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Extending J walking to quantum systems: Applications to atomic clusters

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The J-walking (or jump-walking) method is extended to quantum systems by incorporating it into the Fourier path integral Monte Carlo methodology. J walking can greatly reduce systematic errors due to quasiergodicity, or the incomplete sampling of configuration space in Monte Carlo simulations. As in the classical case, quantum J walking uses a jumping scheme to overcome configurational barriers. It couples the usual Metropolis sampling to a distribution generated at a higher temperature where the sampling is sufficiently ergodic. The J-walker distributions used in quantum J walking can be either quantum or classical, with classical distributions having the advantage of lower storage requirements, but the disadvantage of being slightly more computationally intensive and having a more limited useful temperature range. The basic techniques are illustrated first on a simple onedimensional double well potential based on a quartic polynomial. The suitability of J walking for typical multidimensional quantum Monte Carlo systems is then shown by applying the method to a multiparticle cluster system consisting of rare gas atoms bound by pairwise Lennard-Jones potentials. Different degrees of quantum behavior are considered by examining both argon and neon clusters. Remarkable improvements in the convergence rate for the cluster energy and heat capacity, analogous to those found in classical systems, are found for temperatures near the cluster transition regions.

I. INTRODUCTION

We recently presented a method called J walking (for jump walking)\(^1\) that was shown to greatly reduce systematic errors occurring in random walks in classical systems that arise because of quasiergodicity,\(^2\) the incomplete sampling of configuration space. We demonstrated the technique's utility on small atomic clusters bound by Lennard-Jones potentials. Recently, Tsai and Jordan\(^3\) applied J walking together with histogram methods\(^4\) to a Monte Carlo simulation of the water octomer, and Strozak, Lopez, and Freeman applied J walking to a simulation of argon clusters absorbed on graphite.\(^5\) In this work, we extend the method to quantum systems, incorporating it into the Fourier path integral (FPI) Monte Carlo methodology.\(^6\) Path integral methods based on Feynman's formulation of quantum statistical mechanics\(^7\) provide a prescription for the computer simulation of quantum many-body systems. In the FPI scheme, the quantum paths associated with each coordinate are represented by Fourier series expansions, leading to a generalized configuration space that has been extended to higher dimensions by the inclusion of the auxiliary degrees of freedom from the Fourier coefficients. This enhanced multidimensional space is amenable to the usual methods associated with the Metropolis, Rosenbluth, Rosenbluth, Teller, and Teller algorithm.\(^8\)

As in the classical case, quasiergodicity arises in quantum systems where the sample space contains two or more regions having a very low transition probability between them, resulting in bottlenecks that effectively confine the sampling to only one of the regions.\(^9\) The dichotomy of time scales characterizing the walks produces rapid motion within a confined region and very slow movement between the regions.\(^10\) This results in systematic errors that arise because of the finite length of the walks. They diminish with increased walk length, disappearing in the limit of infinitely long walks. A prototypical example is the double well potential where the wells are separated by a large barrier. For sufficiently low temperatures, the random walker is unable to cross over the barrier within the duration of the walk and hence never samples from the other well.

J walking addresses this problem by coupling the usual small scale Metropolis moves with occasional large scale jumps that move the Metropolis walker to different regions of configuration space, in essence “jumping” over the barriers rather than walking over them. This is accomplished in classical systems by using the Boltzmann distribution obtained at a temperature sufficiently high for the sampling to be ergodic. The distribution generated by the high temperature walker (J walker) becomes the sampling distribution for the low temperature walker’s attempted jumps. Because the distribution’s peaks correspond to the potential minima, the J-walker's motion remains biased about the minima, greatly increasing the likelihood an attempted jump would be accepted. An analogous scheme can be developed for quantum systems simply by sampling from the distribution generated by a high temperature quantum walker. However, because of their larger number of param-
eters, FPI methods offer a greater degree of flexibility than available in the Metropolis simulation of classical systems, allowing several possible variations on the theme. Since the configuration space is often dominated by the coordinate subspace, an alternative scheme is low for the temperature quantum walker to sample from a classical J-walker distribution. Another possibility is to use a quantum J-walker with the same temperature, but having a lower mass, since quasiergodicity becomes less important in high quantum systems where larger zero-point energies and tunneling effectively lower the configurational barriers.

Two complementary implementations for generating the classical J-walker Boltzmann distributions were originally presented. The first ran the J-walker in tandem with the low temperature walker. The low temperature walker occasionally attempted jumps to the current J-walker position by simply using the J-walker coordinates for its trial move. The second scheme ran the J walker beforehand and periodically stored the configurations in an external array. Subsequent jump attempts could be made by accessing the stored configurations via randomly generated indices. The tandem walker scheme required that the J walker be moved an extra number of steps whenever a jump was attempted in order to reduce correlations between the two walkers that caused systematic errors. The extra number of steps needed depended on the temperature difference between the high and low temperature walkers, increasing the computation time greatly as the difference became larger. This scheme is therefore more suited for parallel computation. The external distribution scheme, on the other hand, had only a modest overhead in computation (mostly the time required to generate the distribution), but demanded large storage facilities to handle the distribution arrays. Because of the additional computational overhead inherent in the FPI method (typically up to an order of magnitude), we did not consider the tandem walker method to be feasible for the simulation of quantum systems on scalar computers and so we limited our investigation to externally stored J-walker distributions.

We begin in Sec. II with a brief review of the FPI method and the essential concepts that pertain to J walking. We then develop two quantum J-walking methods based on the use of quantum and classical distributions, and contrast them with the more familiar classical J-walking method. As in our original study of classical systems, we have tested the quantum J-walker method on a number of simple systems. For heuristic purposes, we begin in Sec. III with the simple model of the prototypical one-dimensional double well potential defined by a quartic polynomial. This potential consists of one well of fixed depth containing the global minimum and a second well of variable depth separated by a fixed barrier. The variable depth of the second well allows us to examine in a systematic manner quasiergodic behavior as a function of well depth as well as a function of temperature. A solution of high accuracy can be obtained easily using standard harmonic oscillator basis set expansion techniques so that errors arising from quasiergodicity can be quantified. In Sec. IV, we apply the J-walker method to quantum multiparticle cluster systems consisting of either Ne or Ar atoms bound by pairwise Lennard-Jones potentials and compare it to the usual FPI methods. Finally, in Sec. V we summarize our findings.

II. THEORY

A. Fourier path integral Monte Carlo methods

Path integral formulations of quantum statistical mechanics have become increasingly useful in recent years as continuing developments in computer technology have provided the computational power necessary for their application to realistic systems. There have been two major numerical algorithms developed to evaluate path integrals, discretized path integral methods,11 and Fourier path integral methods,6 although at a fundamental level, the two prescriptions are essentially the same.12 We describe here the application of J walking to the Fourier methods only and note in passing that, in principle, J walking can also be extended to discretized methods.

1. Action integrals

Recent reviews of FPI methods give a comprehensive description of their development and application.6 We review briefly here the major concepts that are instrumental to the application of J walking. For notational simplicity, we limit the discussion to one-dimensional systems; the extension to multidimensional systems is straightforward and is described in the reviews.

Path integral methods enable the calculation of the quantum density matrix, in coordinate representation

$$\rho(x,x';\beta) = \langle x' | \exp(-\beta H) | x \rangle,$$

by employing Feynman path integrals:

$$\rho(x,x';\beta) = \int Dx(u) \exp\{-S[x(u)]\},$$

where

$$S[x(u)] = \frac{1}{\hbar} \int_0^{\beta \hbar} du \left[ \frac{m \dot{x}^2(u)}{2} + V[x(u)] \right]$$

is in the form of a classical action integral in imaginary time. The temperature parameter is $\beta = 1/k_B T$, where $T$ is the temperature and $k_B$ is the Boltzmann constant. The integral $\int Dx(u)$ implies a summation over all paths connecting $x$ to $x'$ in imaginary time $\beta \hbar$. FPI methods represent each path parametrically in a Fourier series. Using $\hbar$ in units of $\beta \hbar$ gives expressions for the path

$$x(u) = x + (x' - x) u + \sum_{k=1}^{\infty} a_k \sin(k \pi u),$$

and for the density

$$\mu(x,x';\beta) = \mu_{fp}(x,x';\beta)$$

$$\times \frac{\int dx \exp\{-S_{\beta=1}^{\infty}(a_k^2/2\sigma_k^2) - \beta \hbar V[x(u)] du\}}{\int dx \exp\{-S_{k=1}^{\infty}(a_k^2/2\sigma_k^2)\}},$$

where

\[ (\text{continued}) \]
\[ \sigma_k^2 = \frac{2\beta \hbar^2}{m(k\pi)^2} \]  
(6)

is the width of the Gaussian-like Fourier coefficient distribution and \( \rho_{fp} \) is the free particle density

\[ \rho_{fp}(x,x';\beta) = \left( \frac{m}{2\pi\beta \hbar^2} \right)^{1/2} \exp\left[ -\left( \frac{m}{2\beta \hbar^2} \right)(x-x')^2 \right]. \]  
(7)

The Gaussian width \( \sigma_k \) provides a natural length scale, growing as the temperature is decreased; like the thermal deBroglie wavelength, it provides a measure of the extent of quantum contributions in the system.

The quantum density given by Eq. (5) can be used in the canonical ensemble to evaluate the expectation value of any operator depending only on the coordinates.\(^{13}\) For example, the expectation value of the potential energy is

\[ \langle V \rangle = \frac{\int \, dx \, da \, w(x,x,a;\beta) V(x)}{\int \, dx \, da \, w(x,x,a;\beta)} , \]  
(8)

where

\[ w(x,x',a;\beta) = \rho_{fp}(x,x';\beta) \exp \left[ -\sum_{k=1}^{\infty} \left( \frac{a_k^2}{2\sigma_k^2} \right) \right] \]

\[ -\beta \int_0^1 V(x(u)) \, du \]  
(9)

is the action weight factor. Since the quantum paths begin and end at the same point for properties dependent only on the coordinates, the free particle density reduces to a constant

\[ \rho_{fp}(\beta) = \left( \frac{m}{2\pi\beta \hbar^2} \right)^{1/2} . \]  
(10)

and so we denote the weight factor simply as \( w(x,a;\beta) \).

The effects of the higher-order Fourier coefficients on the expectation value decrease with increasing \( k \), and in practice, the infinite sums are truncated at some suitably selected maximum value of \( k \) denoted \( k_{\text{max}} \), giving an approximate path

\[ x_k(u) = x + \sum_{k=1}^{k_{\text{max}}} a_k \sin(k\pi u). \]  
(11)

The number of Fourier coefficients required for a given calculation depends on the system being studied, the temperature, and the level of accuracy desired. For notational convenience, we represent the average of some function along an approximate path as

\[ \langle F[x(u)] \rangle_a \equiv \int_0^1 F[x_k(u)] \, du . \]  
(12)

The expectation value for the potential then becomes

\[ \langle V \rangle_{k_{\text{max}}} = \frac{\int \, dx \, da \, w_{k_{\text{max}}}(x,a;\beta) V(x)}{\int \, dx \, da \, w_{k_{\text{max}}}(x,a;\beta)} , \]  
(13)

where

\[ w_{k_{\text{max}}}(x,a;\beta) = \rho_{fp}(\beta) \exp \left[ -\sum_{k=1}^{k_{\text{max}}} \left( \frac{a_k^2}{2\sigma_k^2} \right) \right] \]

\[ -\beta \langle V[x(u)] \rangle_a \]  
(14)

The quantum average is obtained in the limit that all the Fourier coefficients are included

\[ \lim_{k_{\text{max}} \to \infty} \langle V \rangle_{k_{\text{max}}} = \langle V \rangle . \]

For notational simplicity, we will omit the \( k_{\text{max}} \) subscript from subsequent averages. Equation (13) has the form of a classical Monte Carlo average, differing only in the increased dimensionality of the integrations due to the introduction of the Fourier coefficients \( a_k \) as additional integration variables. This addition of auxiliary degrees of freedom in the quantum expressions is characteristic of path integral treatments in general.

### 2. Kinetic energy estimators

The calculation of the kinetic energy expectation value is complicated by the nondiagonality of the kinetic operator in coordinate representation. Several kinetic energy estimators have been developed, two of which are the \( T \) method\(^ {14} \) (for temperature differentiation) and the \( H \) method\(^ {15} \) (for Hamiltonian operating on the density matrix). The \( T \) method evaluates the internal energy \( U \) in its entirety using the statistical mechanical expression

\[ \langle U \rangle_T = -\left( \frac{\partial \ln Q}{\partial \beta} \right)_V , \]  
(15)

where \( Q \) is the standard canonical partition function

\[ Q(T,V,N) = \int \, dx \, \rho(x;\beta) . \]  
(16)

Using Eq. (5) gives

\[ \langle U \rangle_T = \frac{k_{\text{max}} + 1}{2B} + \left( \langle V[x(u)] \rangle_a - \sum_{k=1}^{k_{\text{max}}} \frac{a_k^2}{2B\sigma_k^2} \right) . \]  
(17)

Since \( \langle U \rangle = \langle T \rangle + \langle V \rangle \) for the systems we consider, the \( T \)-method kinetic energy can be written explicitly as

\[ \langle T \rangle_T = \frac{1}{2B} \left[ \frac{k_{\text{max}} + 1}{2} - \sum_{k=1}^{k_{\text{max}}} \frac{a_k^2}{2B\sigma_k^2} \right] - \langle V(x) \rangle . \]

(18)

Another kinetic energy estimator closely related to the \( T \)-method estimator can be obtained by noting that in the limit \( k_{\text{max}} \to \infty \), the term \( \langle V[x(u)] \rangle_a \) in Eq. (18) (where the average is taken over all the paths) vanishes.\(^ {16} \) The expression resulting from the neglect of this term at finite \( k_{\text{max}} \) can also be obtained directly from the mass differentiation of \( Q \) and hence is denoted the \( M \) method (for mass differentiation).
The $H$-method estimator can be obtained directly as the expectation value of the kinetic operator $\hat{T} = (\hat{p}^2 / 2m)\nabla^2$:

$$\langle T \rangle_H = \frac{1}{2\beta} - \langle k_1^2 - K_2 \rangle,$$

which gives

$$\langle T \rangle_H = \frac{1}{2\beta} - \langle k_1^2 - K_2 \rangle,$$  \hspace{1cm} (21)

where

$$K_1 = \frac{\beta \rho^2}{2m} \langle (1 - u) V'[x(u)] \rangle, \hspace{1cm} (22)$$

$$K_2 = \frac{\beta \rho^2}{2m} \langle (1 - u)^2 V''[x(u)] \rangle. \hspace{1cm} (23)$$

Both the $T$ and $M$ methods have the advantage of being computationally simple to evaluate, but suffer the disadvantage of having a variance that grows linearly with $k_{\text{max}}$. The $H$ method is more computationally intensive, but its variance growth is usually much weaker, even decreasing with increasing $k_{\text{max}}$ for some systems.

3. Heat capacity

As with kinetic energy estimators, there are several estimators for the constant volume heat capacity $C_v$. The estimators we used are analogous to the $T$-method kinetic estimator—they are derived by temperature differentiation of the energy

$$\langle C_v \rangle_T = -k_B \beta^2 \left( \frac{\partial^2 \langle U \rangle}{\partial \beta^2} \right) \langle F \rangle,$$  \hspace{1cm} (24)

and hence are called $T$-method heat capacities. Inserting the $T$-method energy expression (17) into Eq. (24) gives the $T$-method heat capacity

$$\langle C_v \rangle_T = k_B \left[ \frac{k_{\text{max}} + 1}{2} - 2 \left( \sum_{k=1}^{k_{\text{max}}} \frac{\alpha_k^2}{2\sigma_k^2} \right) + \beta^2 \langle F^2 \rangle - \langle F \rangle^2 \right], \hspace{1cm} (25)$$

where

$$F = - \sum_{k=1}^{k_{\text{max}}} \frac{\alpha_k^2}{2\sigma_k^2} + \beta \langle V[x(u)] \rangle, \hspace{1cm} (26)$$

while using the sum of the potential energy and the $H$-method kinetic energy given in Eq. (21) gives a hybrid $TH$-method heat capacity

$$\langle C_v \rangle_{TH} = k_B \left[ \frac{k_{\text{max}} + 1}{2} - 2 \left( \sum_{k=1}^{k_{\text{max}}} \frac{\alpha_k^2}{2\sigma_k^2} \right) + \beta^2 \langle F^2 \rangle - \langle F \rangle^2 \right] + 2\langle k_1^2 - K_2 \rangle. \hspace{1cm} (27)$$

4. Partial averaging

Since the number of Fourier coefficients required in the FPI calculations increases rapidly with decreasing temperature or as the system becomes highly quantum, it is useful to employ techniques that improve the convergence of the calculations with respect to $k_{\text{max}}$. One such method is partial averaging,\(^17\) which takes into account the effects of the truncated Fourier coefficients of order higher than $k_{\text{max}}$ that are neglected by the direct Fourier method. The system potential is replaced with an appropriate effective potential obtained as the Gaussian transform of the bare potential

$$V_{\text{eff}}[x_k(u), u] = \frac{1}{\sqrt{2\pi} \sigma(u)} \int_{-\infty}^{\infty} dp \exp \left[ - \frac{p^2}{2\sigma^2(u)} \right] \langle x'_k(u) + p \rangle,$$  \hspace{1cm} (28)

where

$$\sigma^2(u) = \sum_{k=1}^{k_{\text{max}}} \sigma_k^2 \sin^2(k\pi u)$$

$$= (\beta \rho^2 / m) [u(1 - u)] - \sum_{k=1}^{k_{\text{max}}} \sigma_k^2 \sin^2(k\pi u). \hspace{1cm} (29)$$

This leads to an analogous partial average action weight factor

$$\omega_{PA}(x, u, \beta) = \rho \exp \left[ - \sum_{k=1}^{k_{\text{max}}} \frac{\alpha_k^2}{2\sigma_k^2} - \beta \langle V_{\text{eff}}[x(u)] \rangle \right].$$  \hspace{1cm} (30)$$

For polynomial potentials, the Gaussian transform is analytic and the effective potential becomes simply the bare potential plus correction terms comprised of potential derivatives and functions of $\sigma^2(u)$,

$$V_{\text{eff}}[x_k(u), u] = V[x_k(u)] + f_{PA}[x_k(u), \sigma^2(u)]. \hspace{1cm} (31)$$

A similar separability is found in the kinetic energy estimators

$$\langle T_{PA} \rangle_T = \frac{1}{\beta} \left[ \frac{k_{\text{max}} + 1}{2} - \left( \sum_{k=1}^{k_{\text{max}}} \frac{\alpha_k^2}{2\sigma_k^2} \right) - \beta \langle V_{\text{eff}}[x(u)] \rangle \right],$$  \hspace{1cm} (32)

$$\langle T_{PA} \rangle_M = \frac{1}{\beta} \left[ \frac{k_{\text{max}} + 1}{2} - \left( \sum_{k=1}^{k_{\text{max}}} \frac{\alpha_k^2}{2\sigma_k^2} \right) - \beta \langle V_{\text{eff}}[x(u)] \rangle \right],$$  \hspace{1cm} (33)

$$\langle T_{PA} \rangle_H = \frac{1}{2\beta} - \langle k_{PA}^2 - K_{2PA} \rangle,$$  \hspace{1cm} (34)

and in the heat capacity estimators,

$$\langle C_{\text{vPA}} \rangle_T = k_B \left[ \frac{k_{\text{max}} + 1}{2} - 2 \left( \sum_{k=1}^{k_{\text{max}}} \frac{\alpha_k^2}{2\sigma_k^2} \right) + \beta^2 \langle k_{PA}^2 \rangle - \langle F_{PA} \rangle^2 \right] - \beta \langle h_{PA} \rangle, \hspace{1cm} (35)$$
\[ \langle C_{\gamma\alpha} \rangle_{TH} = k_B \left[ \frac{1}{2} \beta \left( \langle F_{PA} (V - K_{1PA}^2 + K_{2PA}^2) \rangle - \langle F_{PA} \rangle (V - K_{1PA}^2 + K_{2PA}^2) \right) + \langle 2K_{1PA}^2 - K_{2PA}^2 \rangle \right] + \langle k_{PA} \rangle , \]  

(36)

where

\[ F_{PA} = - \sum_{k=1}^{\infty} \frac{\sigma_k^2}{2c_k^2} + \beta \langle V_{eff} [x(u)] \rangle \right] \]  

\[ K_{1PA} = \frac{\beta \hbar^2}{2m} \left( 1 - u \right) V_{eff} [x(u)] \right] \]  

\[ K_{2PA} = \frac{\beta \hbar^2}{2m} \left( 1 - u \right)^2 V_{eff} [x(u)] \right] \]  

The terms \( g_{PA} \), \( h_{PA} \), and \( k_{PA} \) are partial average terms that depend on the specific potential.

### B. J walking

For classical systems, the usual Metropolis method uses a random walker to sample the configuration space by making moves from an initial coordinate \( x_i \) to a final coordinate \( x_f \) with a probability of acceptance

\[ p = \min \{ 1, q(x_f|x_i) \} , \]  

(37)

where

\[ q(x_f|x_i) = \frac{T(x_f|x_i)p(x_f)}{T(x_f|x_i)p(x_i)} \]  

(38)

is the acceptance ratio, \( p(x) = Z^{-1} \exp[-\beta V(x)] \) is the Boltzmann distribution with \( Z \) the standard configuration integral, and \( T(x'|x) \) is the transition matrix or sampling distribution.

The sampling distribution is usually generated from uniform deviates \( \xi \) over a finite stepsize range \( \Delta \) to give

\[ T(x'|x) = \begin{cases} 1/\Delta & \text{for } |x' - x| < \Delta/2 \\ 0 & \text{otherwise} \end{cases} \]  

(39)

and

\[ q(x'|x) = \exp(-\beta [V(x') - V(x)]) \]  

(40)

Attempted moves are generated according to \( x' = x + (\xi - \frac{1}{2}) \Delta \) with the maximum stepsize \( \Delta/2 \) usually adjusted to give acceptance probabilities of approximately 50%. The required size decreases with increasing \( \beta \). This temperature dependence can lead to quasiergodic behavior whenever the step size becomes too small relative to the potential barrier heights and widths; the walker becomes effectively trapped within a region of configuration space and is unable to sample the whole space in a representative manner.

The J-walker method is based on the observation that the Boltzmann distributions governing the Metropolis walker are largely dependent on the form of the potential with distribution maxima corresponding to potential minima. The widths of the distribution peaks are inversely dependent on \( \beta \) with higher temperatures resulting in wider distributions. In essence, higher temperature walkers are less constrained since their larger step sizes allow them to overcome the potential barriers more effectively. For a Metropolis walk of a given length, there is a threshold temperature such that walks undertaken above the temperature are ergodic. J-walking then occasionally replaces the usual Metropolis moves with an attempted jump to the position occupied by such a higher temperature walker (J walker). This is equivalent to replacing the usual Metropolis transition matrix given by Eq. (39) with the Boltzmann distribution at the higher temperature whenever a jump is attempted

\[ T_J(x_J|x) = Z_J^{-1} \exp[ -\beta_J V(x_J) ] \]  

for \( 0 < \xi_J < P_J \),

(41)

where \( P_J \) is the jump attempt probability. This gives an acceptance ratio of

\[ q_J(x_J|x) = \exp \left( (\beta_J - \beta) [V(x_J) - V(x)] \right) \]  

(42)

where \( \beta_J \) is the J-walker temperature parameter. In the high temperature limit \( \beta_J \rightarrow 0 \), the acceptance ratio reduces to the standard Metropolis expression given in Eq. (40).

The Boltzmann distribution broadens as the temperature increases, J-walking in this limit reduces to simple jumping with a large stepsize \( \Delta_J \) and so the likelihood of a jump being accepted decreases sharply. In the limit \( \beta_J \rightarrow \beta \), \( q_J(x_J|x) \rightarrow 1 \) since the low temperature walker is now effectively sampling from its own distribution.

For quantum systems, the extra degrees of freedom associated with the Fourier coefficients result in an acceptance ratio

\[ q(x',a'|x,a) = \frac{T(x,a|x',a')} {w(x,a|x')} \]  

(43)

where \( w(x,a|\beta) \) is given by Eq. (14), or by Eq. (30) if partial averaging is used. Since the coordinate moves are usually more important than the Fourier coefficient moves, the usual implementation of the transition matrix is to move a randomly selected Fourier coefficient simultaneously with each attempted coordinate move. Usually the selection scheme for the Fourier coefficients weights the lower-order coefficients more highly than the higher-order coefficients. While the coordinate moves use the typical Metropolis scheme of uniform deviates scaled by a suitable box size \( \Delta_x \), the Gaussian-like distribution of the Fourier coefficients allows attempted coefficient moves to be sampled either from a Gaussian distribution of width \( \sigma_k \) or from uniform deviates scaled by box sizes \( \Delta_{ak} \). Both schemes work well, although sampling from a Gaussian distribution results in more rejected moves at lower temperatures.

Quantum J-walking then couples this scheme with attempted jumps to a higher temperature distribution of \( (x_J,a_J) \) configurations in a manner analogous to the classical case. Whenever a jump is attempted, the transition matrix used is simply the action weight factor for the higher temperature...
\[ T_f(x,a|x,a) = \left( \frac{m}{2\pi \beta J k_B^2} \right)^{1/2} \exp \left\{ -\sum_{k=1}^{k_{\text{max}}} \frac{a_k^2}{2\sigma_k^2} \right\} \]  
\[ -\beta J \langle V(x_j(u)) \rangle_{\sigma_k} \]  
(44)

The subscript on temperature-dependent parameters such as \( \sigma_k \) indicates they have had \( J \) replaced with \( \beta J \). Inserting Eq. (44) together with Eqs. (10) and (14) into Eq. (43) gives

\[ q_f(x,a|x,a) = e^{\Delta S_f}, \]  
(45)

where \( \Delta S_f \) is the change in the action in moving from \( (x,a) \) to \( (x,a_J) \). For the direct FPI formulation,

\[ \Delta S_f = -\left( \langle f[x(u);\sigma_J^2(u)]\rangle_{\sigma_j} - \langle f[x(u);\sigma^2(u)]\rangle_{\sigma_j} \right), \]  
(46)

while if partial averaging is used,

\[ \Delta S_{fPA} = \Delta S_f + \beta \langle f[x(u);\sigma_J^2(u)]\rangle_{\sigma_j} - \langle f[x(u);\sigma^2(u)]\rangle_{\sigma_j} \]  
\[ -\langle f[x(u);\sigma^2(u)]\rangle_{\sigma_j}. \]  
(47)

In the high temperature limit \( \beta J \to 0 \), the Fourier coefficient distribution width \( \sigma_J \) vanishes and the acceptance ratio reduces to the Metropolis expression (with large step sizes implicit), analogous to the classical case, so that the jump acceptance becomes negligibly small. In the limit \( \beta J \to \beta \), \( \sigma_J \to \sigma_J \), and \( \sigma_J^2(u) \to \sigma^2(u) \), giving \( q_f(x,a_J|x,a) \to 1 \), so all attempted jumps are accepted since the low temperature walker is again sampling from its own distribution.

The inverse dependence between \( \beta \) and the mass \( m \) as evident from the ratio \( \beta/m \) appearing in the expressions for parameters such as \( \mu \), \( \omega \), and \( \nu(u) \) suggests alternative formulations of \( J \) walking where the temperature is held constant and the \( J \) walker mass decreased, or where both the \( J \)-walker temperature and mass are varied. We have not investigated these formulations, but only note in passing that they might be considered in those situations where \( J \) walking at higher temperatures is problematic.

The disadvantage of using quantum \( J \)-walker distributions is that the storage requirements are increased by a factor \( k_{\text{max}} \). Since quasiergodicity results primarily from the large barriers associated with coordinate space, it seems plausible to use a distribution that is mostly dependent on the coordinates. Hence an alternative to using the quantum distribution for \( J \) walking is to use a classical \( J \)-walker distribution obtained from the Boltzmann distribution for coordinate moves and to then sample the Fourier moves from a normalized Gaussian distribution. The composite transition matrix for attempted jumps is then

\[ T_f(x,a_J|x,a) = Z_f^{-1} \exp \left\{ -\sum_{k=1}^{k_{\text{max}}} \frac{a_k^2}{2\sigma_k^2} - \beta_j V(x_j) \right\}. \]  
(48)

The prime on \( \sigma_k \) signifies that the coordinate and Fourier coefficient \( J \)-walker temperatures need not be the same, and in fact the resultant expression for the change in the action can be greatly simplified by setting the Fourier coefficient \( J \)-walker temperature equal to that of the low temperature walker. This results in the cancellation of the Fourier coefficient contributions, giving for the direct Fourier path integral formulation

\[ q_f(x,a_J|x,a) = e^{\Delta S_c}, \]  
(49)

where

\[ \Delta S_c = \beta J \langle V(x_j) - V(x) \rangle - \beta \langle \langle V[x_j(u)] \rangle_{\sigma_j} \]  
\[ -\langle V(x(u)) \rangle_{\sigma_j} \], \]  
(50)

and for the partial average formulation

\[ \Delta S_{cPA} = \Delta S_c + \beta \langle f[x(u);\sigma_J^2(u)]\rangle_{\sigma_j} - \langle f[x(u);\sigma^2(u)]\rangle_{\sigma_j}. \]  
(51)

The limiting behavior of the acceptance ratio is not as straightforward as in the purely classical and purely quantum cases discussed previously. In the high temperature limit \( \beta J \to 0 \), the \( J \)-walker effective potential \( \langle V[x_j(u)] \rangle_{\sigma_j} \) reduces to the potential \( V(x) \). Again because the distribution width increases with increasing temperature, the configuration \( x_j \) will not likely correspond to a minimum and the probability of the jump being accepted is small. In the limit \( \beta J \to \beta \), the change in the action \( \Delta S_c \) (or \( \Delta S_{cPA} \) ) will not be zero as was the case for the quantum distribution, and consequently the jump acceptance will not be unity. Physically, the low temperature walker is not sampling from its own distribution if it is sampling from a classical distribution at the same temperature. For higher temperatures, the quantum and classical distributions converge since the effective potentials approach the bare potentials and the partial average contributions become small, but for lower temperatures, these differences become appreciable. So in the limit \( \beta J \to \beta \), \( q_f(x,a_J|x,a) \to 1 \), but in the low temperature limit \( \beta J \to \infty \), \( q_f(x,a_J|x,a) \to 0 \). Thus \( J \) walking from a classical distribution is ultimately limited in its use at low temperatures (or low particle masses).

### III. Quasiergodicity in the Quantum Double Well Potential

We illustrate the effects of quasiergodicity on quantum systems by considering a one-dimensional double well potential defined by the quartic polynomial

\[ V(x) = \sum_{m=0}^{4} a_m x^m, \]  
(52)
where \( a_0 = h \), \( a_1 = 0 \), \( a_2 = -6\delta abh \), \( a_3 = 4\delta (a - b) h \), and \( a_4 = 3\delta h \), with \( \delta = [b^3(b + 2a)]^{-1} \). This function has a minimum located at \( x = b \) with \( V(b) = 0 \), a barrier of height \( h \) located at \( x = 0 \), and a second minimum located at \( x = -a \). By restricting \( a \) to the range \( 0 < a < b \), the \( x = b \) well is the global minimum and the potential varies from a single well and barrier for \( a = 0 \) to a symmetric double well for \( a = b \). We can parametrize the degree of asymmetry in the double well for a given \( b \) and \( h \) by recasting \( a \) in terms of the ratio of the well depths

\[
y = \frac{V(0) - V(a)}{V(0) - V(b)} = \left( \frac{b}{a} \right)^3 \left( \frac{a + 2b}{b + 2a} \right).
\]

(53)

For a given \( b \) and \( y \), \( a \) can be obtained easily by iteration. This potential is a generalization of the double well potential we examined in our study of \( J \) walking on classical systems, where the barrier height and global minimum were fixed at \( h = b = 1 \). For a particle of given mass \( m \), the degree of quantum mechanical behavior in terms of spacings between quantized energy levels can be varied by adjusting the “box size” \( b \) and barrier height \( h \). Alternatively, for a given potential, the degree of quantum behavior can be adjusted by varying the system mass. In order to compare the quantum results with the classical results obtained earlier, we have chosen the latter scheme and have used the previous classical potential with \( h \) and \( b \) fixed at unity and \( a \) variable. We have arbitrarily selected \( m = 1, 0.1, \) and 0.01 amu to model low, moderate, and high degrees of quantum behavior, respectively. Figure 1 shows the potentials for some representative values of \( y \).

### A. Exact results

The problems arising from quasiergodicity in these potentials can be seen by comparing their average energies obtained by FPI methods with the numerically exact solutions obtained directly from the eigenenergies. For systems at a defined temperature and volume, the average of a Hamiltonian operator \( \hat{P} \) in energy representation is

\[
\langle \hat{P} \rangle = \frac{\sum_n \rho_n e^{-\beta E_n}}{\sum_n e^{-\beta E_n}}.
\]

(54)

Because the potentials are quartic polynomials, their eigenenergies can be obtained easily by expanding the eigenfunctions in a harmonic oscillator basis set

\[
\psi_n(x) = \sum_j c_{j+\frac{1}{2}} \phi_j(x)
\]

and using standard numerical techniques on the resulting matrix equations. A suitable oscillator frequency can be obtained in terms of the barrier height \( h \) with \( \omega = \sqrt{2h/m} \). Since FPI methods can provide averages for both the potential and kinetic energies, we calculated expectation values for both the potential and kinetic operators (in amu)

\[
\langle \hat{P} \rangle = \sum_{m=0}^4 c_m \sum_{k} c_{j\pm\frac{1}{2}} \langle \phi_j | x^m | \phi_k \rangle,
\]

(56)

\[
\langle \hat{T} \rangle = \frac{m \omega^2}{2} \sum_{j \pm \frac{1}{2}} c_j c_{j\pm\frac{1}{2}} \langle \phi_j | x^2 | \phi_k \rangle.
\]

(57)

As a check, the eigenenergies were also obtained directly from the differential equations using common two point boundary value numerical methods. Figure 2 shows the energy levels for the \( y = 0.8 \) potential for each of the three masses, while Fig. 3 shows the average internal energy as a function of \( \beta \) for the same potential, as well as the average classical energy for comparison.

### B. FPI results

Using the FPI formulation, the average potential energy for a given potential at some temperature defined by \( \beta \) is given by Eq. (13), while kinetic energies can be obtained with Eqs. (32)–(34). Applying partial averaging to the double well potential given by Eq. (52) gives the effective potential correction term for Eq. (31) in terms of the polynomial coefficients

\[
f_{PA}[x_n(u), \sigma^2(u)] = [\alpha_2 + 3\alpha_3 x_n(u) + 6\alpha_4 x_n^2(u)] \sigma^2(u) + 3\alpha_4 \sigma^4(u).
\]

(58)

Likewise, the \( T- \) and \( M- \) method correction terms in Eqs. (32) and (33) are

\[
g_{PA}[x_n(u), \sigma^2(u)] = [\alpha_2 + 3\alpha_3 x_n(u) + 6\alpha_4 x_n^2(u)] \sigma^2(u) + 6\alpha_4 \sigma^4(u).
\]

(59)
Partial averaging provided significant improvements in convergence for this potential and all results reported subsequently were obtained with partial averaging invoked. Hence for the sake of notational simplicity, the partial average subscripts will be omitted. All of the double well results were obtained as averages of 100 independently initialized FPI Metropolis walks, each consisting of $10^4$ warm-up moves followed by $10^4$ moves with data accumulation. Each attempted move consisted of a coordinate move together with a Fourier coefficient move (sampled from a Gaussian distribution of width $\sigma^2_k$, with the coefficient index selected from a Gaussian distribution of width $k_{\max}/2$ to give higher weight to the lower-order coefficients. The action integrals were evaluated using Simpson's rule with $n_{\text{quad}} = 16$ quadrature points.

In our study of quasiergodic behavior in classical systems we noted that the direction of the error resulting from the incomplete sampling depended on the walk initialization and that this property could be used as an indicator of quasiergodicity. For quasiergodic walks originating at the global minimum, the average value of the potential energy $\langle V \rangle$ is too low since the higher energy wells are insufficiently sampled, while walks originating from higher wells result in values that are too high. This is the case for quantum systems as well. Figure 4 shows the results obtained for $\langle V \rangle$ and for the $H$-method kinetic energy $\langle T \rangle_H$ for quantum particles with masses of $m = 1, 0.1, \text{and } 0.01 \text{amu.}$ The plots show the variation with $\beta$ for a fixed potential having $\gamma = 0.9,$ and the variation with $\gamma$ for a fixed temperature corresponding to $\beta = 10.$ As in the classical case, there is a threshold temperature that marks the onset of quasiergodicity for walks of a given length. It occurs at the corresponding $\beta$ point where the $\langle V \rangle$ curves representing global initialization diverge from the $\langle V \rangle$ curves representing random initialization. The threshold temperature depends slightly on the mass, being lowest for the $m = 0.01 \text{ system.}$ This can be attributed to the effective lowering of the energy barrier by the higher quantum zero point energy with the lower mass and by the increased importance of tunneling. The effects are particularly noticeable in the
FIG. 4. Quasiergodicity in Metropolis FPI walks for the double well potential for particles of mass 1, 0.1, and 0.01 amu. The left-hand plots depict the average potential energy $\langle V \rangle$ and $H$-method kinetic energy $\langle T \rangle_H$ for the $\gamma=0.9$ potential as functions of $\beta$, while the right-hand plots depict $\langle V \rangle$ and $\langle T \rangle_H$ as functions of $\gamma$ for $\beta=10$. In each case, the smooth curves are the numerically exact solutions, while the Metropolis curves were obtained from 100 independently initialized walks consisting of $10^4$ warm-up moves followed by $10^4$ moves with data accumulation, all with partial averaging invoked. Each attempted move consisted of a coordinate move sampled from a uniform distribution, together with one Fourier coefficient move sampled from a Gaussian distribution. The Fourier coefficient moved was also selected randomly from a Gaussian distribution. The number of Fourier coefficients needed to achieve convergence in the kinetic energy at $\beta=10$ is given by $k_{\text{max}}$. Some representative single standard deviation error bars have been included. The potential energy curves exhibit the bifurcation characteristic of quasiergodicity in classical systems. In each plot, the lower branch corresponds to walks initialized at the global minimum $x=1$, while the upper branch corresponds to walks randomly initialized. The kinetic energy curves exhibit no bifurcation.
converges faster than the $T$-method estimator and has a variance that is significantly better for the $H$-method kinetic estimator than for the $T$-method estimator. Figure 5 compares the $H$- and $T$-method kinetic energies as functions of $\beta$ for various $k_{\text{max}}$ for the $\gamma=0.9$ potential for $m=0.01$. As expected, the convergence in $k_{\text{max}}$ is significantly better for the $H$-method kinetic estimator than for the $T$-method. While the $T$ method has a lower variance at low $k_{\text{max}}$, its variance increases much more rapidly with increasing $k_{\text{max}}$ than does the $H$-method variance, so that its variance is comparable to the $H$ method's in the convergence limit. The larger variance at higher $k_{\text{max}}$ can make quasiergodicity less of a problem for highly quantum systems since the longer walks required to reduce the variance in the kinetic energy will also increase the likelihood of the walker passing through the bottlenecks in configuration space.

C. $J$-walking results

The ability of $J$ walking to eliminate the systematic errors arising from quasiergodicity in the quantum double well potential can be seen in Fig. 6, which shows results obtained using $J$ walking from classical and quantum distributions. The plots depict the internal energies as functions of $\gamma$ for $\beta=10$ for masses $m=1$ and $m=0.1$. The jump attempt probability was $P_J=0.1$. Comparison with the analogous FPI Metropolis results depicted in Fig. 4 shows that the large systematic errors in the average potential energy have been eliminated. The results were obtained in a similar manner as those in Fig. 4. Although we have depicted only four combinations, we ran calculations for $k_{\text{max}}=1, 2, 4, 8, 16$, and 32 for all three masses for $J$ walking from both quantum and classical distributions. All combinations gave similarly good results, differing only with the increased noise at higher values of $k_{\text{max}}$. The effects of various $J$-walking parameters on the quantum results are very similar to the effects on the analogous classical results we obtained in our earlier work, and we refer the reader to that paper for a detailed discussion of the effects of varying the jump attempt frequency and distribution parameters, and instead concentrate here on the differences between $J$ walking from quantum distributions and classical distributions.

The quantum distributions consisted of $10^5$ configurations sampled every ten moves from individual high temperature FPI Metropolis walks at $\beta_J=3$. Each configuration consisted of the coordinate and its associated $k_{\text{max}}$ Fourier coefficients. An attempted jump simply required selecting one of the configurations randomly and evaluating the acceptance ratio given by Eq. (45). The computational overhead for the evaluation of the acceptance ratio was kept very low by also storing the Fourier coefficient term $\sum_k |\tilde{a}_k|^2/2\sigma_k^2$ the path averaged effective potential $\langle V(x(u)) \rangle_{\tilde{a}_k}$ and the partial average term $\langle f(x(u);\sigma_k^2(u)) \rangle_{\tilde{a}_k}$ for each configuration in the distribution. Consequently, the extra computational cost required to implement $J$ walking from a quantum distribution was due mostly to the generation of the distribution from a long FPI Metropolis walk (the length being the product of distribution size and the sampling frequency).

The classical distributions similarly consisted of $10^5$ configurations sampled every ten moves from individual high temperature classical Metropolis walks at $\beta_J=3$. However, each classical configuration consisted only of the coordinate, reducing the storage requirements by a factor of $k_{\text{max}}$ compared to the quantum distributions. An attempted jump using a classical distribution consisted of randomly selecting a configuration from the coordinate distribution and generating each of the Fourier coefficients $a_k$ randomly from a Gaussian distribution of width $\sigma_k$. The attempted jump would then be accepted or rejected according to the acceptance ratio given by Eq. (49). The evaluation of the acceptance ratio was made more efficient by storing the potential energy for each configuration, but because all the Fourier coefficients were generated with each jump attempt, the effective potential and partial average terms also had to be generated. This resulted in slightly longer walk times compared to $J$ walking from quantum distributions. However, since the classical distributions could be generated much more quickly than the quantum distributions, the use of classical distributions was more efficient overall than the use of quantum distributions.

Although useful for heuristic purposes, one-dimensional potentials are of limited use as tests since methods that give good results in one-dimensional systems can fail badly when applied to multidimensional systems. For example, simply using step sizes larger than the barrier width gives good results for the classical one-dimensional double well potential since a relatively large
FIG. 6. Typical FPI $J$-walking results for the average internal energy $\langle U \rangle = \langle V \rangle + \langle T \rangle$ as a function of $\gamma$ for the double well potential for $\beta = 10$. The plots at left are for jumps attempted from classical distributions, while the plots at right are for jumps attempted from quantum distributions. The upper plots are for $m=1$ amu and $k_{\text{max}}=1$, the lower plots for $m=0.1$ and $k_{\text{max}}=4$. As in Fig. 4, the smooth curves represent the numerically exact solutions, while the $J$-walker curves were obtained from 100 randomly initialized walks consisting of $10^4$ warm-up moves followed by $10^4$ moves with data accumulation; partial averaging was used throughout. The $J$-walker distributions used consisted of $10^3$ configurations sampled every ten moves from high temperature Metropolis walks at $\beta=3$. Classical distributions contained configurations obtained from individual classical Metropolis walks, while quantum distributions contained configurations and their associated $k_{\text{max}}$ Fourier coefficients generated from individual Metropolis FPI walks. The jump attempt frequency was $P_J=0.1$.

The number of attempted steps will land close to the other minimum, resulting in higher acceptance rates. The likelihood of landing near a minimum in a multidimensional space with a large random step is quite small though, and so the method is useless for the higher-dimensional spaces typically encountered in most Metropolis simulations. The great advantage of $J$ walking is that the locales near the various potential minima are visited representatively by the
high temperature walker and their corresponding configurations stored for future use. The major limitation in classical systems arises from the acceptance ratio depending exponentially on the difference between the temperature parameters [as is evident in Eq. (42)], so that the likelihood of a jump being accepted decreases for larger temperature differences. For quantum systems, the dependence of the jump acceptance probability on the temperature is more complicated, with \( J \) walking from classical distributions inherently limited at lower temperatures and \( J \) walking from quantum distributions practically limited at low temperatures because of the increased storage requirements with increased \( k_B T \). These limitations did not arise in the double well potential, and both quantum \( J \)-walking schemes gave equally good results, although \( J \) walking from classical distributions was still the preferred method for this system because of the lower storage requirements and shorter computational times required to generate the distributions. The differences between the use of classical and quantum distributions are more evident in multidimensional systems and a more thorough comparison of the two is given in the next section where we apply the methods to atomic clusters.

IV. J WALKING IN QUANTUM CLUSTERS

In our initial study of classical \( J \) walking,\(^1\) we tested the method on rare gas clusters bound by pairwise additive Lennard-Jones potentials. Evidence of quasiergodic behavior had been noted in previous Monte Carlo calculations\(^2\) and molecular dynamics studies,\(^2\) and had been postulated to account for differences between Monte Carlo and molecular dynamics results for \( \text{Ar}_{13} \).\(^3\) The clusters' small sizes and consequent modest storage requirements and computational times made them ideal candidates for a realistic test of \( J \) walking. We found that quasiergodicity was indeed evident in the constant volume heat capacity (and to a lesser extent in the internal energy) over a significant part of the temperature range encompassing the transition from "solid-like" behavior to "liquid-like" behavior,\(^2\) and we showed that \( J \) walking was very successful in eliminating the problem.

The large majority of theoretical studies of the equilibrium and dynamical behavior of clusters to date have been classical. While classical mechanical methods are appropriate for many cluster studies, there are many interesting problems where quantum mechanical effects are significant enough to render classical mechanical studies inadequate.\(^6\) The low temperatures typically encountered in rare gas cluster simulations suggest that quantum effects can be significant in the transition regions. Recent studies\(^6\) on small argon and neon clusters found moderate quantum effects in the cluster melting region for argon and large quantum effects for neon. In particular, the large zero-point fluctuations found for neon make the validity of even a qualitative classical description of the dynamical behavior of small neon clusters suspect. Contrast this with the bulk melting temperature of neon (23.48 K), which is quite close to the value obtained from applying the law of corresponding states to the melting temperature of bulk argon (83.95 K). The recent introduction of quantum simulation methods such as path integral Monte Carlo combined with the increased computational performance of modern workstation computers has made the study of many-body quantum mechanical systems feasible. Thus clusters are also ideal for testing quantum \( J \)-walking methods.

A. Metropolis simulations of classical \( \text{Ne}_7 \) and \( \text{Ar}_7 \)

The comparison of quantum and classical Metropolis calculations for small clusters can help provide insight into the limitations of the classical studies, as well as indicate possible quasiergodic behavior in the quantum systems and so we begin with the classical results. The clusters were modeled by the usual pairwise additive Lennard-Jones potential

\[
V = \sum_{i<j} V_{\text{LJ}}(r_{ij}),
\]

\[
V_{\text{LJ}}(r_{ij}) = 4\epsilon \left[ \left( \frac{\sigma}{r_{ij}} \right)^{12} - \left( \frac{\sigma}{r_{ij}} \right)^{6} \right]
\]

with \( \epsilon = 119.4 \) K and \( \sigma = 3.405 \) Å for argon, and \( \epsilon = 35.60 \) K and \( \sigma = 2.749 \) Å for neon.\(^6\) Small clusters are known to become unstable beyond a threshold temperature \( T_B \) that varies with the cluster size \( n \).\(^2\) For the Lennard-Jones potential under free volume conditions, the average energy vanishes in the limit of an infinite number of configurations. Consequently, the choice of boundary conditions can have a pronounced effect on some of the properties of small clusters.\(^24\) We have followed Lee, Barker, and Abraham\(^3\) and confined the clusters by a perfectly reflecting constraint at \( R_c \) centered on the cluster's center of mass. The constraining radius was set to \( R_c = 4\sigma \) for all runs.

The classical internal energy and heat capacity for an \( n \)-atom cluster are given by

\[
\langle U \rangle = \frac{3n k_B T}{2} + \langle V \rangle,
\]

\[
\langle C_V \rangle = \frac{3n k_B}{2} + \frac{\langle V^2 \rangle - \langle V \rangle^2}{k_B T^2}.
\]

Curves for \( \langle U \rangle \) and \( \langle C_V \rangle \) as functions of \( T \) were generated for argon over the temperature range from \( 1 < T < 60 \) K with a mesh size of \( \Delta T = 1 \) K using the usual Metropolis methods. The lowest energy configuration, the pentagonal bipyramid,\(^31\) was used to initialize the \( T = 1 \) K walk; subsequent temperatures used the final configuration from the preceding walk as their initial configuration. Each data point consisted of \( 10^5 \) warm-up passes followed by \( 10^7 \) passes with data accumulation. Because classical Lennard-Jones clusters obey the principle of corresponding states, results for \( \text{Ne}_7 \) were obtained by simply scaling the \( \text{Ar}_{13} \) results.

B. \( J \) walking for classical \( \text{Ne}_7 \) and \( \text{Ar}_7 \)

As we found in our earlier study of classical \( \text{Ar}_{13} \), a single \( J \)-walker distribution generated at a temperature of
sharp contrast to our earlier study of $\text{Ar}_{13}$, which showed pronounced effects of quasiergodicity in the solid–liquid coexistence region. Quasiergodic effects are evident in the slightly higher temperature range from 20 to 40 K. When we graphically displayed samples of the Metropolis configurations from this region, we also noticed the compact isomers and transition forms appearing in the shoulder region, but in addition, we also saw a large percentage of loose random forms that could no longer be related to recognizable isomers. We also saw occasional dissociations consisting of a compact $\text{Ar}_0$ core and a nearby lone atom. This was not the case in the high temperature side of the $C_V$ peak. Here, both Metropolis and $J$ walking are well converged and in agreement with each other. This temperature range corresponds to the cluster dissociation region and many of the Metropolis configurations that we examined appeared consistent with a dense gas enclosed in a spherical container. The low temperature side of the $C_V$ peak appears then to be another coexistence region containing a wide variety of configurations. The discrepancy between Metropolis and $J$-walking results in this region also occurred in the analogous quantum curves and we present a more detailed examination later when we discuss the quantum results.

C. Metropolis FPI for quantum $\text{Ne}_7$ and $\text{Ar}_7$

The FPI treatment of Lennard-Jones clusters has been detailed before\(^6\) and so we merely recap the pertinent points. The computational overhead relative to analogous classical Metropolis simulations increases rapidly as the number of Fourier coefficients $k_{\text{max}}$ needed for convergence grows. There is a slight increase due to the additional dimensionality associated with the extra Fourier coefficients, but since a typical FPI move consists of a coordinate vector move together with one Fourier coefficient vector move (with a bias favoring the lower order coefficients), the computational cost is only moderate. The major cost is found in the calculation of the action integrals. Using Simpson’s rule to evaluate the integrals, the total computation time has a roughly linear dependence on the number of quadrature points $n_{\text{quad}}$ used. Since the paths corresponding to higher $k_{\text{max}}$ become increasingly irregular,\(^17\) $n_{\text{quad}}$ needs to be increased as the number of Fourier coefficients is increased. We determined the minimum number of quadrature points needed for a given $k_{\text{max}}$ by increasing $n_{\text{quad}}$ until the changes in the average energy and heat capacity were within the standard deviation. The values of $n_{\text{quad}}$ that were found to be sufficient for each $k_{\text{max}}$ were

$$k_{\text{max}}-1, 2, 4, 8, 16, 32, 64,$$

$$n_{\text{quad}}-4, 4, 8, 8, 16, 32, 64.$$  \hfill (64)
Taylor series expansion of the Lennard-Jones potential gives for the partial average terms for Eqs. (31)-(36),

\[ f_{\text{PA}} = g_{\text{PA}} = \frac{1}{2} g(u) \nabla^2 V. \]

Although we found the H-method kinetic energy estimator to be superior to the T-method estimator for the double well potential, we primarily used the T-method estimator for the cluster simulations. The computational requirements for the H method (especially for the heat capacity) were substantially greater than those for the T method, and since values of \( k_{\text{max}} \) of 1 or 2 were found to be adequate for the partial averaged T-method kinetic energy and heat capacity over much of the temperature range investigated for each cluster, the faster convergence of the H method with \( k_{\text{max}} \) was not an issue. Another technical difficulty with the H method that disfavored its use at higher temperatures concerns the constraining potential. Unlike the T-method case, the evaluation of the H-method integrals requires a finite constraining potential, such as a high power polynomial \( V_c(r) = \left(\frac{1}{2} (r/R_{\text{cm}}) / R_{\text{cl}}\right)^{20} \), where \( R_{\text{cm}} \) is the center of mass of the cluster. The evaluation of such a potential adds even more computational overhead. We checked some of our T-method results by comparing them with their H-method counterparts and found excellent agreement.

We obtained FPI Metropolis results for the potential, kinetic, and total energies, and for the heat capacity for Ar\(_7\) over a temperature range from 1 to 60 K on a mesh size of \( \Delta T = 1 \) K, and for Ne\(_7\) over a range of 0.5-20 K on a mesh size of \( \Delta T = 1 \) for higher temperatures and \( \Delta T = 0.5 \) for lower temperatures. Each result was obtained from 10^7 passes, initialized from previously warmed configurations. We ran simulations for each point for \( k_{\text{max}} = 1, 2, 4, \ldots \) until the total energy and heat capacity converged, up to a maximum of \( k_{\text{max}} = 32 \), with the number of quadrature points for each \( k_{\text{max}} \) given by Eq. (64). Attempted moves for each pass were made by randomly displacing an atom’s coordinate vector together with one of its Fourier coefficient vectors, with a bias in favor of the lower order coefficients. Both coordinate and Fourier coefficient displacements were sampled from uniform distributions with box sizes fixed to give approximately 50% acceptance ratios for the combined moves.

### D. FPI J walking for quantum Ne\(_7\) and Ar\(_7\)

We ran quantum J-walking calculations using both quantum and classical distributions for both Ar\(_7\) and Ne\(_7\). The two methods gave statistically identical results for each cluster over the common temperature ranges and they both gave good agreement with FPI Metropolis over most of the temperature range, differing only in the low temperature side of the \( C_V \) peak, as in the classical case. Classical Ar\(_7\) distributions were generated in stages at temperatures of \( T_s = 50, 38, 24 \), and 14 K as described in Sec. IV A, with classical J walking used to generate the lower temperature distributions from the preceding ones. Quantum Ar\(_7\) distributions with increasing \( k_{\text{max}} \) were generated at temperatures of \( T_q = 60, 40, 25, 15 \), and 9 K. Similarly, classical Ne\(_7\) distributions were generated at \( T_c = 15, 11 \), and 8 K, and quantum distributions at \( T_q = 20, 15, 11, 9 \), and 6 K. Table I lists the particulars for each distribution.

The data for each cluster were obtained in the same manner as was the Metropolis FPI data, except that the total walk length was only 10^6 passes instead of 10^7 passes; we found the J-walking data to typically have a much lower variance than the Metropolis data, despite the shorter walk length. The jump attempt probability was the same as in the classical J walking \( P_J = 0.1 \). For J walking from classical distributions, an attempted jump consisted of randomly selecting a cluster coordinate configuration from the distribution and randomly generating each of the Fourier coefficients for each coordinate from a Gaussian distribution. This allowed us to run simulations for each point with a complete \( k_{\text{max}} \) series for full comparison with the Metropolis results. For J walking from quantum distributions, an attempted jump consisted of randomly selecting a cluster configuration containing both the coordinates and their attendant \( k_{\text{max}} \) Fourier coefficients. This required the low temperature walker to also have the same value of \( k_{\text{max}} \) and so the convergence in \( k_{\text{max}} \) could not be compared to Metropolis and J walking using classical distributions without generating separate quantum distributions for each \( k_{\text{max}} \). Because the Metropolis FPI and the J walking from classical distributions showed similar convergence and because generating quantum distributions was very time consuming, we simply took the results obtained using the classical distributions to determine the converged value of \( k_{\text{max}} \) needed for each quantum distribution. We

### Table I. J-walker distributions.

<table>
<thead>
<tr>
<th>( \text{Ne}_7 )</th>
<th>( \text{Ar}_7 )</th>
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<tbody>
<tr>
<td>( T(K) )</td>
<td>( T_c(K) )</td>
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<td>15</td>
<td>15</td>
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<td>11</td>
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<td>14</td>
<td>24</td>
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<tr>
<td>9</td>
<td>16</td>
</tr>
</tbody>
</table>

*The J-walker distributions were each generated from a single walk with configurations stored every 100 passes. Those distributions generated using J walking have their corresponding J-walker temperature listed in the \( T_s \) column (quantum distributions generated with J walking used classical J-walker distributions). The columns labeled Size contain the total number of configurations stored. Classical configurations consisted of coordinates and the cluster potential energy, while quantum configurations consisted of coordinates, Fourier coefficients, and various cluster energies.
FIG. 8. A comparison of FPI J-walking and Metropolis results for the $T$-method internal energy (in reduced units) for quantum Ne$_7$ and Ar$_7$. The open symbols were obtained from FPI Metropolis walks consisting of $10^7$ passes initialized from previously warmed configurations. Single standard deviation error bars have been included for those points where the standard deviation is larger than the symbol size. The curves were generated using J walking from externally stored J-walker distributions with jumps attempted with a frequency $P_{J}=0.1$. The distribution parameters are listed in Table I. The J-walking walk length was only $10^5$ passes. The plot at left is for Ne$_7$ using J walking from quantum distributions; the plot at right is for Ar$_7$ using J walking from classical distributions. For J walking using classical distributions, the values of $k_{\text{max}}$ were the same as for the corresponding FPI Metropolis values, but for J walking using quantum distributions, the values were fixed by the distribution $k_{\text{max}}$ values (listed in Table I).

also checked the sensitivity of J-walking results on the distributions by extending the temperature ranges to overlap with other distributions. For example, the Ar$_7$ $T_{c}=50$ classical distribution had a useful range of $35<T<60$ K, while the $T_{c}=38$ K distribution had a useful range of $20<T<45$ K. Data obtained using each distribution agreed over the common range $35<T<45$ K.

A comparison of J-walking and Metropolis results for the total energy, kinetic energy, and heat capacity is shown in Figs. 8, 9, and 10, respectively. Because Ne$_7$ shows larger quantum effects than Ar$_7$, we have shown Ne$_7$ results obtained using quantum distributions and Ar$_7$ results obtained using classical distributions. The results for Ne$_7$ obtained using classical distributions and for Ar$_7$ obtained using quantum distributions were similar. The plots show very good agreement between J walking and Metropolis over most of the temperature range. As in the cases of classical Ar$_7$ and Ne$_7$, discrepancies occur in the temperature range corresponding to the low temperature side of the heat capacity peak. The Metropolis data in this region is very noisy for both the total energy and the heat capacity, and generally lies above the J-walker curves. In an attempt to account for these discrepancies, we checked the convergence for both classical and quantum Ar$_7$ at 35 K for total walk lengths of $10^4$ to $10^6$ passes, using a classical distribution at $T_{c}=38$ K for each. The results are summarized in Table II. For both the classical and quantum J-walking results, the heat capacity and energy both show the $1/N$ decrease in the standard deviation expected for asymptotically convergent stochastic processes, while the Metropolis results are clearly not converged. Quantum J-walking results obtained using a quantum distribution at $T_{q}=40$ K with $k_{\text{max}}=1$ gave good agreement with the results obtained using the classical distribution. These results also agree with J-walking results obtained using the $T_{c}=50$ K classical distribution. A similar analysis of quantum Ne$_7$ at $T=10$ K gave similar results—the Metropolis results were noisy and unconverged, while the J-walker results from both classical and quantum distributions at different temperatures were properly converged and gave good agreement with each other.

The $T$-method kinetic energies obtained using Metropolis did not show the same convergence difficulties, indicating the problem is isolated mostly in the potential energy. There is a slight discrepancy between Ne$_7$ Metropolis and J-walking results at about 9–11 K, indicating perhaps some effects of quasiergodicity in the kinetic energy as well, but there could also be effects arising from incomplete convergence in $k_{\text{max}}$. Different properties converge with $k_{\text{max}}$ at different rates, and the kinetic energy is more slowly con-
### Table II. Metropolis and J walker convergence for Ar, at $T=35$ K.

<table>
<thead>
<tr>
<th>Passes</th>
<th>Metropolis $\langle U \rangle$</th>
<th>J walker $\langle U \rangle$</th>
<th>Metropolis $\langle C_p \rangle$</th>
<th>J walker $\langle C_p \rangle$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10^4$</td>
<td>$-9.1136 \pm 0.1325$</td>
<td>$8.3301 \pm 0.0220$</td>
<td>$26.44 \pm 1.54$</td>
<td>$51.86 \pm 1.62$</td>
</tr>
<tr>
<td>$10^5$</td>
<td>$-8.3772 \pm 0.2277$</td>
<td>$-8.3416 \pm 0.0118$</td>
<td>$40.80 \pm 3.96$</td>
<td>$54.77 \pm 0.63$</td>
</tr>
<tr>
<td>$10^6$</td>
<td>$-8.4370 \pm 0.1147$</td>
<td>$-8.3317 \pm 0.0047$</td>
<td>$50.22 \pm 5.87$</td>
<td>$53.94 \pm 0.20$</td>
</tr>
<tr>
<td>$10^7$</td>
<td>$-8.0757 \pm 0.2519$</td>
<td>$-8.3312 \pm 0.0011$</td>
<td>$57.05 \pm 3.66$</td>
<td>$53.65 \pm 0.14$</td>
</tr>
<tr>
<td>$10^8$</td>
<td>$-8.0405 \pm 0.1426$</td>
<td>$-8.3350 \pm 0.0007$</td>
<td>$65.14 \pm 5.31$</td>
<td>$54.62 \pm 0.04$</td>
</tr>
<tr>
<td>$10^9$</td>
<td>$-7.7926 \pm 0.0549$</td>
<td></td>
<td>$76.49 \pm 2.75$</td>
<td></td>
</tr>
</tbody>
</table>

The $J$-walker distribution consisted of $5 \times 10^5$ classical configurations sampled every 100 passes from a Metropolis walk at $T_0=38$ K. For quantum Ar, $J$-walking results obtained using a quantum $J$-walker distribution of the same size at $T_0=40$ K gave similar results.

The $J$-walker curve was fixed at $k_{\text{max}}=2$ because the quantum distribution had been generated at that value (which was adequate for the total energy and heat capacity over that temperature range). The $k_{\text{max}}=4$ J-walking results obtained using a classical distribution gave good agreement with the Metropolis results.

Figure 11 compares the jump acceptance rates for $J$ walking using quantum and classical distributions. For $J$ walking from quantum distributions, the jump acceptance for each distribution decreases from 100% (for $T=T_q$), approaching 0% as the temperature difference becomes large. The steepness of the decrease is greater at lower temperatures, reflecting the narrowing of the distributions with decreasing temperature. The useful range of the $T_q=15$ K distribution ($k_{\text{max}}=1$) is from 15 down to about 10 K, while the useful range of the $T_q=6$ K distribution ($k_{\text{max}}=8$) is only from 6 to 4 K. In addition, the number of Fourier coefficients required for convergence increases rapidly at low temperatures, further limiting the useful temperature range of a low quantum $J$-walker distribution since the low temperature walker $k_{\text{max}}$ is fixed at the same value as the $J$-walker distribution. Because the distribution size and the time required to generate the distribution grow rapidly with increasing $k_{\text{max}}$, $J$ walking using quantum distributions becomes practically unfeasible at very low temperatures. For $J$ walking from classical distributions, the maximum jump acceptance for a given distribution was less than 100%, even when $T-T_0$ since the classical configurations and generated Fourier coefficients are different than the corresponding quantum configurations. As the temperature becomes lower and the system becomes more quantum, the difference becomes larger. The $T_c=15$ K classical distribution had a useful temperature range of about 20–11 K, with a maximum jump acceptance of about 62%, but the $T_c=11$ K distribution had a useful range of 12–8 K with a maximum jump acceptance of only 7%. Hence classical distributions have fundamental limitations at low temperatures where the Fourier coefficients become strongly coupled to the coordinates. Classical distributions have the advantage of allowing variable $k_{\text{max}}$. Although only curves for converged values of $k_{\text{max}}$ are shown in Fig. 11, the jump acceptance did not vary much with $k_{\text{max}}$.

Figure 12 compares the quantum and classical energies and heat capacities for Ne and Ar. The curves are comprised mostly of $J$-walking data, with Metropolis results...
used for those temperature regions where \( J \)-walking data had not been obtained. The quantum neon \( C_V \) curve is shifted substantially to lower temperatures, with the peak much reduced relative to the classical curve, and the \( U \) curve is much higher than the classical energy throughout. This indicates that quantum effects for Ne, are considerable over the entire temperature range, including the cluster dissociation region. The shoulder in the classical \( C_V \) curve corresponding to the liquid solid coexistence region is completely absent in the quantum curve. This is a consequence of the quantum zero-point motion and tunneling, which effectively raise the minima and lower the saddles of the multidimensional potential surface and allow for easier isomerization.\(^{26}\) These results are consistent with classical and quantum quench studies,\(^{27}\) which showed significant differences between classical and quantum quenched isomer population distributions. The shoulder is also largely diminished in the quantum Ar, \( C_V \) curve, and there is a small, but significant difference between the classical and quantum argon curves for higher temperatures up to the cluster dissociation region. Clearly, it is necessary to incorporate quantum effects for a proper understanding of dynamical and equilibrium behavior of small argon and neon clusters.

V. DISCUSSION

Path integral methods form an important set of tools for extending Monte Carlo methods to quantum many-body systems. Because of the greater computational requirements incurred by quantum simulations relative to similar classical Metropolis simulations, ensuring proper convergence and developing methods that increase the rate of convergence are especially important. Two sources of slow convergence are inherent in FPI Metropolis simulations. The first is due to the increased number of Fourier coefficients needed for highly quantum systems at low temperatures. This increases the effective dimensionality of the simulation, as well as the computational overhead associated with the numerical calculation of the action integrals. The effect is purely quantum; classical simulations converge more quickly as the temperature becomes very low, but quantum simulations become increasingly difficult. The second source of slow convergence is due primarily to bottlenecks in configurational space and is common to classical simulations as well. While the effects of quasiergodicity in quantum systems appear to be diminished somewhat in comparison to analogous classical systems, they can still be quite formidable. Quantum \( J \)-walking is a very useful technique for substantially reducing errors arising from quasiergodicity and increasing the rate of convergence, thus extending the power and scope of FPI methods. The method is related closely to classical \( J \)-walking and shares many of its features. \( J \)-walking is essentially a variant of the usual Metropolis algorithm, and so is very easy to incorporate into existing code. Quantum \( J \)-walking is slightly more complicated to implement than classical \( J \)-walking, but it offers more flexibility. Two variations—\( J \) walking using fully quantum distributions and \( J \) walking using classical coordinate distributions together with Gaussian distributed Fourier coefficients—were shown to be successful over temperature ranges where both quasiergodicity was evident and quantum effects were substantial. Both variations are completely compatible with various FPI techniques such as partial averaging, as well as the many kinetic energy and heat capacity estimators that have been developed. Although \( J \)-walking showed better convergence than FPI Metropolis even at lower temperatures where quasiergodicity was not an issue, its limitations at very low temperatures preclude its general use in this region for overcoming the slow convergence that is a consequence of including large numbers of Fourier coefficients; the method is best suited for handling quasiergodicity in configurational space (these limitations are inherent in scalar computers—this would not be the case with parallel computers).

Because of the close kinship between quantum and classical \( J \)-walking, many of the caveats we originally discussed remain valid. Care must be taken in properly selecting the \( J \)-walking temperature. It must be high enough to ensure a fully ergodic distribution, but low enough to provide sufficient jump acceptance rates. For the multidimensional systems we have examined to date, both criteria cannot be met with a single distribution and multistage distributions generated at several temperatures over the required range are necessary. Quasiergodicity in the lower temperature distributions can be eliminated with \( J \)-walking, using the previously generated distributions for sampling. The walks used to generate the distributions need to be sufficiently lengthy to obtain representative samples of the configuration and Fourier coefficient spaces. This is especially important if the \( J \)-walker distribution is generated at a temperature where quasiergodicity is substantial. The results obtained using such distributions can be checked for consistency by generating additional distributions at the same temperature or at nearby temperatures. Quantum \( J \)-walking results can be checked further by us-

![Graphs showing energy and heat capacity for Ne and Ar](image-url)
ing both quantum and classical distributions for sampling.

To keep distribution sizes within the limitations imposed by finite computer resources, 33 configurations can be sampled periodically, say every ten or 100 passes. This also helps reduce correlations in the distributions. Since most computers have much larger disk storage than memory storage, the large distributions can be stored as a collection of several files, each of which can then be loaded randomly into the computer's primary memory in a periodic manner as required. Writing the files in a parallel manner while the distribution is being generated also helps reduce the correlations within each file. Optical storage devices with capacities of hundreds of megabytes are well suited for J walking. Fully read-writeable devices are commercially available at prices roughly comparable to fixed magnetic disk drives and with access times only slightly slower. 34 Although each cartridge can typically store only a few hundred megabytes per side, very large distributions can be stored on several cartridges, greatly increasing the effective storage capacity of the system. At present, we still do not know the necessary criteria for determining the minimum size required for a given distribution to ensure that it is a representative sample of the configuration space. Consequently, we have tended to make the distributions as large as possible to minimize systematic errors due to unrepresentative distributions. 35

For the systems we examined, J walking using classical distributions was preferable to the use of quantum distributions. These distributions could be generated much more quickly than their quantum counterparts, and since they consumed far less storage per configuration, we could include more configurations in each distribution. The classical distributions can of course be used for J-walking studies of the corresponding classical systems, allowing for comparison between the classical and quantum systems. J walking from classical distributions also has the advantage of being capable of handling arbitrary \( k_{\text{max}} \) so that \( k_{\text{max}} \) convergence can be checked easily for each temperature. J walking from quantum distributions fixes the low-temperature walker's \( k_{\text{max}} \) value to the distribution's value, requiring that the converged value of \( k_{\text{max}} \) over the useful temperature range of the distribution be ascertained before the distribution is generated. Classical distributions are more limited than quantum distributions for handling very low temperatures. For example, for Ne, classical distributions were useful down to about 8 K, while quantum distributions were useful down to 4 K. However, much of the quasiergodic behavior evident in multiparticle systems such as clusters is a consequence of bottlenecks in the coordinate subspace and becomes evident only at higher energies. Consequently, the corresponding temperatures are usually high enough to allow the use of classical distributions. Even in the case of Ne, where quantum effects were quite large over much of the temperature range, J walking using classical distributions was just as effective as J walking using quantum distributions in eliminating quasiergodicity. The use of quantum distributions is generally preferable for handling very low temperatures or for providing an independent check of the results obtained from classical distributions. For Ne, J walking using quantum distributions gave good results for the solid-liquid coexistence region (corresponding to the shoulder in the \( C_v \) curve), a temperature range not accessible using classical distributions, but this was also a region where standard FPI Metropolis gave sufficiently good results.

**ACKNOWLEDGMENTS**

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13. In contrast to classical Monte Carlo methods that are applicable to a variety of ensembles, path integral methods have primarily used the canonical ensemble because of the formal correspondence between the quantum density matrix and the quantum propagator.
27. S. W. Rick, D. L. Leitner, J. D. Doll, D. L. Freeman, and D. D. Frantz,


23 All calculations were performed on DEC3100 and DEC5000 UNIX workstations.
25 Distribution sizes typically totalled 100 to 200 Mb and were stored in 50 separate files.