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First principles semiclassical calculations of vibrational eigenfunctions

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Vibrational eigenfunctions are calculated on-the-fly using semiclassical methods in conjunction with ab initio density functional theory classical trajectories. Various semiclassical approximations based on the time-dependent representation of the eigenfunctions are tested on an analytical potential describing the chemisorption of CO on Cu(100). Then, first principles semiclassical vibrational eigenfunctions are calculated for the CO2 molecule and its accuracy evaluated. The multiple coherent states initial value representations semiclassical method recently developed by us has shown with only six ab initio trajectories to evaluate eigenvalues and eigenfunctions at the accuracy level of thousands trajectory semiclassical initial value representation simulations. © 2011 American Institute of Physics. [doi:10.1063/1.3599469]

I. INTRODUCTION

Quantum properties of molecular systems are encoded in the eigenvalues and eigenfunctions of the molecular Hamiltonian. Many fundamental properties of molecules can be inferred from features such as their eigenfunctions’ nodes and their eigenvalue spectra. For example, the nodal shapes for higher vibrational eigenstates in a normal mode set of coordinates reveal whether and how the modes are coupled, i.e., if the coupling is resonant. Furthermore, by comparing the nodes of the wave function to the histories of ensembles of classical trajectories, one can understand the different types of resonances between eigenmodes and gain insight about the nature of these resonances, for example, how “classical” these resonances are. Unfortunately exact quantum mechanical methods for computing eigenfunctions and eigenvalues are limited to systems comprising a few degrees of freedom and most of the methods rely on having a pre-computed analytical potential energy surface (PES).1–4 The computation of PES is usually a trade-off between human effort and accuracy. This is especially true when one is dealing with complex or floppy systems or bond-breaking processes. On the other hand, while one can employ classical methods to study large many-particle systems, these methods are unable to reproduce intrinsically quantum features: for example, they cannot predict the form of the quantum eigenfunctions. For these reasons, a desirable method should not depend on the analytical development of PES but nevertheless provide access to quantum properties.5–7 Such as the vibrational eigenfunctions. First-principles molecular dynamics (FPMD) approaches have evolved as an alternative to the pre-computation of PES and these allow for the inclusion of quantum effects in the nuclear dynamics.8–17 Although there has been great development in this field, to our knowledge, calculations of molecular eigenfunctions using on-the-fly approaches have not been explored.

In order to reach this goal, several semiclassical approximations to quantum nuclear dynamics will be adopted. An appealing semiclassical method for the computation of eigenfunctions should be able to provide accurate quantized solutions for polyatomic systems and be easy to apply, preferably in a black-box fashion. Furthermore, as mentioned before, if the method can be cast into an FPMD approach, one can avoid the explicit construction of the PES. Some of the original semiclassical methods were based on the Einstein-Brillouin-Keller (EBK) theorem.18 According to these methods an eigentrajectory with specific EBK quantization conditions is searched and a uniform approximation to be corrected with the Maslov phase at the caustic point discontinuity is used.19 This is equivalent to a static Jeffreys-Wentzel–Kramers–Brillouin (JWKB)-like formalism,20 in which the task of finding the multidimensional eigentrajectories seems to be quite involved.

As an alternative to EBK and JWKB approaches, as we expose below, the spectral quantum features can be better understood in terms of the underlying dynamics, in particular within the semiclassical approximation, since periodic orbits set a correspondence between stationary states and their corresponding dynamics. For example, Heller and co-workers have shown strong evidence of this correspondence between classical motion and quantum effects even for spacings of combinations of overtones.21 In particular, in studies also pioneered by Heller, the vibrational eigenfunctions have been calculated on a model potential using a single classical trajectory.22 These calculations used Gaussian wavepackets of frozen width, which is generally, a computationally desirable feature: The Fourier transform of a simple Gaussian integrand

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following a quasi-periodic trajectory leads directly to an approximation to the eigenfunctions of the system of study. In short, semiclassical methods are based on a linear superposition of Gaussian coherent states that lie along the quantized classical trajectories. Such approach is free of any caustic problems and the agreement with the exact calculations is good in the tunneling region. In principle, one can use any classical trajectory to calculate eigenfunctions. However as will be shown below, this will affect the eigenfunctions’ accuracy. Analytical considerations have shown that semiclassical methods can reproduce eigenfunctions of the two dimensional rigid rotor and of a particle in a box exactly, while for the Morse potential eigenstates, the agreement is accurate. Still analytically, the three-dimensional isotropic harmonic oscillator eigenfunctions are calculated by representing the eigenfunctions as the integral of a semiclassical wavepacket over the Lagrangian manifold corresponding to the desired state. In this formulation, a parametric dependence of the wavepacket on the variable describing the Lagrangian manifold corresponding semiclassically to the state of interest is used instead of a time dependence. This formulation is uniform, i.e., it is free of caustic singularities, and it is semiclassical since it approaches the exact wave function uniformly for small \( \hbar \) values (\( \hbar \rightarrow 0 \)). Exact eigenstates have been derived for several systems, including the hydrogen atom, where the semiclassical expression reproduced exactly a given electronic orbital state by integrating over the radial and angular parameters. Nevertheless, one should keep in mind that semiclassical theories remain an approximation to the full quantum mechanical picture and in general, only yield approximate eigenfunctions. To our knowledge, the only general procedure to estimate the error arising from semiclassical approximations other than a direct comparison with the exact quantum eigenfunctions, is a perturbative series correction of the propagator. In this paper, a set of semiclassical molecular dynamics methods will be employed to calculate vibrational eigenfunctions. First, the methods will be tested on analytical potentials where exact calculations can be performed, and then coupled with a first principles approach, in which the potential energy surfaces are computed on-the-fly using density functional theory (DFT). We employ a recent implementation of the original time-averaging filtering of the semiclassical initial value representation method (SC-IVR) (Ref. 36) and extend it for the purpose of calculating vibrational eigenfunctions. This method uses a suitable set of delocalized coherent states that resembles a linear combination of the eigenfunctions of the system to reproduce quantum spectral features. In this way, the multiple coherent states SC-IVR (MC-SC-IVR) mimics the multiple vibrational components and successfully reproduce well-defined spectra for the several systems which have been tested. The major advantages of the method are that very few trajectories can accurately reproduce SC-IVR spectra obtained with thousands of trajectories and further that it can easily be implemented into a first principles molecular dynamics calculation. More recently Roy et al. also used such delocalization of coherent states for the calculation of the vibrational power spectrum of formaldehyde. This is another example of first-principles SC-IVR, as well as it is the work of Pollak and Tatchen on the absorption spectrum of the same molecule. In Sec. II, we will review the expression for computing the eigenfunctions using wavepackets and semiclassical approximations. In Sec. III, we discuss the computation of eigenfunctions with different methods for an analytical potential. In Sec. IV, first principles semiclassical eigenfunctions are calculated for the carbon dioxide molecule and, finally, conclusions are drawn in Sec V.

II. SEMICLASSICAL APPROXIMATION FOR EIGENFUNCTION CALCULATIONS

A. Time-dependent formulation for eigenfunctions calculation

Usually, the time-independent approaches to eigenfunction and eigenvalue calculation are based on the diagonalization of the Hamiltonian, which is represented in a suitable basis set. The computational cost of these types of approaches grows exponentially with the dimensionality of the system. A time-dependent representation could be a valid alternative, since it opens new avenues to quantum approximation methods. Following, we review the basics of the time-dependent method for computing eigenfunctions. Consider a general non-stationary state represented by the wavepacket \( \Psi(x, t) \) with Schrödinger dynamics,

\[
i\hbar \frac{\partial}{\partial t} \Psi(x, t) = \hat{H} \Psi(x, t),
\]

the time-evolved wavepacket can be expressed in terms of a superposition of eigenstates \( \psi_n(x) \),

\[
\Psi(x, t) = \sum_n c_n(t) \psi_n(x) = \sum_n \langle \psi_n(x) | \Psi(x, t) \rangle \psi_n(x).
\]

By combining Eq. (2) with Eq. (1), the time-evolution can be described by the expansion coefficients \( c_n(t) \),

\[
\sum_n \frac{d}{dt} c_n(t) \psi_n(x) = \sum_n c_n(t) E_n \psi_n(x).
\]

By multiplying from the left by \( \langle \psi_m(x) \rangle \), one obtains the dynamical equations for the coefficients \( c_m(t) \),

\[
\frac{d c_m(t)}{dt} = E_m c_m(t)
\]

or, equivalently,

\[
c_m(t) = e^{-iE_m t/\hbar} c_m(0).
\]

By substituting Eq. (5) into Eq. (2), the formulation of a generic non-stationary state as the evolution in terms of (stationary) eigenstates is obtained,

\[
\Psi(x, t) = \sum_n e^{-iE_n t/\hbar} \langle \psi_n(x) | \Psi(x, 0) \rangle \psi_n(x).
\]

Equation (6) is inverted by applying a Fourier transform to both sides and using the Fourier representation for Dirac’s delta function in the energy domain. This results in the time-dependent representation of the \( n \)th eigenfunction,

\[
\psi_n(x) = \frac{1}{\langle \psi_n(x) | \Psi(x, 0) \rangle} \frac{1}{T} \int_{-T}^{+T} e^{iE_n t/\hbar} \Psi(x, t) dt,
\]
where $E_n$ are the eigenfunction energies, and $T$ is the total time the wavepacket is propagated for. By using time inversion symmetry, Eq. (7) can be computed more conveniently since it can be written as

$$\psi_n(x; T) \propto \frac{2}{T} \int_0^T dt \Re(\Psi(x, t)e^{iE_n t}). \quad (8)$$

By either guessing or having the eigenvalue $E_n$, and integrating the time-dependent wave function, the value at $x$ of the eigenfunction for that given eigenvalue can be retrieved. As one integrates up to larger $T$, the global eigenfunction is increasingly refined. If the trajectory is quasiperiodic, it is possible to obtain sufficiently converged eigenfunctions for a reasonable value of $T$.

Equation (8) expresses that any eigenstate can be obtained from a superposition of wavepackets, i.e., the same wavepacket at different times, when integrated using a Fourier phase corresponding to its eigenvalue. The Fourier phase is obtained from a superposition of wavepackets, i.e., the same wavepacket at peak frequencies of the power spectrum leads to constructive interference of the Fourier transforming the wavepacket at peak frequencies of $\omega_j$. The Fourier phase is represented by the total number of cycles $(\gamma_j)$ that this trajectory makes for each degree of freedom, as originally suggested by Heller and co-workers. Then, the expression for the frequency is

$$\omega_j = \frac{2\pi}{T_c} M_j. \quad (11)$$

The criteria for trajectory closure was set so strict that it closes only once within the total simulation time. The frequencies obtained were comparable to the harmonic estimate coming from the Hessian diagonalization at the global minimum. A suitable simulation time and trajectory closure criteria are very important since the correct phase calculation is crucial for the coherent superposition. The trajectory used to generate the coherent state should be taken long enough to adequately explore the entire manifold on which it lies. We found this semiclassical recipe for calculating the eigenfunctions simple and practical. It is, however, limited to a single classical trajectory.

**B. De Leon-Heller semiclassical eigenfunctions**

The original semiclassical quantization of vibrational Hamiltonians has been expressed in terms of action quantization. The Einstein-Brillouin-Keller quantization (EBK) corrected by a Maslov index is one of the seminal approaches. The EBK method implies searching for a trajectory whose action integrals are properly quantized. The energy of such a trajectory is the semiclassical eigenvalue. However, this approach results to be too cumbersome when a multidimensional trajectory search is needed.

The time-dependent formulation presented in Eq. (8) is suitable for implementation into a semiclassical molecular dynamics perspective. As shown by Heller and Davis, it is convenient to replace a trajectory that quantizes the system with a linear superposition of Gaussian coherent states that are generated from that trajectory. This strategy is suggested by Eq. (8) itself, where the eigenfunction is a convolution of a time evolved wavepacket. In our simulations, the wavepacket is composed of a coherent state part,

$$\langle x|p(t), q(t)\rangle = \prod_{j=1}^F \left(\frac{\gamma_j}{\pi}\right)^{1/4} \exp\left[-\frac{\gamma_j}{2}(x_j - q_j(t))^2 + ip_j(t)(x_j - q_j(t))\right], \quad (9)$$

where $F$ is the number of degrees of freedom, $q_j(t)$ and $p_j(t)$ are the position and momentum of the classical trajectory for the $j$th dimension, and $\gamma_j$ are the coherent state widths usually chosen to match the widths of the harmonic oscillator approximation to the wave function at the global minimum. The term in Eq. (9) is multiplied by a time-dependent phase term $e^{i\gamma_j t/\hbar}$ to give the semiclassical representation of the wavepacket. The phase term is crucial for the coherent buildup of the eigenfunction and it has the following form:

$$\gamma_j(t) = \int_0^T dq \cdot dp - E \cdot t + \frac{T}{2} \sum_{j=1}^F \omega_j, \quad (10)$$

where the first two terms are the classical action, a common ingredient in every semiclassical approximation. The time-evolved wavepacket is easily obtained by generating the phase space Gaussian center’s coordinates $(p(t), q(t))$ according to Newton’s equations, the classical action and the phase $\gamma_j(t)$ as in Eq. (10). Initial conditions are taken from the harmonic approximation. The frequency terms in Eq. (10) resemble the semiclassical Maslov phase correction. The frequency values $\omega_j$ were estimated by taking a multidimensional trajectory long enough to close on itself in a time $T_c$ and by counting the total number of cycles ($M_j$) that this trajectory makes for each degree of freedom, as originally suggested by Heller and co-workers. Then, the expression for the frequency is

$$\omega_j = \frac{2\pi}{T_c} M_j. \quad (11)$$

**C. SC-IVR eigenfunctions**

In the same spirit as the above discussion, one can formulate the SC-IVR expression for eigenfunction calculation by invoking the Heller-Herman-Kluk-Kay (HHKK) propagator. In the SC-IVR method, the propagator in $F$ dimensions is approximated by the phase space integral,

$$e^{-iS\hat{t}/\hbar} = \frac{1}{(2\pi\hbar)^F} \int dp(0) \int dq(0) C_i(p(0), q(0)) e^{iS(p(0), q(0))/\hbar}\langle p(t), q(t)|p(0), q(0)\rangle, \quad (12)$$

where $(p(t), q(t))$ are the set of classically evolved phase space coordinates, $S_i$ is the classical action,

$$S_i(p(0), q(0)) = \int_0^t dt' \left(\frac{p^2(t')}{2m} - V(q(t'))\right), \quad (13)$$

and $C_i$ is a pre-exponential factor. In the Heller-Herman-Kluk-Kay version of the SC-IVR, the prefactor is

$$\frac{1}{\sqrt{2}} \left| \frac{\partial q(t)}{\partial q(0)} + \frac{i\hbar}{\gamma} \frac{\partial q(t)}{\partial p(0)} + \frac{i}{\gamma} \frac{\partial p(t)}{\partial q(0)} \right| \quad (14)$$

Reference:

and the basis set is a product of $F$ one-dimensional coherent states of Eq. (9). For bound systems, no significant dependency on width variation was found. By using a $2F \times 2F$ symplectic (monodromy) matrix $M(t) \equiv (\partial(p, q)/\partial(p_0, q_0))$, one can calculate the pre-factor of Eq. (14) from $F \times F$-sized blocks and we monitored the accuracy of the classical approximate propagation by containing the deviation of its determinant from unity to be less than $10^{-6}$. Thus, the SC-IVR representation of the time evolved wavepacket is

$$
\Psi(x, t) = \frac{\int dp \int dq}{(2\pi\hbar)^2} C_i(p(0), q(0)) e^{iS(p(t), q(t))/\hbar} \langle p(t), q(t) | p(0), q(0) \rangle.
$$

Inserting Eq. (15) into Eq. (8), one finds the final SC-IVR expression for the eigenfunction used in this paper. The zero-time coherent-state overlap in Eq. (15) is the employed density for the Monte Carlo phase space integration. In order to gain further numerical stability for the calculation of the prefactor in Eq. (14), one can introduce a phase approximation along the lines of the separable approximation for the double times prefactor used in the time-averaging filtering. This way, with few trajectories we obtained accurate results for the H$_2$O molecule on a model potential and for the CO$_2$ molecule using an on-the-fly approach. Additionally, we obtained accurate spectra for the model potential describing the chemisorption process of CO on Cu(100) copper surface that will be used below. The agreement was excellent not only when dimensionality was reduced to the two stretching motions of a single molecule, but also when a set of four dipole coupled molecules were arranged in a monolayer fashion. Thus, the MC-SC-IVR method has proved to be really advantageous since the number of trajectories can be substantially reduced, while preserving an accuracy comparable to that obtainable with thousands of trajectory calculations. The MC-SC-IVR formulation of the eigenfunction is

$$
\psi_n(x) \propto \frac{\|e^{-iE_n t/\hbar} \sum_{i=0}^{N_{states}-1} e^{iS(p_i(0), q_i(0))/\hbar}}{T} \int_0^T dt e^{iE_n t/\hbar} C_i(p(0), q(0)) |p(t), q(t)\rangle dt,
$$

which is much less computationally demanding and still different from the De Leon-Heller expression. However, both methods avoid any divergence of the semiclassical propagator at the caustic or turning point, which has been one of the major problems of the application of the WKB approximation.

D. Multiple coherent states semiclassical initial value representation eigenfunctions

In a typical SC-IVR simulation, the Monte Carlo phase space integration has been tested to converge with a number of trajectories of the order of thousands. If one requires to propagate trajectories using first-principles dynamics, this number of trajectories is prohibitive. Recently, we developed a method called multiple coherent states SC-IVR (MC-SC-IVR) which reduces significantly the number of classical trajectories to only a few while still preserving good accuracy. The method was developed for power spectra calculations and consists in a SC-IVR strategy that enhances as much as possible the overlap between the reference state (whose time-evolution is Fourier transformed into the power spectrum) and the exact quantum eigenfunctions. In fact, one can think of projecting the power spectrum into the phase space portrait, where it is represented by a set of multidimensional closed “eigen-trajectories” with energy equal to the peaks of the spectrum (see Fig. 1 in Ref. 35). Then, it is clear that a set of trajectories which somehow mimics this distribution is more representative than a single ground state trajectory. For this method, it was found that it is not crucial to know the exact location of the “eigen-trajectories” turning points at the values of potential energy equal to the peaks of the power spectrum. This is so because the Gaussian spreading of each coherent states lying on the top of each “eigen-trajectory” is generally wide enough to include the peaks’ energy shell. In this way, with few trajectories we obtained accurate results for the $\text{H}_2\text{O}$ molecule on a model potential and for the CO$_2$ molecule using an on-the-fly approach. Additionally, we obtained accurate spectra for the model potential describing the chemisorption process of CO on Cu(100) copper surface that will be used below. The agreement was excellent not only when dimensionality was reduced to the two stretching motions of a single molecule, but also when a set of four dipole coupled molecules were arranged in a monolayer fashion. Thus, the MC-SC-IVR method has proved to be really advantageous since the number of trajectories can be substantially reduced, while preserving an accuracy comparable to that obtainable with thousands of trajectory calculations. The MC-SC-IVR formulation of the eigenfunction is

$$
\psi_n(x) \propto \frac{\|e^{-iE_n t/\hbar} \sum_{i=0}^{N_{states}-1} e^{iS(p_i(0), q_i(0))/\hbar}}{T} \int_0^T dt e^{iE_n t/\hbar} C_i(p(0), q(0)) |p(t), q(t)\rangle dt,
$$

which the initial phase space conditions are the equilibrium geometry $q(0)$ for the positions and the harmonic approximated momenta, namely, $p_{j,i}^2/2m = \hbar \omega_j (i + 1/2)$ for each $j$th degree of freedom of the $i$th trajectory. Eq. (18) reduces the phase space integral to a sum over a set of $N_{states}$, which we called “eigen-trajectories,” which are harmonically spaced in the energy domain. An analogous expression holds when the separable approximation is applied. In this way MC-SC-IVR includes the quantum mechanical delocalization by using a set of coherent states placed in a non-local fashion, while their centers are kept fixed during the entire simulation time.
III. MEASURING THE QUALITY OF THE EIGENFUNCTIONS

The most straightforward metric for the accuracy of the semiclassical eigenfunctions, is their overlap with numerically exact solutions,

\[ O = \int \langle \psi_{SC} (x) | \psi_{DVR} (x) \rangle \, dx, \quad (19) \]

where \( \psi_{SC} (x) \) is the normalized semiclassical eigenfunction calculated according to one of the approximations described above, while \( \psi_{DVR} (x) \) is what we consider the numerically exact quantum eigenfunction calculated with the discrete variable representation method in the Sinc function basis.\(^{55}\) Thus, the integral of Eq. (19) becomes a sum over all Discrete Variable Representation (DVR) grid points \( N_{DVR} \),

\[ O = \sum_{i=1}^{N_{DVR}} | \psi_{SC}^* (x_i) \psi_{DVR} (x_i) \Delta x_i |. \quad (20) \]

However, in case of degeneracy the comparison between eigenfunctions of the same eigenvalue calculated with different methods is not so straightforward. In this case, any combination of degenerate eigenfunctions is still an eigenfunction and in principle one does not know how a set of degenerate semiclassical eigenfunctions can be matched with the exact ones. One is tempted to introduce a mixing angle that combines the eigenfunctions with sine and cosine coefficients and represent the semiclassical eigenfunctions as rotations of the exact ones, however we found this procedure to be too cumbersome when dimensionality is increased. A simpler way to overcome this impasse is on one hand, to measure to which extent the semiclassical eigenfunction is an eigenfunction of the DVR Hamiltonian matrix and on the other to measure the completeness of the basis of each group of degenerate functions within the degenerate subspace.

To reach the first goal, the following expression was introduced:

\[ \varepsilon_1 = \frac{|\hat{H}_{DVR}|\psi_{SC} - E_{DVR}|\psi_{SC}|}{E_{DVR}}, \quad (21) \]

where \( \hat{H}_{DVR} \) is the DVR matrix representation of the Hamiltonian operator, \( |\psi_{SC}\rangle \) is the semiclassical eigenfunction evaluated at the DVR grid points and \( E_{DVR} \) is the exact quantum eigenvalue. In Eq. (21) the norm of the vector measuring the deviation of \( |\psi_{SC}\rangle \) from being the exact ket is calculated in the numerator and weighed respect to the value of the eigenvalue. If \( |\psi_{SC}\rangle \) is an eigenket, then \( \varepsilon_1 = 0 \). Moreover, if \( |\psi_{SC,1}\rangle \) and \( |\psi_{SC,2}\rangle \) are two degenerate precise eigenfunctions, then they should both have small values of \( \varepsilon_1 \), even if their overlap with \( |\psi_{DVR}\rangle \) is arbitrary. In order to appreciate how much small \( \varepsilon_1 \) should be for an eigenfunction to be accurate, one can use Eqs. (21) together with Eq. (20) for the ground state (which is not degenerate) and then compare this with the values of \( \varepsilon_1 \) in the case of degenerate states on the same grid points.

To reach the second goal, i.e., to prove that the set of degenerate semiclassical eigenfunctions is complete within the degenerate subspace, the following quantity was introduced:

\[ \varepsilon_2 = \sqrt{\frac{\langle \psi_{SC} | \hat{H}_{DVR} | \psi_{SC} \rangle}{E_{DVR}}}, \quad (22) \]

where the notation is the same as in Eq. (21). Further comments on the \( \varepsilon_2 \) parameter for a given eigenfunction \( |\psi_{SC,i}\rangle \) are necessary to appreciate the meaning of Eq. (22) better. One first notes that

\[ \varepsilon_{2,i}^2 = \frac{\langle \psi_{SC,i} | \hat{H}_{DVR} | \psi_{SC,i} \rangle}{E_{DVR,i}} \]

\[ = \frac{\sum_{l,n} \langle \psi_{SC,i} | \psi_{DVR,l} \rangle \langle \psi_{DVR,l} | \hat{H}_{DVR} | \psi_{DVR,n} \rangle \langle \psi_{DVR,n} | \psi_{SC,i} \rangle}{E_{DVR,i}}, \quad (24) \]

where the identity in terms of DVR complete basis set has been inserted twice. Then, by using the properties that \( |\psi_{DVR,i}\rangle \) is the exact \( i \)th eigenfunction of \( \hat{H}_{DVR} \) of eigenvalue \( E_{DVR,i} \), Eq. (24) is simplified to

\[ \varepsilon_{2,i}^2 = \frac{\sum_{n} |\langle \psi_{DVR,n} | \psi_{SC,i} \rangle|^2 E_{DVR,n}}{E_{DVR,i}}. \quad (25) \]

For non-degenerate cases, Eq. (25) is equal to Eq. (19) plus the contributions coming from the overlap of the \( i \)th semiclassical eigenfunctions with the exact eigenfunctions orthonormal to the \( i \)th one. Thus, if the semiclassical approximation is really accurate, Eq. (25) should give the same value as Eq. (19) does for non-degenerate eigenvalues.

Instead, when two states \( |\psi_{DVR,1}\rangle \) and \( |\psi_{DVR,2}\rangle \) are degenerate, they can both give a significant overlap with the semiclassical eigenfunction \( |\psi_{SC,i}\rangle \) which approximates one of the degenerate eigenstate. In Eq. (25), let us assume that \( |\psi_{SC,i}\rangle \) is good enough to be orthogonal to all exact eigenkets which do not belong to the considered degenerate subspace. In this case, if \( |\psi_{DVR,1}\rangle \) and \( |\psi_{DVR,2}\rangle \) are two eigenfunctions that span the degenerate subspace then we can represent the semiclassical eigenfunction as follows:

\[ |\psi_{SC,i}\rangle \approx \cos \omega |\psi_{DVR,1}\rangle + \sin \omega |\psi_{DVR,2}\rangle, \quad (26) \]

where \( \omega \) is the mixing angle. By inserting this approximation for \( |\psi_{SC,i}\rangle \) into Eq. (25) and considering that \( E_{DVR,1} = E_{DVR,2} = E_{DVR,i} \), one obtains \( \varepsilon_{2,i} = 1 \). Thus, the sum of the squares of the overlaps with the semiclassical
where \( r_{eq} \) is the equilibrium distance, and the interaction of CO with each copper atom is described by the following modified Morse C-Cu interaction potential:

\[
V_i(r_{\text{C}}, r_{\text{O}}, r_i) = A \exp(-\alpha |r_i - r_{0}|) + B \exp(-2\beta (|r_i - r_{\text{C}}| - r_{eq})) - 2B \cos^2(\beta (|r_i - r_{\text{C}}| - r_{eq})).
\]

(29)

The first term in Eq. (29) describes the oxygen-copper repulsion. The remainder of the terms in Eq. (29) represent the carbon-copper interaction. The molecular orientation is taken into account by the angle \( \phi \) between the C-O and the Cu-C bonds. The metal is represented by three layers of 36 (6x6) copper atoms arranged according to a fcc lattice structure. In this work, the molecular axis is fixed perpendicular to the copper surface and the resulting bidimensional molecular motion describes the internal CO mode at 2084 cm\(^{-1}\) and the stretching mode perpendicular to the surface at 353 cm\(^{-1}\). Potential parameters are reported in Ref. 37. In this system there is no degeneracy induced by symmetry considerations. However, given the small value of the external stretch mode, it can happen that overtones of different quanta are located very close in energy space.

The De Leon-Heller method described in Sec. II is applied to the calculation of the overlap according to Eq. (20). For each eigenfunction a classical trajectory is chosen with initial conditions given by the positions at the equilibrium geometry and momenta such that \( \sum p_i^2/2m = E \), where the sum is over each dimension. Whether the set of values \( E \) of the eigenvalues, used to calculate the initial momenta and the phase of Eq. (10), were calculated using a SC-IVR with a single trajectory or MC-SC-IVR, is indicated by the subscripts. For \( E_{\text{DVR}} \) the “exact” eigenvalues’ set was chosen, while for \( E_{\text{1traj}} \) the single trajectory (without separable approximation) spectra values were chosen.\(^{34} \) In Table I one can also find the values of the frequencies evaluated for each trajectory using Eq. (11). These are comparable with the harmonic values and they do not change significantly by changing the trajectories’ initial velocities. If the potential had been harmonic, the prefactor would indeed be the complex exponentiation of the mode frequency over \( \hbar \). Finally, in Table I we report an index labeled as “sum” which is indicative of the overall performance of the method. A closer look at Table I shows how the frequencies are the same within less than 1%, using both the set \( E_{\text{1traj}} \) and \( E_{\text{DVR}} \) as initial conditions. Instead, significant differences can be observed for the overlap values. Clearly, the exact set of energies gives more accurate results. This shows that the De Leon-Heller method can be considered accurate enough for most eigenfunction calculation purposes, when accurate eigenvalues are known.

In Table II single-trajectory results are reported, using Eq. (17). In the second column, the trajectory used for the eigenfunction calculations for all states is the same, i.e., the ground state harmonic trajectory. It is evident that this...
crude approach gradually fails as higher vibrational levels are reached, and already is quite inaccurate for the (2,2) state where the overlap is 90% too small. This shows how the error increases if one attempts to use a unique trajectory to calculate states which are further apart from the energy shell of the single trajectory (see Appendix A of Ref. 59). However, this trajectory shows to be quite accurate for the first vibrational state taken in this set of calculations, i.e., the (2,2) state, is exact within 1% of accuracy. In the following two columns, the overlaps for trajectories with harmonic and anharmonically corrected initial momenta at the energy eigenvalues are reported. For all these columns as for the one labeled “ground,” the energy eigenvalues were taken from the single trajectory power spectrum.\(^\text{34}\) From the second to the fourth column, the accuracy gradually increases, as indicated by the index “sum.” Nevertheless, for the (2,2) eigenstates the missed part of the overlap decreases from 57% to 40% when using the anharmonic correction. This result is still not satisfying. In the last column, the same calculations as in the fourth column are performed, but using a consistent (i.e., with the eigenvalue at the same semiclassical level of calculation) set of separable approximation results for the power spectra and eigenfunction evaluation according to Eq. (16). The separable approximation clearly generate poorer results than the ones reported on the fourth column, where this approximation was not invoked.

Finally Eqs. (18) and (16) are employed for eigenfunction calculations. The results are reported in Table III for different sets of trajectories and for different levels of approximation. These calculations are evidence of the substantial increase of accuracy due to the use of multiple trajectories. In the second column, we report the results of using the MC-SC-IVR approach, where the separable approximation was used both for the eigenfunctions and power spectrum calculations, in order to have a set of consistent results. These results are accurate for our purposes, as shown by the “sum” index. In particular the higher vibrational state taken in this set of calculation, i.e., the (2,2) state, is exact within 1% of accuracy. Exactly the same considerations are valid for a second set of MC-SC-IVR calculations reported on the following column. Here the separable approximation is not employed and the accuracy of the results is slightly less. In the column labeled “4000 traj.s,” the fully converged semiclassical limit is reached using a 4000-classical trajectory calculation for Eq. (15) with 4000 time steps of 10 a.u. each. Here the eigenvalues coming from MC-SC-IVR calculations\(^\text{35}\) are used, i.e., the same ones as in the previous column. Thus, the comparison between these two columns shows the accuracy of our MC-SC-IVR method for eigenfunction calculations respect to the original HHKK semiclassical formulation. Differences between the two sets of overlaps are very small and the MC-SC-IVR method, both with or without the separable approximation, can be used with only few trajectories (six in this case) as a good estimate of the actual semiclassical calculation for vibrational eigenfunctions. These results confirm the previous power spectra calculation ones,\(^\text{35}\) showing that the MC-SC-IVR can properly mimic quantum properties for this kind of bound states at the fully converged SC-IVR level of calculation, with much less effort. In the

<table>
<thead>
<tr>
<th>States</th>
<th>(\omega_1), (\omega_2)</th>
<th>(O(E_{\text{straj}}))</th>
<th>(O(E_{\text{straj}}))</th>
<th>(O(E_{\text{straj}}))</th>
<th>(O(E_{\text{straj}}))</th>
<th>(O(E_{\text{straj}}) - O(E_{\text{DMC}}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZEP</td>
<td>2061.8, 335.6</td>
<td>0.99890</td>
<td>2061.8, 335.6</td>
<td>0.99921</td>
<td>(0.1)</td>
<td>0.99703</td>
</tr>
<tr>
<td>(0.1)</td>
<td></td>
<td>0.98565</td>
<td>2064.2, 319.2</td>
<td>0.98565</td>
<td>(0.2)</td>
<td>0.99393</td>
</tr>
<tr>
<td>(0.2)</td>
<td></td>
<td>2060.2, 304.5</td>
<td>0.99282</td>
<td>2067.3, 306.3</td>
<td>(1.0)</td>
<td>0.99423</td>
</tr>
<tr>
<td>(1.0)</td>
<td></td>
<td>2046.3, 341.1</td>
<td>0.99764</td>
<td>2046.7, 341.1</td>
<td>(1.1)</td>
<td>0.99792</td>
</tr>
<tr>
<td>(1.1)</td>
<td></td>
<td>2044.6, 322.8</td>
<td>0.97876</td>
<td>2044.8, 322.9</td>
<td>(1.2)</td>
<td>0.99344</td>
</tr>
<tr>
<td>(1.2)</td>
<td></td>
<td>2039.8, 302.2</td>
<td>0.96982</td>
<td>2031.2, 282.1</td>
<td>(2.0)</td>
<td>0.97475</td>
</tr>
<tr>
<td>(2.0)</td>
<td></td>
<td>2017.2, 300.9</td>
<td>0.99855</td>
<td>2015.0, 314.8</td>
<td>(2.1)</td>
<td>0.99849</td>
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<tr>
<td>(2.1)</td>
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<td>0.97529</td>
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<td>(2.2)</td>
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<tr>
<td>(2.2)</td>
<td>2018.0, 299.0</td>
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<td>2019.1, 300.7</td>
<td>0.96297</td>
<td>Sum</td>
<td>8.05020</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>8.3935</td>
</tr>
</tbody>
</table>
last two columns of Table III the same calculations as in the second and third columns are, respectively, reported, but using the exact eigenvalues $E_n$ in Eq. (18). Once again, one can observe that the overlaps are not significantly different from those in the second and third columns and they are all very accurate, up to the (2,2) vibrational state.

A graphical comparison is even more striking than the numerical one presented in the previous tables. In Fig. 1 the eigenfunction methods used for calculating the overlaps reported previously are depicted. On panel (a)-(f) the (2,2) eigenfunctions are represented on contour plots, where the dashed color lines are the negative contour of the eigenfunctions. Panel (a) is the exact DVR eigenfunction. Panel (b) eigenfunction is calculated using the ground state trajectory and one can observe how it completely misses the correct nodal arrangement. On panel (c) the De Leon-Heller method wave function is plotted using a single trajectory and one can observe how it completely misses the correct nodal arrangement. On panel (d) Eq. (17) is used with the same set of eigenvalues as for the previous panel. Finally on panels (e) and (f) the MC-SC-IVR eigenfunctions are reported, respectively, with and without the separable approximation. These last plots of eigenfunctions are clearly more accurate than other ones, showing that the MC-SC-IVR method offers a consistent formulation for eigenvalue and eigenfunction calculations.

In order to find the origin of the consistent accuracy of the methods on panel (b) to (c), on Fig. 2, the eigenfunctions of the MC-SC-IVR and De Leon-Heller semiclassical methods at the exact (2,2) vibrational energy level are reported, together with the exact one represented on panel (a). This time all semiclassical methods faithfully reproduce the nodal behavior of the exact eigenfunction. This clearly shows that the performance of the method depends mostly on the accuracy of the calculation of the eigenvalues and that MC-SC-IVR in Eq. (18) and De Leon-Heller method in Eqs. (9)–(11) are almost numerically equivalent.

Now, let us turn to the estimate of the eigenfunctions’ accuracy via the coefficients $\epsilon_1$ and $\epsilon_2$ presented in Sec. III. Even if the CO on Cu(100) vibrational system is not degenerate, we can understand how accurate these parameters are by comparing their values in this non-degenerate case to other degenerate cases. In Table IV, the values of $\epsilon_1$ are calculated for each vibrational state. Since $\epsilon_1$ is the sum of the absolute value of the deviation of the approximate method from the exact one at each DVR point, there are no error compensations and it represents an upper bound for the error estimate. On the second and third columns the single trajectory results are reported, while on the following columns the multiple trajectory results are presented. Once again the indicative index “sum” shows that overall the multiple-trajectories perform best, with an improvement of accuracy of almost 50%. This means that the MC-SC-IVR eigenfunctions mimic the DVR ones better than the single trajectory calculations. However, given such small values of $\epsilon_1$ for all methods, one can consider the single-trajectory methods as highly accurate. In Table V, the same as in Table IV but for $\epsilon_2$.

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FIG. 3. Bidimensional vibrational eigenfunction contour plot cuts for the state \((0, 3^1, 0)\) of CO\(_2\), i.e., the triple bending excitation. The nodes are significantly tilted in the bending and symmetric stretch subspaces, while asymmetric stretch has a small coupling with the bending stretches subspace.

FIG. 4. Bidimensional plots for the “eigen-trajectory” with initial conditions associated to the \((0, 3^1, 0)\) vibrational state of CO\(_2\). The vibrational eigenfunctions of Fig. 3 are visibly on correspondence with this classical trajectory.

IV. FIRST-PRINCIPLES CALCULATION OF VIBRATIONAL EIGENFUNCTIONS

The full dimensional on-the-fly calculation of the vibrational eigenfunctions of the CO\(_2\) molecule is a challenging test for our semiclassical methods. The four vibrational modes are strongly coupled and numerous Fermi resonances originate from the inter-play between the symmetric stretch and the bending modes. In these cases, there is a specific link between classical trajectories and the corresponding quantum phenomenology, as shown by Heller\(^{21}\) in model resonating systems. In other words, there is a quantum to classical correspondence that directly parallels the strong resonance effects observed classically to the quantum properties. This suggests that semiclassical methods are able to reproduce resonating quantum features.

Previous on-the-fly calculations of the vibrational eigenvalues have proved to be very accurate if compared to exact grid methods, even when few trajectories were employed.\(^{34, 35}\) In this section vibrational eigenfunctions will also be calculated using Born-Oppenheimer classical molecular trajectories, where all the semiclassical propagator components are calculated directly from the electronic structure at the level of DFT. For example, the Hessian matrix which is required at each time step for the prefactor calculation of Eq. (14), is calculated from the second derivative operator in the Kohn and Sham formalism. In this way, the vibrational eigenfunction depicted in Fig. 3 has been calculated.

This eigenfunction corresponds to a triple-bending excited state. The distortion of nodal planes and caustic envelopes (the outer edge of the wave functions) are clear signs of non-linear strong couplings. The nodal line shapes are equally tilted in the bending subspace, which reveals the degeneracy between these two modes. Instead, in the bending and symmetric stretch subspace, the three nodes are arranged...
in order to be perpendicular to a U shaped line connecting the maximum of the eigenfunction. In the bending versus asymmetric stretch subspace the nodal lines are straight and perpendicular to the bending mode showing that no significant coupling is present between the two modes.

As anticipated above, this inspection of nodal patterns and their distortion can be explained in terms of classical trajectories.59

By looking at the first panel on the left of Fig. 4, one can see how this periodic trajectory is closing onto itself, an evidence of the synchronization between the two bending modes. An opposite behavior is reported on the last panel where the hallmark of the Fermi resonance is given by the U shape resulting from the convolution of the classical trajectory. By comparing the central panels of Figs. 3 and 4, one can see how the vibrational eigenfunction corresponds to that of the classical trajectory convolution. Further, quantum delocalization effects are indeed taken into account in our semiclassical approximation. In fact, let us compare the right panels of these figures: in Fig. 4 the classical trajectories range from −10 to +10 (in normal modes mass scaled units) along the asymmetric subspace, while in Fig. 3 the eigenfunction is delocalized to occupy almost double the interval.

In order to evaluate the accuracy of the semiclassical eigenfunctions, a set of DVR calculations were performed. In particular, the Hamiltonian operator was diagonalized in coordinate space using a sparse matrix representation interfaced with the ARPACK library diagonalization routines.60 Only the first 20 eigenvalues and eigenfunctions of the matrix were computed to reduce the computational memory request. To obtain higher eigenvalues more grid points are needed and hence more memory space is requested to store the matrix to be diagonalized.

The Hamiltonian operator is expressed in the sinc-DVR formalism in four dimensions (one for each mode).55 The potential energy surface employed is that of the analytical potential obtained by fitting the data on the fly.34 After various tests the converged set of eigenvalues and eigenfunctions were obtained with 52,000 points grid. A ten times denser grid provided the same eigenvalues up to the fourth significant figure and the error for the exact eigenfunctions (i.e., the value of parameter ε₁) is of the order of machine precision. In the diagonalization routine employed, the accuracy of the eigenvectors was checked to be less than 10−12 for the given eigenvalues, i.e., once the eigenvalues have converged up to the fourth decimal place the eigenfunctions corresponding to such eigenvalues are correct to within machine precision.

Such exact eigenfunctions were used to determine the accuracy of the semiclassical ones. The direct overlap comparison was possible only for the ground state, because most of the higher states are degenerate.

In Table VI the overlaps for different levels of semiclassical calculations are reported. One can conclude that all methods are accurate.

In order to have a complete picture of how all methods perform for the first few vibrational states calculated by DVR, a set of calculations for ε₁ and ε₂ are reported in Tables VII and VIII.

On the first row the different semiclassical methods are labeled and on the first column the exact eigenvalues of the vibrational levels are given. We remember that an exact eigenfunction will have a value of ε₁ = 0 and the values of ε₁ reported in Table VII are all of the order of hundredths.

![FIG. 5. Bidimensional vibrational eigenfunction contour plot cuts for the CO₂ vibrational state (1, 0², 0), i.e., the single symmetric stretch excitation. The nodal line in the bending/symmetric subspace is U shaped, in correspondence with the 2:1 Fermi resonance. The asymmetric stretch has a small coupling with the bending stretch.](image_url)
However, in order to quantify the overall accuracy of each method on the last row of Table VII the sums of the values in each column are reported. Even these sums are still of the order of hundredths and any distinction between the semiclassical methods is quite superfluous. The same considerations are valid for \( \varepsilon_2 \), where the exact value is \( \varepsilon_2 = 1 \).

From Table VIII, we note that all semiclassical methods are very accurate. Instead, when calculating the overlap of a single degenerate semiclassical eigenfunction with the DVR eigenfunction we found errors even of the order of 50%. As discussed previously, the introduction of the coefficients \( \varepsilon_1 \) and \( \varepsilon_2 \) is indeed necessary to evaluate accuracy of the degenerate eigenfunctions.

The most interesting vibrational states are the Fermi resonating states. The states labeled by a single symmetric stretch excitation and a double bending are represented, respectively, in Figs. 5 and 6.

Since twice the bending frequency is about a single symmetric stretch excitation, these states resonate and split into the Fermi states reported in Figs. 5 and 6. In these plots, the bending subspaces are as usual synchronized as shown by the tilted nodal lines. Instead, in the symmetric/asymmetric (Fig. 5) and in the bending/asymmetric (Fig. 6) subspace the nodal lines are perpendicular to the excited mode, showing that no significant coupling is present between the asymmetric stretch mode and the other ones. More interesting is the bending/symmetric stretch subspace in both figures, where the nodal line is U-shaped. In Fig. 5 the U shaped nodal line is pointing upwards, while it points downwards in Fig. 6. This shape is typical of a 2:1 Fermi resonance and one can also detect it from the envelope of the corresponding classical trajectories’ subspace.

V. CONCLUSIONS

In this paper a semiclassical initial value representation method (MC-SC-IVR) has been implemented for the calculation of vibrational eigenfunctions. It has been compared with other semiclassical methods and they all result equally good. The fact that these methods, implemented using an on-the-fly approach, are accurate enough to correctly describe a complex quantum system such as the CO2 molecule is noteworthy. In conclusion, we have observed that none of the methods employed can be considered superior to the others and that the accuracy of the semiclassical eigenfunction is mainly dictated by the accuracy of the corresponding eigenvalue calculation. To explore the role of the choice of trial eigenvalue, we scanned the eigenvalue of Eq. (8) and observed how the overlap of Eq. (20) changed as a function of the scanned eigenvalue. The numerical results show that a resolution in the spectrum better than 10 cm\(^{-1}\) is required to have a few percent of eigenfunction overlap error. Finally, these eigenfunctions can be used as guiding functions for more accurate Monte Carlo calculations. Work in this direction is in progress in our groups.

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### References
