics, so that the observation of a metallic state in two-dimensional Si MOFSETs in 1994 [9] came as an important surprise. It gave rise to a new interest in the subject [10, 11, 12] and raised the question of the possibility that electron-electron interaction could stabilize a metallic phase. Recent progresses in the weak disorder limit seem to indicate that it could indeed be the case [13].

Numerical methods have proved to be very useful in putting the scaling theory of localization for non-interacting particles on very firm grounds. It is therefore very tempting to try to develop similar approaches for the many-body problem in spite of the intrinsic difficulties in dealing with correlations. Indeed, a number of technical problems need to be overcome. (i) Obtaining the ground state of a decently large number $N$ of correlated particles is already a challenging task. (ii) In order to study Anderson localization (and not a mere trapping of the electrons which is found for very strong disorder) one needs the localization length $\xi$ to be rather large yet smaller than the system size $L$ which must hence be rather large itself. (iii) One needs to calculate a physical observable sensitive to localization which must hence be some sort of correlation function [14]. Indeed, thermodynamic quantities such as the electronic density do not show localization in average.

In this letter, we propose a practical scheme to study numerically many-body localization. We use zero temperature Green Function quantum Monte-Carlo (GFMC) technique to study the ground state of the system. This technique takes full advantage of the fact that the non-

![Figure 1](image-url)
log $g$ as a function of the system size $L$, in square systems with $\nu = 1/25$. Disorder and interactions are increased from top to bottom as indicated by the arrow. The different curves are: $r_w = 0.005, r_s = 0$ (△), $r_w = 0.1, r_s = 0$ (×), $r_w = 0.15, r_s = 0$ (●), $r_w = 0.175, r_s = 0$ (◇), $r_w = 0.2, r_s = 0$ (□), $r_w = 0.225, r_s = 0$ (□), $r_w = 0.25, r_s = 0$ (□). The collapse of the linear fits (dashed lines) at $L = 0$ is the signature of one parameter scaling in the localized regime.

interacting ground state can be found exactly (by diagonalization of the one-body problem) so that upon switching on the interaction the GFMC simulations are done with a very good starting point. Our tool to measure the localization properties is the “Thouless conductance” of the system, which can be related to the distribution of the winding numbers in the imaginary time path integral. We apply our method to polarized (spinless) electrons in two dimensions for which both theory and experiments agree that the system is insulating. The main point of scaling theory of localization is that $\beta(g) \equiv d \log g / d \log L$ which depends on disorder, interaction, density and size is in fact a function of $g$ only. Our chief result is presented in Fig. 1 where we establish this scaling for interacting electrons. We find that $\beta(g)$ is unaffected by the presence of the correlations due to Coulomb repulsion in agreement with what is expected from the weak disorder limit [12]. Upon increasing the interaction strength, the system flows toward the insulating limit and the system localization length $\xi$ (shown in Fig. 4) decreases.

Model and method. We consider a system of $N$ spinless electrons in a rectangular $L_x \times L_y$ lattice with periodic boundary conditions. The Hamiltonian reads,

$$ H = -t \sum_{\langle \vec{r}, \vec{r}' \rangle} c_{\vec{r}}^\dagger c_{\vec{r}'} + \sum_{\vec{r}} v_r n_{\vec{r}} + \frac{U}{2} \sum_{\vec{r} \neq \vec{r}'} V(|\vec{r} - \vec{r}'|) n_{\vec{r}} n_{\vec{r}'} + \lambda, $$

(1)

where $c_{\vec{r}}^\dagger$ et $c_{\vec{r}}$ are the usual creation and annihilation operators of one electron on site $\vec{r}$, the sum $\sum_{\langle \vec{r}, \vec{r}' \rangle}$ is restricted to nearest neighbors and $n_{\vec{r}} = c_{\vec{r}}^\dagger c_{\vec{r}}$ is the density operator. The disorder potential $V_{\vec{r}}$ is uniformly distributed inside $[-W/2, W/2]$. $U$ is the effective strength of the two body interaction $V(\vec{r})$. To reduce finite size effects, $V(\vec{r})$ is obtained from the bare Coulomb interaction using the Ewald summation technique. The expressions for $V(\vec{r})$ has been given in [10]. At small filling factor $\nu \equiv N/(L_x L_y) \ll 1$, we recover the continuum limit and we are left with two dimensionless parameters, the usual $r_s = m^* e^2/(\hbar^2 \epsilon \sqrt{\pi n})$ ($m^*$ effective mass, $\epsilon$ electron charge, $\epsilon$ dielectric constant and $n$ electronic density) interaction parameter which for our model reads $r_s = U/(2t \sqrt{\pi n})$ and a parameter $r_w = W/(t \sqrt{\pi n})$ controlling the strength of the disorder. In the diffusive limit without interaction, the product of Fermi momentum $k_F$ by the mean free path $l$ is given by $k_F l = 192 \pi / r_s^2$.

The GFMC method and our particular implementation has been given in [10] to which we refer for details and references. GFMC is a lattice version of the standard zero-temperature quantum Monte-Carlo methods (like diffusive quantum Monte-Carlo) that have enjoyed important success for both bosonic and fermionic systems [17]. Its principle is to project an initial variational guiding wave-function (GWF) $|\Psi_G\rangle$ onto the exact ground state $|\Psi_0\rangle$ by applying the projector operator $e^{-\beta H}$ in a stochastic way. Quantum Monte-Carlo methods suffer from the so called sign problem when dealing with fermionic statistics. One way out of the sign problem which has been quite successful is the fixed node approximation [17] where upon projection onto $|\Psi_0\rangle$ the sign of the wave-function is kept fixed. The method is variational and calculates the best wave-function compatible with the nodal structure of the GWF. Important effort is usually spent looking for a GWF as close to the real ground state as possible. In the present case however, we can obtain the ground state without interaction exactly by diagonalizing the corresponding one-body problem. As we are interested in the evolution of the localization properties upon switching on the interaction, we have an excellent starting point to begin with. The general form of our GWF is a Slater determinant multiplied by a Jastrow function,

$$ \Psi_G(\vec{r}_1, \vec{r}_2, ..., \vec{r}_N) = \text{Det} \left[ \phi_i(\vec{r}_j) \right] \times \prod_{i < j} J(|\vec{r}_i - \vec{r}_j|). $$

(2)

The Jastrow part introduces some correlation and account for Coulomb repulsion. We use modified Yukawa functions [18] : $J(r) = \exp\left[ -a \frac{\sqrt{r}}{\Delta} \left( 1 - e^{-B(r)/\rho} \right) \right]$ where $a = 1/\sqrt{\Delta}$ is the average distance between electrons. $A(r_s)$ and $B(r_s)$ are variational parameters that we optimize while imposing the cusp condition $B = \sqrt{r_s} / \Delta$ to reproduce the short distance behaviour. The nodal structure of the GWF depends only on the Slater determinant $\text{Det} \left[ \phi_i(\vec{r}_j) \right]$ which enforces the antisymme-
try. The Slater determinant is constructed out of one-body orbitals $\phi_i$ that are obtained in two different ways leading respectively to $\Psi_{\text{liq}}$ and $\Psi_{\text{Har}}$ GWF. The orbitals of $\Psi_{\text{liq}}$ are calculated by exact diagonalization of the one-body (disordered) problem so that $\Psi_{\text{liq}}$ coincides with the exact ground state without interaction ($r_s = 0$). The calculation of the orbitals of $\Psi_{\text{Har}}$ proceeds in a similar way but we include iteratively the (Hartree) mean field potential due to the density $\langle n_r \rangle$ of electrons in the one-body problem. The Hartree potential tends to screen the disorder leading to an increase of the GWF's localization length. We shall verify however that the GFMC results are not sensitive to the choice of GWF.

Measuring the localization properties. The idea to use the sensibility $D$ of the system to a tilt in its boundary conditions as a criteria of localization was introduced very early by Edwards and Thouless [13]. Indeed, in a periodic system, the position of the boundary can be moved by a simple gauge transformation so that all sites are equivalent with respect to the boundary and a localized state is expected to have an exponentially small sensitivity to the boundary. More precisely, in presence of a small Aharonov-Bohm flux $\phi$, a current $I = -\partial E/\partial \phi$ flows in the system ($E$ is the total energy). When the flux is small, we have $I = -D\phi$ where $D = \partial^2 E/\partial \phi^2|_{\phi=0}$ is the curvature of the energy. This quantity $D$ is referred as the “Thouless conductance”, the Drude weight, the conductivity stiffness or the superfluid stiffness depending on the context. For bosonic systems, $D$ is simply related to the superfluid fraction [20]. For fermions, it is related to the low frequency limit of the imaginary part of the conductivity [21]. For disordered system the product of $D$ with the density of states is proportional to the conductance of the system [19, 22]. In most cases, $D$ is positive and the system is diamagnetic but in some instances (one dimensional systems with even number of spinless electrons [23], or two dimensional systems with degenerate ground states [24]) a paramagnetic ($D < 0$) response can been found so that the widely used interpretation of $D$ as a conductance can sometimes be problematic. In any case, it is a good measure of the localization properties of the system.

Following [20], we calculate the diffusive constant $g$ of the motion of the center of mass of the system in imaginary time along the $x$ direction, $g = \lim_{\beta \to 1} N \langle R^2_s(\beta) \rangle / (t/\beta)$, where $\langle R^2_s(\beta) \rangle$ is the second moment of the center of mass along $x$. An example of the calculation of $g$ is shown in the left panel of Fig. 3. $g$ is simply related to $D$ as $g = DL^2/\langle Nt \rangle$ yet $g$ is easier to access in the simulations. We note that in the fixed node approximation, $g$ is always positive by construction. In particular, in the absence of disorder $r_w = 0$ and interaction $r_s = 0$, we find $g = 2$, i.e. the sum of the curvature of the individual one-body levels [21].

**Numerical results: square samples.** We now turn to the numerics and show that $g$ is an appropriate measure of localization. In Fig. 2 we plot $\log g$ (averaged upon disorder) as a function of system size $L$ for square samples at fixed filling factor. Without interaction, scaling theory of localization [2] predicts that $\log g$ is independent of $L$ for small disorder (Ohm’s law) while for strong disorder, $g$ decreases exponentially with $L$ so that $\log g = \log g_0 - L/\xi$. The existence of a universal scaling law $g_L$ in this case means that $\log g_0$ is just a constant, independent of the disorder strength. Indeed, the numerics are fully consistent with this picture: for $r_w \leq 15$, $\log g$ is roughly constant. Upon increasing disorder further, $\log g$ starts to decrease linearly with $L$ while all curves intercept at a single point at $L = 0$. We find $g_0 = -0.47 \pm 0.08$. Further check of the method can be done by comparing the localization length $\xi$ with results $\xi_{ex}$ of an exact diagonalization of the one-body Hamiltonian [25]. The result is shown in the lower right panel of Fig. 3. We find both methods in good agreement as $\xi \propto (\xi_{ex} - 1)$ [20].

We are now ready to switch on the interaction in our system. We find (Fig. 2 full symbols) that upon increasing $r_s$, the localization length decreases so that the system becomes more insulating. More importantly, all the curves still intercept at the same single point at $L = 0$ indicating that the universal scaling function $g(L)$ is unaffected by the electron-electron correlations. To confirm this important point, we have performed simulations with our two different wave-functions $\Psi_{\text{liq}}$ and $\Psi_{\text{Har}}$ and find that the resulting $\xi$ agree very well as shown in the low-
find that the universal scaling function $\beta(g)$ is unaffected by the interactions. Yet, upon increasing the interaction strength, the system flows toward the insulating limit. This picture is in agreement with what is expected in the weak disorder and weak interaction limit. A natural extension of this work would be the study of non polarized electrons where the existence of an intrinsic metal-insulator transition remains a controversial issue. We note that in Fig. 1, the localization length $\xi$ extrapolates to zero at $r_s \approx 10$ for the two studied values of disorder. This could be the signature of a transition toward some sort of disordered Wigner crystal. Remarkably, $r_s \approx 10$ also corresponds to the density at which the metal-insulator transition was observed for all but the cleanest samples.

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