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Joseph E. Lawrence, and David E. Manolopoulos

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Joseph E. Lawrence and David E. Manolopoulos

AFFILIATIONS
Physical and Theoretical Chemistry Laboratory, Department of Chemistry, University of Oxford, South Parks Road, Oxford OX1 3QZ, United Kingdom

Email: joseph.lawrence@chem.ox.ac.uk

ABSTRACT
We analyze the golden rule limit of the recently proposed isomorphic ring polymer (iso-RP) method. This method aims to combine an exact expression for the quantum mechanical partition function of a system with multiple electronic states with a pre-existing mixed quantum-classical (MQC) dynamics approximation, such as fewest switches surface hopping. Since the choice of the MQC method adds a degree of flexibility, we simplify the analysis by assuming that the dynamics used correctly reproduces the exact golden rule rate for a nonadiabatic (e.g., electron transfer) reaction in the high temperature limit. Having made this assumption, we obtain an expression for the iso-RP rate in the golden rule limit that is valid at any temperature. We then compare this rate with the exact rate for a series of simple spin-boson models. We find that the iso-RP method does not correctly predict how nuclear quantum effects affect the reaction rate in the golden rule limit. Most notably, it does not capture the quantum asymmetry in a conventional (Marcus) plot of the logarithm of the reaction rate against the thermodynamic driving force, and it also significantly overestimates the correct quantum mechanical golden rule rate for activationless electron transfer reactions. These results are analyzed and their implications discussed for the applicability of the iso-RP method to more general nonadiabatic reactions.

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I. INTRODUCTION

The ability to simultaneously treat non-Born-Oppenheimer dynamics and nuclear quantum effects in atomistic simulations is important in a wide variety of contexts, ranging from proton coupled electron transfer reactions in proteins to energy transport in photovoltaics. Recently, Tao, Shushkov, and Miller have proposed a new method, called isomorphic ring polymer molecular dynamics (iso-RPMD or iso-RP),1,2 which aims to extend the well known Born-Oppenheimer ring polymer molecular dynamics (RPMD) method3–8 to systems with multiple electronic states. The iso-RP ansatz is to construct an “isomorphic” multistate Hamiltonian for a ring polymer of classical nuclei that is consistent with the quantum mechanical partition function, and then to combine this with a pre-existing mixed quantum-classical (MQC) dynamics approximation such as Tully’s fewest switches surface hopping.

One potential application for iso-RPMD is the study of thermal reactions that involve transitions between two different electronic states.9 The most widely used theory for describing these reactions is Marcus theory,10 which assumes that the nuclear degrees of freedom in each state can be modeled with classical harmonic free energy surfaces and that the transitions between the states can be treated using perturbation theory (Fermi’s golden rule). Marcus theory often works well at high temperatures, but it fails to capture nuclear quantum effects, such as tunneling and zero-point energy, which become important at lower temperatures. These nuclear quantum effects can be included in simple analytical theories,11 but in order to understand real chemical systems, it is desirable to have a simulation method that includes nuclear quantum effects and is capable of treating fully atomistic models of multidimensional systems with anharmonicity. One might hope that iso-RPMD would provide a practical way to do this.

In the adiabatic limit, in which excited electronic states do not make any contribution to the canonical quantum mechanical partition function, it is clear that iso-RPMD will work well, since it reduces in this limit to the standard RPMD approximation (in which
the dynamics of the ring polymer takes place on the ground adiabatic Born–Oppenheimer potential energy surface). However, how well it can be expected to work in the opposite nonadiabatic limit, in which the rate of a transition between two diabatic electronic states is governed by the strength of the perturbative electronic coupling between these states as it is in Marcus theory, is less clear. This is what we shall investigate in the present paper.

II. ISOMORPHIC RPMD

To set the scene for this investigation, let us begin by introducing the iso-RPMD formalism.\textsuperscript{1,2} Although this formalism is in principle applicable to systems with many electronic states, it will be sufficient for our purposes to consider just a two level system, for which the Hamiltonian can be written in the diabatic representation as

\[ \hat{H} = \sum_{i=1}^{f} \frac{\hat{p}_i^2}{2m_{i}} + V(\hat{q}), \]  

(1)

where \( 1 \) is the \( 2 \times 2 \) identity matrix and \( V(\hat{q}) \) is the diabatic potential matrix,

\[ V(\hat{q}) = \begin{pmatrix} V_{1}(\hat{q}) & \Delta(\hat{q}) \\ \Delta(\hat{q}) & V_{2}(\hat{q}) \end{pmatrix}. \]

(2)

The “isomorphic” RPMD for this two level system is based on writing the exact quantum partition function

\[ Q = \text{Tr}[e^{-\beta \hat{H}}] \]

in the form

\[ Q = \lim_{n \to \infty} \left( \frac{n}{2\pi \hbar} \right)^{n} \int d^{n}q \int d^{n}p \text{tr}_e[\hat{e}^{-\beta n \hat{H}_e(p,q)}]. \]

(4)

where \( \text{tr}_e[\cdots] \) denotes a trace over the electronic degrees of freedom. The isomorphic ring polymer Hamiltonian \( \hat{H}_n^{iso}(p,q) \) is

\[ \hat{H}_n^{iso}(p,q) = h_n(p,q) 1 + V_n^{iso}(q), \]

(5)

where

\[ h_n(p,q) = \sum_{j=1}^{n} \sum_{\nu=1}^{f} \left\{ \frac{\hat{p}_j^2}{2m_{\nu}} + \frac{1}{2} m_{\nu} \omega_{n}^2 (q_{j+1}-q_{j})^2 \right\}. \]

(6)

is the free ring polymer Hamiltonian, with \( m_{\nu} = m_{\nu}/n, \omega_{n} = n/\beta h \), and \( q_{j+1} = q_{j+1} \). The isomorphic diabatic potential energy matrix is

\[ V_n^{iso}(q) = \begin{pmatrix} V_{1n}^{iso}(q) & \Delta_n^{iso}(q) \\ \Delta_n^{iso}(q) & V_{2n}^{iso}(q) \end{pmatrix}. \]

(7)

where the diagonal elements are chosen to be

\[ V_{ii}^{iso}(q) = \frac{1}{n} \sum_{j=1}^{n} V_{i}(q_{j}). \]

(8)

Here, \( q = (q_{1}, q_{2}, \ldots, q_{n}) \) is a vector whose elements are the position vectors for each ring polymer bead and \( p = (p_{1}, p_{2}, \ldots, p_{n}) \) are the conjugate momenta.

In order to ensure that the iso-RP partition function exactly reproduces the quantum partition function in the infinite \( n \) limit, Tao et al.\textsuperscript{1} required that \( \Delta_n^{iso}(q) \) is chosen so that

\[ \text{tr}_e[e^{-\beta h_n(p,q)}] = e^{-\beta h_n(p,q)} \mu(q), \]

(9)

where

\[ \mu(q) = \text{tr}_e \left[ \prod_{\nu=1}^{n} e^{-\beta h_{\nu}(q)} \right]. \]

(10)

with \( \beta_n = \beta/n \). Here, \( \mu(q) \) orders the matrix product in order of increasing bead index \( j \), in accordance with a standard path integral discretization of Eq. (3). This leads to the following expression for the isomorphic coupling:

\[ \Delta_n^{iso}(q)^{2} = \cosh \left( \frac{\beta_n(q) e^{V_{1n}^{iso}(q)}/2}{\beta_n} - V_{2n}^{iso}(q)/2 \right)/4, \]

(11)

where

\[ V_{\pm n}^{iso}(q) \]

(12)

Note that in general \( \mu(q) \) can become negative,\textsuperscript{13} which leads to a complex \( \Delta_n^{iso}(q) \) and a non-Hermitian \( V_{n}^{iso}(q) \). However, when \( \Delta_n^{iso}(q) \) has a constant sign, one can show that \( \mu(q) > 0 \), so \( \Delta_n^{iso}(q) \) is real and \( V_{n}^{iso}(q) \) is real and symmetric for all values of \( q \). We shall therefore assume that \( \Delta(q) \) has a constant sign from this point on.

All of the above is simply a rearrangement of a formally exact expression for the quantum mechanical partition function. The isomorphic RPMD ansatz is to take the classical multistate Hamiltonian \( \hat{H}_n^{iso}(p,q) \) and use it with a mixed quantum classical (MQC) dynamics method, such as Tully’s fewest switches surface hopping. This is an appealingly simple idea, but how well does it work? In the following, we will consider the more specific question of whether or not iso-RPMD can give accurate reaction rates in the nonadiabatic (golden rule) limit, and in particular, whether it can capture nuclear quantum effects such as tunneling and zero point energy in this limit.

III. GOLDEN RULE LIMIT

The exact quantum mechanical rate constant for transfer from state \( |0\rangle \) to state \( |1\rangle \) in the golden rule limit can be written as\textsuperscript{1}

\[ k = \frac{1}{Q_{1}} \int_{0}^{\infty} \sigma(c) d\tau, \]

(13)

where \( Q_{i} \) is the reactant partition function and

\[ \sigma(c) = \text{tr}_e \left[ e^{-\beta h_{1}\hat{e}^{-\beta h_{1}}}(\Delta(\hat{q}) e^{\beta h_{1}})(\Delta(\hat{q})). \]

(14)

Here,

\[ \hat{H}_{i} = \sum_{j=1}^{f} \frac{\hat{p}_j^2}{2m_{j}} + V_{i}(\hat{q}) \]

(15)

is the nuclear diabatic Hamiltonian on state \( i \) and \( \text{tr}_e[\cdots] \) denotes a trace over the nuclear coordinates.

The classical limit of Eq. (13), which becomes exact at high temperatures where the ring polymer can be replaced by a single bead, is simply\textsuperscript{15}
\[ k_{\text{id}} = \left( \frac{2\pi}{\hbar} \right) \bar{A}(q)^2 \delta(V_-(q)) \Bigg|_0, \tag{16} \]

where \( V_-(q) = V_0(q) - V_1(q) \) is the diabatic energy gap and the thermal average is defined as

\[ \langle A(q) \rangle_0 = \frac{\int d^4p \int d^4q e^{-\beta h(p,q)} A(q)}{\int d^4p \int d^4q e^{-\beta h(p,q)}}. \tag{17} \]

Unfortunately, it is well known that many MQC dynamics methods do not give the correct rate in the high-temperature, nonadiabatic limit [i.e., are not consistent with Eq. (16)]. In particular, the original formulation of Tully’s fewest switches surface hopping suffers from “over coherence.” There has been a concerted effort to fix this problem with decoherence corrections, \[^{16–24}\] but that is not the focus of the present study. Here, we shall simply sidestep the issue by assuming that the MQC method used with iso-RPMD is consistent with Eq. (16) in the one bead limit.

Having made this assumption, we need not consider the specific details of the MQC dynamics and can simply make use of the isomorphism to write down the iso-RP expression for the golden rule rate,

\[ k_{\text{iso}} = \lim_{n \to \infty} \left( \frac{2\pi}{\hbar} \right) \bar{A}(q)^2 \delta(V_{\text{iso}}(q)) \Bigg|_{0,n}, \tag{18} \]

where the ring polymer thermal average is defined as

\[ \langle A(q) \rangle_{0,n} = \frac{\int d^4p \int d^4q e^{-\beta h(p,q)} A(q)}{\int d^4p \int d^4q e^{-\beta h(p,q)}}. \tag{19} \]

with \( H_{0,n}(p,q) = h_n(p,q) + V_{\text{iso}}(q) \).

We can further simplify Eq. (18) by taking the golden rule limit of the iso-RP coupling \( \Delta h_{\text{iso}}(q)^2 \). To keep the notation as simple as possible, we shall do this only in the special case where \( \Delta A(q) = \Delta \) is a constant (the Condon approximation). The more general case of a coordinate-dependent \( \Delta \) can also be treated, but it does not add anything useful to our discussion. The golden rule limit of \( \Delta h_{\text{iso}}(q)^2 \) is worked out for constant \( \Delta \) in the Appendix, where it is shown to lead to a golden rule iso-RP rate in the infinite \( n \) limit that can be written in path integral notation as

\[ k_{\text{iso}} = \frac{2\pi}{\sqrt{\hbar Q \tau}} \int_0^\beta d\lambda \bar{A}(\lambda)^2 \delta(V_{\text{iso}}(q(\tau)))) e^{-\beta \lambda^2}. \tag{20} \]

Here, the \( \lambda \)-dependent path-integral expectation value is defined as

\[ \langle A(q(\tau)) \rangle_\lambda = \frac{\int \mathcal{D}q(\tau) e^{-\beta S(\tau)(q(\tau))} A[q(\tau)]}{\int \mathcal{D}q(\tau) e^{-\beta S(\tau)(q(\tau))}}. \tag{21} \]

with the action

\[ S^{(1)} = S^{(1)} + S^{(1)}_1. \tag{22} \]

\[ S^{(1)} = \int_0^\beta \frac{1}{2} \sum_{j=1}^2 m_j \ddot{q}_j + V_0(q(\tau)) d\tau, \tag{23} \]

\[ S^{(1)}_1 = \int_0^\beta \frac{1}{2} \sum_{j=1}^2 m_j \dot{q}_j^2 + V_1(q(\tau)) d\tau, \tag{24} \]

and the \( \lambda \)-dependent Boltzmann factor is

\[ e^{-\beta \lambda^2} = \int \frac{\mathcal{D}q(\tau) e^{-\beta S(\tau)(q(\tau))}}{\mathcal{D}q(\tau) e^{-\beta S(\tau)(q(\tau))}}. \tag{25} \]

The isomorphic diabatic energy gap in Eq. (19) can be written in the same notation as

\[ V_{\text{iso}}(q(\tau)) = V_0^{\text{iso}}(q(\tau)) - V_1^{\text{iso}}(q(\tau)). \tag{26} \]

again with constant \( \Delta \). The effect of the nuclear motion on the electronic dynamics of this model (and hence the rate of transfer from state 0 to state 1) is fully characterized by the spectral density,

\[ J(\omega) = \frac{\pi}{2} \sum_{i=1}^\infty \frac{\delta(\omega - \omega_i)}{2} \tag{30} \]

in which we have chosen to work in mass-scaled coordinates such that \( m_j = 1 \).

Due to the simplicity of the problem, it is possible to obtain closed form expressions for each of the terms appearing in Eq. (20).

The \( \lambda \)-dependent Boltzmann factor can be written as

\[ e^{-\beta \lambda^2} = e^{-\beta \lambda^2} + \lambda \beta, \tag{31} \]

with

\[ F(\lambda) = \frac{\lambda}{2} \int_0^\beta d\lambda \left( 1 - \frac{\cosh(\omega \beta)}{\sinh(\omega \beta/2)} \right) \omega. \tag{32} \]

Furthermore, using the fact that \( V_{\text{iso}}(q(\tau)) \) only depends on the average (centroid) of the imaginary time path, \( q = \frac{\hbar}{\beta} \int_0^\beta q(\tau) d\tau \), it is straightforward to show that

\[ \left\{ \delta(V_{\text{iso}}(q(\tau))) \right\}_\lambda = \frac{1}{2} \int_0^\beta \frac{\beta^2 \lambda^2}{\pi^2} e^{-\beta \lambda^2} \Delta q(1/2) d\lambda, \tag{33} \]

where

\[ \lambda = \frac{1}{\pi} \int_0^\infty f(\omega) d\omega \tag{34} \]

is the Marcus theory reorganization energy.
Note that in the high temperature limit, $F(\lambda) \rightarrow \Lambda^2 \left(1 - \frac{1}{\beta} \right)$, and hence, the $\lambda$ dependence of Eq. (31) exactly cancels the $\Lambda$ dependence of Eq. (35). In this limit, the isomorphic rate correctly reduces to the rate given by Marcus theory: $k_{iso} \rightarrow k_{MT}$ as $\beta \rightarrow 0$, where

$$k_{MT} = \frac{\Delta^2}{h} \sqrt{\frac{\beta \epsilon}{\Lambda}} e^{-\frac{\Delta^2}{\beta \Lambda} - \frac{\beta \epsilon}{\Lambda}}. \tag{35}$$

Unfortunately, however, it is already clear that there is an issue with the iso-RP rate at lower temperatures. Since $F(\lambda)$ is independent of $\epsilon$, we see that the iso-RP rate retains the Marcus theory dependence on $\epsilon$ at all temperatures. Hence, the ratio of the iso-RP rate to the Marcus rate is independent of the driving force, $k_{iso}/k_{MT} = \Lambda$, and so, on a plot of the logarithm of the rate as a function of the driving force, the iso-RP rate is simply a vertically shifted version of the famous Marcus parabola. This clearly does not capture the behavior of the correct quantum mechanical rate at low temperatures.

To demonstrate this, we shall consider two models with Brownian oscillator spectral densities,

$$J(\omega) = \frac{\Lambda}{2} \frac{\gamma \Omega^2 \omega}{(\omega^2 - \Omega^2)^2 + \gamma^2 \omega^2}, \tag{36}$$

and with parameters chosen to be representative of typical condensed phase electron transfer reactions. Model 1 is weakly quantum mechanical with $\beta \Omega = 2$, $\gamma = \Omega$, and $\beta \Lambda = 60$, corresponding to a reaction coordinate frequency of $\Omega \approx 400$ cm$^{-1}$ and a Marcus reorganization energy of $\Lambda \approx 1.5$ eV at 300 K. Model 2 is more strongly quantum mechanical with $\beta \Omega = 6$, $\gamma = \Omega$, and $\beta \Lambda = 60$, corresponding to a reaction coordinate frequency of $\Omega = 1200$ cm$^{-1}$. These parameters are roughly comparable to those obtained in the classic experimental paper in the Marcus inverted regime by Miller et al., who extracted 1500 cm$^{-1}$ as the characteristic inner sphere frequency, along with a total reorganization energy of 1.2 eV, for a series of organic electron transfers in tetrahydrofuran. The approximate 40%–60% split of the reorganization energy into inner-sphere and outer-sphere contributions they obtained experimentally is also broadly compatible with the Brownian oscillator spectral density for model 2, in which 60% of the reorganization energy comes from modes with frequencies less than $\omega \approx 1000$ cm$^{-1}$ at 300 K.

Figure 1 compares the iso-RP rate for model 1 as a function of the driving force, $\epsilon$, with both the exact quantum mechanical rate and the classical rate (given by Marcus theory). The isomorphic rate is unable to capture the dependence of the exact rate on the driving force, retaining the parabolic form seen in Marcus theory and failing to reproduce the large increase in the rate in the inverted regime due to nuclear quantum effects. The largest error is in the inverted regime: when the bias is twice the reorganization energy, the iso-RP rate is too small by nearly 2 orders of magnitude. In contrast, it is reasonably accurate in the case of zero bias ($\epsilon = 0$), where $k_{iso}/k_{MT} = 1.7$ compared to $k/k_{MT} = 2.6$ for the exact rate. Perhaps the most worrying aspect of the results in Fig. 1, however, is that the iso-RP rate seems to be significantly less accurate than the Marcus rate in the activationless case ($\epsilon = 0$), where it predicts a quantum enhancement, whereas the exact rate is actually slightly smaller than the Marcus rate.

The overestimation of the rate in the activationless case is even more striking in model 2, as shown in Fig. 2. Here, the iso-RP rate is over 30 times larger than the exact rate at $\epsilon = \Lambda$. Moreover, the error is again largest deep in the inverted regime, with the iso-RP rate underestimating the exact rate by nearly 3 orders of magnitude. In contrast, it is reasonably accurate in the case of zero bias ($\epsilon = 0$), where $k_{iso}/k_{MT} = 1.7$ compared to $k/k_{MT} = 2.6$ for the exact rate. Perhaps the most worrying aspect of the results in Fig. 1, however, is that the iso-RP rate seems to be significantly less accurate than the Marcus rate in the activationless case ($\epsilon = 0$), where it predicts a quantum enhancement, whereas the exact rate is actually slightly smaller than the Marcus rate.

V. ANALYSIS

To understand the behavior of the iso-RP rate, it is helpful to consider another approximate method for calculating reaction
rates in the golden rule limit, namely, Wolynes theory.\textsuperscript{14} This theory makes a steepest descent approximation to the time integral in Eq. (13) to give

\[ k_{WT} = \frac{\Delta^2}{Q_0 \hbar} \frac{2\pi}{-\beta \epsilon(\lambda_{sp})} e^{-\beta \epsilon(\lambda_{sp})}, \]  

(37)

where the saddle point condition is \( \epsilon'(\lambda_{sp}) = 0 \). Wolynes' theory is well known to be very accurate for the spin-boson model. It correctly recovers the Marcus theory rate in the high-temperature limit (\( k_{WT} \rightarrow k_{SM} \) as \( \beta \rightarrow 0 \)), and it is also remarkably accurate at lower temperatures, where the path-integral distribution corresponding to \( e^{-\beta \epsilon(\lambda_{sp})} \) is dominated by imaginary time paths close to the golden-rule instanton.\textsuperscript{3} For example, for our strongly quantum mechanical model 2 at zero bias, \( k_{WT}/k = 0.98 \), and in the activationless case \( k_{WT}/k = 1.00 \).

It is clear that one of the main differences between \( k_{WT} \) and \( k_{iso} \) is that in Wolynes theory, the Boltzmann factor \( e^{-\beta \epsilon(\lambda)} \) is evaluated at \( \lambda = \lambda_{sp} \), whereas the iso-RP rate [Eq. (20)] involves an integral over all values of \( \lambda \) in the interval \([0, \beta]\). This difference can be used to explain the behavior of the iso-RP rate. We begin by noting that the (\( \epsilon \) independent) ratio of the iso-RP rate to the Marcus theory rate is given by

\[ \frac{k_{iso}}{k_{MT}} = \frac{1}{\beta} \int_0^\beta d\lambda \frac{e^{-\beta \epsilon(\lambda)}}{e^{-\beta \epsilon(\lambda_{sp})}}, \]  

(38)

where \( \tilde{F}_{MT}(\lambda) = \Lambda \beta \left(1 - \frac{1}{\beta}\right) \) is the high temperature limit of \( \tilde{F}(\lambda) \).

Then, since \( k_{WT} \rightarrow k_{SM} \) as \( \beta \rightarrow 0 \), we can write

\[ \frac{k_{WT}(\epsilon)}{k_{MT}(\epsilon)} = \left(\frac{F''_{MT}(\lambda_{MT}(\epsilon))}{F''(\lambda_{sp})}\right) e^{-\beta \epsilon(\lambda_{sp})} e^{\beta \epsilon(\lambda_{MT}(\epsilon))}, \]  

(39)

where

\[ \lambda_{MT}(\epsilon) = \frac{\beta(\Lambda - \epsilon)}{2\Lambda}. \]  

(40)

is the high temperature limit of \( \lambda_{sp} \). Now, assuming that both \( \lambda_{sp}(\epsilon) \approx \lambda_{MT}(\epsilon) \) and \( F''(\lambda_{sp}) \approx F''_{MT}(\lambda_{MT}(\epsilon)) \) continue to hold at lower temperatures (at least approximately), we see that

\[ e^{-\beta \epsilon(\lambda_{MT}(\epsilon))} \approx \frac{k_{WT}(\epsilon)}{k_{MT}(\epsilon)} \approx \frac{k(\epsilon)}{k_{iso}(\epsilon)}. \]  

(41)

Finally, making the substitution \( \lambda = \lambda_{MT}(\epsilon) \) to convert the integration variable from \( \lambda \) to \( \epsilon \) in Eq. (38) and using the approximation in Eq. (41), we can rewrite the quantum enhancement in the rate predicted by iso-RPMD as

\[ \frac{k_{iso}}{k_{MT}} \approx \frac{1}{2\Lambda} \int_{-\Lambda}^\Lambda d\epsilon \frac{k(\epsilon)}{k_{MT}(\epsilon)}. \]  

(42)

For both models considered above (models 1 and 2), this approximation is within 1% of the actual ratio, indicating that the assumptions in Eq. (41) are valid and that this does indeed provide a useful way to understand the behavior of the iso-RP rate.

From Eq. (42), we see that the quantum enhancement predicted by iso-RPMD can be interpreted as the average of the true quantum enhancement over the entire Marcus normal regime, \( \epsilon \in [-\Lambda, \Lambda] \). This can be used to understand why, for example, iso-RPMD underestimates the rate in the symmetric case and overestimates the rate in the activationless case. Symmetric reactions (with \( \epsilon = 0 \)) have a larger quantum enhancement than other driving forces in the normal regime, due to nuclear tunneling. The iso-RP rate for a symmetric reaction is therefore contaminated by imaginary time paths for which the tunneling is weaker, which leads to an underestimation of the quantum mechanical rate. Conversely, in the activationless case, where the exact rate is typically very close to the Marcus theory rate, iso-RPMD considerably overestimates the rate as it is dominated by the large nuclear tunneling seen around \( \epsilon = 0 \). In both cases, iso-RPMD is dominated by unphysical imaginary time paths, which leads to an inaccurate rate. This is in contrast to the situation in the adiabatic limit, where RPMD rate theory accurately captures deep tunneling due to its connection to the semiclassical instanton approximation.\textsuperscript{18} Modifying iso-RPMD so that it is dominated by the correct (nonadiabatic) instanton path in the golden rule limit may well therefore provide a useful route to improving its accuracy for nonadiabatic reactions.

VI. CONCLUSION

Our analysis has shown that the iso-RP ansatz is unable to quantitatively describe reaction rates in the golden rule limit, when nuclear quantum effects are important. We have shown that the ansatz leads to a uniform quantum enhancement over the Marcus theory rate, independent of the thermodynamic driving force. Hence, it does not capture the quantum asymmetry that is seen in a plot of the logarithm of the rate constant against the thermodynamic driving force. (However, since it correctly describes the high temperature limit, iso-RP will capture the classical asymmetry seen in anharmonic systems.) Furthermore, we have shown that the quantum enhancement predicted by the iso-RP ansatz is approximately the average of the true quantum enhancement over the entire Marcus normal regime. This explains why iso-RP tends to underestimate the rate for symmetric reactions and to overestimate the rate for activationless reactions. Since the average quantum enhancement in the normal regime is typically dominated by reactions near the symmetric limit, iso-RPMD may prove qualitatively useful in describing these reactions. However, it is clear that applying the method to reactions with driving forces nearer the activationless limit, or those in the Marcus inverted regime, would be inadvisable.

By design, we have only considered the golden rule limit of iso-RPMD and it is clear that the failings seen in this regime will not be universal to all coupling strengths. In particular, provided a sensible MQC method is used, iso-RPMD should still accurately describe systems which are close to being electronically adiabatic. Furthermore, even for systems in which there are significant nonadiabatic effects, provided these effects are not rate limiting in the process considered, iso-RPMD may provide accurate results. It does share the single most important feature of standard adiabatic RPMD, in that it obeys quantum detailed balance by construction (provided a MQC method with this property is used).\textsuperscript{2,3} We also expect that iso-RPMD will be accurate for systems in which the nonadiabatic transition is effectively classical and nuclear quantum effects are important for crossing a barrier away from this transition. In fact, the accuracy of iso-RPMD has already been demonstrated in such a situation by Tao...
et al. in their calculation of state-resolved reaction rates for a one-
dimensional two-state model of the F + H₂ reaction. Application
of the method to similar but more complex systems will no doubt
provide physical insight in the future.

For systems in which nuclear quantum effects are important
and the rate limiting step involves passage through the diabatic
crossing, such as many biological and inorganic electron transfer
reactions, more work is clearly needed to make iso-RPMD an accurate
method for calculating reaction rates. Provided such systems are
in the golden rule limit, however, Wolynes theory already provides
a quantitatively accurate way to calculate reaction rates in both the
normal and inverted Marcus regimes. Moreover, one can connect
Wolynes theory in the golden rule limit to conventional RPMD rate
theory in the Born-Oppenheimer limit using a simple interpolation
formula, which enables the calculation of accurate reaction rates
for arbitrary electronic coupling strengths.

In order to apply iso-RPMD to such systems, one would need to modify
the isomorphic Hamiltonian or the dynamics to make it more closely related
to an accurate approximation to the quantum rate in Eq. (13) in the
golden rule limit. This might be achieved by making a connection to
Wolynes theory, or to a newer path integral based approach such as
the recently proposed general grand quantum transition state theory
of Thapa et al., which already has a very similar functional form
to Eq. (20). However, whether or not such an improvement is possible
remains to be seen.

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APPENDIX: GOLDEN RULE LIMIT OF Δⁿiso[q]²

Here, we evaluate the golden rule limit of the iso-RP coupling,
Δⁿiso(q). In order to keep the argument as simple as possible, we
will consider a position independent Δ(q) = Δ. It is also simpler to
work in path integral notation, where ring polymer configurations
q (in the infinite n limit) become cyclic paths q(τ) parameterized by
imaginary time τ. Consequently, functions of a single ring polymer
configuration, A(q), become functionals of the imaginary time path,
A[q(τ)]. Making this notational change, we can recast Eq. (18) in the
infinite n limit as

\[ k_{\text{iso}} = \left( \frac{2\pi}{\hbar} \Delta_{\text{iso}}[q(\tau)] \right)^2 \delta(V_{\text{iso}}[q(\tau)]) \],

where

\[ V_{\text{iso}}[q(\tau)] = V_0^\text{iso}[q(\tau)] - V_1^\text{iso}[q(\tau)], \]

\[ V_0^\text{iso}[q(\tau)] = \frac{1}{\beta \hbar} \int_0^{\beta \hbar} V(q(\tau)) \, d\tau \],

\[ V_1^\text{iso}[q(\tau)] = \int \frac{S_0^\text{iso}[q(\tau)]}{\beta}, \]

and the expectation value is given by

\[ \langle A[q(\tau)] \rangle_{\text{iso}} = \frac{\int Dq(\tau) e^{-\frac{1}{2} S_0^\text{iso}[q(\tau)]/\hbar} A[q(\tau)]}{\int Dq(\tau) e^{-\frac{1}{2} S_0^\text{iso}[q(\tau)]/\hbar}}, \]

with

\[ S_0^\text{iso}[q(\tau)] = \int_0^{\beta \hbar} \sum_{i=1}^1 \frac{1}{2} m_i \dot{q}_i^2(\tau) + V_0(q(\tau)) \, d\tau. \]

Note that the integrals over the positions of the ring polymer beads have become a "path integral" over all possible paths q(τ). The circle
indicates that the paths are cyclic, q(τ + βh) = q(τ), and the measure
\[ Dq(\tau) \] contains the integrals over the ring polymer momenta (which
have become a "path integral" over all possible paths).

In order to evaluate each term in this expansion, one can expand the
trace as

\[ \mu[q(\tau)] = \lim_{n \to \infty} \prod_{j=1}^n \left( \sum_{\sigma_j} \langle \sigma_j | e^{-\beta \hbar V(q(\tau))} | \sigma_{j+1} \rangle \right). \]

where \( \sigma_j \) and \( \sigma_{j+1} = \sigma_1 \). Then, noting that

\[ \langle 0 | e^{-\beta \hbar V(q)} | 0 \rangle = e^{-\beta \hbar V(q) + \mathcal{O}(\beta^2)}, \]

\[ \langle 1 | e^{-\beta \hbar V(q)} | 1 \rangle = e^{-\beta \hbar V(q) + \mathcal{O}(\beta^2)}, \]

and

\[ \langle 0 | e^{-\beta \hbar V(q)} | 1 \rangle = -\beta \Delta_{\text{iso}} e^{-\beta \hbar V(q) + V(\tau)} + \mathcal{O}(\beta^2), \]

we see that each off-diagonal matrix element in the product in Eq. (A10)
contributes one power of \( \Delta \). Equivalently, noting that the set \( \{ \sigma \} \)
in the infinite n limit corresponds to a stochastic path \( \sigma(\tau) \), we see
that every time \( \sigma(\tau) \) changes from 0 to 1 (known as a kink) we
pick up one power of \( \Delta \). Since the path \( \sigma(\tau) \) has cyclic symmetry,
the must have an even number of kinks and hence \( \mu[q(\tau)] \) is an even
function of \( \Delta \).
It is immediately obvious that the only paths that contribute to the zeroth order term are those which do not contain any kinks

$$\mu_0[q(r)] = e^{-\beta V_+[q(r)]} + e^{-\beta V_-[q(r)]},$$  
(A14)

corresponding to $\sigma(\tau) = 0$ and $\sigma(\tau) = 1$ for all $\tau$ in the track over diabatic states in Eq. (A8). Similarly, the only paths that contribute to $\mu_1[q(r)]$ are those which contain two kinks. Each of these paths can be characterized by two parameters, the total time which the path spends on diabat 1, which we denote $\lambda h$,

$$\lambda h = \int_0^h \sigma(\tau) \, d\tau,$$  
(A15)

and the time at which $\sigma(\tau)$ changes from 0 to 1, which we label $\tau'$. We can thus write the second order term as

$$\mu_2[q(r)] = \frac{1}{h} \int_0^h d\lambda \left[ \int_0^h d\tau' e^{-(\beta - \lambda) V_+[q(\tau + \tau')] - \lambda V_-[q(\tau + \tau')]} \right],$$  
(A16)

where

$$V_0^+(q(\tau)) = \frac{1}{(\beta - \lambda) h} \int_0^h V_0(q(\tau)) \, d\tau,$$  
(A17)

$$V_1^+(q(\tau)) = \frac{1}{\lambda h} \int_0^h V_1(q(\tau)) \, d\tau.$$  
(A18)

Now, by first recognizing that

$$\mu_0[q(r)] e^{\beta V_+^0[q(r)]/2} = \cosh(\beta V_+^0[q(r)]/2),$$  
(A19)

and then making use of

$$\cosh^2[a + b \Delta^2] - a^2 = \frac{2b \Delta^2}{\sinh(a)} + O(\Delta^4),$$  
(A20)

it is straightforward to show that we can write the coupling as

$$\Delta^i [q(r)]^2 = \Delta^2 \frac{\mu_1[q(r)]}{\mu_2[q(r)]} + O(\Delta^4),$$  
(A21)

where

$$\mu_1[q(r)] = \frac{2}{h} \int_0^h d\lambda \left[ \int_0^h d\tau' e^{-(\beta - \lambda) V_+^0[q(\tau + \tau')] - \lambda V_-^0[q(\tau + \tau')]} \right].$$  
(A22)

Next recognizing that

$$\frac{\delta[V_-^0[q(r)]]}{\mu_2[q(r)]} = \frac{\delta(V_+^0[q(r)])/e^{-\beta V_+^0[q(r)]}}{\mu_2[q(r)]},$$  
(A24)

and also that

$$\frac{e^{-\beta V_+^0[q(r)]}/\mu_2[q(r)]}{e^{-\beta V_-^0[q(r)]}/\mu_1[q(r)]} = \frac{1}{h} \int_0^h d\lambda \left[ \int_0^h d\tau' e^{-\beta V_+^0[q(\tau + \tau')]/h} \right],$$  
(A25)

where

$$S^{(1)}[q(r)] = S^{(1)}_0[q(r)] + S^{(1)}_1[q(r)],$$  
(A26)

$$S^{(1)}_i[q(r)] = \int_0^h \sum_{\tau = 1}^2 \frac{1}{2} m_\tau q_\tau^2(\tau) + V_\tau(q(\tau)) \, d\tau,$$  
(A27)

we see that we can rewrite Eq. (A1) in the golden rule limit as in Eq. (25),

$$k_{iso} = \frac{2\pi}{\hbar Q_i} \int_0^\beta d\lambda \left\{ \Delta^2 \delta(V_-^-[q(\tau)]) \right\} e^{-\beta \lambda / h},$$  
(A29)

with

$$\langle A[q(r)] \rangle = \frac{\oint V(q(r)) e^{-\beta V_-^0[q(r)]/h} A[q(r)]}{\oint V(q(r)) e^{-\beta V_+^0[q(r)]/h}}$$  
(A30)

and

$$e^{-\beta \lambda / h} = \text{tr}_\lambda e^{-\beta V_-^0[h_\lambda - L_\lambda]} = \oint V(q(r)) e^{-\beta V_-^0[q(r)]}/h,$$  
(A31)

and

$$Q_i = e^{-\beta V_-^0(0)} = \text{tr}_0 e^{-\beta h_\lambda},$$  
(A32)

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