Mean first-passage times for solvated LiCN isomerization at intermediate to high temperatures

Micha M. Schleeh,1,a Johannes Reiff,1,b,∗ Pablo L. García-Müller,2 Rosa M. Benito,3 Florentino Borondo,4,5 Jörg Main,1 and Rigoberto Hernandez6,7,b

1) Institut für Theoretische Physik I, Universität Stuttgart, 70550 Stuttgart, Germany
2) Departamento de Tecnología, Centro de Investigaciones Energéticas Medioambientales y Tecnológicas, Avda. Complutense 40, 28040 Madrid, Spain
3) Grupo de Sistemas Complejos, Escuela Técnica Superior de Ingeniería Agronómica, Alimentaria y de Biosistemas, Universidad Politécnica de Madrid, 28040 Madrid, Spain
4) Instituto de Ciencias Matemáticas (ICMAT), Cantoblanco, 28049 Madrid, Spain
5) Departamento de Química, Universidad Autónoma de Madrid, Cantoblanco, 28049 Madrid, Spain
6) Department of Chemistry, Johns Hopkins University, Baltimore, Maryland 21218, USA
7) Departments of Chemical & Biomolecular Engineering, and Materials Science and Engineering, Johns Hopkins University, Baltimore, Maryland 21218, USA

(Dated: 5 January 2022)

The behavior of a particle in a solvent has been framed using stochastic dynamics since the early theory of Kramers. A particle in a chemical reaction reacts slower in a diluted solvent because of the lack of energy transfer via collisions. The flux-over-population reaction rate constant rises with increasing density before falling again for very dense solvents. This Kramers turnover is observed in this paper at intermediate and high temperatures in the backward reaction of the LiNC $\longrightarrow$ LiCN isomerization via Langevin dynamics and mean first-passage times (MFPTs). It is in good agreement with the Pollak–Grabert–Hänggi (PGH) reaction rates at lower temperatures. Furthermore, we find a square root behavior of the reaction rate at high temperatures and have made direct comparisons of the methods in the intermediate- and high-temperature regimes; all suggesting increased ranges in accuracy of both the PGH and MFPT approaches.

I. INTRODUCTION

The mean first-passage time (MFPT) is a useful estimate of an inverse reaction rate constant (for simplicity called rate in this paper) whether it is determined numerically or analytically.17 For example, the inverse MFPT can be used to numerically obtain the Kramers rate through the direct observation of an ensemble of trajectories of a given system. Its use in resolving rates in chemical reactions has been demonstrated in several systems including the isomerization reaction of lithium cyanide (LiNC/LiCN). The latter has received a lot of attention in part because of the early introduction of a useful model potential by Essers et al.21 Recently, for example, some of us obtained the decay rate of trajectories near the normally hyperbolic invariant manifold (NHIM) for this reaction so as to demonstrate the use of nonrecrossing dividing surfaces in time-dependent transition state theory (TST).22

In turn, the objective of this paper is to demonstrate a stronger connection between rates obtained using MFPT to those from flux-over-population formulas. Pollak, Grabert and Hänggi (PGH)25 employed the latter in constructing the theory describing the Kramers turnover23 in the rates from low to high friction. The turnover was observed in the LiNC reaction25 and has subsequently been seen to correlate well with the Pollak–Grabert–Hänggi (PGH) theory.20,11,19

Here, we obtain the MFPT and the PGH theory explicitly for the backward reaction LiCN $\longrightarrow$ LiNC. Perhaps surprisingly, we found that the MFPT can be useful in describing reactions of particles undergoing Brownian motion at much higher temperature than the barrier height. Throughout this work, we use the terms low, intermediate, and high to qualify our temperature range in reference to the barrier height. Intermediate temperatures are on the order but lower than the barrier height, whereas low temperatures are much lower than the barrier height by at least a factor of 10 such that the system is mostly bound and the reactions must be activated. Notably, we benchmarked the rates obtained from the MFPT to several alternative methods so as to confirm the relative accuracies across a broad range of friction and temperature.

At high temperatures, the rate of escape and the corresponding rate constant have to be carefully defined. Here, a significant proportion of the initial conditions are at initial energies above the barrier. These are not trapped and simply escape across the barrier without return (as long as an absorbing boundary condition is used on the other side). These trajectories do not contribute to a steady-state flux or the corresponding rate constant. However, there is still a small population—
LiCN involves the breaking and making of weak bonds between a lithium cation and a cyanide anion. The LiNC is the stable conformation of the isomerization reaction, and hence the backward reaction, Li–C≡N → C≡N–Li, (1) is exothermic. Figure 1 shows the LiCN molecule in a Cartesian body-fixed frame. Through the approximation of separating the translation and rotation of the center of mass of the whole molecule, it is possible to reduce the equations of motion to two internal variables.

The reaction saddle is marked with a cyan cross. At \( \vartheta = 0 \) the isomer exists in its LiNC configuration \( V(4.7947, 0) = -0.23421 \text{ a.u.} \), whereas at \( \vartheta = \pi \) it is in its LiNC configuration \( V(4.3487, \pi) = -0.24461 \text{ a.u.} \).

These variables, \( R \) and \( \vartheta \), are known as the Jacobi coordinates: \( R = |R| \) is the distance between the NC center of mass and the lithium atom, and \( \vartheta = \angle (\mathbf{r}_{\text{NC}}, \mathbf{R}) \) is the angle between \( \mathbf{R} \) and \( \mathbf{r}_{\text{NC}} \), where \( |\mathbf{r}_{\text{NC}}| = r_c = 2.186 \mu_0 \) and \( \mu_0 \) is the Bohr radius. At \( \vartheta = 0 \), the potential is at a local minimum that corresponds to the metastable conformation of the LiCN reactant in the backward reaction. The global minimum of the potential lies at \( \vartheta = \pi \), and corresponds to the stable conformation of the LiNC product.

The Hamiltonian can now be specified using the Born–Oppenheimer potential energy surface and the usual approximations as

\[
\mathcal{H} = \frac{p_R^2}{2\mu_1} + \frac{1}{R^2} \left[ \frac{p_\vartheta^2}{2m^*(R)} \right] + V(R, \vartheta) \tag{2}
\]

containing the kinetic and potential energy \( V(R, \vartheta) \) energy in \( R \) and \( \vartheta \) where the reduced masses are \( \mu_1 = [1/m_{\text{Li}} + 1/(m_C + m_N)]^{-1} \), \( \mu_2 = (1/m_C + 1/m_N)^{-1} \), and

\[
m^*(R) = \left( \frac{1}{\mu_1} + \frac{R^2}{\mu_2 \mu_0^2} \right)^{-1}. \tag{3}
\]

We assume pure \(^7\text{Li}, ^{12}\text{C}, \) and \(^{14}\text{N} \) isotopes for the atomic masses \( m_{\text{Li}}, \text{m}_C, \text{and } m_N \), respectively. The potential energy surface (PES) in Jacobi coordinates is shown in Fig. 2. The reaction saddle is marked with a cyan cross and lies at \( R = 4.2197 \text{ a.u.} \) and \( \vartheta = 0.2922\pi \).

In this paper, we use atomic units (a.u.) for all of our figures and calculated values. For example, the distances are given in terms of the Bohr radius \( \alpha_0 \), the atomic time unit is given by \( h/E_0 \), where \( E_0 \) is the Hartree energy, and the mass is given in electron masses \( m_e \). In these units, we found high-temperature rates \( k \) on the order of \( 10^{-4} \text{ a.u.} \) (cf. Fig. 5), corresponding roughly to \( k = 4 \times 10^{12} \text{ s}^{-1} \) in SI units. The inverse of such a rate is
comparable to the time scale of the isomerization dynamics reported for high excitation energies in Ref. [26] which are also the dynamics accessed at high temperatures.

B. Langevin implementation

Solvent effects on the LiNC $\rightleftharpoons$ LiCN isomerization reaction have previously been addressed through the introduction of an argon bath. [16,17,25] The interaction with the bath can be reduced to the Langevin equation though a mapping to the characteristic friction and random noise. [16,17,25] With the approximation $(d/dt)/(p_\alpha/(m^* R)) \approx \dot{p}_\alpha/(m^* R)$, the equations of motion follow from Hamiltonian [2] as

\begin{align*}
\dot{\vartheta} &= \left(\frac{1}{\mu_1 R^2} + \frac{1}{\mu_2 R^2}\right)p_\vartheta, \quad (4a) \\
\dot{p}_\vartheta &= -\frac{dV(R, \vartheta)}{d\vartheta} - \gamma p_\vartheta + R \xi_\vartheta, \quad (4b) \\
\dot{R} &= \frac{p_R}{\mu_1}, \quad (4c) \\
\dot{p}_R &= \frac{p_R^2}{\mu_1 R^3} - \frac{dV(R, \vartheta)}{dR} - \gamma p_R + \xi_R, \quad (4d)
\end{align*}

where the stochastic forces $\xi_\vartheta$ and $\xi_R$ satisfy the respective fluctuation-dissipation theorems [24]

\begin{align*}
\langle \xi_{\vartheta_i}(t) \xi_{\vartheta_j}(t') \rangle &= 6 \gamma k_B T m^* \delta_{ij} \delta(t - t') \quad (5a) \\
\text{and} \quad \langle \xi_{R,i}(t) \xi_{R,j}(t') \rangle &= 6 \gamma k_B T m_1 \delta_{ij} \delta(t - t') \quad (5b)
\end{align*}

for uniformly distributed noise. The canonical momentum $p_\vartheta$ is an angular momentum. It should therefore not be surprising that the last term in Eq. [4b] includes a product with the radial coordinate $R$ as it leads to the correct units. The random forces are generated at the beginning of the calculation and use a fixed $R$ (= 4.2196 a.u.) at the barrier with the reduced mass $m^*$ (= 2406 a.u.). This is a nontrivial approximation because the reduced mass varies as much as 25% across the positions in $R$, but it is consistent with prior work and the error is least when the trajectories are near the barrier. A numerical consequence of this approximation is that the random forces do not vary with $R$, thus allowing a significant simplification in coding the equations of motion and in implementing the theory. Thus the price of this approximation is that the results may be affected in so far as the effective temperature is renormalized.

The results have been calculated with a fourth order Langevin Runge–Kutta keeping the random force constant during each time step.

C. Pollak–Grabert–Hänggi theory

The PGH theory for activated processes driven by Markovian forces has been seen to be very effective in the low- and intermediate-temperature regimes across the Kramer turnover of the rates with respect to friction. [23] Among several examples, [23,25] it was shown to be effective for characterizing the dynamics across the potential model of Straub–Borkovec–Berne (SBB). [25,26] The Langevin implementation is memoryless, which is formally described by the friction kernel,

$$
\gamma(t, t') = \gamma_0 \delta(t - t').
$$

(6)

The SBB approximation uses parabolic functions attached to each other to create a single potential well followed by a saddle as an inverse parabolic function, and implemented in the generalized Langevin equation with a memory friction. It was used previously by some of us to model the minimum energy path of the LiNC $\rightleftharpoons$ LiCN isomerization reaction. [25] In the SBB model, memory is introduced through a single exponentially decaying term in the friction kernel,

$$
\gamma(t, t') = \alpha^{-1} \exp\left(-\frac{|t - t'|}{\alpha \gamma}\right).
$$

(7)

The propagation of particles inside this potential is strongly dependent on the memory time scale $\tau = \alpha \gamma$ used in the friction kernel and the form of the friction kernel $\gamma(t, t')$ itself. For those relaxation processes that occur at times much longer than $\tau$, the response looks ohmic as in the Langevin case. However, the SBB model now allows for dynamical responses from the solvent that can compete with the dynamics in the system. This leads to an effective friction which arises from the mean of the modulate frictions from the previous times. We found earlier [13] that in practice, this led to an increase of observed rates by about a factor of 5 when using the Langevin equation (LE) rather than the SBB model.

D. Mean first-passage time rates

The first-passage time is the time a particle needs to reach a certain region for the first time given some initial state. In case of a reaction, the first-passage time is defined as the time the particle is propagated from a point in the reactant region to a point on some characteristic surface at or beyond a dividing surface (DS) [11,16,43]. In reactive systems characterized by one-dimensional barriers, the DS reduces to a point. It is naively taken to be the saddle point, but other choices are available, just like for variational TST, for example. [25,12] The first-passage times for a series of trajectories from different initial points in the reactive regime experiencing varying thermal forces vary stochastically. Averaged together they lead to the mean first-passage time $t_{MFPT}$, whose inverse is the rate of escape $k_{MFPT}$ [12,14,15]

$$
k_{MFPT} = \frac{1}{t_{MFPT}}.
$$

(8)

In the limit of a harmonic barrier, $k_{MFPT}$ has been seen to be precisely equal to the TST rate, and both are equal to the correct Kramers rate $k$ [13].
Generally, the rate problem is treated exclusively in the activated regime. Therein, the typical energies of the system are characteristic of an average temperature that is well below the energetic barrier. The smooth turnover in the Kramers rates with friction was resolved by Mel’nikov and Meshkov\cite{15} and Pollak, Grabert and Hänggi (PGH).\cite{16} They found a mathematical expression for the rate connecting the low-friction regime—where the rate increases with $\gamma$—and the high-friction regime—where the rate decreases with $1/\gamma$. PGH imposes a rate-determining region (or DS in phase space) which requires temperatures to be low enough that the reacting system is somehow thermalized. Since our initial work\cite{17,19,21} demonstrating the applicability of the PGH theory at surprisingly higher temperatures, Pollak and coworkers have extended it for temperatures near threshold\cite{22}

In the present problem, however, we must also consider much higher temperatures in which the reactive system usually accesses energies much higher than the barrier along the reaction coordinate. The process is consequently effectively barrierless, and the rate problem reduces to the determination of a steady-state flux for a given thermal molecular beam. That is, the typical energies of those states accessing and crossing the barrier at high temperatures—in the sense that they are much larger than the barrier—correspond to states that cross the barrier freely (or ballistically) at their typical velocity. As a consequence, the rates reduce to a simple power law at high temperature,

$$k \propto \sqrt{\langle v^2 \rangle} \propto \sqrt{T}. \quad (9)$$

This behavior is well known since the early work on molecular beams\cite{23,24} Alternatively, we can recover Eq. (9) from the mean-squared displacement observed in small and long times from the Langevin equation,\cite{25} where the mean-squared displacement is given by

$$\langle \Delta x^2 \rangle = \frac{2ND}{\gamma} \left( \gamma t - 1 + e^{-\gamma t} \right) \approx 2ND\gamma t^2 \approx 2Nk_B T t^2. \quad (10)$$

Thus the velocity dependence of Eq. (9) is again recovered when

$$t \ll 1/\gamma \quad (11)$$

is satisfied\cite{26} and the reactants have a short MFPT (and associated small $t$) due to the fast barrierless crossing.

### III. RESULTS

#### A. Mean first-passage time rates of LiCN isomerization

The PES of LiNC $\rightarrow$ LiCN (cf. Fig. 3) is replotted in Fig. 5 in terms of the Cartesian coordinates, $x$ and $z$, of Li relative to the $C_{\infty v}$ axis of CN. The minimum energy path of the potential nearly follows a semi-circle with a radius of $R \approx 4.5$ a.u. The minimum energy of the LiCN molecule ($E_{\text{min,LiCN}} = -0.23421$ a.u.) is not as deep as that of LiNC ($E_{\text{min,LiNC}} = -0.24461$ a.u.). Consequently, a trajectory starting at the LiCN state has to overcome a smaller barrier height of $E^f = 0.00535$ a.u. (corresponds to $E^f/k_B = 1690$ K) than one starting from the LiNC state. At room temperature ($T = 300$ K), for example, the barrier of this backward reaction is thus low enough that such trajectories are activated to above threshold energy frequently enough that they can be observed numerically. This is also true even for the forward reaction despite its higher barrier.

To ensure that selected trajectories are properly identified as reactive, they must first cross the saddle and then reach the product side without turning around. This condition is satisfied using an absorbing boundary\cite{27,28} defined by an angle of $\varphi = 0.67\pi$, and shown as the dotted

![Fig. 3. Potential energy $V$ as a function of body-fixed Cartesian coordinates ($x, z$). The origin is located at the cyanide compound’s center of mass. Filled circles illustrate the positions of the individual atoms in our model. The circles’ radii are chosen proportional to the atomic masses. For lithium, an arbitrary position on an example trajectory at $T = 300$ K is shown. The trajectory starts at the local minimum $x = 0$, $z = 4.7947$ corresponding to the LiCN configuration. It crosses the saddle located at $x = 3.3520$, $z = 2.5631$ and ends at the absorbing boundary, indicated by the dashed line. The potential’s global minimum $x = 0$, $z = -4.3487$ corresponds to the LiNC isomer.](image-url)
white line in the Cartesian PES plot of Fig. 3. Indeed, if the trajectory reaches this line, then it generally has enough momentum in the direction of the product side to make it very unlikely for it to turn around and climb back over the saddle to the reactant side. Use of absorbing boundary conditions along space and/or energy to address low to mid friction also has significant precedent in the literature.\textsuperscript{22–25} In the present case focusing on high temperatures, we did not add an energy condition to the absorbing boundary because the spatial constraint is sufficient to remove the high-energy trajectories once they have reached the product basin (cf. Fig. 3) before their long-time return to reactants. Its use enables us to better compare the MFPT calculations to the AAMD results obtained earlier in Ref. 16 where direct rates were determined based on the flux through a DS along the reaction coordinate.\textsuperscript{22}

The representative trajectory shown in Fig. 3 is only one of 1500 propagated trajectories from the ensemble used to calculate the MFPT rate $k_{\text{MFPT}}$. Trajectories of this thermal ensemble are initialized at a specific temperature $T$ and located near the reactive well. For simplicity in the current implementation, they are all placed at the LiCN minimum at $R = 4.7947$ a.u. and $\vartheta = 0$, which in Cartesian coordinates corresponds to $x = 0$ a.u. and $z = 4.7947$ a.u. Their velocity is sampled from a Maxwell–Boltzmann distribution with temperature $T$. The assumption here is that the redistribution to a quasi-bound distribution in space is fast compared to the escape times when the system is trapped. Meanwhile, in the barrierless cases, there would be no such trapping and we are merely considering the steady-state flux from the injection of particles at this origin. This case could presumably be accessible through increasingly exquisite spectroscopic methods such as transition state (TS) spectroscopy.\textsuperscript{52,53}

At higher bath temperatures, the distributions in energy—total, potential and kinetic—of the initial reactant ensemble necessarily shift to higher values. Consequently, a larger percentage of the states in the reactant ensemble will possess the required energy to surmount the barrier at any given time. More of these states escape over the barrier and are removed from the simulation. The remaining and decreasing population of reactive states is re-thermalized according to the bath temperature, and thereby continues to contribute to the steady-state rate. As the temperature increases further, this activation is faster leading to increasingly larger kinetic energies in the trajectories contributing to the steady-state rate. This, in turn, causes smaller MFPTs and higher rates. Specifically, ensembles thermalized at some temperature $T$ and propagated at a certain friction value $\gamma$ lead to rates that depend not on the fast inertial trajectories but rather those that endure several traversals of the reaction region—at energies below activation threshold—before escaping.

![FIG. 4. Mean first-passage time rates $k_{\text{MFPT}}/5$ of the LiCN $\rightarrow$ LiNC backward reaction as a function of friction $\gamma$ at temperatures $T = 300$ K (blue circles), $T = 450$ K (orange triangles), and $T = 600$ K (green diamonds). For comparison, the corresponding PGH rates obtained for a GLE model—with friction kernel specified by a bath parameter $\alpha = 1.5625$ a.u. and decay time $\tau (= \alpha \gamma)$—are shown as solid blue, dashed orange, and dotted green lines, respectively. The reported rates $k_{\text{MFPT}}/5$ are obtained from the MFPT rates for the LE model (with ohmic friction), and scaled by a factor of 5 to make the shapes of the Kramers turnovers comparable.](https://example.com/fig4.png)

**B. Kramers turnover**

We now report the MFPT rates $k_{\text{MFPT}}$ across the friction domain at several intermediate temperatures in Fig. 4. As before,\textsuperscript{16,19,25} we observe a clear Kramers turnover, and the shape of $k_{\text{MFPT}}$ is in very good agreement with the corresponding PGH rate formula. The rate maxima are always around $\gamma \approx 4 \times 10^{-4}$ a.u.

To obtain the rates in Fig. 4 at different ensembles at $T = 300$ K, $T = 450$ K, and $T = 600$ K are thermalized on each friction value $\gamma$. Each mean rate $k_{\text{MFPT}}$ is calculated using 1500 trajectories propagated by the Langevin equation. At low friction ($1 \times 10^{-5}$ a.u. $< \gamma < 2 \times 10^{-4}$ a.u.), we find the expected linear increasing behavior of the rates. Similarly, at high friction, the rates decrease strongly with $1/\gamma$. These two limits have been known since the work of Kramers. In combination, they give rise to the eponymous Kramers turnover. In absolute values, however, the MFPT rates overestimate the expected rates by a factor of 5. This is unfortunate, but not unexpected, because the MFPT rates—just like those from flux over population\textsuperscript{25}—are known to be approximate due to several factors. In the present implementation, these MFPTs are uniformly shorter, leading to faster rates than measured through the other approaches, presumably because there is an acceleration from the initial distribution that is not being dissipated by the bath.

The rates calculated using trajectories propagated via the LE are also overestimated by about a factor of 5 compared to the SBB model and AAMD calculations. Specifically, the Kramers turnover rate maxima $k_{\text{max}}$ in the LE and generalized Langevin equation (GLE) from
FIG. 5. Kramers turnover rate maxima $k_{\text{max}}$ as functions of temperature $T$. The rates $k_{\text{MFPT}}$ (blue triangles), calculated with an LE friction kernel, are shown in the low- and high-temperature regimes, which are separated by the effective temperature of the barrier $E^\ddagger/k_B = 1690\,\text{K}$. The reported PGH rates, marked with purple crosses and cyan plusses, are obtained for the GLE model with bath parameters $\alpha = 1.5625\,\text{a.u.}$ (corresponding to short-time memory) and $\alpha = 1.313 \times 10^6\,\text{a.u.}$ (corresponding to long-time memory), respectively. The AAMD- (red diamonds), LE- (orange circles), and GLE-based (green squares) reactive-flux rates are calculated via the flux-over-population method, and are available only in the high-temperature regime. For reference, rate calculations based on two TST formulas (cf. Appendix [B]) are shown with dashed and solid gray lines.

Ref. [19] are in agreement only if the rates obtained for the reactive flux calculations for the LE were to be divided by a factor of 5. Meanwhile, the GLE rates obtained using PGH theory for the near ohmic limit (at very small $\alpha$) do not suffer from this overestimation. This suggests that the LE rates calculated using reactive flux suffer from a numerical error not present in the GLE dynamics. We conjecture, though have not proven, that this discrepancy arises from a slower energetic redistribution among the quasi-bound states allowing for a faster escape of the initial higher energy configurations.

To confirm the effect of friction in the Kramers turnover of $k_{\text{MFPT}}$ seen in the PGH theory in Fig. 4, the bath parameter $\alpha$ in the PGH friction kernel is set to a small value $\alpha = 1.5625\,\text{a.u.}$ The maximal rates for fixed $\alpha$ in this case are still found at friction values $\gamma_0$ in a similar range and hence this case ($\alpha = 1.5625\,\text{a.u.}$) indeed leads to a short memory time scale of $\tau \approx 6.25 \times 10^{-4}\,\text{a.u.}$ This places it in the memoryless dynamics regime characteristic of the LE, as noted above. Meanwhile, the PGH rates obtained at the longer memory time $\tau \approx 5.25 \times 10^2\,\text{a.u.}$—corresponding to a larger $\alpha$—as was observed in the AAMD simulations still exhibits similar maximal rates in Fig. 5. At intermediate temperatures, for example, the memory time scale is still much lower than the MFPT ($t_{\text{MFPT}} \gtrsim 1 \times 10^4\,\text{a.u.}$), and thus the system remains in a nearly memoryless regime. As the PGH theory—which is also known to be in good agreement with TST[20] at the appropriate limits—describes the low- and intermediate-temperature regime well, it is not surprising that the MFPT rates are effective as long as we ignore a temperature-independent uniform factor.

C. High-temperature regime

The MFPT rates can also be used to describe the high-temperature regime, where the typical energies are well above the barrier height $E^\ddagger/k_B = 1690\,\text{K}$. Therein the Kramers turnover maxima in the MFPT rates are compared to the corresponding reactive-flux rates[19] in Fig. 5. Specifically, we compare against LE-, GLE-, and AAMD-based calculations, where we consider AAMD to yield the most accurate results. Notably, the latter includes cavity reorganization effects which may not be fully described within the LE-based reactive-flux and MFPT approaches.

Figure 5 also includes TST benchmark calculations based on the Eyring–Polanyi (cf. Appendix [B] and the Polanyi–Wigner (cf. Appendix [B]) rate expressions. While the Eyring–Polanyi rate describes the reaction rate reasonably well for $k_B T \ll E^\ddagger$, as is well known[8][9][10][11], it quickly deviates from the other results as the temperature is increased. Indeed, the usual rule of thumb is that there is a breakdown in rate theories—because they begin to violate the separation between reactant and transition state regions—above this threshold. Thus the result in the all-atom LiCN dynamics in an argon solvent seen in Refs. [18] and [25] was surprising because it suggested the existence of an observable rate at such high temperatures, and their turnover with apparent friction obeyed the Kramers turnover seen through the lens of PGH theory. Here, we see that part of the origin of the breakdown in the Eyring–Polanyi theory is that it suggests an infinite rate which is not possible when the particles move at velocities distributed around a Boltzmann distribution for a given $k_B T$. Instead, at high enough temperatures, the reactants move at apparent rates that are proportional to their average velocity because at such energies the barrier is not visible. The results in Fig. 5 demonstrate that the rates, as also captured by Polanyi–Wigner, move smoothly from the well-known low-temperature regime through intermediate temperatures in which the barrier plays a role, albeit a decreasing one with increasing temperature.

Thus at increasing temperatures, the reaction is no longer limited by the activation process that is inher-
ent in the Eyring–Polanyi formula and rises much faster leading to a very large upper bound of the rate. On the other hand, the Polanyi–Wigner formalism appears to capture the slowing down in the rates due to the decreasing quasi-bound population in the reactive basin with increasing temperature. Specifically, the prefactor in the Polanyi–Wigner rate is no longer temperature dependent but rather determined by the vibrational frequency $\omega_0/2\pi \approx 3.2 \times 10^{12} \text{s}^{-1}$ of the reactant well and, therefore, leads to a flattening in the rate with increasing high temperatures. Nevertheless, the interaction with the solvent manifests itself by way of the decreased rates through the intermediate temperature regime.

The comparison between the maxima of the PGH rates and the AAMD rates for higher temperatures in Ref. 16 is also shown in Fig. 4. The agreement shows that the PGH theory can also be applied to high temperatures. In this work, we further found that the PGH rates are able to describe the intermediate- and high-temperature regime, both with the memory time scale appropriate to the system (cyan pluses) and through the approximate Langevin approach (purple crosses). This also justifies the use of the $k_{\text{MFPT}}$ obtained from the LE friction kernel at the high temperatures.

The AAMD, GLE reactive-flux, and PGH results reported in Fig. 5 are in good agreement at high temperatures. As already observed in Ref. 19, the reactive-flux results based on the LE grossly overestimate the rates by a factor of 4.5 to 4.8 and 5.1 to 9.3 compared to those seen in the GLE and AAMD reactive-flux calculations, respectively. The MFPT rates for the LE are lower than those from the reactive-flux calculations on the LE but not enough to match the other methods. Specifically, $k_{\text{MFPT}}$ overestimates by a factor between 2.9 at $T = 2500$ K and 2.2 at $T = 5500$ K compared to the GLE reactive flux. The overestimation factor is larger when compared to AAMD calculations, monotonically rising from 3.3 at $T = 2500$ K to 4.6 at $T = 5500$ K. There is, however, no such simple trend when considering the whole temperature range via comparison with PGH rates. Instead, the overestimation factor is particularly high at very low temperatures (9.7 at $T = 300$ K), quickly drops until it reaches a local minimum at $T = 2500$ K, and finally rises again slowly with temperature.

Compared to the LE reactive-flux approach, the MFPT appears to better capture the escape of the quasi-trapped initial states at high temperatures. These low-probability states contribute to the steady-state flux obtained from the AAMD, GLE reactive flux and PGH leading to an even lower rate. The disagreement between the latter and those obtained using the LE is surprising because the PGH results for a GLE with a very small $\alpha$ is formally the exact answer for an LE and should be in direct agreement with the reactive flux obtained on the LE. Thus, both the MFPT and reactive-flux calculations of the LE may be suffering from numerical errors due to the fact that the needed contributing trajectories are rare. The better agreement obtained with the MFPT may reflect the fact that it does not suffer as much from the constraints related to the selection of the initial states from the quasi-bound population.

Despite some of the loss in accuracy arising from the strong localization of the initial distribution of states, the advantage of the MFPT rates compared to AAMD rates is that they can be calculated for the whole temperature regime in less time. Furthermore, a square root function shape is observable for $k_{\text{MFPT}}$, as described in Sec. 1D. This shape arises because the rate is known to depend on the square root of temperature in this regime.

The difference between $k_{\text{MFPT}}$ and reactive-flux-based approaches in Fig. 5 is a consequence of the assumptions inherent in each method. For the reactive-flux rate, an extended ensemble is initialized at the minimum of LiCN and the rates are obtained by the flux-over-population method across numerically integrated trajectories. In the GLE, the rates are obtained through a direct (and analytical) analysis of the reactive ensemble across the barrier. In principle, long-time returns to the barrier may be absent from the latter leading to an overestimate in the transition state theory calculation. Such long-time sojourns are long only in the sense that they are longer than the effective reactive timescales but may be short in an absolute sense because the temperatures (and the kinetic energies) are large. A related consequence of the high-temperature regime is that the kinetic energy tends to be dominated by equipartition leading the effective average rate across the barrier to simply be described by the barrierless average velocity. The ensuing square root behavior is seen correctly in the MFPT rates in Fig. 5 at high temperatures whereas the PGH rates are unfortunately linear.

**IV. CONCLUSION AND OUTLOOK**

In this work, we have calculated the rates of the LiCN → LiNC backward reaction using mean first-passage times. We benchmarked them relative to the results of earlier AAMD calculations and the PGH rates across several temperatures and friction regimes. The MFPT rates are effective in describing the Kramers turnover in the intermediate temperature regime as shown by comparisons to the PGH rates calculated with a short memory time scale. At very high temperatures when the reactants motion effectively becomes ballistic, the MFPT approach was seen to correctly capture the square-root dependence of the rates on temperature.

At low temperatures, there are several examples of near-perfect unadjusted agreement between methods like the reactive flux and GLE-based theories like Grote–Hynes. See, for example, reports of the rates in cyclohexane interconversion, NaCl dissociation, $S_N2$ reactions, and even some enzymes. However, for the LiCN isomerization reaction at intermediate and high temperatures, we found, as before that methods based on Langevin dynamics overestimate the rate roughly by a
factor of 5 compared to AAMD or GLE simulations. The program used for our paper was written independently of the one used in Ref. [21]. It therefore seems unlikely that this factor is merely caused by a bug in some code. Instead, this suggests a physical origin which we conjecture here, as done earlier in Ref. [19] that it originates in otherwise unaccounted mean field dissipation in the determination of the stochastic trajectories. Thus the MFPT approach pursued in this paper was found to provide no worse agreement than other methods while being relatively simple to implement in systems with arbitrary dimensionality. This overestimation nevertheless poses a challenge for future work.

Furthermore, comparisons of the computed rate constants to experiment would clarify which theoretical approach is most suitable to describe the LiCN isomerization reaction. The experimental measurement of these rates especially in the high-temperature regime, also poses a challenge for future work. Specifically, for LiCN, it may be difficult to construct an initial distribution of states localized at the reactant well that is weakly solvated by a thermalized argon bath. Perhaps, spectroscopic scalpels such as those offered by TS spectroscopy [22,23] could be employed, and we look forward to seeing such advances.

In addition, the MFPT approach provides a deeper view of the structure of the LiCN isomerization reaction. The footprint (or projection) of the ensemble of trajectories that contribute to the rate onto the domain of reactant coordinates appears to spread across the intrinsic reaction coordinate (IRC). [24,25] Though not addressed quantitatively here, we conjecture that this spread can be used to characterize the manifold (or tube) that envelopes the possible reaction pathway [26] but now constructed organically without the assumptions required of the IRC.

This work also opens at least two new possible directions to address rates in chemical reactions and other activated systems: The first lies in the need for more direct comparison between NHIM rates [20] with MFPT rates. The second lies in the use of these approaches to obtain the rates for a time-dependent LiCN system driven by an electrical field so as to fully demonstrate the computational advantages of this approach in chemical reactions generally. Finally, reproducing the calculations of the MFPT on the Langevin equation with those for the general Langevin equation (with the observed memory time of the solvent) could also confirm the generality of our finding.

Appendix A: LiCN potential surface

The LiNC $\rightarrow$ LiCN PES according to Ref. [21] consists of two parts: a damped long-range energy plus a short-range energy.

The long-range part is composed of the electrostatic energy

$$ E_{\text{el}}(R, \vartheta) = \sum_{L=0}^{\infty} R^{-L-1} P_L(\cos \vartheta) \langle Q_{L,0} \rangle $$

and the induction energy

$$ E_{\text{ind}}(R, \vartheta) = \sum_{l_1,l_2=0}^{\infty} R^{l_1-l_2-2} \sum_{L=|l_1-l_2|}^{l_1+l_2} P_L(\cos \vartheta) C_{l_1,l_2,L} $$

Here, $P_L$ is the Legendre polynomial of order $L$, $\langle Q_{L,0} \rangle$ denotes the expectation value of the order-$L$ CN multipole moment, and $C_{l_1,l_2,L}$ are the induction coefficients. Numerical values for $\langle Q_{L,0} \rangle$ and $C_{l_1,l_2,L}$ are given in Table I. The damping is represented by

$$ F(R) = 1 - \exp \left[ -a(R - R_0)^2 \right] $$

with fit parameters $a = 1.5156$ a.u. and $R_0 = 1.9008$ a.u.

The short-range term can be written as

$$ E_{\text{SR}}(R, \vartheta) = \sum_{L=0}^{\infty} D_L(R) P_L(\cos \vartheta) $$

### Table I. Expectation values $\langle Q_{L,0} \rangle$ of the CN$^-$ multipole moments used in Eq. (A1) and induction energy coefficients $C_{l_1,l_2,L}$ used in Eq. (A2). Originally published in Refs. [21] and (partially) [23]. The bold value $C_{2,1,3}$ differs from the original publication.

| $L$ | $\langle Q_{L,0} \rangle$ | $C_{1,1,L}$ | $C_{2,1,L}$ | $C_{2,2,L}$ | $C_{3,1,L}$ | $C_{3,2,L}$ | $C_{3,3,L}$ |
|-----|-----------------|-------------|-------------|-------------|-------------|-------------|-------------|
| 0   | $-1.00$         | $-10.53$    | $-57.49$    | $-458.2$    |             |             |             |
| 1   | $-2.2151$       | $-10.31$    | $-101.45$   |             |             |             |             |
| 2   | $-3.414$        | $-3.17$     | $-35.71$    | $-35.56$    | $-353.7$    |             |             |
| 3   | $-3.819$        | $1.866$     | $-37.62$    |             |             |             |             |
| 4   | $-15.84$        | $5.23$      | $5.95$      |             | $-112.6$    |             |             |
| 5   | $-14.29$        |             | $-14.23$    |             |             |             |             |
| 6   | $-43.82$        |             |             |             |             |             | $-108.3$    |

### Table II. Parameters $A_L$, $B_L$, and $C_L$ found for the analytical expression of the short-range interaction in Eq. (A5) as fitted to the potential originally published in Ref. [21]. The bold value $C_2$ differs from that in the original publication, but all of the other values are the same.

| $L$ | $A_L$ | $B_L$ | $C_L$ |
|-----|-------|-------|-------|
| 0   | $-1.38321$ | $0.14001$ | $0.207892$ |
| 1   | $-2.95719$ | $1.479777$ | $-0.011613$ |
| 2   | $-4.74203$ | $1.81199$ | $-0.017181$ |
| 3   | $-1.88853$ | $1.28750$ | $0.027728$ |
| 4   | $-4.41433$ | $2.32297$ | $-0.070693$ |
| 5   | $-4.02565$ | $2.77538$ | $-0.137720$ |
| 6   | $-5.84259$ | $3.48085$ | $-0.186331$ |
| 7   | $-2.61681$ | $2.65559$ | $-0.005882$ |
| 8   | $-6.34466$ | $4.34498$ | $-0.152914$ |
| 9   | $15.2023$  | $-6.54925$ | $1.302568$  |
LiCN isomerization

This curve differs from what can be obtained using the parameters published in the same article (dashed line). Modifying the parameters as detailed in Sec. 3 yields much better agreement, as illustrated by the solid line.

\[ D_L(R) = \exp(-A_L - B_L R - C_L R^2) \]  

Numerical values for the fit parameters \( A_L, B_L, \) and \( C_L \) are given in Table I.

Combining long and short-range energies, the final PES reads

\[ V(R, \vartheta) = \left[E_{\text{ol}}(R, \vartheta) + E_{\text{ind}}(R, \vartheta)\right]F(R) + E_{\text{SR}}(R, \vartheta). \]  

We noticed a discrepancy while comparing the minimum energy path calculated using the parameters published in Ref. [21] with Fig. 2 from Ref. [21] As shown in Fig. 6, the two curves differ visibly. Two parameters in the original source code[28] differ significantly from the originally published values, possibly due to errors introduced during the paper’s production process. The correct values—shown in bold in Tables I and II—yield a much better agreement with Fig. 2 from Ref. [21].

An implementation of the potential in the Python programming language using the same parameters as the original source code[29] can be found on GitHub.[30]

**Appendix B: Approximative rate formulas**

Most approximative rate formulas follow the Arrhenius form[31]

\[ k(T) = \nu(T) \exp\left(-\frac{E^\ddagger}{k_B T}\right) \]  

where \( \nu \) is a possibly temperature-dependent prefactor, \( E^\ddagger \) is the reaction’s barrier height or activation energy, \( k_B \) is the Boltzmann constant, and \( T \) is the temperature. In the following we present two important TST variants of this equation.

### 1. Polanyi–Wigner rate

One of the earliest results of TST is the unimolecular rate equation derived, amongst others, by Polanyi and Wigner in 1928[32] It follows the Arrhenius rate law (B1) with the pre-exponential factor \( \nu(T) \) given by the vibrational frequency \( \omega_0/(2\pi) \) of the reactant well[33]

\[ k(T) = \frac{\omega_0}{2\pi} \exp\left(-\frac{E^\ddagger}{k_B T}\right), \]  

where \( E^\ddagger \) is the internal energy difference between the barrier and the reactant of the isolated system. This equation has been derived in various contexts. It can, e.g., be recovered from the underdamped regime \( \omega^2 k_B T / E^\ddagger \ll \gamma \ll \omega^\ddagger \) of Kramers’s medium-to-high-viscosity rate[34]

\[ k(T) = \left[\sqrt{\left(\frac{\gamma}{2\omega^\ddagger}\right)^2 + 1} - \frac{\gamma}{2\omega^\ddagger}\right] \frac{\omega_0}{2\pi} \exp\left(-\frac{E^\ddagger}{k_B T}\right), \]  

where \( \gamma \) is the friction and \( \omega^\ddagger \) is the inverse barrier frequency. Equation (B2) can therefore be seen as a TST rate which is an upper bound for the rate at the turnover in Kramers’s original theory for a solvated reaction.[35]

### 2. Eyring–Polanyi rate

The usual (or modern) form for the classical TST rate equation is given by the Eyring–Polanyi equation[36,37]

\[ k(T) = \frac{\kappa k_B T}{\hbar} \exp\left(-\frac{\Delta G^\ddagger}{k_B T}\right) \]  

with \( \kappa = 1 \), where \( \kappa \) is the transmission coefficient, \( \hbar \) is Planck’s constant, and \( \Delta G^\ddagger \) is the Gibbs energy of activation in the context of a solvent.

The Gibbs energy of activation can be approximately determined from the enthalpy of activation \( \Delta H^\ddagger \) via

\[ \Delta G^\ddagger = \Delta H^\ddagger - T \Delta S^\ddagger, \]  

where \( \Delta S^\ddagger \) is the entropy of activation. In turn, the enthalpy of activation for an unimolecular gas-phase reaction can be written as

\[ \Delta H^\ddagger = E^\ddagger - k_B T. \]  

The energy of activation \( E^\ddagger \) and the entropy of activation \( \Delta S^\ddagger \) can finally be determined from the minimum energy path and the potential of mean force of the reaction[31,38] by equating the latter with the Gibbs energy.

The transmission coefficient \( \kappa \) in the Eyring–Polanyi equation describes the fraction of states that cross the DS between reactants and products at most once, i.e., those that do not recross. This quantity cannot be determined from straightforward statistical mechanics and is therefore of great interest[39] in the general case. In TST, it is assumed to be approximately one[40] This approximation is valid if the temperature is not too high or if the friction is sufficiently strong.
ACKNOWLEDGMENTS

Useful discussions with Thomas Bartsch are gratefully acknowledged. The German portion of this collaborative work was partially supported by the Deutsche Forschungsgemeinschaft (DFG) through Grant No. MA1639/14-1. The Spanish portion was supported by the Spanish Ministry of Science, Innovation and Universities (MICIU) under Contract No. PGC2018-093854-B-100. The US portion was partially supported by the National Science Foundation (NSF) through Grant No. CHE 1700749. This collaboration has also benefited from support by the European Union’s Horizon 2020 Research and Innovation Program under the Marie Sklodowska-Curie Grant Agreement No. 734557.

DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

REFERENCES

1P. Hänggi and P. Talkner, “Non-markov processes: the problem of the mean first passage time,” Z. Physik B 45, 79–83 (1981)
2R. Müller, P. Talkner, and P. Reimann, “Rates and mean first passage times,” Physica A 247, 338–356 (1997)
3P. Reimann, G. J. Schmid, and P. Hänggi, “Universal equivalence of mean first-passage-time and Kramers rate,” Phys. Rev. E 60, R1 (1999)
4S. Redner, A Guide to First-Passage Processes (Cambridge University Press, 2001).
5A. L. Vega, R. Giannas, and S. Miret-Artés, “Mean first passage time and the Kramers turnover theory in activated atom-surface diffusion,” Phys. Chem. Chem. Phys. 4, 4885 (2002).
6P. D. Shepherd and R. Hernandez, “An optimized mean first passage time approach for obtaining rates in activated processes,” J. Chem. Phys. 117, 9227–9233 (2002).
7S. Park, M. K. Sener, D. Lu, and K. Schulten, “Reaction paths based on mean first-passage times,” J. Chem. Phys. 119, 1313-1319 (2003).
8G. Brocks and J. Tennyson, “Ab initio rovibrational spectrum of LiNC and LiCN,” J. Mol. Spectrosc. 99, 263–278 (1983).
9R. M. Benito, F. Borondo, J-H. Kim, B. G. Sumpter, and G. S. Ezra, “Comparison of classical and quantum phase space structure of nonrigid molecules, LiCN,” Chem. Phys. Lett. 161, 60–66 (1989).
10F. Borondo, A. A. Zembekov, and R. M. Benito, “Quantum manifestations of saddle-node bifurcations,” Chem. Phys. Lett. 246, 421 (1995).
11F. Borondo, A. A. Zembekov, and R. M. Benito, “Saddle-node bifurcations in the linlc system: Classical aspects and quantum manifestations,” J. Chem. Phys. 105, 5068 (1996).
12A. A. Zembekov, F. Borondo, Zembekov, and R. M. Benito, “Semiclassical quantization of fragmented tori: Application to saddle-node states of linlc,” J. Chem. Phys. 107, 7934 (1997).
13J. C. Losada, R. M. Benito, and F. Borondo, “Frequency map analysis of the 3D vibrational dynamics of the LiCN/LiNC molecular system,” Eur. Phys. J. Spec. Top. 165, 183–193 (2008).
14D. Prado, E. G. Vergini, R. M. Benito, and F. Borondo, “Supercars in the LiNC-LiCN isomerization reaction,” Europhys. Lett. 88, 40003 (2009).
15G. E. Murgiá, D. A. Wisniacki, P. I. Tamborenea, and F. Borondo, “Control of chemical reactions using external electric fields. The case of the LiNC-LiCN isomerization,” Chem. Phys. Lett. 490, 356–361 (2010).
16P. L. García-Müller, R. Hernandez, R. M. Benito, and F. Borondo, “Detailed study of the direct numerical observation of the Kramers turnover in the LiNC=LiCN isomerization rate,” J. Chem. Phys. 137, 204301 (2012).
17P. L. García-Müller, R. Hernandez, R. M. Benito, and F. Borondo, “The role of the CN vibration in the activated dynamics of LiNC=LiCN isomerization in an argon solvent at high temperatures,” J. Chem. Phys. 141, 074312 (2014).
18A. Vergel, R. M. Benito, J. C. Losada, and F. Borondo, “Geometrical analysis of the LiNC vibrational dynamics: A stability geometrical indicator,” Phys. Rev. E 89, 022901 (2014).
19A. Junginger, P. L. García-Müller, F. Borondo, R. M. Benito, and R. Hernandez, “Solvated molecular dynamics of LiCo isomerization: All-atom argon solvent versus a generalized Langevin bath,” J. Chem. Phys. 144, 024104 (2016).
20M. Feldmaier, J. Reif, R. M. Benito, F. Borondo, J. Main, and R. Hernandez, “Influence of external driving on decays in the geometry of the LiCN isomerization,” J. Chem. Phys. 153, 084115 (2020).
21P. Essers, J. Tennyson, and P. E. S. Wormer, “An SCF potential energy surface for lithium cyanide,” Chem. Phys. Lett. 89, 223-227 (1982).
22Y. Nagahata, R. Hernandez, and T. Komatsuzaki, “Phase space geometry of isolated to condensed chemical reactions,” J. Chem. Phys. 155, 210901 (2021).
23E. Pollak, H. Grabert, and P. Hänggi, “Theory of activated rate processes for arbitrary frequency dependent friction: Solution of the turnover problem,” J. Chem. Phys. 91, 4073–4087 (1989).
24H. A. Kramers, “Brownian motion in a field of force and the diffusional model of chemical reactions,” Physica (Utrecht) 7, 284–304 (1940).
25P. L. García-Müller, F. Borondo, R. Hernandez, and R. M. Benito, “Solvent-induced acceleration of the rate of activation of a molecular reaction,” Phys. Rev. Lett. 101, 178302–01-04 (2008).
26A. P. Pollak and E. J. Reed, “Coherent chemistry with THz pulses: Ultrafast field-driven isomerization of LiNC,” Phys. Rev. A 91, 052706 (2015).
27R. Kubo, “The fluctuation-dissipation theorem,” Rep. Prog. Phys. 29, 255–284 (1966).
28P. Hänggi, “Escape from a metastable state,” J. Stat. Phys. 42, 105–148 (1986).
29J. E. Straub and B. J. Berne, “Energy diffusion in many-dimensional Markovian systems: The consequences of competition between inter- and intramolecular vibrational energy transfer,” J. Chem. Phys. 85, 2999–3006 (1986).
30P. Zwanig, “Comments on a paper by Straub, Borkovec, and Berne,” J. Chem. Phys. 86, 5801–5803 (1987).
31B. J. Berne, M. Borkovec, and J. E. Straub, “Classical and modern methods in reaction rate theory,” J. Phys. Chem. 92, 3711–3725 (1988).
32P. Hänggi, P. Talkner, and M. Borkovec, “Reaction-rate theory: Fifty years after Kramers,” Rev. Mod. Phys. 62, 251–341 (1990) and references therein.
33J. E. Straub, M. Borkovec, and B. J. Berne, “Shortcomings of current theories of non-Markovian activated rate processes,” J. Chem. Phys. 83, 3172–4 (1985).
34J. E. Straub, M. Borkovec, and B. J. Berne, “Non-Markovian activated rate processes: Comparison of current theories with numerical simulation data,” J. Chem. Phys. 84, 1788–1794 (1986).
35P. L. Garciá-Muller, J. Hill, Free Energy Transduction and Biochemical Cycle Kinetics (Springer New York, 1989).
36H. Eyring, “The activated complex in chemical reactions,” J. Chem. Phys. 3, 107–115 (1935).
