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A path integral study of the role of correlation in exchange coupling of spins in double quantum dots and optical lattices

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Abstract

We explore exchange coupling of a pair of spins in a double dot and in an optical lattice, using the frequency of exchanges in a bosonic path integral, evaluated using Monte Carlo simulation. The algorithm gives insights into the role of correlation through visualization of two-particle probability densities, instantons, and the correlation hole. We map the problem to the Hubbard model and see that exchange and correlation renormalize the model parameters, dramatically reducing the effective on-site repulsion at larger separations.

(Some figures in this article are in colour only in the electronic version)

Lattice models are popular in solid state physics and often serve as simple models for atomic orbitals, especially in the theory of magnetism [1]. Quantum dot arrays and optical lattices are new realizations of lattices. These artificial lattices are candidates for quantum computers, where spins on exchange-coupled dots are qubits for universal quantum computation [2, 3]. A fundamental concept is intersite exchange, in which virtual hopping leads to spin coupling of neighboring sites. A two-site model is one of the simplest quantum problems, yet the quantitative mapping from a three-dimensional model of a double dot or optical lattice experiment to an effective two-site model has many subtleties requiring careful treatment of exchange and correlation [1, 3–6].

In this paper we use path integral Monte Carlo (PIMC) to extract accurate singlet–triplet splitting from a spatial model. Similar PIMC algorithms have been used to study spin dynamics in ³He [7–9] and Wigner crystals [8, 10], and the approach is particularly simple for two-site models. This two-particle problem has been previously solved with direct diagonalization (DD) methods with a careful choice of basis functions [4, 6] and is amenable to variational or diffusion quantum Monte Carlo (QMC) [11]. However, the simple and elegant PIMC approach is a more direct solution without variational bias or basis set issues and offers theoretical insights into this important problem. We first show that the splitting energy, $J$, is easily extracted from the average permutation of the two-particle path integral, even when $J \ll k_B T$. This PIMC algorithm is a black-box calculator, providing accurate numerical estimates of $J$ for technologically relevant models of dots or optical lattices with arbitrary interactions and confinement potentials. More importantly, PIMC allows us to ask questions about quantum correlation. For example, do the particles exchange across the barrier simultaneously, or do they briefly double occupy the dot? Or, does the motion of one particle over the barrier correlate with the location of the other particle? We answer these questions by viewing representative trajectories (instantons) for a double dot and calculating pair correlation functions. Magnetic fields are known to modulate $J$ [3, 4, 6], and we show how to include them in PIMC with a simple Berry’s phase [12]. Finally we model recent experiments of exchange coupled atoms in an optical trap, demonstrating broader utility [13].

The mapping from a continuous model with interacting particles to a lattice model introduces subtle complications. For a non-interacting system it is reasonable to reduce the Hilbert space to just one orbital per site, coupled by a hopping matrix element, $t$. The non-interacting many-body ground
state is a product state of these single-particle orbitals. Low energy excited states are spanned by this basis, so an effective lattice model is an excellent approximation. Interactions are typically added to this lattice model as on-site energies, $U$, or intersite terms, $V$. For small $t$, this gives the well-known $J = 4t^2/(U - V)$.

There can be a serious flaw when considering interactions in this order. When interactions are added to the continuum Hamiltonian, correlation enters as virtual excitations to higher energy orbitals. At first this seems insignificant, since there may be still a one-to-one mapping to an effective lattice model. But, when choosing effective lattice parameters, one must remember that many-body states in the continuum model have quantum fluctuations that are simply not present in the lattice model.

As a specific example, consider two electrons in a double quantum dot. This system is often represented as a two-site Hubbard model, where the sites represent the 1s ground states of the dots. Correlation terms involve virtual excitation of the electron to the $2p_x$ and $2p_y$ states of the dots. These quantum fluctuations create van der Waals attraction, in addition to mean-field repulsion. Van der Waals attraction and other correlations renormalize the interaction parameters to new values, $U_t$ and $V_t$.

When we consider hopping between sites, more complications emerge. The hopping barrier has contributions from both the external potential and electron–electron interactions. While the mean-field Hartree contribution can simply be added to the effective potential, the fluctuating part is not so trivial. In the transition state, an electron passes over a barrier whose height has quantum fluctuations. Thus we expect interactions to renormalize the hopping constant, $t$. At the Hartree–Fock level, Hund–Mulliken theory already predicts a renormalized $t$ and $U_t$ due to long-range exchange [1, 3]. However, neglect of correlation in Hund–Mulliken theory can lead to catastrophic failure at intermediate dot separations [6]. PIMC includes all correlations, and illuminates their role in barrier hopping with the concept of instantons.

We start with the two-dimensional model for the GaAs double quantum dot studied in [6],

$$H = \frac{p_x^2}{2m^*} + \frac{p_y^2}{2m^*} + \frac{e^2}{\epsilon |r_1 - r_2|} + V_{\text{ext}}(r_1) + V_{\text{ext}}(r_2),$$

with $m^* = 0.067 m_e$ and $\epsilon = 12.9$. The external potential comes from two piecewise-connected parabolic potentials,

$$V_{\text{ext}}(r) = \frac{1}{2} m_o \omega^2 \{\min[(x-d)^2, (x+d)^2] + y^2\},$$

with minima at $x = \pm d$. We report $d$ relative to the oscillator length $r_0 = \sqrt{\hbar/m_o}$. The two lowest energy two-electron states are spatially symmetric and antisymmetric under exchange, with energies $\epsilon_+$ and $\epsilon_-$, respectively. The exchange coupling, $J = \epsilon_+ - \epsilon_-$, has been calculated previously using DD on a basis of Fock states built from seven single particles states [6]. Much care was taken to test convergence with the number of states and careful evaluation of Coulomb matrix elements. We note that the same quality of DD calculation in three dimensions would typically take more single-particle states.

QMC techniques give essentially exact answers to many problems without basis set convergence issues, and often work just as easily in multiple dimensions. PIMC is nice for quantum dot problems [14] because it does not require a trial wavefunction. However, direct calculation of either $\epsilon_+$ or $\epsilon_-$ with PIMC often have large statistical errors in energy ($\sim 1$ meV in dots). Instead, we use particle exchange statistics to estimate energy differences $J$ to high accuracy ($\sim 1\mu$eV) in PIMC.

To calculate $J$, we split the partition function into terms that are spatially symmetric and antisymmetric under exchange, $Z = Z_+ + Z_-$. These terms can be expressed as symmetrized or antisymmetrized imaginary time path integrals [15, 16], (see appendix),

$$Z_{\pm} = \frac{1}{Z} \sum_{r,\tau} (\pm 1)^{\beta} \int DR(\tau) e^{-\beta S_{\pm}[R(\tau)]}. \quad (3)$$

This is a sum over all two-particle paths $R(\tau)$ with the boundary condition $R(\beta) = P R(0)$, where $P$ swap particle positions and $Z$ is the identity. The symbol $(\pm 1)^{\beta}$ takes the values $(\pm 1)^{\beta} = 1$ and $(\pm 1)^{\beta} = \pm 1$ at low temperature, only one state contributes to each partition function, so $Z_{\pm} = e^{-\beta \epsilon_{\pm}}$. Thus,

$$e^{-\beta J} = \frac{Z_+}{Z_-} = \sum_{\rho} \int DR(-\beta) e^{-\beta S_{\pm}} \sum_{\rho} \int DR e^{-\beta S_{\pm}} \equiv \langle (-1)^{\rho} \rangle_{\beta}, \quad (4)$$

or $J = -k_B T \ln\langle (-1)^{\rho} \rangle_{\beta}$. Thus the exchange coupling can be calculated by sampling a symmetric (bosonic) path integral [16] and taking the average of $\langle (-1)^{\rho} \rangle$, which is $+1$ for identity paths and $-1$ for exchanging paths.

We ran PIMC simulations [16] with our open-source p\textsuperscript{3}c code for the dots studied in [6], with the results shown in figure 1(a). To aid other researchers, we have made the simulation available as a tool on nanoHUB [17]. Coulomb interactions are included with a pair action that correctly handles the cusp condition. We observed convergence of the path integral results with 6400 discrete slices, but a higher-quality pair action [16] could require fewer slices. We see near perfect agreement with DD, and speculate that small deviations may be due to the finite basis in the DD calculation or approximations in the evaluation of Coulomb matrix elements at larger $d$ [6].

To learn more, we collect the two-particle density, $\rho(x_1, x_2)$, which is the probability to find one electron at $x_1$ and the other at $x_2$, integrated over all values of $y_1$ and $y_2$, and shown in figure 2(b). We calculate double occupation, $x_D$, which we define as the probability for the electrons to lie on the same side of the $x = 0$ plane. From $J$ and $x_D$ we deduce renormalized values for $t$ and $U_{\text{td}} - V$, figures 1(c) and (d). Interactions renormalize $t$ to smaller values, consistent with Hund–Mulliken theory or a larger renormalized mass. The larger $J$ arises from the dramatic decrease in $U_{\text{td}} - V$ at larger dot separations, as correlation enables more virtual hopping.

There are two minima, $(x_1, x_2) = (\pm d, \mp d)$, in the total potential, marked ‘+’ in figure 2(b). For non-zero $J$, some paths must go between these minima. In a semiclassical
Figure 1. PIMC results. (a) Exchange couplings $J$ for $\hbar \omega = 4$ meV (●) and $\hbar \omega = 6$ meV (○) double dots with a piecewise parabolic potential (inset). Dashed lines are direct diagonalization results from [6]. (b) The double dot occupation probability $x$. Using $J$ and $x$ we fit (c) $t$ and (d) $V_r - U_r$ parameters for an effective two-site Hubbard model. Dashed lines in (c) show the bare hopping $t$ for one electron in the double dot. Dashed lines in (d) are $V - U$ with $V = e^2/2\epsilon_d$ and $U$ taken from a PIMC calculation on a single dot.

Figure 2. Paths and pair densities for a double dot. (a) Simplified instanton with double occupation of the right dot. (b) Pair densities $\rho(x_1, x_2)$ with the lowest density contour line that connects both potential minima (+ markers) at $(\pm d, \mp d)$. (c) Simplified instanton with simultaneous exchange. (d) Actual path showing double occupation, sampled from $\hbar \omega = 4$ meV, $d = 1.5r_0$ dots. (e) Actual path showing simultaneous exchange, sampled from $\hbar \omega = 6$ meV, $d = r_0$ dots. Trajectories (d) and (e) are also plotted in (b).

Figure 3. Crossing density, $\rho_2(x, x)$, equivalent to the diagonal of the pair densities in figure 2(b).

In the path integral, a magnetic field is easily implemented as a Berry’s phase $q\Phi_B$, where $q$ is the electron charge and $\Phi_B$ is the total magnetic flux enclosed by the path of the two electrons. The exchange splitting is then $J(B) = -k_B T \ln(\langle e^{q\Phi_B} (\pm 1)^2 \rangle / \langle e^{q\Phi_B} \rangle)$.

For charged particles, magnetic fields can be used to tune the exchange coupling and even change its sign [3, 18].
Figure 4. Conditional density of one electron when the other electron is in the white box, showing the correlation hole during an instanton. Panels (a)–(c) are the $\hbar \omega = 4 \text{meV}, d = 1.5 \sigma_0$ dots and (d)–(f) are the $\hbar \omega = 6 \text{meV}, d = 1.0 \sigma_0$ dots. Numerical factors are the likelihood of the first electron being in the white box. PIMC results are shown on the left of each image, with DD results [6] on the right.

Figure 5. Magnetic field dependence included with a Berry’s phase for several double quantum dots. The denominator is small and Monte Carlo sampling errors are catastrophic. In practice, we find that fields up to 4 T in strength are practical for the geometries we study, yielding the results in figure 5.

To test the applicability of the method to an optical lattice, we consider the exchange of two $^{87}\text{Rb}$ atoms in a double-well optical trap [13]. This system resembles the double dot, only with much heavier particles, a much lower temperature, short-range interactions, and a different confining potential. While the experiments in [13] have very little correlation, we present results to demonstrate feasibility for such systems, which can be made more strongly interacting. The experiment has a double-well potential, $V(x) = V_{\text{long}} \sin^2(\pi x/\lambda) + V_{\text{short}} \cos^2(2\pi x/\lambda)$, with $\lambda = 765 \text{nm}$ and $V_{\text{long}} = 10E_r$, where $E_r = \hbar^2/2M_{\text{Rb}}\lambda^2$ [13]. We model interactions as $V(r) = V_0 \text{sech}^2 \kappa r$ with $V_0 = 50.5 \mu\text{K}$ and $\kappa = 0.1 \text{nm}^{-1}$ to reproduce the $^{87}\text{Rb}$ scattering length. Figure 6 shows $J$ as the barrier $V_{\text{short}}$ is varied, confirming agreement with experiment.

In conclusion, we have demonstrated a PIMC algorithm for computing exchange splitting in double quantum dots and optical lattices. The exchange splitting arises from instantons in the path integral, and we have collected data on these path crossings, including double occupation and the correlation hole. Correlations renormalize $t_i$ and $U_i - V_i$, with a dramatic decrease in $U_i - V_i$ at large separation. We find that simultaneous crossing occurs more often with closely spaced dots, while further separated dots are more likely to have instantons with double occupations. Finally, we have demonstrated the versatility of the algorithm with the inclusion of magnetic fields and applications to laser-trapped atoms.

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**Appendix. Form of the discretized path integral and the action**

The partition function for the effective mass Hamiltonian, equation (1), can be written as an imaginary time path integral [14–16],

$$Z = \int \mathcal{D}R(\tau) e^{-\frac{i}{\hbar}S_{\text{E}}[R(\tau)]}.$$  \hspace{1cm} (A.1)

The path integral $\int \mathcal{D}R(\tau)$ and Euclidean action $S_{\text{E}}$ are easiest to define in the discretized form we used in the Monte Carlo integration. By dividing imaginary time into $N_T$ discrete steps, each of length $\Delta \tau = \beta \hbar/N_T$, the path $R(\tau)$ becomes an array of positions (‘beads’) $r_i^j$, where $i$ indicates the slice number ($0 \leq i < N_T$) and $j = 1, 2$ labels the two electrons. Then the path integral becomes a multiple integral over all bead
positions,
\[ \int D\mathbf{r}(\tau) \rightarrow \prod_{i=0}^{N_T-1} \int d\mathbf{r}_{1j} \int d\mathbf{r}_{2j}. \quad (A.2) \]

The action \( S_E \) represents the terms in the effective mass Hamiltonian and is a function of the bead coordinates,
\[
S_E = \sum_{i=0}^{N_T-1} \left[ \frac{m^* |\mathbf{r}_{i+1} - \mathbf{r}_i|^2}{2\Delta \tau} + \frac{m^* |\mathbf{r}_{i+2} - \mathbf{r}_i|^2}{2\Delta \tau} 
+ 2 \ln (2\pi \Delta \tau / m^*) + V_{\text{ext}}(\mathbf{r}_i) \Delta \tau + V_{\text{ext}}(\mathbf{r}_{i+1}) \Delta \tau 
+ u_{\text{coul}}(\mathbf{r}_{i+1}, \mathbf{r}_{i+2}, \mathbf{r}_i, \mathbf{r}_i; \Delta \tau) \right]. 
\quad (A.3)

The first three terms (which explicitly contain \( m^* \)) are the kinetic action and are derived from the free particle propagator in two dimensions. The next two terms are the action for the confining potential, \( V_{\text{conf}}(\mathbf{r}) \), evaluated in the primitive approximation [16]. The last term is the pair Coulomb action [16], which we have fit to a short time approximation \[16\]. The last term is the pair Coulomb propagator for the imaginary time interval \( \Delta \tau \). Because of special symmetry of the Coulomb potential, this propagator is only a function of two coordinates, \( q_i = (|\mathbf{r}_{i+1} - \mathbf{r}_{i+2}| + |\mathbf{r}_{i} - \mathbf{r}_{i+2}|)/2 \) and \( s_i^2 = (|\mathbf{r}_{i+1} - \mathbf{r}_{i+2}| - (\mathbf{r}_{i} - \mathbf{r}_{i+2}))^2 \). For simplicity, we have dropped the \( s_i^2 \) dependence; this approximation is exact as \( \Delta \tau \rightarrow 0 \). We evaluated the short time Coulomb propagator using the high-accuracy Trotter method of [19] and stored tabulated values of \( u_{\text{coul}}(q_i; \Delta \tau) \) on a grid for efficient evaluation during our Monte Carlo simulations.

To perform the trace implicit in equation (A.1), we identify slice \( N_T \) with slice 0 in equation (A.3) by setting \( \mathbf{r}_{N_T+1} = \mathbf{r}_{01} \) and \( \mathbf{r}_{N_T+2} = \mathbf{r}_{02} \). The division of the partition function into spatially symmetric \( Z_+ \) and antisymmetric \( Z_- \) parts may be accomplished by summing over permutations \( P = \mathcal{I}, \mathcal{P} \), as in equation (3). Permuting configurations \( (P = \mathcal{P}) \) are handled by setting \( \mathbf{r}_{N_T+1} = \mathbf{r}_{01} \) and \( \mathbf{r}_{N_T+2} = \mathbf{r}_{01} \) in equation (A.3).

References