Non-adiabatic effects within a single thermally averaged potential energy surface: Thermal expansion and reaction rates of small molecules

J. L. Alonso, A. Castro, J. Clemente-Gallardo, P. Echenique, J. J. Mazo et al.

Citation: J. Chem. Phys. 137, 22A533 (2012); doi: 10.1063/1.4747699
View online: http://dx.doi.org/10.1063/1.4747699
View Table of Contents: http://jcp.aip.org/resource/1/JCPSA6/v137/i22
Published by the American Institute of Physics.
Non-adiabatic effects within a single thermally averaged potential energy surface: Thermal expansion and reaction rates of small molecules

J. L. Alonso,1,2,3 A. Castro,2,3,4,5 J. Clemente-Gallardo,1,2,3,4 P. Echenique,1,2,3,4,6 J. J. Mazo,7 V. Polo,2,3,8 A. Rubio,9,10 and D. Zueco5,7

1Departamento de Física Teórica, Universidad de Zaragoza, Pedro Cerbuna 12, E-50009 Zaragoza, Spain
2Instituto de Biocomputación y Física de Sistemas Complejos (BIFI), Universidad de Zaragoza, Mariano Esquillor s/n, Edificio I+D, E-50018 Zaragoza, Spain
3Unidad Asociada IQFR-BIFI, Mariano Esquillor s/n, Edificio I+D, E-50018 Zaragoza, Spain
4Zaragoza Scientific Center for Advanced Modeling (ZCAM), Universidad de Zaragoza, Mariano Esquillor s/n, Edificio I+D, E-50018 Zaragoza, Spain
5Fundación ARAID, Paseo María Agustín 36, E-50004 Zaragoza, Spain
6Instituto de Química Física Rocasolano, CSIC, Serrano 119, E-28006 Madrid, Spain
7Instituto de Ciencia de Materiales de Aragón, and Departamento de Física de la Materia Condensada, CSIC-Universidad de Zaragoza, E-50009 Zaragoza, Spain
8Departamento de Química Orgánica y Química Física, Universidad de Zaragoza, Pedro Cerbuna 12, E-50009 Zaragoza, Spain
9Nano-Bio Spectroscopy Group and ETSF Scientific Development Centre, Departamento de Física de Materiales, Universidad del País Vasco, E-20018 San Sebastián, Spain
10Centro de Física de Materiales CSIC-UPV/EHU-MPC and DIPC, E-20018 San Sebastián, Spain

(Received 29 May 2012; accepted 9 August 2012; published online 31 August 2012)

At non-zero temperature and when a system has low-lying excited electronic states, the ground-state Born–Oppenheimer approximation breaks down and the low-lying electronic states are involved in any chemical process. In this work, we use a temperature-dependent effective potential for the nuclei which can accommodate the influence of an arbitrary number of electronic states in a simple way, while at the same time producing the correct Boltzmann equilibrium distribution for the electronic part. With the help of this effective potential, we show that thermally activated low-lying electronic states can have a significant effect in molecular properties for which electronic excitations are oftentimes ignored. We study the thermal expansion of the Manganese dimer, Mn₂, where we find that the average bond length experiences a change larger than the present experimental accuracy upon the inclusion of the excited states into the picture. We also show that, when these states are taken into account, reaction-rate constants are modified. In particular, we study the opening of the ozone molecule, O₃, and show that in this case the rate is modified as much as a 20% with respect to the ground-state Born–Oppenheimer prediction. © 2012 American Institute of Physics. [http://dx.doi.org/10.1063/1.4747699]

I. INTRODUCTION

In the ab initio study of the behaviour of molecular systems, it is common to perform the Born–Oppenheimer separation of the nuclear and electronic parts of the molecular wavefunction. This approximation is based on the large difference between the masses of the electrons and the nuclei,1,2 and therefore becomes exact only in the classical limit for the nuclei. Computing the electronic energy spectrum for different positions of the nuclei, one obtains the different so-called potential energy surfaces (PESs)—one for the electronic ground state, and one for each excited state. Now the question arises, how may we use these PESs in order to produce accurate and convenient physical models? The simplest option is ground-state Born–Oppenheimer (gsBO), or typically just Born–Oppenheimer (BO). One may then consider the nuclei to be quantum particles and solve their corresponding Schrödinger equation, or take the classical limit and perform BO molecular dynamics. In any case, a model based on only one PES (usually the ground-state one) is an adiabatic approximation, based on the neglect of the non-adiabatic couplings.

However, at non-zero temperature and when a system has low-lying excited electronic states, these electronic states are involved in any chemical process, and their influence produces the so-called non-adiabatic effects. In this paper, we use a thermodynamically accurate generalization, introduced in Ref. 3, of the gsBO potential, built with the PESs of the lowest lying electronic states appropriately weighted to produce the correct Boltzmann equilibrium distribution of the electronic part, to specifically study the thermal expansion and reaction rates of small molecules. For both phenomena, a substantial amount of research is constructed on the implicit assumption of one electronic potential, which may be fitted to experimental results or computed ab initio. The possibility of electronic excitations is typically either ignored or handled by independently considering each of the excited PES (although we can mention Ref. 4 as a very recent exception to this general trend, with similar aims to the ones we pursue here). The thermally
it is shown that tunneling below the barrier only occurs for studies of hot molecules and which will be used throughout the document. In Sec. III A, we present the first application of the effective-potential technique: the study of the thermal expansion of the Managese dimer, Mn2; in Sec. III B, we discuss the influence of the inclusion of low-lying excited electronic states on the reaction rate of small molecules. Reaction rates are affected on the one hand by the energy barrier reduction. On the other hand, the curvature at the top of the barrier is smaller for our effective potential than for the gsPES. In this section we will also study the case of the opening of the ozone molecule, O3. Finally, in Sec. IV, we comment on the most important conclusions of this work and highlight some possible implications and future lines of research.

II. THEORETICAL FRAMEWORK

Let us have a quantum-classical system formed by N classical particles described by their Euclidean coordinates \(R := (R_1, \ldots, R_N)\) and momenta \(P := (P_1, \ldots, P_N)\), and n quantum particles described by an n-body wavefunction \(|\psi\rangle\). The starting point of our formalism is the assumption that the following is an accurate formula to compute canonical equilibrium expected values of quantum-classical observables \(\hat{O}(R, P)\):

\[
\langle \hat{O}(R, P) \rangle = \frac{\int dR dP \text{tr}(\hat{O}(R, P) e^{-\frac{H_{\text{th}}(R, P)}{k_B T}}} )}{\int dR dP \text{tr}(e^{-\frac{H_{\text{th}}(R, P)}{k_B T}}) },
\]

(2.1)

where \(k_B\) is the Boltzmann constant and \(T\) the temperature, \(dRdP\) denotes integration over all position and momenta in the appropriate ranges. The object \(\hat{H}(R, P)\) is the quantum-classical Hamiltonian, which, in the case of molecular systems, takes the form

\[
\hat{H}(R, P) := \hat{1} \sum_{K=1}^{N} \frac{\hat{p}_K^2}{2M_K} + \hat{H}_e(R),
\]

(2.2)

where \(\hat{1}\) denotes the identity matrix, \(M_K\) is the mass of the \(K\)th nucleus, and the electronic Hamiltonian, \(\hat{H}_e(R)\), contains all particle interactions and the electronic kinetic term (see Ref. 2 for an explicit expression).

It is also convenient to write Eq. (2.1) as

\[
\langle \hat{O}(R, P) \rangle = \int dR dP \text{tr}(\hat{O}(R, P) \hat{\rho}_{\text{eq}}(R, P)),
\]

(2.3)

in terms of a \((R, P)\)-dependent equilibrium density matrix, defined by

\[
\hat{\rho}_{\text{eq}}(R, P) := e^{-\frac{H_{\text{th}}(R, P)}{k_B T}} \int dR' dP' e^{-\frac{H_{\text{th}}(R', P')}{k_B T}}.
\]

(2.4)

As shown in Ref. 3, the justification of Eq. (2.1) for computing equilibrium expected values in quantum-classical
models stems on the one hand from plausibility arguments related to the classical limit of the behaviour of the nuclei, such as the ones given in Ref. 27. However, it can also be obtained as the zeroth order approximation (in the quantum-classical mass ratio) to the canonical equilibrium associated with a, in principle, more rigorous quantum-classical formulation based on the Wigner formalism, as shown by Kapral and Ciccotti28 and Nielsen et al.29 It may also be rationalized by entropic arguments.30 In any case, and as it can be seen in the references in Ref. 3, irrespective of how good an approximation Eq. (2.1) is for any given application—always a difficult question—it is often the desired, target expectation value when designing quantum-classical schemes.

The main realization in which the effective potential that we will use in this work is based is that, for observables which do not depend explicitly on the electronic degrees of freedom, i.e., which are of the form

\[ \hat{O}(R, P) = \hat{\bar{O}}(R, P), \]  

where \( O(R, P) \) is a number, the target expected value in Eq. (2.1) can be rewritten as

\[ \langle O(R, P) \rangle = \frac{\int dRdP \, O(R, P) e^{-\frac{\hat{H}(R, P)}{k_BT}}}{\int dRdP \, e^{-\frac{\hat{H}(R, P)}{k_BT}}} \]  

Now, \( H_{\text{eff}}(R, P, T) \) is a purely classical, \( T \)-dependent, effective Hamiltonian defined as

\[ H_{\text{eff}}(R, P, T) := -k_BT \ln \text{tr} e^{-\frac{\hat{H}(R, P)}{k_BT}} = \sum_{k=1}^{N} \frac{\hat{p}_{k}^2}{2M_k} - k_BT \ln \text{tr} e^{-\frac{\hat{H}(R)}{k_BT}} =: \sum_{k=1}^{N} \frac{\hat{p}_{k}^2}{2M_k} + V_{\text{eff}}(R; T), \]  

where we have used Eq. (2.2). In the last line, we have finally defined the promised, purely classical, \( T \)-dependent, \( P \)-independent, effective potential

\[ V_{\text{eff}}(R; T) := -k_BT \ln \text{tr} e^{-\frac{\hat{H}(R)}{k_BT}}, \]  

which can be used to describe the behaviour of the nuclei in a classical mechanical setting, producing the correct target equilibrium in Eq. (2.1) for classical observables—by construction—and including the influence of all the electronic excited states.

Indeed, if we consider the adiabatic basis \( \{|\psi_k(R)\rangle\} \), which diagonalizes the electronic Hamiltonian \( \hat{H}_e(R) \) for each fixed position of the nuclei,

\[ \hat{H}_e(R)|\psi_k(R)\rangle = E_k(R)|\psi_k(R)\rangle, \]  

being \( \{E_k(R)\} \) the corresponding PESs, i.e., the eigenvalues of the electronic Hamiltonian as a function of the nuclear positions, then we can rewrite the effective potential in Eq. (2.8) as

\[ V_{\text{eff}}(R; T) = -k_BT \ln \sum_k e^{-\frac{E_k(R)}{k_BT}}, \]  

\[ = E_0(R) - k_BT \ln \left[ 1 + \sum_{k>0} e^{-\frac{\Delta E_{10}(R)}{k_BT}} \right], \]  

where we have defined

\[ \Delta E_{10}(R) := E_k(R) - E_0(R). \]  

The expression in Eq. (2.10) explicitly shows the difference between the ground-state PES, \( E_0(R) \), i.e., the gsBO potential, and the effective potential \( V_{\text{eff}}(R; T) \) introduced in Ref. 3 and used in this work. In particular, it is worth remarking that

- At \( T = 0 \), our effective potential becomes the gsBO one. Indeed, it is easy to see from Eq. (2.10) that

\[ \lim_{T \to 0} V_{\text{eff}}(R; T) = E_0(R), \quad \forall R. \]  

- The same reasons that produce the previous result allow us to include in the sum in Eq. (2.10) only the lowest lying electronic states and still get a good enough approximation to the exact expression if the temperature is low compared to the states we are neglecting, i.e., if \( k_BT \ll \Delta E_{10}(R) \) for them. This fact will be used in the practical cases presented in Sec. III.

- It can be seen from Eq. (2.10) that an exact property of the effective potential is that it is strictly lower than the gsPES, i.e., \( V_{\text{eff}}(R; T) \leq E_0(R), \forall R, T \). However, since an additive constant in a potential energy is not measurable, it must be noticed that this inequality is relevant only inasmuch the difference \( E_0(R) - V_{\text{eff}}(R; T) \) actually depends on \( R \). See Secs. III A and III B for concrete examples of this situation.

- As we discussed in Ref. 3, from Eq. (2.10), we see that, if the second derivative of \( \Delta E_{10} \) at a barrier top \( q_B \) verifies

\[ \frac{\partial^2 \Delta E_{10}}{\partial q^2}(q_B) \left( 1 + e^{-\frac{\Delta E_{10}(q_B)}{k_BT}} \right) > \frac{1}{k_BT} \left( \frac{\partial \Delta E_{10}}{\partial q}(q_B) \right)^2, \]  

then we can prove

\[ \left| \frac{\partial^2 V_{\text{eff}}}{\partial q^2}(q_B; T) \right| < \left| \frac{\partial^2 E_0}{\partial q^2}(q_B) \right|, \]  

\[ \text{i.e., the curvature of the effective potential at the barrier top is smaller than the one associated with the gsPES. In avoided crossings—a very interesting general case, and the one actually studied in Sec. III B—since the first excited state approaches the gsPES and then recedes from it, we have that the derivative \( (\partial \Delta E_{10}/\partial q)(q_B) \) will be approximately zero and the condition in Eq. (2.13) will be approximately satisfied, together with Eq. (2.14).} \]
non-adiabatic couplings. This is a consequence of the fact that the purpose of the effective potential is the computation of averages at the canonical equilibrium given by Eq. (2.4), and not the dynamics that may lead to it. The non-adiabatic couplings are necessary to carry the system to equilibrium—by providing the necessary channels to couple the various electronic states. Because of this, we are considering that the difference between these averages and the ones that would be obtained by considering the gsPES only is a non-adiabatic effect. However, the equilibrium averages predicted by Eq. (2.1) can be obtained without explicit consideration of the couplings. The magnitude of those couplings might be very relevant to compute the speed of the thermalization: small couplings may necessitate long thermalization times, but those analyses are beyond the scope of the present work.

In this respect, also note that all excited electronic states are included in the definition of the effective potential—because all states are included in Boltzmann’s equilibrium formula, regardless of how they may be coupled by external fields or, in the quantum-classical case, by the non-adiabatic couplings. The weight of each state is entirely determined by its energy. Any state is present in the averaging, even if its couplings to the ground state and to any other accessible state are zero. This is an example of ergodic difficulties, and obviously, a dynamical averaging would not include such a state unless it is already included in the initial state sampling.

The straightforward application of our scheme would then be inadequate if one is interested in computing a “restricted equilibrium” average, in which a state (or full subspace of states) is known to be absent, due to symmetry rules. The “experimental average” would not contain those states, even if the true canonical ensemble average does. However, in this situation it would be easy to correct our scheme by simply not including the forbidden states in the formulas.

III. RESULTS
A. Non-adiabatic effects on the thermal expansion of the \( \text{Mn}_2 \) dimer

Theoreticians usually identify the “molecular structure” with the equilibrium structure, i.e., the point determined by the absolute minimum of the ground-state Born-Oppenheimer potential energy surface (gsPES). This point in \( \mathbb{R}^{3N} \) space (\( N \) being the number of atoms) is well defined and has an easy intuitive meaning: the position occupied by the nuclei at equilibrium, if they had infinite mass (in which case they would be classical point particles). This geometry corresponds to a motionless molecule, that does not exist because molecules vibrate and move even at zero temperature. Therefore, this equilibrium structure is a theoretical concept that is not provided—at least not directly—by the experimental techniques utilized to probe molecular structure.

In fact, different experimental techniques yield different averaged results, whose value and meaning depend on the physical process involved in the measurement. For example, x-ray diffraction provides distances between the electronic charge distribution centroids. Gas-phase electron diffraction (ED), on the other hand, provides internuclear distances. Microwave spectroscopy measures moments of inertia, which can be directly related to nuclear distances to the molecular center of mass. Infrared, Raman, and ultraviolet spectrometries can also be used for complementary analysis. In all cases, the results are averaged over the populated rotational and vibrational molecular states—and, as we will show below, possibly over different electronic potential energy surfaces. Those techniques achieve a remarkable precision (of the order of 0.001 Å), but nevertheless each one of them provides different numbers.

In order to compare the results obtained in different experiments and in precise ab initio calculations, it is necessary to use a “common denominator” representation, which can very well be the equilibrium structure mentioned above, usually called the \( r_e \) structure. One must therefore know how to relate the experimental result to this concept. In an ED experiment at a given temperature, for example, one obtains the so-called \( r_g \) structure, an operational concept with no clear physical meaning. It can be related, however, to the thermally averaged internuclear distance, or \( r_g \) structure, which is of physical significance. It is not equivalent to the \( r_e \) structure, not even at 0 K, because of the vibrational and rotational (also called centrifugal) distortions. The relationship between \( r_e \) and an averaged structure such as \( r_g \) is not straightforward, but must be considered if we want to validate high precision theoretical \( ab \) initio calculations with experimental results or vice versa. This relationship was first considered by Bartell,, and has later been developed by several authors. It was soon realized that, in general, molecules expand as temperature increases, due to both the anharmonicity of the vibrations and to the centrifugal “force” that rotations exert on the structure.

In all these studies, an assumption is implicitly made: the electrons are adiabatically tied to their ground state, and therefore the analysis is performed by considering the gsPES only. However, as discussed in the Introduction, the existence of non-zero non-adiabatic couplings permits the system to visit electronic excited states, and the thermodynamic averaging should account for this possibility if the energy gap between the gsPES and the excited ones is not large in comparison with the thermal energy, \( k_B T \). If calculations and experiments are to be successfully compared, it is therefore necessary to consider the possible effect of the excited electronic states. We propose the use of our thermally averaged potential energy surface introduced in Sec. II for this purpose.

In principle, the rotational-vibrational averaging necessary to compute \( r_g \) must be performed assuming quantum-mechanical nuclei. This is obvious at 0 K, where the zero-point vibration would be completely absent in a classical treatment. However, we are mostly interested in the high-temperature regime—where the excited electronic states are expected to play a larger role and the system becomes classical. This fact can be exemplified by looking at closed—albeit approximate—theoretical formulae that exist for the simplest cases on which we concentrate in this study:
diatomic molecules. By truncating the Taylor expansion of the PES around the equilibrium bond length, i.e.,
\[ V(r) = \frac{1}{2} k_2 (r - r_e)^2 - k_3 (r - r_e)^3, \]  
(3.1)

Toyama et al.\(^{34}\) computed, in the quantum case, the temperature-induced variation of the average internuclear distance (with respect to \(r_e\)) as
\[ ⟨Δr⟩(T) := ⟨r⟩(T) - r_e = \frac{2k_B T}{k_2 r_e} + \frac{3k_3ω_e}{2k_2^2} \coth \left( \frac{ω_e}{2k_B T} \right), \]  
(3.2)

where \(ω_e\) is the vibrational frequency at equilibrium (\(ω_e = \sqrt{k_2/μ}\), where \(μ\) is the reduced mass of the two nuclei).

The first term in Eq. (3.2) is the centrifugal distortion; it arises from the harmonic potential term in Eq. (3.1), and can be considered as the variation in the average bond length caused by the overall molecular rotations. Interestingly, a classical treatment leads to the same expression for this term. The second term, on the contrary, has a genuinely quantum behaviour at low temperatures. In fact, it does not approach zero as \(T \to 0\) K, producing a zero-point vibration variation of the bond length with respect to \(r_e\). We can now take the classical limit by considering \(μ \to ∞\), in which case this second term becomes linear in \(T\): \(3k_3 k_B T / k_2^2\). But note that the same behaviour occurs if we take \(k_B T\) large with respect to \(ω_e\)—i.e., the system becomes classical for large enough temperatures.

In view of this, and since we are interested in the high-temperature regime in which our effective potential may significantly differ from the gsPES, we have used the classical approximation to compute the average bond length. To this end, we used Eq. (2.6), which for the dimer case (and considering that the function \(O(R, P)\) is in this case nothing else than the internuclear distance \(r = |R_1 - R_2|\)) reads
\[ ⟨r⟩_{eff}(T) = \int_0^L dr r^3 e^{-\frac{V_{gsPES}(r)}{k_B T}} \right] \int_0^L dr r^2 e^{-\frac{V_{gsPES}(r)}{k_B T}}. \]  
(3.3)

Note the presence of a somewhat arbitrary upper limit of integration \(L\). This value cannot be made arbitrarily large, since in the limit \(L \to ∞\), the equilibrium bond length is also infinite (assuming that, as it is always the case, the potential function remains finite at large internuclear distances). In fact, at equilibrium and at any non-zero temperature, a dilute gas of diatomic molecules in an infinite space does not contain dimers but isolated atoms. In the real world, dimers exist because there is always some form of container, or they are in a very long-lived metastable state. In practice, one must choose a value of \(L\) such that the results are almost insensitive to small variations of it—acknowledging that, if \(L\) is increased to very large values, the large value of \(⟨r⟩(T)\) will start growing.

The question that we want to answer here is whether or not the use of the effective potential in Eq. (3.3) leads to significantly different results with respect to the results obtained using only the gsPES, i.e.,
\[ ⟨r⟩_0(T) = \int_0^L dr r^3 e^{-\frac{V_{gsPES}(r)}{k_B T}} \]  
(3.4)

The answer cannot of course be universal, and depends on the chosen system and the temperature regime observed. In order to illustrate the issue, we have concentrated on the Mn\(_2\) molecule; a Van der Waals weakly bound molecule and a specially difficult theoretical case,\(^{31,38,39}\) for which a good feedback between experiments and theory could help validate the conflicting theoretical results. As we shall show, the existence of very low-lying excited electronic states in the vicinity of the equilibrium distance has an important effect on \(⟨r⟩(T)\), and therefore it is crucial to consider them to make proper comparisons.

To build our averaged potential defined in Eq. (2.10), we depart from the potential energy curves provided by Tzeli et al.,\(^{31}\) computed from first principles with very accurate multireference variational calculations. We only consider the six almost-degenerate, lowest lying states, displayed in Fig. 1. We have adjusted these curves to Morse functions, i.e.,
\[ V(r) = D[e^{-2αr} - 2e^{-αr} + 1] + V_∞. \]  
(3.5)

By finding the best match for the parameters \(D, α, r_e\), and \(V_∞\), an almost perfect fit can be obtained with respect to the results of Tzeli et al. The plot clearly shows the reasons for choosing Mn\(_2\) in this study: the lowest lying states are extremely close to each other, and the potential well is very shallow.

The results are shown in Fig. 2. The two key curves are the ones denoted by \(⟨Δr⟩_{eff}\) and \(⟨Δr⟩_0\), which are the results obtained with Eqs. (3.3) and (3.4), respectively. The difference due to the use of the effective potential instead of the ground state one is notable. It is larger than the resolution of modern experimental techniques, even in the lower temperature range. One may then conclude that any assessment of the quality of a theoretical model based on a comparison...
This fact is illustrated in Fig. 3, where we have considered a much the position of the minima of the two curves differs. The gap should not be larger than a few mHa—which is easily understood since, at 300 K, $k_BT$ is approximately 1 mHa. But also note that, even if the gap is small, there is no change with respect to $\langle \Delta r \rangle_0$ if the positions of the minimum of the two curves do not differ ($\Delta r_e$ is small). In other words, if the two PES are merely a rigid vertical shift of each other, the thermally averaged PES is also a rigid vertical shift and nothing change.

B. Non-adiabatic effects on the reaction rate and the opening of ozone

Reaction-rate theory focuses on studying the behaviour at long times of systems with different equilibrium states. This is a subject of great interest in many biological, chemical, and physical problems. As noted by Arrhenius in 1889, the cornerstone of the theory is the $e^{-A/T}$ temperature dependence of the reaction rates. Such dependence can be understood in the framework of a transition-state theory where the system evolves as a function of a given reaction coordinate $q$ from the metastable state $A$ to the metastable state $C$ through an energy barrier $B$, being the activation constant related with the barrier energy $U = E_B - E_A$ (see Fig. 4).

FIG. 2. Calculated variation of the average internuclear distance as a function of temperature for Mn$_2$. $\langle \Delta r \rangle_{\text{eff}}$ is the result computed using, in Eq. (3.3), the effective potential including all six lowest lying states in Fig. 1, while $\langle \Delta r \rangle_0$ is only computed with the gsPES. $\langle \Delta r \rangle_{\text{eff}}$ is the third-order approximation to the quantum result, as calculated using Eq. (3.2), and $\langle \Delta r \rangle_{\text{class}}$ is its classical limit. $\langle \Delta r \rangle_{\text{arm}}$ is the centrifugal term, common to all cases.

FIG. 3. Variation of the average internuclear distance, at 300 K, for a fictitious dimer with two relevant PES, as schematically shown in the inset. The results are given as a function of the displacement of the excited PES with respect to the gsPES, in two directions: variation of the energy minimum $\Delta E_{10}$, and difference in the position of this minimum, $\Delta r_e$.

FIG. 4. Sketch for the usual reaction-rate problem, in terms of a generic reaction coordinate $q$: Two metastable states at $q_A$ and $q_C$ are connected by a barrier with height $U = E_B - E_A$ and maximum at $q_B$. The curvatures are also shown.
In general the reaction rate can be written as

\[ r = k \, e^{-U/k_B T}, \]  

(3.6)

where the prefactor \( k \) depends on the temperature \( T \), the friction coefficient or "damping" of the system, and the details of the potential energy function. An accurate estimation of this prefactor has been shown to be a formidable task and many articles have been devoted to this end—deserving an special mention the celebrated one by Kramers.\(^{41}\) See also Refs. 23, 42, and 43.

As it has been shown in Ref. 3 and summarized in Sec. II, the inclusion of excited electronic levels becomes important at certain temperatures for obtaining suitable PESs of molecular systems, different from the gsBO one. The object of this section is to consider the effect of these excited states on the thermal-activation rate calculations.

Let us denote by \( r_{\text{eff}} \) the activation rate for the effective potential \( V_{\text{eff}}(q; T) \) in Eq. (2.10), and by \( r_0 \) the rate for the gsPES, \( E_0(q) \) (both of them expressed as a function of the reaction coordinate \( q \)). The main effect of the inclusion of the new terms associated with the excited electronic states is a reduction of the energy barrier from \( U_0 = E_0(q_B) - E_0(q_A) \) to \( U_{\text{eff}} = V_{\text{eff}}(q_B^e) - V_{\text{eff}}(q_A^e) \). We will have in general \( q_A^e \approx q_A \) and \( V_{\text{eff}}(q_B^e) \approx E_0(q_A) \), and thus the change in the energy barrier is given by the change at the potential maximum. Regarding the activation rate, in the simplest picture, we find that \( r_{\text{eff}} \approx e^{-U_{\text{eff}}/k_B T} \) and \( r_{\text{eff}}/r_0 = e^{-(U_{\text{eff}}-U_0)/k_B T} \). Therefore, if \( U_{\text{eff}} \approx U_0 \) is large enough, the effect of the energy barrier reduction on the activation rate will be important. In addition to this effect, it is also important to realize that the activation rate will also show a deviation from the expected \( e^{-U/k_B T} \) temperature dependence due to the fact that \( U_{\text{eff}} \) itself depends on \( T \). With this in mind, the importance of the excited electronic states can be studied looking for deviations of \( r(T) \) from its expected dependence.

A more detailed analysis must take into account the influence of the prefactor \( k \) too. To this end, some approximations need to be done. Let us place our problem in the context of the large-barrier (\( U/k_B T \gg 1 \)) and strong-friction limit. In such a situation, we learnt from Kramers that\(^{41}\)

\[ r_{\text{KHD}} \approx \dfrac{\omega_A}{\gamma} \sqrt{\dfrac{k_B T}{2\pi}} \left\{ \int_{q_A}^{q_B} e^{U(q)/k_B T} dq \right\}^{-1}, \]  

(3.7)

where \( U(q) := V(q) - V(q_A) \), being \( V(q) \) the appropriate potential. Here, \( \omega_A \) is associated with the curvature of the potential at point \( A \), \( \gamma \) is the friction coefficient, and KHD stands for Kramers high-damping limit.

This formula can be used directly to calculate the reaction rate by performing the integral numerically, and this is what we will do in this section. However, before doing that, let us introduce two simple approximations that are instructive and give some insights about the general features of the problem.

For a large barrier, a narrow region around the maximum gives most of the contribution to the integral. In many cases we can use the so-called parabolic barrier approximation:\(^{41}\) \( U_{\text{near B}} \)

\[ \simeq U_B - \dfrac{1}{2} \omega_B^2 (q - q_B)^2. \]  

Then

\[ r_{\text{KHD}}^{\text{pb}} \simeq \dfrac{\omega_A \omega_B}{2\pi \gamma} e^{-U_B/k_B T}. \]  

(3.8)

In this situation, we have

\[ \dfrac{r_{\text{eff}}}{r_0} \approx \dfrac{\omega_B^{\text{eff}}(T)}{\omega_B} e^{-(U_{\text{eff}}(T)-U_0)/k_B T}. \]  

(3.9)

Now, if \( V_{\text{eff}} \sim E_0 \) beyond the barrier and \( V_{\text{eff}} < E_0 \) at the barrier, we expect \( \omega_B^{\text{eff}} < \omega_B \) (see also the end of Sec. III). Hence, the prefactor effect opposes the exponential one: The rate will typically diminish because of the changes in the prefactor, but it will typically increase due to the changes in the barrier. In any case, given its exponential dependence, the effect of the barrier reduction is expected to be dominant in most cases.

Special care must be taken if the region of the potential close to the barrier (i.e., the one that contributes significantly to the integral in Eq. (3.7)) cannot be accurately approximated by a parabolic function around its maximum. Another common approximation to compare with is given by a cusp barrier

\[ U_{\text{near B}} \approx \begin{cases} U_B + m_1 (q - q_B) & \text{if } q \leq q_B, \\ U_B - m_2 (q - q_B) & \text{if } q > q_B, \end{cases} \]  

(3.10)

where \( m_1, m_2 > 0 \) are the slopes at each side of the barrier. In this case, the activation rate can be written as

\[ r_{\text{KHD}}^{\text{cb}} \approx \dfrac{\omega_A}{2\pi \gamma} \sqrt{\dfrac{2\pi}{k_B T}} \lambda e^{-U_B/k_B T}. \]  

(3.11)

where \( \lambda := m_1m_2/(m_1 + m_2) \), and we have

\[ \dfrac{r_{\text{eff}}}{r_0} \approx \dfrac{\lambda^{\text{eff}}(T)}{\lambda} e^{-(U_{\text{eff}}-U_0(k_B T)}, \]  

(3.12)

In conclusion, we can write in both cases

\[ \dfrac{r_{\text{eff}}}{r_0} \approx b(T) e^{\delta(T)/k_B T}, \]  

(3.13)

where \( \delta(T) := U_0 - U_{\text{eff}}(T) > 0 \) is the barrier reduction, and \( b(T) \) accounts for the changes in the prefactor. For the simplest case where \( E_0(q) \) and \( V_{\text{eff}}(q) \) show a maximum at the same coordinate \( q_B \) (the position of the maximum is not affected by the new terms), it is easy to see that \( e^{\delta(T)/k_B T} \approx (1 + e^{-\Delta/k_B T}) \) with \( \Delta := E_1(q_B) - E_0(q_B) \), where we have only considered the first excited state \( E_1 \).

As a working case, we will consider now the case of the ring opening reaction of ozone, which has been previously studied in Ref. 3. Figure 5 shows the potential profile for ozone as a function of the opening angle \( \phi \) at different temperatures. Since this is enough for our purposes, we consider only the gsBO PES, \( E_0(\phi) \), and the first excited state one, \( E_1(\phi) \)—taken from Ref. 3, where they where computed using the complete active-space self-consistent field method.\(^{44}\) The effective potential \( V_{\text{eff}}(\phi) \) is constructed with them. As we can clearly see in the bottom graph, the potential profile is modified by the inclusion of the new term corresponding to \( E_1(\phi) \) only in a narrow region close to the barrier. Also, in this case, the potential barrier is close to 9000 K and thus \( U/k_B T \gg 1 \) for the temperature range of interest, which justifies the basic
approximations behind the formulae in this section. However, there exist a clear asymmetry between $E_0(\phi)$ and $E_1(\phi)$. As a consequence, the barrier location moves with $T$, and the barrier energy reduction $\delta(T)$ does not follow the expected $T$ dependence. In addition, as we can see in Fig. 5, neither the parabolic nor the linear cusp approximations will be suitable to fit the barrier profile close to the maximum. Hence, we have directly used Eq. (3.7) to numerically compute the $r_{\text{eff}}/r_0$ ratio of the system. The results are presented in Fig. 6, where we can see that, in spite that the small barrier reduction observed, the activation rate changes up to a 20% upon the inclusion of the excited electronic states. This indicates that the non-adiabatic effects associated with low-lying states should be included in any analysis of this kind if one aims for high accuracy in the predictions. Besides, in the same Fig. 6, we also plot the contributions to the ratio $r_{\text{eff}}/r_0$ coming from the changes both in the potential barrier height and shape. As expected the increase of the rate motivated by the barrier reduction is moderated by the prefactor change. The two effects work against each other, and the exponential dependence on the barrier overcomes the influence of the curvature.

IV. CONCLUSIONS

In this work, we have shown that the ground-state PES, as defined in the Born–Oppenheimer approximation and typically used for many applications in chemistry, physics, and materials science, is not the only electronic state that significantly contributes to the theoretical prediction and calculation of thermodynamic observables at non-zero temperature already for small molecular systems. Although this fact was probably expected by the reader, we provide actual numerical examples of relevant observables being significantly modified by the inclusion of thermally activated low-lying excited electronic states in real systems at not-too-high temperatures: The average bond length of the manganese dimer is shown to change in an amount which is accessible to modern experimental techniques, while the reaction rate of the ring opening of ozone is shown to change up to a 20%. Moreover, our compact effective potential, which includes any number of such states and which, despite its simplicity, is able to produce the correct Boltzmann weight for the electronic subsystem—a long sought property in the field of quantum-classical models.

Also, as discussed in Sec. II, a general result when using our effective potential is that, in any avoided-crossing situation, the curvature on the top of the barrier is smaller than the gsPES curvature. As shown in Ref. 25, if the curvature is smaller, the tunneling below the energy barrier will occur at lower temperature. Therefore, the inclusion of low-lying electronic states is important if one wants to adequately discriminate possible quantum-like behaviour of the nuclei from simple classical effects due to the direct influence of temperature on the potential—for example, in biological systems.25,26

Additionally, although the effective potential used in this work has been derived in Ref. 3 assuming classical nuclei and equilibrated electrons, it could also be used as an exter-
nal potential to perform calculations on quantum nuclei. This procedure would allow to study low temperatures, while still including a possible correction due to electronic excitations. Notice, however, that, although the use of the gsPES in the BO approximation as a potential for quantum nuclei can be rigorously justified (see, e.g., Eq. (17) in Ref. 45), in the case of the effective potential used in this work, its use in this manner is not justified in principle because of its intrinsic non-adiabatic origin.

Finally, it is also reasonable to expect that the use of our temperature-dependent effective potential provides new insights that could lead to answer the intriguing question of the thermodynamical stability of some diatomic dications, an issue we plan to address in the next future.

ACKNOWLEDGMENTS

We thank A. Rey for insightful comments and illuminating discussions that have greatly contributed to the work we present here. This work has been supported by the research grants E24/1 and E24/3 (DGA, Spain), FIS2008-01240, FIS2009-13364-C02-01, FIS2009-12648, and FIS2011-25167 (MICINN, Spain, cofinanced by FEDER funds), and ARAID and Ibercaja grants for young researchers (Spain). A.R. acknowledges funding from European Research Council Advanced Grant DYNamo (ERC-2010-AdG - 2181036), European Community project 217 del Gobierno Vasco (IT-319-07), ACI-Promociona (ACI2009-2181036), European Community project THEMA (Contract 219 No. 228539), Ikerbasque, and SGiker ARINA (UPV/EHU).

1M. Born and J. R. Oppenheimer, “Zur Quantentheorie der Molekeln,” Ann. Phys. (Leipzig) 84, 457 (1927).
2P. Echenique and J. L. Alonso, “A mathematical and computational review of Hartree–Fock SCF methods in quantum chemistry,” Mol. Phys. 105, 3057 (2007).
3J. L. Alonso, A. Castro, P. Echenique, V. Polo, A. Rubio, and D. Zueco, “Ab initio molecular dynamics on the electronic Boltzmann equilibrium distribution,” New J. Phys. 12, 083064 (2010).
4G. Mazzola, A. Zen, and S. Sorella, “Finite temperature electronic simulations beyond the Born-Oppenheimer approximation,” e-print arXiv:1205.4526.
5C. Zhu, A. W. Jasper, and D. G. Truhlar, “Non-Born-Oppenheimer Liouville-von Neumann dynamics. Evolution of a subsystem controlled by linear and population-driven decay of mixing with decoherent and coherent switching,” J. Chem. Theory Comput. 1, 527 (2005).
6D. G. Truhlar, “Decoherence in combined quantum mechanical and classical mechanical methods for dynamics as illustrated for non-Born–Oppenheimer trajectories,” in Quantum Dynamics of Complex Molecular Systems, edited by D. A. Micha, and I. Burghardt (Springer, Berlin, 2007), pp. 227–243.
7J. R. Schmidt, P. V. Parandekar, and J. C. Tully, “Mixed quantum-classical equilibrium: Surface hopping,” J. Chem. Phys. 120, 044104 (2008).
8G. A. Worh and L. S. Cebberbaum, Annu. Rev. Phys. Chem. 55, 127 (2004).
9T. J. Martinez, M. Ben-Nun, and R. D. Levine, J. Phys. Chem. 100, 7884 (1996).
10H. Nakamura, Nonadiabatic Transitions: Concepts, Basic Theories and Applications (World Scientific, 2002).
11M. Baer, Beyond Born–Oppenheimer: Conical Intersections and Electronic Nonadiabatic Coupling Terms (Wiley, 2006).
12O. Prezhdo, J. Chem. Phys. 111, 8366 (1999).
13J. L. Alonso, A. Castro, J. Clemente-Gallardo, J. C. Cuchí, P. Echenique, and F. Falceto, “Statistics and Nosé formalism for Ehrenfest dynamics,” J. Phys. A 44, 395004 (2011).
14It is also worth remarking that some common criticisms that stated that Ehrenfest is a fully coherent method, and thus it cannot account for such important effects as mixing and decoherence, have recently been challenged: J. L. Alonso, J. Clemente-Gallardo, J. C. Cuchí, P. Echenique, and F. Falceto, “Ehrenfest dynamics is purity non-preserving: a necessary ingredient for decoherence,” J. Chem. Phys. 137, 054106 (2012); e-print arXiv:1205.0885.
15L. S. Bartell, “Effects of anharmonicity of vibration on the diffraction of electrons by free molecules,” J. Chem. Phys. 23, 1219 (1955).
16Z. Varga, M. Hargittai, and L. S. Bartell, “On the thermal expansion of molecules,” Struct. Chem. 22, 111 (2010).
17Z. Varga, M. Hargittai, and L. S. Bartell, “On the thermal expansion of molecules: a sequel,” Struct. Chem. 22, 1065 (2011).
18L. S. Bartell, “Bond lengths of vibrationally hot AXn molecules. A simple treatment,” J. Chem. Phys. 70, 4581 (1979).
19L. S. Bartell, S. K. Doun, and S. R. Goates, “Inference of vibrational anharmonicity in hot SF6: An electron diffraction study,” J. Chem. Phys. 70, 4585 (1979).
20S. R. Goates and L. S. Bartell, “Electron diffraction studies of hot molecules. I. Observed and calculated thermal expansions of SF6, CF4, and SF6+,” J. Chem. Phys. 77, 1866 (1982).
21S. R. Goates and L. S. Bartell, “Electron diffraction studies of hot molecules. II. Anharmonic shrinkage effects in SF6, CF4, and SF6+,” J. Chem. Phys. 77, 1874 (1982).
22L. S. Bartell, W. Vunce, and S. R. Goates, “Electron diffraction studies of hot molecules. III. Stretching and bending anharmonicity in CF4Cl,” J. Chem. Phys. 80, 3923 (1984).
23P. Hänggi, P. Talkner, and M. Borkovec, “Reaction-rate theory: fifty years after Kramers,” Rev. Mod. Phys. 62, 251 (1990).
24D. G. Fleming, D. J. Arsenault, O. Sukhorukov, J. H. Brewer, S. L. Mielke, D. G. Truhlar, G. C. Schatz, B. C. Garrett, and K. A. Peterson, “Kinetics of the reaction of the heaviest hydrogen atom with H2, the + H2 + H + → He + H2 reaction: Experiments, accurate quantum calculations, and variational transition-state theory, including kinetic isotope effects for a factor of 36.1 in isotopic mass,” J. Chem. Phys. 135, 184310 (2011).
25J. P. Bothma, J. B. Gilmore, and R. H. McKenzie, “The role of quantum effects in proton transfer reactions in enzymes: quantum tunneling in a noisy environment?” New J. Phys. 12, 055002 (2010).
26S. S. Iyengar, I. Summer, and J. Jakowski, “Hydrogen tunneling in an enzyme active site: a quantum wavepacket dynamical perspective,” J. Phys. Chem. B 112, 7601 (2008).
27F. Mauri, R. Car, and E. Tosatti, “Canonical statistical averages of coupled quantum-classical systems,” Europhys. Lett. 24, 431 (1993).
28R. Kapral and G. Cicotti, “Mixed quantum-classical dynamics,” J. Chem. Phys. 110, 8919 (1999).
29S. Nielsen, R. Kapral, and G. Cicotti, “Statistical mechanics of quantum-classical systems,” J. Chem. Phys. 115, 5805 (2001).
30Let us assume that the probability distribution function of a hybrid quantum-classical system can indeed be represented by a “parameterized” density matrix ρ(R, P), where the parameters (R, P) are the classical variables. Likewise, any observable can be represented by a parameterized operator ˆO(R, P). The average of any observable can be computed as:

\[ \langle O \rangle = \text{Tr} \int dR \int dP \hat{O}(R, P) \rho(R, P). \]

Then, the \( \hat{\rho}(R, P) \) that maximizes the entropy, defined as

\[ S[\hat{\rho}] = -k_B \int dR \int dP \text{Tr} \left[ \hat{\rho}(R, P) \log \hat{\rho}(R, P) \right], \]

subject to the constraints \( \int dR \int dP \text{Tr} \hat{\rho}(R, P) = 1 \) and \( \langle H \rangle = E \), is precisely given by Eq. (2.4) in the paper.
31D. Tzeli, U. Miranda, I. G. Kaplan, and A. Mavridis, “First principles study of the electronic structure and bonding of mno2,” J. Chem. Phys. 129, 154310 (2008).
32K. Kuchitsu and L. S. Bartell, “Effect of anharmonic vibrations on the bond lengths of polyatomic molecules. I. Model of force field and application to water,” J. Chem. Phys. 36, 2460 (1962).
K. Kuchitsu and L. S. Bartell, “Effects of anharmonicity of molecular vibrations on the diffraction of electrons. II. Interpretation of experimental structural parameters,” J. Chem. Phys. 35, 1945 (1961).

M. Toyama, T. Oka, and Y. Morino, “Effect of vibration and rotation on the internuclear distance,” J. Mol. Spectrosc. 13, 193 (1964).

D. R. Herschbach and V. W. Laurie, “Influence of vibrations on molecular structure determinations. I. General formulation of vibration–rotation interactions,” J. Chem. Phys. 37, 1668 (1962).

V. Spiridonov, N. Vogt, and J. Vogt, “Determination of molecular structure in terms of potential energy functions from gas-phase electron diffraction supplemented by other experimental and computational data,” Struct. Chem. 12, 349 (2001).

B. S. Butayev, V. P. Spiridonov, A. S. Saakjan, A. Y. Nasarenko, and A. G. Gershikov, “Effects of centrifugal distortion on the distance and amplitude parameters of diatomic molecules,” J. Mol. Struct. 119, 295 (1985); online at http://dx.doi.org/10.1016/0166-1280(85)87007-X.

B. Wang and Z. Chen, “Magnetic coupling interaction under different spin multiplets in neutral manganese dimer: Caspt2 theoretical investigation,” Chem. Phys. Lett. 387, 395 (2004).

S. Yamamoto, H. Tatewaki, H. Moriyama, and H. Nakano, “A study of the ground state of manganese dimer using quasidegenerate perturbation theory,” J. Chem. Phys. 124, 124302 (2006).

S. Arrhenius, Z. Phys. Chem. (Leipzig) 4, 226 (1889).

H. Kramers, Physica (Utrecht) 7, 284 (1940).

E. Pollak and P. Talkner, Chaos 15, 026116 (2005).

V. I. Melnikov, Phys. Rep. 209, 1 (1991).

B. O. Roos, “The complete active space self-consistent field method and its applications in electronic structure calculations,” Adv. Chem. Phys. 69, 399 (1987).

N. L. Doltsinis and D. Marx, J. Theor. Comput. Chem. 1, 319 (2002).

I. Corral, A. Palacios, and M. Yanez, “Electronic structure and lifetimes of GaX2+ (X = N, O, F) in the gas phase. Unraveling stability trends,” Phys. Chem. Chem. Phys. 13, 18365 (2011).