**Ab initio** Lattice Results for Fermi Polarons in Two Dimensions

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We investigate the attractive Fermi polaron problem in two dimensions using nonperturbative Monte Carlo simulations. We introduce a new Monte Carlo algorithm called the impurity lattice Monte Carlo method. This algorithm samples the path integral in a computationally efficient manner and has only small sign oscillations for systems with a single impurity. As a benchmark of the method, we calculate the universal polaron energy in three dimensions in the scale-invariant unitarity limit and find agreement with published results. We then present the first fully nonperturbative calculations of the polaron energy in two dimensions and density correlations between the impurity and majority particles in the limit of zero-range interactions. We find evidence for a smooth crossover transition from fermionic quasiparticle to molecular state as a function of the interaction strength.

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One of the most interesting and fundamental problems in quantum many-body physics is the polaron problem, where a mobile impurity interacts with a bath of particles. With the advent of trapped ultracold atomic gases, the polaron problem can now be realized for both bosonic and fermionic baths and also in the universal limit where the range of the particle interactions is negligible [1]. In a fermionic medium, the impurity can undergo a transition and change its quantum statistics by binding fermions from the surrounding Fermi gas [2,3]. The impurity is dressed by fluctuations of the Fermi sea forming a quasiparticle or polaron state. But with increasing particle interaction strength, molecules will form by capturing one or even two particles from the Fermi sea, and this behavior has been shown to depend on the mass ratio of the two components of the Fermi gas for the 3D case [2–10]. In 1D, the exact analytical solution for equal masses shows that the polaron-molecule transition is a smooth crossover [11,12].

In 2D, the Fermi polaron properties have been studied by using different theoretical and experimental approaches, and these have predicted various scenarios for the existence or absence of a polaron-molecule transition [13–21]. The Fermi polaron system has been studied by using diagrammatic Monte Carlo (diag MC) calculations [20,21]. The diag MC method uses a worm algorithm to stochastically sample Feynman diagrams to high orders in the coupling constant. In this work, we introduce a nonperturbative ab initio approach called the impurity lattice Monte Carlo (ILMC) method [22] to investigate highly imbalanced Fermi gases. Unlike diag MC calculations, the impurity lattice Monte Carlo calculation directly samples the path integral and so is a fully nonperturbative calculation. We present calculations for the energy of the 2D polaron and density correlations between the impurity and majority particles as a function of the interaction strength. Our results show evidence for a smooth crossover from a polaron to a molecule.

Impurity lattice Monte Carlo method.—The impurity lattice Monte Carlo method is a hybrid of two Monte Carlo algorithms, namely, worldline and auxiliary-field Monte Carlo algorithms. In the ILMC approach, the spacetime worldlines of each impurity are sampled explicitly, while all other particles are handled by using the auxiliary-field formalism. The impurity worldlines also function as local auxiliary fields felt by the other particles in the system. We illustrate the method by considering a d-dimensional many-body system of two-component fermions with equal masses $m_\uparrow = m_\downarrow = m$ with attractive interactions in the zero-range limit. In our many-body system, $N$ up-spin particles fill the Fermi sea, and one down-spin particle is an impurity immersed in this Fermi sea. In the zero-range limit, the interaction potential can be replaced by a delta function interaction. Using the lattice spacing to regularize the short-distance physics, we can write our lattice Hamiltonian as

$$H = H_0 + C_0 \sum_{\vec{n}} a_\uparrow(\vec{n}) a_\downarrow(\vec{n}),$$

where $\vec{n}$ denote spatial lattice points on a d-dimensional $L^d$ periodic cube. The free lattice Hamiltonian $H_0$ is given by

\[ H_0 = \sum_{\vec{n}} \left( \frac{p_{\vec{n}}^2}{2m} - \frac{U_d}{2} n_{\uparrow}(\vec{n}) n_{\downarrow}(\vec{n}) \right), \]
\[ H_0 = H_0^{\uparrow} + H_0^{\downarrow} \]
\[ = -\frac{\hbar^2}{2m^2} \sum_{\mu=1}^{d} \sum_{n,i=\uparrow,\downarrow} a_i^{\dagger}(\vec{n}) a_i(\vec{n} + \hat{\mu}) - 2a_i(\vec{n}) \]
\[ + a_i(\vec{n} - \hat{\mu}) \],

where \( a \) is the spatial lattice spacing. We fix the coupling constant \( C_0 \) in order to reproduce either the desired two-particle scattering length or the binding energy of a shallow dimer at infinite volume. The partition function of the system can be written as

\[ Z = \text{Tr}(M_{L_t}) \tag{3} \]

where \( M \) is the normal-ordered transfer matrix operator

\[ M = e^{-a_i H_0 + C_0 \sum n a_i^{\dagger}(\vec{n}) a_i(\vec{n}) a_i^{\dagger}(\vec{n}) a_i(\vec{n})} / \hbar \],

and \( a_i \) is the temporal lattice spacing.

We are interested in the system containing one down-spin particle together with \( N \) up-spin particles. Let us consider one forward time step from \( n_i \) to \( n_i + 1 \). If the down-spin worldline remains stationary at some lattice site \( \vec{n} \) during this time step, then there is no interaction between the impurity and the up spins. We get an effective transfer matrix for the up-spin particles that has the form

\[ M_{\uparrow}(n_i) = \left( 1 - \frac{d a_i \hbar}{m a^2} \right) e^{-a_i H_0^{\uparrow} + C_0 \sum n a_i^{\dagger}(\vec{n}) a_i(\vec{n})} / \hbar \].

We note the local potential generated by the down-spin impurity sitting at lattice site \( \vec{n} \). If the down-spin worldline instead hops from one spatial lattice site to another, then there is no interaction between the impurity and the up spins. Therefore, the effective up-spin transfer matrix is

\[ M_{\uparrow}(n_i) = \left( \frac{a_i \hbar}{2ma^2} \right) e^{-a_i H_0^{\uparrow}} / \hbar \].

For more details on the impurity lattice Monte Carlo formalism, we refer to Ref. [22].

**Polarons in three dimensions at unitarity.**—As a benchmark of the ILMC method, we present simulations of polarons in the 3D unitarity limit. We define \( \epsilon_p < 0 \) as the difference between the ground-state energy of the system with a single impurity compared to the system without the impurity. In the unitarity limit, where the S-wave scattering length diverges, the polaron energy is a universal quantity and scales with the Fermi energy, \( \epsilon_p = \theta \epsilon_F \), where \( \theta \) is a universal dimensionless number.

Using our effective up-spin transfer matrix, we compute Euclidean time projection amplitudes and determine the ground-state energy by taking the ratio of projection amplitudes for \( L_t \) and \( L_t - 1 \) time steps in the limit of large \( L_t \). For more details, see Supplemental Material [23].

In order to determine the polaron energy at unitarity, we have performed simulations for several different lattice volumes \( L^3 \) as well as several different values for \( N \), the number of up-spin particles. Since the scattering length is tuned to infinity, taking the limit of infinite volume at a fixed particle number corresponds to taking the continuum limit with the interaction range going to zero. At a fixed particle number, we determine the polaron energy for each system at lattice volumes 6, 7, 8, 9, and 10. We then apply a linear extrapolation in the inverse lattice spacing, the expected leading lattice-spacing dependence of corrections from the scale-invariant unitarity limit. See, for example, Ref. [24] for similar extrapolations in the unitarity limit. Repeating this procedure for systems with \( N = 15, 20, 25, 30, \) and 35, we also extrapolate the polaron energy to the thermodynamic limit. For the thermodynamic limit extrapolation, we perform a linear fit in \( 1/N \).

In Fig. 1, we show results for the polaron energy in the unitarity limit. Using the impurity lattice Monte Carlo method, we find \( \theta = \epsilon_p / \epsilon_F = -0.622(9) \). This result is in very good agreement with the result \( \theta = -0.618 \) determined in Ref. [2] using diagrammatic Monte Carlo calculations with the variational calculations with one and two particle-hole pair excitations giving \( \theta = -0.6066 \) [25] and \( \theta = -0.6158 \) [7], respectively. All these theoretical calculations are also consistent with the experimental values \( \theta = -0.58(5) \) [26] and \( \theta = -0.64(7) \) [10] measured in ultracold atomic gases.

**Attractive polarons in two dimensions.**—We now consider attractive polarons in two dimensions. There is no analog of the unitarity limit in two dimensions, since the only scale-invariant fixed point is in a weakly interacting limit [27,28]. But there is a very interesting and important question as to whether a polaron-molecule transition occurs in the ground state as a function of the interaction strength.
At this time, experiments are not yet conclusive on the question of a transition [17,18]. The existence and nature of such a transition impact the overall phase diagram for spin-imbalanced 2D Fermi gas [29–33]. Some work using a variational approach did not show any ground-state transition [13]. However, later studies found evidence for a transition when treating molecule and polaron variational wave functions in a similar fashion [15], and similar findings have been obtained in diagrammatic Monte Carlo simulations [20,21]. While these variational and diag MC studies are impressive and informative, one does not gain information about the nature of the transition itself. Separate calculations are needed to describe the fermionic polaron and the molecular state, and there is no overlapping region where both calculations are reliable. In order to remedy this situation, we use impurity lattice Monte Carlo simulations to study the nonperturbative physics of the transition in detail.

We consider again one down-spin impurity and \( N \) up-spin particles in the limit of zero-range attractive interactions. The delta function interaction is tuned according to the two-body bound state energy \( |e_B| \). Using the impurity lattice Monte Carlo calculation, we calculate the polaron energy as a fraction of the up-spin Fermi energy. We tune the coupling constant in order to get the two-body bound states with binding momentum \( k_B = \sqrt{m|e_B|} \) equal to 0.22\( \hbar /a \), 0.31\( \hbar /a \), 0.43\( \hbar /a \), 0.53\( \hbar /a \), and 0.62\( \hbar /a \).

We run simulations for several different lattice areas \( L^2 \) and several different particle numbers \( N \). The lattice sizes go from \( L^2 \times L = 20^2 \times 100 \) to \( L^2 \times L = 80^2 \times 700 \). For each \( L^2 \) and \( N \) we find the ground-state energy by extrapolating to the limit \( L \to \infty \) by fitting the Euclidean time projection amplitude to the asymptotic function \( e_0 + ae^{-\kappa L} \). To magnify the details, we subtract the dimer energy in vacuum, \( e_B \), from the polaron energies and scale by \( e_F \), the majority up-spin particle Fermi energy.

In Fig. 2, we have plotted the subtracted-scaled polaron energy \( (e_p + |e_B|)/e_F \) versus the dimensionless parameter \( \eta \equiv \frac{1}{2} \ln(2e_F/|e_B|) \), which characterizes the strength of the interaction. The simulations are done with \( N = 21 \) and \( N = 20 \) up-spin particles. For comparison, we have plotted the diagrammatic Monte Carlo results from Ref. [20] and variational results from Refs. [15,19]. The dashed line shows polaron variational results including one particle-hole \( (p - h) \) pair [15], and the solid line gives polaron variational results including up to two \( p - h \) pairs [19]. The dot-dashed line shows the improved variational results for the molecule state including one \( p - h \) pair [15]. We also compare with experimental data [34] presented in Ref. [17] and find good agreement for the weak coupling region \( \eta > 1 \). For the strong coupling region \( \eta < 1 \), the experimental uncertainty increases dramatically, and there seem to be nonuniversal systematic effects in the experimental realization that need to be corrected (see, for example, Ref. [35]).

![FIG. 2 (color online). Ground-state energy as a function of the dimensionless parameter \( \eta = \frac{1}{2} \ln(2e_F/|e_B|) \) in comparison with diag MC results [20]. The experimental data are from Ref. [34]. Dashed line: Polaron variational results using one particle-hole \( (p - h) \) pair [15]. Solid line: Polaron variational results using up to two \( p - h \) pairs [19]. Dot-dashed line: Improved molecule state variational results including one \( p - h \) pair [15]. The vertical band represents the region where the crossover transition from a polaron to a molecule occurs.](image-url)
indicator of a transition from fermionic polaron to paired molecular state as a function of the coupling strength.

Despite the computational advantages of one initial state over another, we can also obtain the same ground-state energies within error bars using different initial states. The paired Gaussian wave packet initial state does well for simulations far into the weak coupling limit when we change the size of the wave packet. At strong coupling, the optimal wave packet is fairly compact, indicating a localized pair. At weak coupling, the optimal wave packet is very large, growing as big as the lattice length $L$.

The location of our crossover region is in good agreement with the results of diag MC calculations which have found transitions at $\eta = -0.95 \pm 0.15$ [20] and $\eta = -1.1 \pm 0.2$ [21]. This result is also consistent with variational calculations [19] which obtain a transition in the region $-0.97 < \eta < -0.80$. Our result also compares well with the experimental result $\eta = -0.88(20)$, obtained after converting the experimental data [34] performed in a quasi-2D trap to their corresponding pure 2D values [35].

We have done simulations for a wide range of particle numbers and found rapid convergence to the thermodynamic limit $N \to \infty$. In fact, we find very little dependence on $N$ for $N \geq 8$. The lattice results in Fig. 2 correspond to simulations with the largest numbers of particles: $N = 21$ for weak coupling and $N = 20$ for strong coupling. As we can see in Fig. 2, there is also relatively little spread in the lattice results for different values of the binding momenta. This indicates that we are close to the continuum limit of zero-range interactions, and so corrections to the continuum limit are also numerically small.

Density-density correlations.—In order to uncover the underlying nature of the polaron-molecule transition, we have used ILMC to measure the density-density correlation function between impurity and majority particles:

$$
\rho_{1\downarrow}^{(\vec{r})} = \int d^2 r' \langle \rho_{1\uparrow}(\vec{r} + \vec{r}') \rho_{1\downarrow}(\vec{r}') \rangle.
$$

We have considered $N = 8, 14, 20$ up-spin particles on an $L^2 = 40^2$ periodic lattice and interactions ranging from weak coupling to strong coupling, $\eta = 1.5, 0.5, -0.8, -1.0$. The results are shown in Fig. 3. We express quantities in terms of the natural length scale $h/k_F$ and plot the logarithm of the correlation function to show the behavior both near and far from the central peak.

There are several interesting features apparent in Fig. 3. First, there is little dependence on $N$, which indicates that one is already close to the thermodynamic limit even for the small $N = 8$ system. Second, although the height of the central peak depends on $\eta$, the width of the peak is approximately $h/k_F$ and largely independent of the interaction strength.

Third, at strong coupling we see a tall central peak and a deficit in the wings of the profile that dips below the background value. This can be interpreted as the impurity stripping away an up-spin particle from its immediate surroundings. The fourth and perhaps most important finding is that there is no sign of a sharp phase transition such as a divergence of the correlation function or non-analytic dependence on the interaction strength. This shows that the polaron-molecule transition is a smooth crossover.

Discussion and outlook.—In all of the lattice simulations presented here, we have found that, with a sensible choice of the initial state, the sign oscillations are mild. This is good news for large-scale ILMC simulations of other systems. Some possible examples include alpha particles in dilute neutron gases, $\Lambda$ hyperons in neutron and nuclear matter, and imbalanced cold atomic systems. In these simulations, one uses auxiliary-field Monte Carlo simulations for the majority particles in addition to the impurity worldline updates. If the simulations without the impurity can be done, then simulations with the impurity can also be done with little increase in sign oscillations.

At weak coupling, the impurity worldline has only a minor effect on the eigenvalues of the $N \times N$ matrix of single-particle amplitudes, and so the determinant of the matrix remains positive. At strong coupling, we find one large positive eigenvalue that depends on the impurity worldline. But this large eigenvalue is positive, and the other eigenvalues once again are largely independent of the worldline. So the determinant remains positive in this case also. This reflects the underlying physics that, at strong coupling, we have a tightly bound molecule which interacts weakly with the surrounding Fermi gas. The sign problem is worst in the crossover region when we are in between the two extreme cases. However, even in the crossover region, the sign problem remains quite manageable for the systems we have considered.
By calculating the energy of the impurity and the density-density correlations versus interaction strength, we have found evidence for a smooth crossover from a polaron to a molecule. We are now applying the ILMC method to investigate the polaron-molecule transition in three dimensions as well as impurities in paired superfluid systems, with applications to ultracold atomic systems and alpha particles in neutron gases.

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