The short periodic orbit method for excited chaotic eigenfunctions

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In this paper [published in Phys. Rev. E 102, 042210 (2020)], a new method for the calculation of excited chaotic eigenfunctions in arbitrary energy windows is presented. We demonstrate the feasibility of using wavefunctions localized on unstable periodic orbits as efficient basis sets for this task in classically chaotic systems. The number of required localized wavefunctions is only of the order of the ratio \( t_H/t_E \), with \( t_H \) the Heisenberg time and \( t_E \) the Ehrenfest time. As an illustration, we present convincing results for a coupled two-dimensional quartic oscillator with chaotic dynamics.

I. INTRODUCTION

The accurate computation of the eigenstates of a dynamical system is a central problem in Physics and Computational Chemistry. Usual standard methods [1] are based on the variational principle, which implies that in order to get an approximation to the \( N^{th} \) eigenstate, the \( N - 1 \) lower lying ones should also be calculated. This makes the task particularly demanding when interested in excited states. The existence of a large density of states or classically chaotic dynamics also contribute to significantly increment the computational burden. In the case of systems with hard wall boundaries, such as quantum billiards, eigenstate computations are not performed by the variational procedure, and then efficient methods to calculate only the states within specific energy windows have been developed [2]. This fact dramatically reduces the computational time for arbitrarily excited levels, and that fully opened the possibility of studying these interesting states. Unfortunately, such effective methods are not readily available for systems with continuous potentials.

In this paper, we present a method which extends the procedure recently developed by us [3, 4] for the computation of eigenstates of classically chaotic systems. This procedure is based on the use of wavefunctions localized on unstable periodic orbits (POs), which leads to small basis sizes, of the order of the ratio \( t_H/t_E \) between the Heisenberg, \( t_H \), and the Ehrenfest, \( t_E \), times. Recall that while the former provides the time scale at which the eigenenergies are accurately obtained, which scales as \( t_H \sim O(\hbar^{1-d}) \), being \( d \) the number of degrees of freedom of the system, the latter gives the time lapse where the semiclassical evolution of a wave packet is valid, being \( t_E \sim O(\log \hbar) \). A specific definition of these two important characteristic times for our system will be given below in Sec. [1].

The efficiency of the method is demonstrated by systematic application to small energy windows defined in different energy ranges of a two-dimensional coupled quartic oscillator with highly chaotic dynamics. The key point for the success of our method is the strong energy localization of our basis functions. This interesting property has enabled the calculation of the eigenfunctions of other chaotic systems, such as billiards [2], maps [6], or molecules [4, 6], which makes us to expect the general validity of our approach. Other attempts to use phase space basis functions have been made previously in the literature, especially in relation with molecular systems. For example, Davis and Heller used nonorthogonal complex Gaussians, which render analytical expressions for the elements of the Hamiltonian matrix in the case of polynomial potentials [7]. Reimers and Heller extended that semiclassical basis sets for rigid rotors [8]. Wavelets defined in the phase space have been also successfully applied in a series of papers by Poirier and coworkers [9–11]. Similarly, phase space Gaussians were also applied, on the one hand, by Halverson and Poirier [12], by incorporating some of their ideas previously used with wavelets, and, on the one hand, by Shimshovitz, Bäcić and Tan nor [13, 14], using the von Neumann lattice. More recently, Brown and Carrington [15, 16] have developed an iterative eigensolver which boosts the numerical calculations. Other long standing methods for computing molecular energies are the Discrete Variable Representation (DVR) [17, 18] using localized basis elements, for which the kinetic and potential energy contributions are easily computed, and the use of the powerful Lanczos diagonalization method [19].

Unstable POs are a cornerstone in classical and quantum chaotic dynamics [20]. They are the only remnants of order in the chaotic sea, and their invariant manifolds organize the classical dynamics in their neighbor-
hundreds \[21\]. From the quantum mechanical point of view, POs have also been shown to have a deep impact on the energy levels of classically chaotic systems \[20\], and also on the characterization of many of their eigenfunctions, where they can yield "scars" on the probability distributions, as shown by Heller in his seminal paper \[22\]. Bogo-
molny showed that this localization phenomenon can also arise in the combination of groups of eigenfunctions \[23\], and later Berry developed the corresponding phase space version of this theory \[24\].

Different methods have been reported in the literature for the construction of localized states over unstable POs (hereafter called "scar functions" due to its similarity with Heller’s scarring on eigenfunctions). Some use averages over groups of eigenfunctions around the PO quantization condition \[25\]; others make use of short PO theory \[26,27\], apply the asymptotic boundary layer \[28\], or perform the quantum propagation of wavepackets launched along the PO of interest \[21,29\].

In this work, we discuss the spectral properties of our basis functions, and demonstrate the accuracy of the proposed method by carrying out a systematic study of the convergence, both for eigenenergies and eigenfunctions in a small energy window. We also show the feasibility of using remarkably small basis sizes to calculate arbitrarily excited states. Only a brief description of the most relevant computational details and results are given in the text. Let us remark that the method reported would render even more accurate results in the case of more generic systems that present regular and irregular motion at the same time because of the more simple nodal structure of their eigenstates.

The rest of this work is organized as follows. First, we briefly describe in the next section the system under study, which is an anharmonic potential. Second, we describe in Sec. \[\text{III}\] the method that we have develop to construct efficient basis sets, in the sense that their sizes are small compared to the number of accurately computed eigenfunctions. Third, Sec. \[\text{IV}\] is devoted to the results and the corresponding discussion. Finally, we sum up the paper in Sec. \[\text{V}\] with the conclusions and outlook.

II. SYSTEM

The system we have chosen to study is described by the classical Hamiltonian

\[
\mathcal{H}(p_x, p_y, x, y) = \frac{p_x^2}{2} + \frac{p_y^2}{2} + \frac{x^2y^2}{2} + \frac{x^4 + y^4}{400}. \tag{1}
\]

The dynamics of this system is extremely chaotic, and no signs of regularity in phase space are observed at first sight \[21\].

The eigenfunctions of the associated quantum Hamiltonian belong to the five irreducible representations of the \(C_{2v}\) symmetry group. Nevertheless, we only consider here the totally symmetric (even on the axes, \(x, y\), and the diagonals, \(x = \pm y\)) eigenfunctions, which belong to the \(A_1\) representation. The mean number of \(A_1\) eigenenergies smaller than \(E\) is semiclassically estimated using the method reported in the Ref. \[30\] as \(\mathcal{N}(E) \simeq 0.251E^{3/2}/h^2 + 0.605E^{3/4}/h\). Then, the mean energy density given by \(\rho(E) = d\mathcal{N}(E)/dE \simeq 0.376E^{1/2}/h^2 + 0.454E^{-1/4}/h\), while the Heisenberg time is defined by the relationship

\[
t_H = 2\pi\hbar\rho(E). \tag{2}
\]

For the construction of the semiclassical basis one selects a set of unstable short POs in such a way that the sum of their periods is greater than \(t_H \[31\]

\[
\sum_{\gamma} T_\gamma > t_H, \tag{3}
\]

being \(T_\gamma\) the period of a short PO \(\gamma\). At this point it is worth noticing that the monodromy method \[32,33\] is exceptionally fast for calculating periodic orbits in a general Hamiltonian system, specially in high dimension. Nevertheless, here the situation is much more simple because our Hamiltonian has only two degrees of freedom and it presents the symmetries of the \(C_{4v}\) point group. As a consequence, the shortest periodic orbits, which are symmetrical, are trivially evaluated. Furthermore, the Hamiltonian \[1\] is mechanically similar, and then it presents no bifurcations as a function of the energy. Thus, the phase space structure does not change with the energy, and then one can compute all the periodic orbits at a reference energy, \(E\), and scale them for a different energy, \(E'\), as

\[
x'_{\gamma'} = \left(\frac{E'}{E}\right)^{1/4} x_{\gamma}, \quad p'_{x_{\gamma'}, \gamma'} = \left(\frac{E'}{E}\right)^{1/2} p_{x_{\gamma}}; \\
y'_{\gamma'} = \left(\frac{E'}{E}\right)^{1/4} y_{\gamma}, \quad p'_{y_{\gamma'}, \gamma'} = \left(\frac{E'}{E}\right)^{1/2} p_{y_{\gamma}}. \tag{4}
\]

with \(t' = (E'/E)^{-1/4} t\). Of course the first two periodic orbits of Fig.\[1\] are given without any calculation and consequently let us consider, for instance, orbits 3, 6 and 7. By fixing the starting point with \(y = 0\) and \(p_x = 0\) one looks for the coordinate \(x\) in such a way that the first intersection of the trajectory with the diagonal occurs orthogonal to it.

In this paper, we use the POs displayed in Fig.\[1\] with \(\sum T_\gamma \simeq 32\) and \(t_H \simeq 26\) at \(E = 110\), the maximum energy considered in the calculation. These POs are arranged by the number of bounces with the symmetry lines. Obviously, a criterion of general validity for arranging short POs with the purpose of covering phase space in an efficient way is desirable. We believe that such criterion could be defined in terms of the properties of the first homoclinic orbit of each short PO, particularly in higher dimension. Still, the condition \(3\) and the fact that one only needs the shortest POs of the system strongly limits the time required to search for POs.
Figure 1. Set of desymmetrized periodic orbits used in the calculation (red curves). The equipotential at energy \( E = 1 \) for Eq. (1) is shown in solid black curve.

Moreover, symmetry considerations can further simplify this task.

Associated with an unstable short PO \( \gamma \), one obtains the discrete set of Bohr-Sommerfeld (BS) energies, \( E_n^\gamma \), computed from the relationship

\[
\frac{S_\gamma}{\hbar} - \frac{\mu_\gamma \pi}{2} = 2\pi n,
\]

with \( S_\gamma \) the action of \( \gamma \) and \( \mu_\gamma \) its Maslov index.

We provide a semiclassical construction of wavefunctions with \( n \) excitations along \( \gamma \), the so-called tube functions of \( \gamma \), with mean energy \( E_n^\gamma \) and small energy dispersion given by \( \hbar \lambda_\gamma/\sqrt{2} \); where \( \lambda_\gamma \) is the stability exponent of \( \gamma \). Although the time evolution of these wavefunctions can be evaluated at semiclassical level in terms of homoclinic and heteroclinic orbits \([34, 35]\), here we prefer to use a quantum evolution, following Ref. \([36]\), in order to obtain highly accurate eigenfunctions and eigenvalues.

In this respect, it is worth mentioning that the quantum evolution of tube functions up to times of the order of the mean Ehrenfest time \( t_E \), that in our case is given by

\[
t_E = \frac{1}{2\lambda} \ln \left( \frac{A}{\hbar} \right),
\]

with \( \lambda \simeq 0.385E^{3/4} \) the Lyapunov exponent of the system, and \( A \simeq 11.1E^{3/4} \) the area of a characteristic Poincaré surface of section, is a simple task.

### III. METHOD

For the computation of eigenvalues and eigenfunctions contained in an energy window defined by the central energy \( E_0 \) and width \( \Delta E \), \([E_0 - \Delta E, E_0 + \Delta E]\), we proceed as follows.

First, we select the set of BS energies of the POs existing in the enlarged energy window \([E_0 - (\Delta E + \delta), E_0 + (\Delta E + \delta)]\), with \( \delta \simeq 2(t_H/t_E)/\rho(E) \) an energy window containing \( 2t_H/t_E \) eigenvalues.

Second, we construct tube functions of the selected BS energies, for instance \( E_0^\gamma \). By propagating a Gaussian wave packet over \( \gamma \) using the frozen Gaussian approximation \([37]\), one has

\[
\psi_n^\gamma(x,y) = \int_0^{T_H} e^{iE_n^\gamma t/\hbar} \phi(x,y,t) dt,
\]

where \( \phi(x,y,t) = \exp \{ -\alpha[(x-x_0)^2 + (y-y_0)^2]/\hbar + i[px(x-x_0)/\hbar + p_y(y-y_0)/\hbar + 2\pi nt/T_H] \} \),

\[
(x_0, y_0, p_x, p_y) \text{ the classical evolution over } \gamma.
\]

The energy dispersion of \( \psi_n^\gamma(x,y) \) behaves like \( Cn^2 \) in leading order, with \( C \) a constant that depends on the parameter \( \alpha \) of Eq. (8). By selecting the parameter \( \alpha \) that minimizes the energy dispersion for a finite value of \( \hbar \), we obtain a good estimate of the parameter that minimizes the coefficient \( C \). Of course, the main idea behind this choice is that as the energy dispersion is smaller, the wave function is closer to an eigenfunction. In conclusion, one selects the parameter \( \alpha \) in such a way that the energy dispersion of \( \psi_n^\gamma(x,y) \) results a minimum.

Third,scar functions are constructed by quantum propagation of the tube functions (7) up to \( t_E \),

\[
\langle x,y|n,n \rangle = \int_{-t_E}^{t_E} \cos \left( \frac{\pi t}{2t_E} \right) e^{(E_n^\gamma - \delta)t/\hbar} \psi_n^\gamma(x,y) dt.
\]

This time evolution is the most demanding point of our method, but still it can be efficiently carried out using procedures like the discrete variable representation \([17, 18]\) because of the small energy dispersion of the scar functions \([9]\). Similar results are expected when other basis functions with similar spectral properties are used, e.g. the scar functions constructed using the method reported in Ref. \([26]\). Similarly, Eq. (7) could be also used to construct the semiclassical basis set, but the much smaller spectral dispersion of the scar functions \([9]\) after a relative short time propagation makes these last functions better suited. As a particular example, notice in Fig. 2(a) that the spectra of four of the scar functions involved in our calculations is mostly concentrated in the window of interest. Here, the infinite resolution spectra are shown as vertical thin lines, while its finite-resolution counterpart, which is defined as the
Figure 2. (a) Bohr-Sommerfeld quantized energies (vertical green lines) for the 61 scar functions in the basis set used to obtain the 9 eigenstates of $A_1$ symmetry (vertical black lines) of the quartic oscillator $[1]$ lying in the energy window $[105.5, 107.5]$ (between gray shaded regions). The infinite (vertical colored lines) and low resolution (continuous thick curves) projections on the eigenenergies spectrum of the $|11, 44\rangle$ (red), $|5, 24\rangle$ (blue), $|14, 55\rangle$ (cyan), $|17, 67\rangle$ (brown) scar functions shown in Fig. 4 (see text for notation), which are the ones which are mainly involved in the reconstruction of the eigenfunction 297, have been superimposed. (b) Errors in the computed eigenenergies (orange circles) and eigenfunctions (purple diamonds) given by Eq. (11).

Lorenzian convolution

$$I_\Delta(E) = \frac{\Delta E}{2\pi} \sum_N \frac{\langle \psi_\gamma^N \rangle^2}{(E - E_N)^2 + (\Delta E/2)^2}, \quad (10)$$

with $\Delta E = 0.4$ and $E_N$ is the energy of the eigenstate $|N\rangle$, are shown as vertical black thick lines at the bottom of Fig. 2(a). Notice that this spectra can only be obtained once the eigenfunctions of the system have been calculated.

Fourth, the Gram-Schmidt Selective method [3] is then used to choose the most fit scar functions. The method works as follows. One selects the scar function with minimum energy dispersion as the first function of the basis, for instance $|\gamma, n\rangle$. Next, one eliminates, for the other scar functions, the contribution along $|\gamma, n\rangle$, for instance $|\eta, m\rangle$ (with $\eta$ other PO) is replaced by $|\eta, m\rangle - \langle \gamma, n | \eta, m \rangle |\gamma, n\rangle$. Then, one evaluates the energy dispersion of the new wave functions and selects as the second function of the basis, the one with minimum dispersion. This process continues in a similar way to the Gram-Schmidt method while the minimum energy dispersion is smaller than a given value, for instance 2%. The method has the effect of describing the subspace spanned by the desired eigenfunctions, i.e., those whose energies lie within the energy window under study.

Fifth, once the final basis set is obtained the corresponding Hamiltonian matrix is computed and diagonalized using standard procedures.

Figure 3. Probability density of the eigenfunctions contained in the energy window $E = [105.5, 107.5]$. They are scaled to $E = 1$ for comparison. The corresponding eigenstate number is given in the plots.

IV. RESULTS AND DISCUSSION

Figure 8 presents the results of a computation performed in the energy window $[105.5, 107.5]$ for $\hbar = 1$. There are 66 scar functions contained in the enlarged energy window $[100.2, 112.8]$ [see vertical green lines in Fig. 2(a)], and the Gram-Schmidt Selective method chooses 61 scar functions in order to reproduce with high accuracy the 9 eigenvalues [black lines contained in the central white region of Fig. 2(a)] and eigenfunctions of interest. The figure displays the probability density of
these eigenfunctions in the fundamental domain. As can be seen, they all present a very complex nodal pattern, something characteristic of classically chaotic systems. With respect to the scar phenomenon, we clearly observe that the eigenfunction 293 is localized along the first PO of Fig. 1.

The eigenfunction $|j\rangle$ of our computation is compared with the "exact" result $|j'\rangle$ obtained using a basis set of 5000 harmonic oscillator functions. We compute the error of eigenvalues and eigenfunctions by the dimensionless quantities

$$\rho(E) |E_j - E_{j'}| \quad \text{and} \quad 1 - \langle j'|j\rangle^2,$$

respectively. The mean error of the 9 eigenvalues and eigenfunctions of Fig. 3 is of order $\sim 10^{-4}$ and $\sim 10^{-5}$, respectively, as shown in Fig. 2(b). A similar accuracy based on the variational principle would require much larger full matrix diagonalizations. In particular, for the energy window reported in the manuscript, the same level of accuracy requires a harmonic oscillator basis set formed by $\sim 50$ times more basis elements than the one reported here, which exerts a diagonalization time $\sim 50^3$ times larger. Furthermore, by accepting an error ten times greater than the previous ones, the same calculation provides 20 consecutive eigenfunctions. With respect to the stability of the method, let us remark that in other energy windows of the spectrum up to $E = 110$, we found essentially the same result. Moreover, for smaller energies a reduced number of POs is required as long as Eq. (3) is accomplished. Finally, we find that the dimension of the basis is only 40% greater than the number of eigenfunctions contained in the enlarged energy window.

Let us consider now the representation of the eigenfunctions obtained in our calculation in the basis set of scar functions, this showing the basis elements that contribute the most (largest overlaps) to their reconstruction. Notice that this procedure provides valuable information on the influence that each PO has on the emergence of the properties of individual chaotic eigenfunctions, this representing an important added bonus of our method to the theory of quantum chaos. As an illustrative example, we present in Fig. 4 the reconstruction of the eigenfunction 297 shown in Fig. 3. As already mentioned, it exhibits a complex nodal pattern, as it is not localized over any single PO. Successive steps in the reconstruction are given in the figure. In each one of them we show the result of adding the scar function depicted in the panel above it. Notice that 41.5% of the eigenfunction is reconstructed solely with the scar function $|11,44\rangle$. By adding the scar function $|5,24\rangle$, 65.3% of the eigenfunction is reconstructed. This figure increasingly raises when more basis functions are included, up to a value of 84.4% when four basis functions are combined. In other words, the square modulus of the projection of the eigenstate 297 onto the subspace spanned by the four scar functions of the figure is 0.844. Notice also how the reconstruction procedure, that we have just described, progressively refines the intricate details of the eigenfunction.

The other eigenfunctions of Fig. 3 have a similar partial reconstruction in terms of a small number of scar functions. In particular, we remark the cases where only one scar function is able to reproduce more than 50% of the eigenfunction. The scar function $|1,56\rangle$ reproduces 79.7% of 293, $|8,41\rangle$ reproduces 60.6% of 296, and $|16,56\rangle$ reproduces 58.9% of 300.

In order to clearly show the strong energy localization of our basis functions, we conclude this section by displaying in Fig. 5 the square modulus of the overlap of the first wave function in Fig. 4, i.e., the scar function $|11,44\rangle$, with the eigenfunctions of the system. In
Figure 5. Square modulus of the overlap of the first wave function of Fig. 4, i.e., the scar function |11, 44⟩, with the eigenfunctions of the system, as a function of the eigenenergies in linear (a) and logarithmic (b) scales. The thin green line marks the position of the Bohr-Sommerfeld quantization energy (E ≃ 106.75), which lies within the reference energy window [105.5, 107.5] (between the gray shaded regions).

Panel (a) we use a linear scale to remark the fact that the energy dispersion of the wave function consists only of contributions within a few mean level spacings. Notice here that the highest peak lies very close to the Bohr-Sommerfeld energy of the scar function (thin green line at E ≃ 106.75), and corresponds to the overlap associated with the eigenstate 297. Recall that the scar function |11, 44⟩ is precisely the main contribution to the reconstruction of that eigenstate. Furthermore, in panel (b) we use a logarithmic scale to emphasize that the intensities decay exponentially as we move away from the corresponding Bohr-Sommerfeld energy.

V. CONCLUSIONS AND OUTLOOK

The short periodic orbit method, formulated and tested in this paper, is an extremely powerful technique for calculating very excited chaotic eigenfunctions. We have shown the feasibility of computing eigenfunctions of a classically chaotic system using small basis sets. The key point is the use of scar functions constructed in the neighborhood of a set of short periodic orbits that covers the relevant phase space efficiently. These semiclassical functions have excellent properties regarding localization in energy that make them suitable to describe only the portion of the system’s Hilbert space spanned by the eigenstates of interest. In particular, the accuracy of the method has been demonstrated by performing a systematic calculation of the eigenfunctions of a coupled quartic oscillator using narrow energy windows in excited regions of the spectrum. Let us finally remark that each eigenfunction obtained in the calculation is partially reconstructed using only a few scar functions, of the order of the t_H/t_E ratio. This fact provides a useful way of classifying chaotic eigenfunctions.

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