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Citation: J. Chem. Phys. 130, 234113 (2009); doi: 10.1063/1.3155062
View online: http://dx.doi.org/10.1063/1.3155062
View Table of Contents: http://jcp.aip.org/resource/1/JCPSA6/v130/i23
Published by the American Institute of Physics.

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Multiple coherent states for first-principles semiclassical initial value representation molecular dynamics

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(Received 8 April 2009; accepted 27 May 2009; published online 19 June 2009)

A multiple coherent states implementation of the semiclassical approximation is introduced and employed to obtain the power spectra with a few classical trajectories. The method is integrated with the time-averaging semiclassical initial value representation to successfully reproduce anharmonicity and Fermi resonance splittings at a level of accuracy comparable to semiclassical simulations of thousands of trajectories. The method is tested on two different model systems with analytical potentials and implemented in conjunction with the first-principles molecular dynamics scheme to obtain the power spectrum for the carbon dioxide molecule. © 2009 American Institute of Physics. [DOI: 10.1063/1.3155062]

I. INTRODUCTION

One of the goals of quantum nuclear molecular dynamics simulations is that of simulating the interplay between electronic properties and quantum nuclear effects. Electronic properties are local in nuclear configuration state, whereas nuclear wave functions are nonlocal for the same coordinates. In the first-principles (FP) approach to quantum molecular dynamics, nonlocal properties of the nuclear Schrödinger equation have been to some extent localized and are calculated directly by quantum chemistry methods. The main advantage of this method is that the interplay between the electronic structures and nuclear motion is reproduced within the limits of the Born–Oppenheimer (BO) approximation. From the computational standpoint, the use of FP methods allows avoiding the use of precomputed potential energy surfaces (PESs). Fitting PESs to empirical or to calculated electronic structure data is usually a tradeoff between the desired accuracy and the human and computational effort required to construct them and it is often biased by the functional forms chosen. In FP simulations, the interatomic potentials and derivatives are calculated directly from electronic wave functions, i.e., on-the-fly, thus avoiding any artifact resulting from fitting as well as the cumbersome fitting process, which gets formidable as the degrees of freedom of the system increase. On-the-fly simulations are sometimes advantageous when describing rapidly changing regions of the PES, such as bond-breaking processes, as fitting errors can be larger in these cases. Simulations of the dynamics of floppy molecules can benefit from this approach as well. FP, on-the-fly simulations have been employed very successfully for studying conical intersections.1

To date, there have been several implementations of FP approaches for the treatment of nuclear molecular dynamics with or without quantum effects. In a typical FP molecular dynamics simulation following Newton’s equations and therefore not taking quantum effects into consideration, a velocity autocorrelation function is calculated and it is Fourier transformed to obtain the vibrational frequencies. In this classical dynamics approach, anharmonic effects are properly accounted for, but the zero point energy and tunneling effects are not taken into account in the classical formalism. Furthermore, long simulations are necessary in order to obtain a good frequency resolution.2–4 Several techniques have been developed to speed up classical simulations. For example, in the BO molecular dynamics approach, the electronic structure calculations for a given simulation step are converged based on previous step information. However, this approach can lead to systematic energy drifts.5 Alternatively, extended Lagrangian molecular dynamics approaches (ELMD)6–9 involve the classical propagation of nuclear and electronic degrees of freedom on an equal footing by assigning classical variables to the electronic ones. The electronic degrees of freedom, in turn, are expanded in terms of plane waves,6 Gaussian functions,8 or real-space grids.9 ELMD propagation is computationally efficient, however, the resultant energy surface is not always close to the actual BO PESs, as typically these dynamics oscillate around the BO PESs. Of concern are the dependencies that have been found on the fictitious electronic masses.8,10 In recent years, several avenues have been successfully pursued in order to include quantum effects in a FP approach to quantum molecular dynamics.7,11–13 One can cast the path-integral expression for the quantum propagator into an extended Lagrangian formalism,13 or adopt the differential approach of the variational multiconfiguration Gaussian wave packet method,12 where the potential is approximated to be locally harmonic. Other approaches include a mean field approximation for the
time-dependent self-consistent quantum propagation\textsuperscript{14,15} and multiple spawning methods developed by Ben-Nun and Martinez.\textsuperscript{15}

Semiclassical methods based on classical trajectories\textsuperscript{6–22} are naturally suitable for carrying out on-the-fly molecular dynamics simulations. It is well known from the SC developments in the early 1970s\textsuperscript{28,29} that these kinds of approximations include all quantum effects at least qualitatively and often proved to be quantitatively accurate in describing molecular systems.\textsuperscript{17,18,25,30–39}

More recently we implemented the semiclassical initial value representation (SC-IVR) of the quantum propagator into a FP molecular dynamics scheme\textsuperscript{40} and showed how SC-IVR can be coupled with FP electronic structure approaches for classical molecular dynamics. A very recent study\textsuperscript{41} has shown similar results for vibrational absorption spectra. By employing a single FP classical trajectory, we obtained the vibrational power spectrum for the CO\textsubscript{2} molecule with 1 cm\textsuperscript{-1} spectrum resolution. However, the accuracy of this simulation worsens for the reproduction of anharmonic effects. For this purpose, we suggest reproducing anharmonic effects. For this purpose, we suggest employing the direct product of one dimensional coherent reference states,

\[ \langle q|p(t),q(t) \rangle = \prod_i (\gamma_i/\pi)^{F/4} \exp \left[ -\frac{\gamma}{2} (q_i - q_i(t))^2 + \frac{i}{\hbar} p_i(t) \cdot (q_i - q_i(t)) \right]. \]

In the previous expression, the coherent states have a fixed width, \( \gamma_i \). For bound systems, a reasonable choice for the width parameter is that of employing the width of the harmonic oscillator approximation to the wave function at the minimum of the PES where the dynamics will be carried out. In our previous experience and that of others, we found no significant dependency upon variation in the width parameter.\textsuperscript{19,40,42}

In this paper, the main application of FP-SC-IVR molecular dynamics will be the calculation of spectral density,

\[ I(E) = \langle \chi|\delta(\hat{H} - E)|\chi \rangle = \sum_n \langle \chi|\psi_n \rangle^2 \delta(E - E_n), \]

where \( |\chi\rangle \) is some reference state and \{\( |\psi_n \rangle \)\} and \( \{E_n\} \) are the collections of exact eigenstates and eigenvalues of the Hamiltonian \( \hat{H} \). The spectrum of Eq. (4) embodies all the information coming from the eigenvalues and eigenvectors; the peaks are located at the positions of the eigenvalues and their intensity is proportional to the overlap between the trial state \( |\chi\rangle \) and the actual eigenvectors associated with the PES. Replacing the Dirac delta function of Eq. (4) by its Fourier representation, one can more conveniently achieve the same spectrum with the following time dependent representation:\textsuperscript{26}

\[ I(E) = \frac{1}{2\pi\hbar} \int_{-\infty}^{+\infty} \langle \chi|e^{-iE/t}|\chi \rangle e^{iEt/t} dt \]

Here, the spectrum is written in terms of the Fourier trans-
form of the survival probability. The SC-IVR approximation for the survival probability is
\begin{equation}
\langle \psi | e^{\iota t \hat{H}/\hbar} | \psi \rangle = \frac{1}{(2\pi \hbar)^4} \int dp(0) \int dq(0) C_0(p(0), q(0)) \times e^{i S_q(0, q(0)) \hbar} \langle \psi_t | \psi \rangle \langle \psi | \psi_t \rangle.
\end{equation}
(6)
suggests a suitable choice for the phase-space reference state \( | \psi \rangle = | p_{eq}, q_{eq} \rangle \). Usually, a coherent wave packet, which mimics the ground state wave packet and its motion following Eq. (3), is chosen. The purpose of this paper, however, is to suggest an alternative expression to the single harmonic reference state that improves the accuracy of the results for excited states, while maintaining a reasonable computational effort.

In order to reduce the number of trajectories required for Monte Carlo phase space sampling in Eq. (6), one can introduce a time average integral at the cost of a longer simulation time. Insertion of the time average integral is not an approximation per se and it does not imply any ergodicity property. This is in principle exact, by virtue of Liouville's theorem. Using this approach, Kaledin and Miller, and Ceotto obtained the time-averaging (TA) SC-IVR approximation for the spectral density,
\begin{equation}
I(E) = \frac{1}{(2 \pi \hbar)^2} \int dp(0) \int dq(0) \times \frac{\text{Re}}{\pi \hbar T} \int_0^T dt_1 \int_0^T dt_2 C_t(p(t_1), q(t_1)) \times \langle \psi(t_2) | \psi(t_1) \rangle \times \langle \psi(t_1) | \psi(t_2) \rangle e^{i(S_q(p(0), q(0)) + E t_2)/\hbar} \times \langle \psi(t_1) | \psi(t_2) \rangle e^{i(S_q(p(0), q(0)) + E t_1)/\hbar}^*.
\end{equation}
(7)
Here, the \( p(t_1), q(t_1) \) and \( p(t_2), q(t_2) \) position and momentum variables evolve from the same initial conditions, but to different times, and \( T \) is the total simulation time. The first time integral is the Fourier transform of Eq. (5), while the second time integration corresponds to the time-averaging filter. This double-time integration implies that for any given trajectory from time 0 to time \( T \), all possible intervals from \( t_1 \) to \( t_2 \) are considered. The prefactor of Eq. (2) for each interval now depends on two times, i.e., \( C_t(p(t_1), q(t_1)) \). This approach would be really advantageous for FP applications if the number of trajectories can be reduced to a few. In the following, we will introduce and describe an implementation of the reference state representation aimed to achieve this goal.

In order to make Eq. (7) even less computationally demanding, its prefactor [Eq. (7)] is approximated as phase difference, \( C_t(p(t_1), q(t_1)) = \exp[i(\phi(t_1) - \phi(t_2))/\hbar] \), where \( \phi(t) = \text{phase}[C_t(p(0), q(0))] \). With this "separable approximation" Eq. (7) becomes
\begin{equation}
I(E) = \frac{1}{(2 \pi \hbar)^2} \int dp(0) \int dq(0) \times \frac{\text{Re}}{\pi \hbar T} \int_0^T dt \langle \psi(t) | \psi(t) \rangle \times e^{i(S_q(p(0), q(0)) + E t + \phi(p(0), q(0))/\hbar)}^2,
\end{equation}
(8)
leading to a simplification of the double-time integration to a single- and positive-definite time integral. This approximation offers also the advantage of reducing the instabilities of the complex phase space integrand by means of the time-averaging filter.

The PES, gradients, and Hessians are computed at each nuclear configuration directly from a Gaussian-based density functional theory. The Kohn–Sham orbitals were expanded on a nonorthogonal Gaussian basis. The computationally intense part of our calculation is the evaluation of the potential energy together with its higher derivatives at each time step. The computer time required to complete this calculation on a standard desktop (3.2 GHz Intel CPU) was usually a few hours for a \( 2 \) cm\(^{-1}\) spectra resolution for the small molecules studied here. The classical nuclear dynamics forces,
\begin{equation}
M_j \ddot{R}_j = -\nabla_i E_{\text{DFT}}[\mathbf{R}],
\end{equation}
(9)
have been calculated on the BO PES and the classical propagation is performed according to the velocity-Verlet algorithm, as implemented in the Q-CHEM electronic structure package.

III. MULTIPLE COHERENT STATES REPRESENTATION

By inspection of the definition of the power spectrum given in Eq. (4), one can develop a new strategy for SC-IVR power spectra calculation that would enhance its precision. According to this equation, the reference state should significantly overlap with each eigenstate in order to have a well defined intensity peak. In our previous work, we showed that when a single ground harmonic state is chosen as the reference, the spectral accuracy is poor for the excited states. Instead, by constructing a reference state that is a combination of states that mimic the spectral eigenfunctions with the right eigenvalue spacings, one should obtain more intense spectral peaks. This is of relevance because we found by numerical simulation that usually the more intense the peaks are, the higher the accuracy obtained for a fixed-time simulation.

For illustrative purposes, a representation of the multiple reference state coherent basis is shown on Fig. 1. In Fig. 1(a), the representation of a potential well and its vibrational energy levels are depicted. The power spectrum for this potential is shown on Fig. 1(b). With the goal of obtaining high peak intensities and therefore improved accuracy, a set of carefully, yet systematically chosen, classical trajectories are sought. Figure 1 shows representative classical trajectories that can be associated with each vibrational level. These "eigentrajectories" are chosen to have turning points located at the values of potential energy equal to the peaks of the power spectrum. On panel Fig. 1(c), the phase portrait for the
quantum wave packet. Instead, in the approach proposed for semiclassical applications, where the basis set is updated to be a combination of coherent states, a power spectrum for the potential is sketched on panel (b). The dotted arrow lines on panel (a) represent the classical trajectories associated with each vibrational level. The phase portrait of these trajectories is shown on panel (c). Two choices for the location of the coherent states basis for the MC[n] approach are shown. The dark circles represent trajectories that start at the bottom of the potential well and have enough energy to reach the classical turning point for a harmonic approximation of the potential. An alternative approach depicted in gray is that of initializing a MC[n] coherent state basis located at the classical turning points with zero or negligible initial momentum.

In light of these considerations and inspired by previous coherent state grids calculations by Davis and Heller, we have chosen the reference states $|\chi\rangle$ to be a combination of coherent states,

$$|\chi\rangle = \sum_{j=1}^{N_{states}} |p_{eq}^i, q_{eq}^i\rangle,$$

located at either the black or gray circles on Fig. 1(c). In the first case (black circles), the coherent states are located at the equilibrium molecular position $q_{eq}$, and the initial momentum is prescribed in such a way that the kinetic energy is just enough to reach the classical turning point of the targeted vibrational energy level. The harmonic approximation is employed to set these initial momenta, namely, $p_{eq}^2/2m=\hbar\omega(n+1/2)$. In the second case (gray circles), the positions of the initial coherent states are placed at the turning points $q_{eq}$ and the momenta are set to be a small value $\epsilon$ close to zero.

There have been several examples of coherent state grids for semiclassical applications, where the basis set is updated at each infinitesimal time step and it follows coherently the quantum wave packet. Instead, in the approach proposed above, the coherent states are placed in a nonlocal fashion and their centers are kept fixed all along the simulation time.

To gain numerical insight about the multiple coherent (MC) states approach presented above, one can look at the survival probability expression in Eq. (6) in terms of position and momenta for a single coherent state,

$$\langle\chi|p(t), q(t)\rangle|\chi(0), q(0)\rangle = e^{-\gamma(q_t - q_{eq})^2/4 - q_0^2 - q_{eq}^2(4 - q_0^2 - q_{eq}^2)/4(4 - q_0^2 - q_{eq}^2)/2}\times e^{-(p_t - p_{eq}^0)^2/2 - (p_0 - p_{eq})^2/2}\times e^{-(p_t - p_{eq}^0)^2/2 - (p_0 - p_{eq})^2/2}.$$  

(11)

The previous equation shows that the signal for the Fourier transform is significant only if the classical trajectories happen to be on the neighborhood of $(p_{eq}^0, q_{eq})$. Choosing trajectories around this neighborhood is important since one can facilitate phase space Monte Carlo integration if the coherent basis states and the classical trajectories are chosen to be close to the modes associated with the desired vibrational energy peaks. As stated above, in the case of calculations of bound states, a good guess can be obtained from the evaluation of the nuclear Hessian at the equilibrium geometry of the potential well of interest. By inserting Eq. (10) into Eq. (7) one obtains the final expression for the MC states power spectra calculations

$$I(E) = \frac{1}{(2\pi\hbar)^3} \sum_{j=1}^{N_{states}} \int_{0}^{T} dt_1 \int_{t_1}^{T} dt_2 \sum_{i} \langle p_{eq}^i, q_{eq}^i|p(t_1), q(t_1)\rangle e^{i(S_{i}(p(0), q(0)) + E_{i})/\hbar}$$

$$\times \left[ \sum_{j} \langle p_{eq}^j, q_{eq}^j|p(t_2), q(t_2)\rangle e^{i(S_{j}(p(0), q(0)) + E_{j})/\hbar} \right]^*,$$

(12)

where the phase space integral has been reduced to a sum over the coherent states centers. An analogous expression holds when the separable approximation is applied. In the Appendix we show how one can calculate the full harmonic spectrum starting from Eq. (7) and show how the center of the momentum distribution is associated with the spectral peak intensities.

IV. NUMERICAL TESTS ON MODEL POTENTIALS

In this section, we describe two applications of the proposed method employing model potentials. The systems studied are the carbon monoxide molecule adsorbed in the copper (100) surface and the vibrational spectra of the water molecule. These two applications allow for testing our method in well-known potentials, for which results are readily available.

A. Carbon monoxide adsorbed on the copper (100) surface

To test our ideas on a simple, yet physically relevant model system, we performed SC-IVR simulations to model the interactions of a carbon monoxide (CO) molecule chemisorbed on the copper Cu(100) surface. We chose this model due to the availability of a well-tested analytical potential in the form of

\begin{align*}
\langle \chi | p(t), q(t) | \chi(0), q(0) \rangle &= e^{-\gamma(q_t - q_{eq})^2/4 - q_0^2 - q_{eq}^2(4 - q_0^2 - q_{eq}^2)/4(4 - q_0^2 - q_{eq}^2)/2} \times e^{-(p_t - p_{eq}^0)^2/2 - (p_0 - p_{eq})^2/2} \times e^{-(p_t - p_{eq}^0)^2/2 - (p_0 - p_{eq})^2/2}. \\
&\quad \times e^{-(p_t - p_{eq}^0)^2/2 - (p_0 - p_{eq})^2/2}. \\
&= \exp \left[ -\frac{1}{2} \left( \frac{q_t - q_{eq}}{\gamma} \right)^2 - \frac{1}{4} \left( q_0^2 - q_{eq}^2 \right) - \frac{1}{2} \left( p_t - p_{eq}^0 \right)^2 - \frac{1}{2} \left( p_0 - p_{eq} \right)^2 \right].
\end{align*}
where \( \mathbf{r}_c \) and \( \mathbf{r}_O \) are, respectively, the carbon and the oxygen Cartesian coordinates and \( \mathbf{r}_i \) are that of the \( i \)th coordinates of the copper atom. The intramolecular term \( V_{CO} \) is a standard Morse potential, while the interaction of CO with each copper atom is described as

\[
V(\mathbf{r}_C, \mathbf{r}_O) = \sum_{i=1}^{N_{\text{copper}}} V_i(\mathbf{r}_C, \mathbf{r}_O, \mathbf{r}_i) + V_{CO}(\mathbf{r}_C - \mathbf{r}_O),
\]

and has the appropriate initial momentum \( \mathbf{p}_{eq}, \mathbf{q}_{eq} \) and has the appropriate initial momentum.

The first term of Eq. (14) describes the oxygen-copper repulsion. The remainder of the terms in Eq. (14) represents the carbon-copper interactions. The molecule orientation is taken into account by the angle \( \phi \) between the O–C and C–Cu bonds. The metal is represented by three layers of 36 (6 \( \times 6 \)) copper atoms arranged according to fcc lattice structure. The molecular axis is fixed perpendicular to the surface and the resulting molecular motion is described by two degrees of freedom, the internal CO mode and the stretching mode perpendicular to the surface, as depicted in Fig. 2.

The potential parameters were adjusted to obtain the first normal mode at 2084 cm\(^{-1}\) and the second one at 353 cm\(^{-1}\). Since there is an order of magnitude difference between mode frequencies, the time-averaging filter will more likely perform better on the fast mode, while the accuracy of this filter on the slow and floppy mode remains to be verified.

We performed single and MC states calculations of the CO–Cu system to investigate the accuracy of the MC states approach to SC-IVR dynamics proposed in this work. A single-trajectory simulation using the time-averaging filter yields the data shown in the panel (0,0) of Fig. 3. The corresponding vibrational eigenvalues are reported on Table I in the column denoted as MC state [1], (MC[1]). For comparison, we show the result obtained employing the separable approximation of Eq. (8) as MC state/separable, MC[1]/S. Although these frequencies have values very close to the exact numerical ones, a closer inspection of the vibrational eigenvalue spacings shows that single-trajectory simulations take into account anharmonicity only for the first quantum excitation; higher vibrational energy levels show a harmonic-like spacing as reported in Table II.

In order to find the source of this deficiency, we calculated a single trajectory power spectrum using different reference states. The chosen state is denoted by the quantum numbers in parenthesis in each one of the panel of Fig. 3. This figure highlights how the intensities of the spectral peaks are highly dependent on the initial state. This is of importance because a higher spectral peak usually results in higher accuracy. The value and the spacing of the eigenvalues corresponding to the vibrational reference state (highlighted in each panel of Fig. 3) are reported in Tables I and II under the columns peak-centered (PC), and peak-centered/ separable approximation PC/S. By choosing various reference states, the vibrational anharmonicity is well reproduced and the spacings obtained are almost exact. The percentage error of the calculated vibrational level spacings for these calculations was always less than 1%. Obviously it is desirable to reproduce these data with the same level of accuracy in a single spectrum. Hence, we employed the MC-TA-SC-IVR approach using the following procedure. For every desired state, a trajectory which starts at the equilibrium coordinates \( (\mathbf{p}_{eq}, \mathbf{q}_{eq}) \) and has the appropriate initial momentum associated with the harmonic approximation for the desired mode is propagated. The results of the simulations using this approach are shown in Tables I and II under columns MC[6]/S and MC[6], where the number 6 in brackets indicates the number of trajectories employed for our coherent basis set approach. These six classical trajectories were able to reproduce an accurate power spectrum. The deviation from the exact values for this computational approach was within few wave numbers of the numerical DVR results. A range of frequencies from the ground state energy of 1210 to
TABLE I. Vibrational energy levels for the different internal and external stretching excitations: On the second column the exact grid calculation results. The MC states MC[1]/S column represents a single trajectory time-averaging semiclassical calculation carried out with the separable approximation. The same calculation without the separable approximation is listed under the MC[1] column. The peak centered with separable approximation (PC/S) and peak-centered (PC) columns correspond to the initially chosen coherent states shown in the panels of Fig. 3. The MC states simulations of six trajectories are shown in columns MC[6]/S and MC[6] with and without the separable approximation, respectively. The six-trajectory calculations are able to reproduce the anharmonic spacings associated with each mode, as shown in Table II. The trajectories for each entry of the table were carried out for 6000 time steps.

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\(^{a}\)Exact frequencies in cm\(^{-1}\) by DVR calculations (Ref. 54).

6000 cm\(^{-1}\) was obtained with errors of the order of 2–3 cm\(^{-1}\). It is also worth noting that the MC-SC-IVR approach is able to reproduce vibrational states that are not included into the harmonic combinations of Eq. (10) with the same level of accuracy.

B. Water molecule

A more challenging test for the time-averaging MC-SC-IVR is the calculation of the vibrational spectra of the water molecule. This system has strongly coupled modes and strong anharmonic effects in the higher vibrational levels. For this study, we employed the realistic potential surface of Ref. 55.

We summarize the results of our calculations in Table III. The spectroscopic vibrational terms and their associated symmetry have been reported on the first column. For comparison purposes, the harmonic levels are reported on the second column. The next three columns are TA-SC-IVR calculations performed with the separable approximation of Eq. (8) but at a different computational level. Under the MC[1]/S column, we show the calculated frequencies for a single 8000 time-step trajectory. Under the MC[8]/S label, the results of eight classical trajectories sampled according to the harmonic approximation of the first eight vibrational states and the MC states basis of Eq. (10) is employed. On the column labeled “separable,” the results from a simulation employing 32,000 trajectories using the separable approximation and a single ground harmonic reference state are reported. The data shown in this column are converged at the semiclassical level and we employ this column as a reference for comparison. The purpose of this table is that of comparing the results of our few-trajectory MC states approach to the considerably computationally more expensive 32,000 trajectories simulation at the same level of semiclassical approximation. The results of Table III show that the MC[8]/S simulation reproduces with a certain accuracy the semiclassically converged vibrational levels. The MC[8]/S simulations correctly describe anharmonic effects, a deficiency observed for single-trajectory calculations. As in the case of the CO/Cu(100) simulation shown previously, even if the ZPE is reproduced accurately at all levels of semiclassical dynamics, the vibrational energy levels spacings of higher-energy states

TABLE II. Peak spacings of the data shown in Table I. The MC states simulations with six trajectories with and without the separable approximation (MC[6]/S and MC[6]) are able to reproduce the anharmonic spacings accurately. The single-trajectory calculations MC[1] and MC[1]/S show harmonic-like spacing for the second and third harmonic modes. The peak-centered trajectories (PC/S and PC) are able to reproduce the appropriate level spacings.

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</tr>
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<td>(1,1)</td>
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<td>345</td>
<td>334</td>
<td>332</td>
<td>337</td>
<td>334</td>
</tr>
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<td>315</td>
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<td>345</td>
<td>316</td>
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<tr>
<td>(2,1)</td>
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<td>344</td>
<td>337</td>
<td>334</td>
<td></td>
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<tr>
<td>(2,2)</td>
<td>314</td>
<td>345</td>
<td>346</td>
<td>314</td>
<td>312</td>
<td></td>
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</table>
are different for each type of semiclassical simulation. The single-trajectory MC\[1\]/S calculation gives a harmonic bending spacing of 1607 cm\(^{-1}\). In contrast, the eight-trajectory MC\[8\]/S simulation provides an anharmonic spacing of 1599, 1560, and 1537 cm\(^{-1}\). This is to be compared to the spacing of 1588, 1552, and 1528 cm\(^{-1}\) resulting from the 32 000 trajectories calculation. A similar trend is observed for the symmetric and asymmetric stretch modes. For example, the (1,1,1) vibrational state has an energy of 13 789 cm\(^{-1}\) under the MC\[8\]/S simulation, while the single-trajectory semiclassically converged result of 32 000 trajectories using the TA-SC-IVR approach resulting from Monte Carlo integration of Eq. (8) with a standard sampling width and 2000 time steps for each trajectory. The remainder of the columns shows results employing the full TA-SC-IVR approach of Eq. (7). Uncertain peaks are marked with *.

<table>
<thead>
<tr>
<th>(v_1), (v_2), (p_3)</th>
<th>Harm.</th>
<th>MC[1]/S</th>
<th>MC[8]/S</th>
<th>Separable</th>
<th>MC[1]</th>
<th>MC[8]</th>
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<tr>
<td>000 (A(_1))</td>
<td>4711</td>
<td>4635</td>
<td>4637</td>
<td>4644</td>
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<td>100 (A(_1))</td>
<td>6361</td>
<td>6244</td>
<td>6236</td>
<td>6232</td>
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<td>200 (A(_1))</td>
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<td>7791</td>
</tr>
<tr>
<td>010 (A(_2))</td>
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<td>8363</td>
<td>8353</td>
<td>8364</td>
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</tr>
<tr>
<td>001 (B(_1))</td>
<td>8652</td>
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<td>8447</td>
<td>8442</td>
<td>8447</td>
<td>8450</td>
</tr>
<tr>
<td>300 (A(_1))</td>
<td>9661</td>
<td>9458*</td>
<td>9333</td>
<td>9312</td>
<td>9460*</td>
<td>9323</td>
</tr>
<tr>
<td>110 (A(_2))</td>
<td>10 191</td>
<td>9975</td>
<td>9941</td>
<td>9933</td>
<td>9973</td>
<td>9938</td>
</tr>
<tr>
<td>101 (B(_2))</td>
<td>10 302</td>
<td>10 058</td>
<td>9991*</td>
<td>9974</td>
<td>10 056</td>
<td>9989</td>
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<tr>
<td>210 (A(_1))</td>
<td>11 841</td>
<td>11 583</td>
<td>11 502</td>
<td>11 492*</td>
<td>11 581</td>
<td>11 495</td>
</tr>
<tr>
<td>201 (B(_2))</td>
<td>11 952</td>
<td>11 655</td>
<td>11 578*</td>
<td>11 521*</td>
<td>11 663</td>
<td>11 577*</td>
</tr>
<tr>
<td>020 (A(_1))</td>
<td>12 372</td>
<td>12 098</td>
<td>12 083</td>
<td>12 053*</td>
<td>12 096</td>
<td>12 054</td>
</tr>
<tr>
<td>011 (B(_2))</td>
<td>12 483</td>
<td>12 181</td>
<td>12 131</td>
<td>12 127</td>
<td>12 179</td>
<td>12 131</td>
</tr>
<tr>
<td>002 (A(_2))</td>
<td>12 593</td>
<td>12 265</td>
<td>12 180</td>
<td>12 171</td>
<td>12 277*</td>
<td>12 177</td>
</tr>
<tr>
<td>120 (A(_1))</td>
<td>14 022</td>
<td>13 706</td>
<td>13 647</td>
<td>13 628</td>
<td>13 704</td>
<td>13 660</td>
</tr>
<tr>
<td>111 (B(_2))</td>
<td>14 133</td>
<td>13 789</td>
<td>13 705</td>
<td>13 690</td>
<td>13 788</td>
<td>13 714</td>
</tr>
<tr>
<td>102 (A(_2))</td>
<td>14 244</td>
<td>13 873</td>
<td>13 789</td>
<td>13 750</td>
<td>13 885*</td>
<td>13 829</td>
</tr>
</tbody>
</table>

V. FIRST-PRINCIPLES SC-IVR APPLICATIONS

A challenging test for the MC states FP-SC-IVR method is the calculation of the full dimensional vibrational power spectrum of the CO\(_2\) molecule. A successful method should reproduce spectral features such as degenerate bending modes, strong intermodal couplings, as well as the Fermi resonances. To evaluate the accuracy of the MC-FP-SC-IVR method, the vibrational spectrum of the CO\(_2\) molecule will be compared with numerically exact discrete variable representation (DVR) eigenvalue calculations. These calculations were performed on a potential fitted to a set of FP points obtained at the same level of theory and are reported on Table IV under the DVR heading. On the first column of the same table, we report the experimental values, so one can also evaluate how realistic is the description offered by the DFT functional. In these calculations, Cartesian coordinates have been used to describe the classical molecular motion of Eq. (9) on a PES generated by a B3LYP density functional\(^{48}\) with the cc-pVDZ basis set. On the second column of Table IV one can find the spectroscopic label of each vibrational state and on the third the harmonic estimate for the vibrational energy levels. The main results on Table IV are from the application of the MC states FP-SC-IVR method. The columns labeled MC\[8\] and MC\[8\]/S list the data obtained from calculations where the first eight states are chosen as the MC states reference basis. The MC\[8\]/S column shows results employing Eq. (7) with only eight on-the-fly classical trajectories. The subsequent

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**TABLE III.** Water vibrational eigenvalues at different computational levels. On the first column, the spectroscopic vibrational terms are shown. The second column lists the second vibrational energy levels using normal mode analysis. The column labeled MC\[1\]/S shows semiclassical results obtained with a single trajectory of 8000 time steps. The column labeled MC\[8\]/S lists the results obtained from an eight-trajectory calculation of 4000 time steps each using the MC states approach of Eq. (10). The column labeled “separable” shows a semiclassically converged result of 32 000 trajectories using the TA-SC-IVR approach resulting from Monte Carlo integration of Eq. (8) with a standard sampling width and 2000 time steps for each trajectory. The main results on Table IV are from the application of the MC states FP-SC-IVR method. The columns labeled MC\[8\] and MC\[8\]/S list the data obtained from calculations where the first eight states are chosen as the MC states reference basis. The MC\[8\]/S column shows results employing Eq. (7) with only eight on-the-fly classical trajectories. The subsequent
TABLE IV. Some of the calculated vibrational energy eigenvalues. All data are in wave numbers. Experimental frequencies associated with Fermi resonances are indicated by a distinct superscript symbol for each group of resonant states in the first column. Uncertain peaks are marked with (&). The first column represents the experimental vibrational frequencies associated with the modes listed on the second column. The third column shows the harmonic DFT results. The trajectories listed on the fourth and fifth columns were carried out for 5000 time steps. Exact numerical DVR calculations in the same B3LYP/cc-PVDZ model chemistry used for the FP-SC-IVR calculations are reported on the last column.

<table>
<thead>
<tr>
<th>Exp.</th>
<th>Mode</th>
<th>Harmonic</th>
<th>MC[8]</th>
<th>MC[8]/Sd</th>
<th>DVR</th>
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<td></td>
<td>ZPE</td>
<td></td>
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<tr>
<td>667.4</td>
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<td>656.62</td>
<td>644</td>
<td>638</td>
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<tr>
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<td>1280</td>
<td>1269</td>
<td>1252.91</td>
</tr>
<tr>
<td>1388.2</td>
<td>1,0,0</td>
<td>1363.46</td>
<td>1388</td>
<td>1390</td>
<td>1372.29</td>
</tr>
<tr>
<td>1932.5</td>
<td>0,3,0</td>
<td>1969.86</td>
<td>1922</td>
<td>1902</td>
<td></td>
</tr>
<tr>
<td>2003.2</td>
<td>0,3,0</td>
<td>1969.86</td>
<td>2032</td>
<td>2028</td>
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<tr>
<td>2076.9</td>
<td>1,1,0</td>
<td>2020.08</td>
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<td>2109</td>
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<tr>
<td>2349.1</td>
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<td>2372</td>
<td>2359</td>
<td>2359.51</td>
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<tr>
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<td>2585.0</td>
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<td>2603</td>
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<tr>
<td>2671.7</td>
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<td>2669</td>
<td>2662</td>
<td>2640.15</td>
</tr>
<tr>
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<tr>
<td>2797.2</td>
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<td>2726.92</td>
<td>2817</td>
<td>2788</td>
<td>2757.14</td>
</tr>
<tr>
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<tr>
<td>6972.6</td>
<td>0,0,3</td>
<td>7270.41</td>
<td>6833</td>
<td>6825</td>
<td>6821.35</td>
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</tbody>
</table>

1Experimental frequencies in cm⁻¹ from Ref. 57.
2The first number is the symmetric stretch quantum, the second is the degenerate bendings, and the third one is the asymmetric stretch. The exponent of the second number is the l_i degeneracy index.
3Vibrational levels according to a normal modes harmonic model.
4Using the separable approximation of Eq. (8).
5Exact frequencies in cm⁻¹ from Ref. 40.

VI. CONCLUSIONS

In this paper, a novel semiclassical implementation for power spectrum calculation is presented. The results of this method are comparable with those of Monte Carlo phase space integration employing thousands of trajectories. Our results are valid for bound harmonic-like potentials such as those of molecules with a few degrees of freedom. The proposed MC states, MC[n] method, is a strategy for employing a basis set of a small number of coherent states, which resemble a linear combination of approximate eigenfunctions of the system. Each one of the trajectories for our basis set is chosen to resemble a target vibrational state. In this way, a multiple vibrational components signal is successfully reproduced and results in a well-defined power spectrum. We found that a harmonic MC[n] ansatz provides a great improvement over a single ground harmonic reference state simulation. The method has been developed by analytical considerations that show the importance of the reference state choice in deriving the harmonic spectrum and tested on both precomputed and on-the-fly FP PESs. The proposed methodology can be extended to the use of additional trajectories. A possible extension is the use of trajectories sampled from a Gaussian distribution around the Harmonic approximation to the chosen target vibrational modes. Further investigations along this line are currently underway in our laboratories.

ACKNOWLEDGMENTS

A. A.-G. and S.A. thank the Faculty of Arts and Sciences of Harvard University for financial support. M.C. and G.F.T. thank the University of Milan for funding and CILEA (“Consorto Interuniversitario Lombardo per L’Elaborazione Automatica”) for computational time allocation. A.A.-G. and S.A
thank the FAS Research Computing for cluster computing support. A.A.-G. thanks the Defense Advanced Projects Agency (under award FA9550–08–1–0285 administered by the Air Force of Scientific Research) and the National Science Foundation (under award 0835713) for the support.

APPENDIX: ANALYTICAL CONSIDERATIONS

In order to gain further understanding of the physical meaning of the MC state approach, here we derive the harmonic oscillator power spectrum starting from Eq. (7) in units of $\hbar$ and given an arbitrary coherent state. The phase space equations of motion in mass-scaled form are

$$q(t) = q(0) \cos(\omega t) + \frac{p(0)}{\omega} \sin(\omega t), \quad (\text{A1})$$

$$p(t) = p(0) \cos(\omega t) - \omega q(0) \sin(\omega t), \quad (\text{A2})$$

and the difference of the corresponding actions is

$$S(t_2) - S(t_1) = p(0) q(0) \left( \cos^2(\omega t_2) - \cos^2(\omega t_1) \right)$$

$$+ \left( \frac{p^2(0)}{2\omega} - \frac{\omega^2 q^2(0)}{2} \right) \sin(\omega t_2)$$

$$+ \sin(\omega t_1)), \quad (\text{A3})$$

Another element of Eq. (7) is the generic reference state $|p_{eq}, q_{eq}\rangle$ coherent states overlap.

$$\langle p_{eq}, q_{eq}|p(t_2), q(t_2)\rangle \langle p(t_1), q(t_1)|p_{eq}, q_{eq}\rangle$$

$$= \exp \left[ -\frac{\gamma}{4}(q^2(t_2) + q^2(t_1)) - \frac{\gamma}{2} p_{eq}^2 + \frac{\gamma}{2} q_{eq}(q(t_2) + q(t_1)) \right]$$

$$- \frac{1}{4\gamma} \left[ p_{eq}^2(t_2) + p_{eq}^2(t_1) \right] - \frac{1}{2\gamma} p_{eq}^2 \left[ p_{eq}(p_{eq}^2 + p_{eq}^2(t_1) + p(t_1)) \right]$$

$$\times \exp \left[ -\frac{i}{2} \left[ p_{eq}(q(t_2) - q(t_1)) - q_{eq}(p(t_2) - p(t_1)) \right]$$

$$+ (p(t_2)q(t_2) - p(t_1)q(t_1)) \right\}, \quad (\text{A4})$$

whose exponent can be suitably divided into a real and imaginary part. When setting $\gamma = \omega$ and $q_{eq} = 0$, the following is obtained for the real part of the exponent of Eq. (A4):

$$-\frac{\omega q^2(0)}{2} + \frac{p^2(0)}{2\omega} \left[ \cos(\omega t_2) + \cos(\omega t_1) \right]$$

$$- \frac{p_{eq}^2}{2\omega} \left[ \sin(\omega t_2) + \sin(\omega t_1) \right] \quad (\text{A5})$$

by substituting into the equations of motion (15) and (16). The imaginary contribution after summation with the difference of actions expression [Eq. (A3)] is given by,

$$-\frac{i}{2} \left[ p_{eq}q(0)(\cos(\omega t_2) - \cos(\omega t_1)) \right]$$

$$- \frac{i}{2} \frac{p_{eq}p(0)}{\omega} (\sin(\omega t_2) - \sin(\omega t_1)), \quad (\text{A6})$$

For the harmonic motion, the prefactor of Eq. (2) has the following expression:

$$C_{\omega}(p(t_1), q(t_1)) = \exp^{i\omega t_2 t_1/2}. \quad (\text{A7})$$

By inserting the results of Eqs. (A5)–(A7) into Eq. (7), the following expression for the spectral density is obtained,

$$I(E) = e^{-p^2/2\omega} \frac{Re}{(2\pi)^T} \int_0^T dt_1 \int_0^T dt_2 e^{-i(\tau t_2 - t_1)(E - \omega^2)} \int dq(0)$$

$$\times \exp \left[ -\frac{\omega q^2(0)}{2} - \frac{p_{eq}^2}{2}\left( \sin(\omega t_2) + \sin(\omega t_1) \right)$$

$$+ i \cos(\omega t_2) - i \cos(\omega t_1)) \right] \int dp(0) \exp \left[ -\frac{p^2(0)}{2\omega} \right]$$

$$+ \frac{p_{eq}p(0)}{2\omega} \left( \cos(\omega t_2) + \cos(\omega t_1) - i \sin(\omega t_2) \right)$$

$$+ i \sin(\omega t_1)) \right]. \quad (\text{A8})$$

After solving the Gaussian integrals and changing variables, $\tau = t_2 - t_1$, Eq. (A8) becomes

$$I(E) = e^{-p^2/2\omega} \frac{Re}{\pi} \int_0^{+\infty} d\omega \exp^{+iE(E-\omega^2)} \exp \left[ \frac{p^2}{2\omega} e^{-i\omega t} \right]. \quad (\text{A9})$$

Setting $p_{eq} = 0$ and performing the time integration, Eq. (A9)
leads to obtaining a spectral density composed of a single peak corresponding to the ground-state energy, \( I(E) = \frac{\partial E}{\partial E - \omega/2} / 2\pi \). In summary, we have shown that if the \([0,0]\) reference state is used, the single spectral peak obtained will be that associated with the ground-state energy. In the case of employing an arbitrary reference state for the momentum and by employing a series expansion for the relevant part of the exponent of Eq. (A9), one can derive the final expression for a harmonic power spectrum,

\[
I(E) = \frac{1}{\pi} e^{\frac{-p_{eq}^2}{2\omega}} \sum_{k=0}^{\infty} \left( \frac{p_{eq}^2}{2\omega} \right)^k \int_0^{\infty} d\tau e^{iE\tau - \omega/2 - \omega\tau}
\]

\[
eq \frac{1}{2\pi} e^{\frac{-p_{eq}^2}{2\omega}} \frac{p_{eq}^2}{2\omega} \sum_{k=0}^{\infty} \delta \left( E - \omega \left( k + \frac{1}{2} \right) \right).
\]  

(A10)

Equation (A10) spans the entire power spectrum and it has been obtained with a reference state of nonzero momenta. The same reasoning can be applied to reference states of the form \([0, q_{eq}]\) as described in Fig. 1(c). In other words, the arbitrary reference states of the type \([p_{eq}, q_{eq}], [0, q_{eq}], \) or \([p_{eq}, 0]\) exhibit a full power spectrum since they can be expanded in terms of harmonic oscillator eigenstates. In conclusion, the analytical considerations presented above highlight the relevance of the ansatz suggested in Eq. (10), where the reference state is given in terms of a set of carefully chosen coherent states.