# Orbits in the H<sub>2</sub>O molecule

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We study the forms of the orbits in a symmetric configuration of a realistic model of the  $H_2O$  molecule with particular emphasis on the periodic orbits. We use an appropriate Poincaré surface of section (PSS) and study the distribution of the orbits on this PSS for various energies. We find both ordered and chaotic orbits. The proportion of ordered orbits is almost 100% for small energies, but decreases abruptly beyond a critical energy. When the energy exceeds the escape energy there are still nonescaping orbits around stable periodic orbits. We study in detail the forms of the various periodic orbits, and their connections, by providing appropriate stability and bifurcation diagrams. © 2001 American Institute of Physics. [DOI: 10.1063/1.1356068]

Small molecules are quantum systems, but their classical study reveals very important features. The method of the Poincaré surface of section plays a prominent role in such a study. A difficulty that appears often in realistic systems is that one cannot always choose a flat Poincaré surface of section that intersects all periodic orbits. In this paper we use a curved Poincaré surface of section in order to study a realistic model of the H<sub>2</sub>O molecule. We find the phase plots for different values of the energy of the molecule. Then we find all the periodic orbits of period 1 and 2 and compute their stability and bifurcation diagrams.

## I. INTRODUCTION

Two dimensional Hamiltonian systems are often studied using the technique of the Poincaré map, where one reduces the four dimensional, continuous time flow of the system to an associated *two dimensional* discrete map, by choosing an appropriate surface of section. Usually a flat surface of section is chosen and the orbits that cross this surface with a particular direction are the consequents of the Poincaré map.

For many systems a flat Poincaré surface of section (PSS) is not suitable since there are orbits of these systems that do not cross the surface of section. In these cases another choice must be made. In this paper we study a symmetric model of the  $\rm H_2O$  molecule where we have chosen a curved PSS. This choice for the PSS permits us to study all the periodic orbits of the system for all the values of energy.

The study of periodic orbits and phase plots in realistic models of simple molecules, such as the model we are studying in this paper, gives interesting information that can be compared with the behavior of the corresponding quantum system and with experiments. <sup>1,2</sup> In particular the relation between classical periodic orbits and quantum mechanical eigenfunctions was emphasized by several authors, starting with the work of Gutzwiller. <sup>3</sup> The most surprising result was that in many cases one simple periodic orbit determines the

basic form of the eigenfunction.<sup>4–7</sup> Further details can be found by using more periodic orbits<sup>8</sup> and the asymptotic structures around unstable periodic orbits.<sup>9</sup>

A partial study of periodic orbits in the H<sub>2</sub>O molecule has been done by Lawton and Child, <sup>10,11</sup> Jaffé and Brumer, <sup>12</sup> and by Kellman, <sup>13</sup> who emphasize the bifurcation of the local stretching modes from the normal stretching vibrational mode. These papers contain references to previous work on orbits in the H<sub>2</sub>O molecule.

We describe the model of the  $H_2O$  molecule in Sec. II, and use a convenient set of coordinates in defining the Hamiltonian. We calculate the equipotential surfaces for the symmetric molecule (Sec. III) and in Sec. IV we define an appropriate PSS and discuss its properties.

Then we study in detail the phase plots on such a PSS and find the main islands of stability and the chaotic zones (Sec. V). We calculate the main periodic orbits and their bifurcations and stability in Sec. VI. Finally we summarize our conclusions in Sec. VII.

## II. A MODEL FOR THE H<sub>2</sub>O MOLECULE

For the study of the H<sub>2</sub>O molecule we have chosen a simple model that is well suited for classical calculations.<sup>14</sup> The potential energy has the form

$$\begin{split} V_{\rm HHO} &= V_{\rm O}^{(1)} \cdot f(R_1, R_2, R_3) + V_{\rm OH}^{(2)}(R_1) + V_{\rm OH}^{(2)}(R_2) \\ &+ V_{\rm HH}^{(2)}(R_3) + V_{\rm HHO}^{(3)}(R_1, R_2, R_3), \end{split} \tag{1}$$

where  $R_1$ ,  $R_2$ , and  $R_3$  are the O-H<sub>1</sub>, O-H<sub>2</sub>, and H<sub>1</sub>-H<sub>2</sub> distances, respectively. Distances are measured in Å and energies in eV. The single-body term in Eq. (1) is given by

$$V_{\rm O}^{(1)} \cdot f(R_1, R_2, R_3) = 1.958 \cdot \frac{1}{2} \left[ 1 - \tanh \left( \frac{3\rho_3 - \rho_1 - \rho_2}{2} \alpha \right) \right], \tag{2}$$

where  $\alpha$ =1.9018,  $\rho_1$ = $R_1$ -0.9572,  $\rho_2$ = $R_2$ -0.9572, and  $\rho_3$ = $R_3$ -1.5139. The two-body terms are given by the equations

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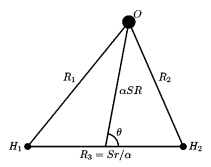


FIG. 1. Coordinates for the H<sub>2</sub>O molecule.

where  $r_i = R_i - 0.9696$ ,  $a_1 = 4.507$ ,  $a_2 = 4.884$ ,  $a_3 = 3.795$ , and  $D_1 = 4.6211$ ,

$$V_{\rm HH}^{(2)}(R_3) = -D_2(1 + a_4r_3 + a_5r_3^2 + a_6r_3^3) \exp\{-a_4r_3\}, \qquad (4)$$

where  $r_3 = R_3 - 0.7414$ ,  $a_4 = 3.961$ ,  $a_5 = 4.064$ ,  $a_6 = 3.574$ , and  $D_2 = 4.7472$ . Finally the three-body term is given by

$$V_{\text{HHO}}^{(3)}(R_1, R_2, R_3) = 0.01892 \cdot P(\rho_1, \rho_2, \rho_3)$$

$$\times \prod_{i=1}^{3} (1 - \tanh(\gamma_i \rho_i / 2)), \tag{5}$$

where  $\gamma_1 = \gamma_2 = 2.6$ ,  $\gamma_3 = 1.5$ , and  $P(\rho_1, \rho_2, \rho_3)$  is the polynomial

$$P(\rho_{1}, \rho_{2}, \rho_{3}) = 1 + \sum_{i=1}^{3} C_{i}\rho_{i} + \sum_{i=1}^{3} \sum_{j=i}^{3} C_{ij}\rho_{i}\rho_{j}$$

$$+ \sum_{i=1}^{3} \sum_{j=i}^{3} \sum_{k=j}^{3} C_{ijk}\rho_{i}\rho_{j}\rho_{k}$$

$$+ \sum_{i=1}^{3} \sum_{j=i}^{3} \sum_{k=j}^{3} \sum_{l=k}^{3} C_{ijkl}\rho_{i}\rho_{j}\rho_{k}\rho_{l}.$$
 (6)

The coefficients  $C_i$ ,  $C_{ij}$ ,  $C_{ijk}$ , and  $C_{ijkl}$  are defined in the paper by Murrell and Carter. <sup>14</sup>

Instead of the original coordinates  $R_1$ ,  $R_2$ , and  $R_3$  we will use the scaled Jacobi coordinates SR, Sr,  $\theta$  (Fig. 1) that are defined by the equations

$$SR = \frac{1}{2\alpha} \sqrt{2R_1^2 + 2R_2^2 - R_3^2},$$

$$Sr = \alpha R_3, \tag{7}$$

$$\cos\theta = \frac{R_2^2 - R_1^2}{2\alpha \, SR \, R_2},$$

where  $\alpha$  is given by the relation

$$\alpha = \left\{ \frac{m_{\rm O} + 2m_{\rm H}}{4m_{\rm O}} \right\}^{1/4}.$$
 (8)

The inverse relations are

$$R_1 = \sqrt{\left(\frac{\text{Sr}}{2\alpha}\right)^2 + (\alpha \text{ SR})^2 - \text{SR Sr cos } \theta},$$

$$R_2 = \sqrt{\left(\frac{\text{Sr}}{2\alpha}\right)^2 + (\alpha \,\text{SR})^2 + \text{SR} \,\text{Sr}\cos\theta},\tag{9}$$

$$R_3 = \frac{\mathrm{Sr}}{\alpha}$$
.

The potential function given by Eqs. (1)–(6) can be written in scaled Jacobi coordinates using Eqs. (9).

The kinetic energy of the H<sub>2</sub>O molecule has the form

$$T = \frac{1}{2\mu} \left[ P_{SR}^2 + P_{Sr}^2 + \left( \frac{1}{SR^2} + \frac{1}{Sr^2} \right) P_{\theta}^2 \right], \tag{10}$$

where  $P_{\rm SR}$ ,  $P_{\rm Sr}$ , and  $P_{\theta}$  are the conjugate momenta of SR, Sr, and  $\theta$ , respectively. The parameter  $\mu$  is defined by the relation

$$\mu = \sqrt{\frac{m_{\rm O} m_{\rm H}^2}{m_{\rm O} + 2m_{\rm H}}} = \frac{m_{\rm H}}{2\alpha^2}.$$
 (11)

The Hamiltonian can now be written as the sum of the kinetic and potential energies

$$H(P_{SR}, P_{Sr}, P_{\theta}, SR, Sr, \theta) = T(P_{SR}, P_{Sr}, P_{\theta}, SR, Sr) + V_{HHO}(SR, Sr, \theta).$$
(12)

#### III. EQUIPOTENTIAL SURFACES

If we consider initial conditions with

$$\theta = \pi/2$$

and

$$P_{\theta} = 0, \tag{13}$$

we can easily see that  $\dot{\theta}=0$  and  $\dot{P}_{\theta}=0$ , meaning that the submanifold of the phase space determined by Eqs. (13) is invariant under the Hamiltonian flow and hence we can perform a standard type reduction in order to get a two degrees of freedom Hamiltonian system on this submanifold.

The reduced Hamiltonian  $\widetilde{H}$  is related to the original Hamiltonian through the relation

$$\tilde{H}(P_{SR}, P_{Sr}, SR, Sr) = H(P_{SR}, P_{Sr}, 0, SR, Sr, \pi/2) = E$$
 (14)

and its numerical value is the energy E of the molecule. This Hamiltonian describes the symmetric water molecule, where the distances  $O-H_1$  and  $O-H_2$  are equal.

The equipotential surfaces for the symmetric molecule are shown in Fig. 2. The potential has two minima at SR =  $\pm 0.676\,065$ , Sr=1.310 82, where it takes the value  $E_0$  = -10.047. Between the two minima there is a saddle point at SR=0, Sr=1.6057, where the potential takes the value  $E_1$ = -8.993. For  $E < E_1$  orbits are confined inside the potential well around a minimum of the potential. For  $E > E_1$  the two wells join and orbits can pass from one well to the other.

For values of energy greater than  $E_2 = -2.7892$  orbits can escape to infinity following the two horizontal channels at the left and the right of Fig. 2. Escape through these channels corresponds to a configuration of the molecule where the oxygen atom has become unbounded and the two hydrogen atoms form an  $H_2$  molecule. Although for  $E > E_2$  most orbits escape there are still bounded orbits as one can see in Fig. 6(c) below.

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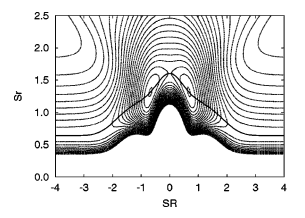


FIG. 2. Equipotential surfaces for the symmetric  $H_2O$  molecule. The solid line is the Poincaré surface of section.

## IV. A POINCARÉ SURFACE OF SECTION

A PSS should intersect almost all the orbits. Such intersections occur naturally if the potential has a plane of symmetry. However this does not happen in the present case. On the other hand many orbits have a tendency to pass through a local minimum of the potential (although this is not always the case). Thus in order to find a suitable Poincaré surface of section we calculated numerically for each value of SR the corresponding value of Sr, where the potential function has a minimum. Then we fitted the numerically determined function  $Sr = f_{num}(SR)$  by the analytic form

$$Sr = f(SR) = \alpha \overline{f}(\alpha SR), \tag{15}$$

where

$$\overline{f}(x) = 0.740350 + 0.482392 \exp(-3.34209x^2) + 0.631258 \exp(-0.112678x^4)$$
 (16)

and  $\alpha$  is given by Eq. (8).

This curve is given in Fig. 2. We determined that the difference between the numerically determined curve  $Sr = f_{num}(SR)$  and the fitted analytic form is very small. For our computations we used the fitted form, given by Eqs. (15) and (16).

Contrary to the textbook approach we have not chosen SR and  $P_{\rm SR}$  as the two coordinates on the PSS. If we had made such a choice we would have the following problem. Consider the orbits of Fig. 3. Both orbits cross the PSS at the

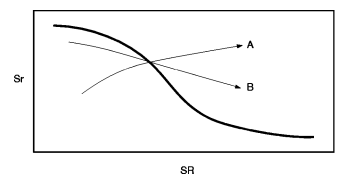


FIG. 3. The thick line gives the PSS. Orbits A and B cross the line at the same point with the same  $P_{SR}$  but opposite  $P_{Sr}$ .

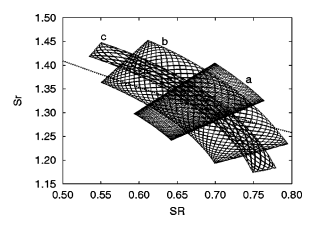


FIG. 4. Orbits a and c belong in the islands of 1a and 1c, respectively. The orbit b lies between these two islands. The thick line represents the PSS.

same point with coordinates SR, Sr with the same momentum  $P_{\rm SR}$  and they both have the same energy E. Then we solve Eq. (14) in order to determine  $P_{\rm Sr}$ , and we get two solutions  $\pm P_{\rm Sr}$ . From Fig. 3 we can see that both solutions are valid, but they correspond to different orbits. This in turn means that on such a PSS different orbits would overlap and this is not permissible.

In order to solve this problem we have chosen as coordinates on the PSS, SR and the component  $P_t$  of the momentum vector  $P = (P_{SR}, P_{Sr})$  tangential to the curve in Eq. (15). Thus the PSS is determined by the coordinate SR along the curve of Fig. 2 and the component  $P_t$  of the momentum. Some typical orbits on the plane (SR,Sr) are shown in Fig. 4.

A variable that is canonically conjugate to  $P_t$  is

$$Q_t = \int_0^{SR} \sqrt{1 + f'(x)} dx. \tag{17}$$

In fact the symplectic form in the variables  $Q_t$ ,  $P_t$  is

$$\Omega = (dQ_t) \wedge (dP_t) \tag{18}$$

and it can be shown that this is preserved. The form Eq. (18) in the variables SR,  $P_t$  is written

$$\Omega = (1 + f'(SR))^{1/2} (dSR) \wedge (dP_t). \tag{19}$$

Thus the two-form  $d\operatorname{SR} \land dP_t$  is not preserved in general. In our study we use the variable SR instead of  $Q_t$  because it is not practical to compute  $Q_t$  numerically.

Although the coordinates SR and  $P_t$  are not canonically conjugate, this does not cause any problems in the study of the system. In fact the integration is performed in the canonical coordinates SR, Sr,  $P_{\rm SR}$ ,  $P_{\rm Sr}$  and the coordinates SR and  $P_t$  are used only on the PSS. The result is that the phase plots on the PSS are only slightly distorted relative to the phase plots that we would get if we had chosen canonically conjugate coordinates on the PSS.

# V. PHASE PLOTS ON THE POINCARÉ SURFACE OF SECTION

In Figs. 5 and 6 we give the distribution of the orbits on the Poincaré surface of section for various values of the energy *E*.

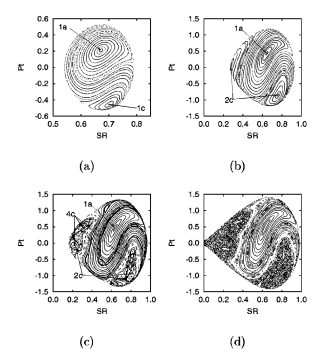


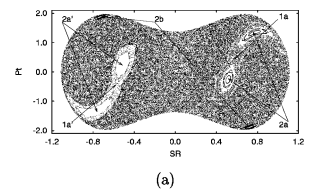
FIG. 5. Phase plots for small values of the energy: (a) E = -9.9, (b) E = -9.3, (c) E = -9.1, and (d) E = -8.99.

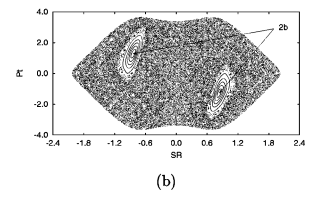
The energy of Fig. 5(a) is close to the minimum of the potential. In this plot practically all the orbits are ordered, defining closed invariant curves. Many invariant curves form two sets around two stable periodic orbits. The upper one is the orbit 1a and the lower one the orbit 1c (see Sec. VI). However there are also invariant curves between these two sets, that start on the left or upper side of the boundary, and terminate on the right side of the boundary. In Fig. 4 we can see why this is happening. In this figure we see three orbits for this value of the energy in configuration space. The orbits labeled as a and c belong to the islands of the periodic orbits 1a and 1c, respectively. We see that the PSS crosses the outline of these orbits at two nonadjacent sides. When this is happening we get closed invariant curves on the PSS. But for the orbit labeled b, which lies between the two islands of Fig. 5(a), the PSS crosses the outline of this orbit at two adjacent sides (Fig. 4), and on the PSS we get an open invariant curve.

Figures 5(b), 5(c), and 5(d) refer to larger energies and are given in a larger scale than Fig. 5(a). Orbit 1c is continued as the double period orbit 2c in Fig. 5(b), the second point being near the left limit of the figure. The islands around both points 2c are now larger. In this case the chaotic orbits are also insignificant.

In Fig. 5(c) the orbit 2c is unstable (and is represented by two points at the centers of two dark regions) and it has produced four stable points by bifurcation that correspond to a stable orbit 4c of period 4, surrounded by islands of stability above and below each point of orbit 2c. In this case there are some small chaotic zones around each unstable orbit.

The chaotic zones have grown considerably in Fig. 5(d). In this case there are still four islands of stability 4c, as in Fig. 5(c), but very small. All the rest of the area previously





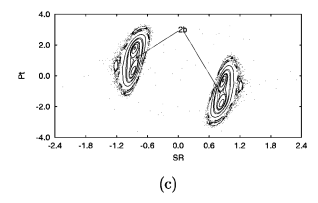


FIG. 6. Phase plots for energies: (a) E=-8.0, (b) E=-3.0, and (c) E=-2.5.

occupied by bifurcations of orbit 2c and their corresponding islands are now chaotic. This case is just above the limiting value of the energy  $E_1 = -8.993$  at which the two potential wells join into one. Thus there is one symmetric figure for SR<0 that joins Fig. 5(d) at the small throat near the point (0,0). The point (0,0) represents orbit 1b, which is unstable and produces a large chaotic region around it.

In Fig. 6(a) we give the phase plot for a larger value of the energy (E=-8) when the two potential wells are well connected at SR=0 (see the equipotentials of Fig. 2). Figure 6(a) is symmetric with respect to the axis SR=0. The apparent asymmetries in the islands of this figure are due to the use of nonsymmetric initial conditions on both sides of the axis SR=0. In this case orbit 1a is unstable and has generated by bifurcation orbit 2a, surrounded by two islands of stability on the right half of Fig. 6(a). A symmetric orbit 1a' and its bifurcations 2a' appear on the left side of the figure.

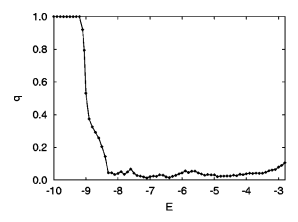


FIG. 7. Proportion of regular orbits, as a function of the energy E.

Orbits 1a, 1a' have generated two chaotic domains that surround the islands 2a (respectively, 2a'). At the lower left and upper right sides of this figure there are two elongated islands that belong to the family 2b. The center of the figure (0,0) represents again the unstable orbit 1b, which generates a large chaotic domain.

Most of Fig. 6(a) is covered by one chaotic orbit for which we have calculated  $6 \cdot 10^4$  points. We see that the points of the chaotic orbit are not distributed uniformly on the PSS. This is a transient phenomenon and is due to stickiness of the orbits near cantori surrounding islands 2a, or islands 2a'. These cantori form partial barriers. Orbits need a long time in order to cross the cantori and for considerable times the regions inside and outside the cantori seem separated. This causes the apparent nonuniformity of the distribution of points on the PSS. But after a larger number of iterations the distribution of the points becomes uniform.

The phase plots for larger values of E have some similarity to Fig. 6(a) but there are also differences. For example, in Fig. 6(b) we see that there are two symmetric islands, one on each side of the axis SR=0. These islands belong to the irregular family 2b, which is symmetric with respect to the point (0,0) (see Sec. VI), and not to two different families like families 1a and 1a' of Fig. 6(a). The chaotic domain in this case is again very large.

For E above the energy  $E = E_2$  orbits can escape. In Fig. 6(c) we can see the islands around an orbit that belongs in family 2b. The orbits that are outside these islands escape and this is why the area between the islands is empty.

The percentage of the area of the PSS covered by organized orbits as a function of the energy is presented in Fig. 7. We see that for values of energy close to  $E_1\!=\!-10.047$  (the minimum value of the potential) almost all orbits are organized. An abrupt change happens at values of energy around  $E\!=\!-9$ . For  $E\!>\!-9$  most of the PSS is filled by chaotic orbits. There are islands of stability for even larger values of the energy, and we notice that for values of the energy close to  $E\!=\!-3$  the percentage of the organized orbits increases, although it remains below 0.2 (or 20%). This increase is due to the increase in the size of the island around family 2b. The islands continue to increase in size even when E becomes larger than the escape energy  $E_2\!=\!-2.79$ , but for still larger E they become smaller.

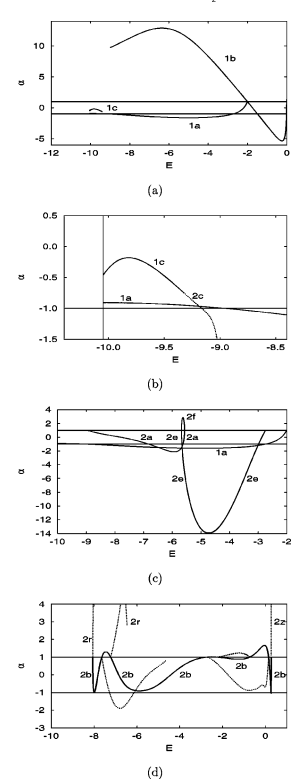


FIG. 8. Stability diagrams: (a) families 1a, 1b, 1c, (b) families 1a, 1c, 2c (continuation of 1c), (c) family 1a and its period 2 bifurcations, and (d) families 2b, 2r, 2z.

## VI. PERIODIC ORBITS

We have computed the most important families of periodic orbits for the symmetric water molecule. The stability diagram of the period-1 families as a function of the energy of the molecule can be seen in Fig. 8(a). Such a diagram gives the Hénon stability parameter  $^{16}$   $\alpha$  of each family as a

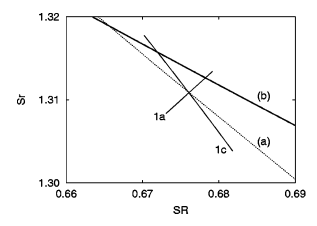


FIG. 9. Periodic orbits 1a and 1c for E = -10.04664 in configuration space. Line (a) represents the numerically determined minima of the potential and the line; (b) represents the selected PSS.

function of E. For a period-k orbit of an area-preserving map  $\sigma$ , the Hénon stability parameter is defined by the relation

$$\alpha = \frac{1}{2} \operatorname{Trace}(D[\sigma^k](x_0)), \tag{20}$$

where  $x_0$  is a point of the periodic orbit and  $D[\sigma^k](x_0)$  is the Jacobian matrix of  $\sigma \circ \cdots \circ \sigma$  (k times) evaluated at  $x_0$ . When the stability parameter of a periodic orbit lies in the interval from -1 to 1 the orbit is stable, otherwise it is unstable.

Since the Poincaré map is not symplectic, as we have explained in Sec. IV, the determinant of the Jacobian matrix  $D\sigma(x)$  is not equal to 1 for arbitrary points x on the PSS. However for a periodic orbit with period k the determinant of  $D[\sigma^k]$  is exactly equal to 1. In fact, using Eq. (19) it can be proven that the determinant of the Jacobian matrix of  $\sigma^k$  at a point  $x_0 = (SR_0, P_{t0})$  is given by the relation

$$\det D[\sigma^{k}](x_{0}) = \sqrt{\frac{1 + f'(SR_{0})}{1 + f'(SR_{k})}},$$
(21)

where  $x_k = \sigma^k(x_0) = (SR_k, P_{tk})$  is the kth image of  $x_0$ . If an orbit is periodic with period k, then  $SR_0 = SR_k$ , hence  $\det D[\sigma^k](x_0) = 1$ .

For values of energy close to the minimum value of the potential  $(E=E_1=-10.047)$  there exist two period-1 families. We call these families 1a and 1c. They correspond to the two normal modes of the water molecule. In Fig. 9 we can see the corresponding periodic orbits at E=-10.04664 in configuration space.

Family 1c has a peculiar behavior that is the result of our choice of the PSS. This family is born at E=-10.047, but beyond the energy E=-9.331 it appears as a period-2 family, which we call 2c [Fig. 8(b)]. We note that no bifurcation appears here. In order to understand what is happening, we should see Fig. 10. In this figure we see one orbit of family 1c that intersects the PSS only at one point, as expected for small energies. An orbit of the same family for a larger value of the energy intersects the PSS at two points and thus it appears as a period-2 orbit. We will call the change of the apparent period of a family that is not related to a bifurcation, a *transition*. We note here that there would be a transition even if we had chosen, as the PSS, the numerically determined curve of the minima of the potential

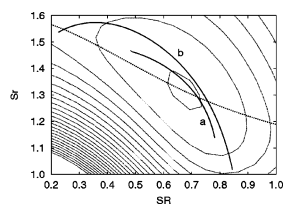


FIG. 10. The transition  $1c \rightarrow 2c$ . Orbit a (E = -9.8) belongs to family 1c. Orbit b (E = -9.2) belongs to family 2c.

instead of the fitted function. Family 2c becomes unstable at E = -9.164 and generates by bifurcation two stable period 4 families.

For values of the energy between  $E = E_1 = -10.047$  and E = -7.2 the largest islands on the PSS are around family 1a and family 2a, which bifurcate from 1a [Figs. 5 and 6(a)]. Family 1a is created at the minimum energy of the system  $E = E_1 = -10.047$  as a stable family and becomes unstable at E = -8.95 [Fig. 8(a)]. At this value of E it generates by bifurcation the period-2 family 2a [Fig. 8(c)]. Notice that when the stability parameter a of a period-1 orbit is equal to a1 the same orbit described twice has a stability parameter a3 the same are a4. Family a5 a starts at a5 with stability parameter a6. Family a8 a starts for larger values of a8. Family a9 are mains unstable until a9 and exists for larger values of a9. Family a9 are mains unstable until a9 and exists for larger values of a9. Family a9 are mains unstable until a9 and exists for larger values of a9. Family a9 are mains unstable until a9 are mains unstable until a9 are mains for smaller values of a9. Fig. a9.

Family 2a for E = -6.82 becomes unstable by crossing the line  $\alpha = -1$  and becomes stable again for E = -5.62. The stability parameter of family 2a takes the value  $\alpha=1$  at E = -5.568. At this energy an unstable period-2 family, which we call 2f, is created. Family 2e becomes unstable (going from larger to smaller values of the energy) for E =-2.987 and becomes stable again for E=-5.661. The stability parameter of family 2e takes the value  $\alpha=1$  at E =-5.675. At this energy the unstable family 2f, which was created at a larger energy from 2a, joins 2e [Fig. 8(c)]. The bifurcation diagram of families 1a, 2a, 2e, 2f, 1b is shown in Fig. 11. This figure gives the value of SR for each energy. We see that family 2f exists only in a small interval of values of E and joins family 1a at its maximum E and family 2e at its minimum E. This explains why families 2a, 2e, and 2f form a loop in Fig. 8(c).

Between E=-2.47 and E=-2.00 family 1a remains stable. At E=-2.00 family 1a joins family 1b, and for larger values of energy family 1a does not exist. We may thus say that family 1a is a bifurcation of family 1b. Family 1b is represented by the point (0,0) on the surface of section. It is created at the energy  $E_2=-8.993$  [Fig. 8(a)], at the saddle point of the potential (Fig. 2), with coordinates SR=0, Sr=1.6057. For energies somewhat larger than  $E_2$ 

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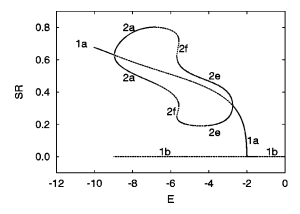


FIG. 11. Bifurcation diagram of family 1a.

family 1b is unstable ( $\alpha$ =9.73). As E increases, the stability parameter of family 1b becomes larger, but later (for larger E) it becomes smaller. At E=-2.00 this family becomes stable. It becomes unstable again at E=-1.50 with stability parameter  $\alpha$ =-1. For larger E family 1b is unstable, but at E=-0.0205 it becomes stable again until E=-0.0110, where it becomes unstable by crossing the line  $\alpha$ =1 and thus generating another period-1 family. As the energy approaches the energy E=0 the stability parameter of family 1b tends to infinity, and the return time (period) of the periodic orbit also tends to infinity.

In Fig. 12 we can see that periodic orbits of family 1b appear as straight lines in configuration space with SR=0 and varying Sr. In physical space this corresponds to a colinear configuration of the molecule with the oxygen atom in the middle and the two hydrogen atoms oscillating symmetrically at each side of the oxygen atom. For large intervals of the energy this configuration is unstable. In Fig. 12 we can also see how the periodic orbits of family 1a appear in configuration space for different values of the energy. We note that there is also a family 1a' that is symmetric to family 1a with respect to the axis SR=0. As the energy increases families 1a and 1a' approach family 1b, and join 1b for E = -2.00.

At energy E=-8.06 two irregular period-2 families are created [Fig. 8(d)]. Irregular families are created in pairs (one stable and one unstable) by a tangent bifurcation. This

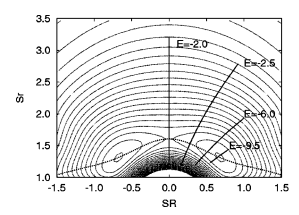


FIG. 12. Periodic orbits of families 1b (axis SR=0) and 1a. The dotted line represents the PSS.

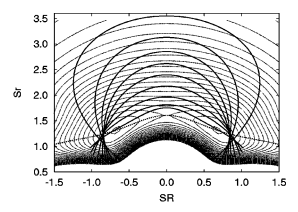


FIG. 13. Orbits of family 2b in configuration space. The dotted line represents the PSS.

means that irregular orbits are not created by the bifurcation of an existing periodic orbit, but from each other. The stable family is called 2b and the unstable one is called 2r. The stability parameter of family 2r grows from  $\alpha=1$  to  $\alpha=17.12$  as the energy increases and then decreases, until it reaches the value  $\alpha=2.77$  at E=-6.454. For this energy family 2r appears as a period-4 family and we have a transition, as in the case of the transition  $1c \rightarrow 2c$ .

The stable family 2b plays an important role in the structure of the phase space. The stability diagram of family 2b and the period-2 families that bifurcate from 2b can be seen in Fig. 8(d). The diagram is very complex, much more than the stability diagram of family 1a. Here we consider only family 2b and its equal period bifurcations. The stability parameter of family 2b crosses (or just reaches) the horizontal line  $\alpha$ =1 eight times. Each time this happens a family of period-2 orbits bifurcates from family 2b. For E=0.27 family 2b joins a new unstable family 2z and they both disappear there, i.e., they do not exist for larger E.

Families 2b, 2r, and 2z are symmetric with respect to the point (0,0). The families that bifurcate from 2b are not symmetric but they come in symmetric pairs. In Fig. 8(d) some families that have bifurcated from family 2b suddenly terminate because they are going through transitions. Some orbits of family 2b can be seen in Fig. 13. In physical space these orbits correspond to oscillations of the oxygen atom between the two symmetric minima of the potential.

The island around the orbits of family 2b is the largest island for values of energy close to E=-3. Generally, from E=-8 to E=-3 the island around 2b occupies a significant measure of the PSS.

In the present paper we studied the stability (and instability) of the various types of orbits on the plane of symmetry  $\theta = \pi/2$  and  $P_{\theta} = 0$ . However orbits that are stable on this plane may be unstable in a direction perpendicular to this plane. An example is shown in Fig. 14, where we have calculated two stability parameters for family 2a. One is the usual stability parameter  $\alpha_{\theta}$  for deviations out of the symmetry plane. We see that while family 2a is stable with respect to deviation on the symmetry plane it becomes unstable with respect to perpendicular deviations for E = -8. At the transition to instability there is a bifurcation of a new period 2

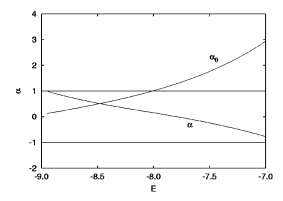


FIG. 14. Stability diagrams of the family 2a for deviations on the symmetry plane  $(\alpha)$  and perpendicular to it  $(\alpha_{\theta})$ .

family, which extends out of the symmetry plane and is initially stable.

The study of the stability of orbits in a direction out of the symmetry plane is important in many cases. Such studies in galactic dynamics have been made for a long time. <sup>17</sup> A detailed study of the stability and the bifurcations of periodic orbits with respect to the third dimension in the H<sub>2</sub>O molecule will be given in a future paper.

#### VII. SUMMARY AND CONCLUSIONS

We have made a systematic study of the classical orbits for the symmetric configuration of a realistic model of the  $H_2O$  molecule. In order to study the distribution of the ordered and the chaotic orbits for this system, we had to choose an appropriate PSS. The usual choice for the PSS is a plane surface in phase space, but for the particular system studied here, this would not be convenient because for any choice of a plane PSS there would be important periodic orbits that would not cross it.

The PSS we chose is a surface passing approximately through the minima of the potential. Most nonescaping orbits cross this PSS. Choosing appropriate coordinates on this PSS, we were able to reduce the study of the dynamics of the symmetric H<sub>2</sub>O molecule to the study of a two-dimensional map. With this particular choice of the PSS we ensure that the Poincaré map describes the complete dynamics of the molecule. A minor problem with this particular choice for the PSS is that some periodic families appear to change period without a bifurcation, due to an extra intersection of a periodic orbit with the PSS.

We found the distribution of the orbits on this PSS by distinguishing between ordered and chaotic orbits. The ordered orbits are represented by isolated points if they are periodic and by invariant curves if they are quasiperiodic. Most orbits are ordered for small energies. But as the energy increases beyond a critical value the proportion of ordered orbits decreases abruptly. When the energy increases beyond the escape energy ( $E_2 = -2.7892$ ), most orbits escape to infinity, but there are still orbits trapped around stable periodic orbits. This remains true even when the energy increases above  $E_3 = 0$ , but for positive energies the regions of ordered motion are very small, and practically insignificant.

We studied the main periodic orbits of periods 1 and 2 for negative values of the energy. We constructed stability diagrams and bifurcation diagrams for the various families of periodic orbits. These diagrams are essential in understanding the role of various stable orbits in trapping nonperiodic orbits around them. In fact for different energies the trapping takes place around different orbits and we can separate the interval between the critical energy and the escape energy into two regions. In the first region (lower energies) the ordered orbits are trapped mainly around families 2a and 2a'. In the second region, close to the escape energy the ordered orbits are trapped mainly around the irregular family 2b.

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<sup>1</sup>S. Farantos and M. Founargiotakis, Chem. Phys. 142, 345 (1990).
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<sup>&</sup>lt;sup>2</sup>S. Keshavamurthy and G. Ezra, J. Chem. Phys. **107**, 156 (1997).

<sup>&</sup>lt;sup>3</sup>M. Gutzwiller, J. Math. Phys. **12**, 343 (1971).

<sup>&</sup>lt;sup>4</sup>E. Heller, E. Steckel, and M. Davis, J. Chem. Phys. **73**, 4720 (1980).

<sup>&</sup>lt;sup>5</sup>M. Davis and E. Heller, J. Chem. Phys. **75**, 3916 (1981).

<sup>&</sup>lt;sup>6</sup>S. Farantos and J. Tennyson, J. Chem. Phys. **82**, 800 (1985).

<sup>&</sup>lt;sup>7</sup>M. Founargiotakis, S. Farantos, G. Contopoulos, and C. Polymilis, J. Chem. Phys. **91**, 1389 (1989).

<sup>&</sup>lt;sup>8</sup>M. Berry, Proc. R. Soc. London, Ser. A **423**, 219 (1989).

<sup>&</sup>lt;sup>9</sup>W. Schweizer, W. Jans, and T. Uzer, Phys. Rev. A 58, 1382 (1998).

<sup>&</sup>lt;sup>10</sup>R. Lawton and M. Child, Mol. Phys. 44, 709 (1981).

<sup>&</sup>lt;sup>11</sup>R. Lawton and M. Child, Mol. Phys. 37, 1799 (1979).

<sup>&</sup>lt;sup>12</sup>C. Jaffé and P. Brumer, J. Chem. Phys. **73**, 5646 (1980).

<sup>&</sup>lt;sup>13</sup>M. Kellman, Annu. Rev. Phys. Chem. **46**, 395 (1995).

<sup>&</sup>lt;sup>14</sup> J. Murrell and S. Carter, J. Phys. Chem. **88**, 4887 (1984).

<sup>&</sup>lt;sup>15</sup>G. Birkhoff, *Dynamical Systems* (American Mathematical Society, New York, 1927).

<sup>&</sup>lt;sup>16</sup>M. Hénon, Ann. Astrophys. 28, 992 (1965).

<sup>&</sup>lt;sup>17</sup>G. Contopoulos and P. Magnenat, Celest. Mech. **37**, 387 (1985).