MANY-DIMENSIONAL HAMILTONIAN SYSTEMS

by

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I. INTRODUCTION

It is well known that autonomous Hamiltonian systems with one degree of freedom \( H(p,q) \) are integrable. For two or more degrees of freedom, integrability is exceptional [Without loss of generality, we consider autonomous systems for which the Hamiltonian is explicitly independent of time. Nonautonomous systems in \( N \) degrees of freedom can be made autonomous in \( N+1 \) degrees of freedom by introducing an extended phase space; thus \( H(p,q,t) \) is equivalent to \( H(p_1, q_1, p_2, q_2) \); see Lichtenberg and Lieberman, 1983, Sec. 1.2b].

Practically all that is known about nonintegrable Hamiltonian systems is for the near integrable case having two degrees of freedom [Near integrable Hamiltonians have the form \( H=H_0+\epsilon H_1 \) with \( H_0 \) integrable, \( H \) not integrable, and the perturbation strength \( \epsilon \ll 1 \). Yet, such systems are special in many respects and do not exhibit the generic behavior of higher dimensional systems. In this manuscript, we compare systems having two degrees of freedom with systems having more than two degrees of freedom.

For systems having one degree of freedom, the motion is on the smooth level curves \( H(p,q) = \text{const} \) in the \( (p,q) \) phase space, as illustrated in Fig. 1a for the pendulum Hamiltonian

\[
H = \frac{p^2}{2} - \cos q. \tag{1}
\]

The elliptically-shaped orbits inside the singular separatrix curve correspond to oscillations of the pendulum, and the orbits outside the separatrix correspond to rotations. The motion can also be represented in
the action-angle variables \((I, \theta)\) of the pendulum, where

\[
I = \frac{1}{2\pi} \int \phi \, dp \, dq
\]  

(2)

is the canonical action and \(\theta\) is the canonical angle. In these coordinates, the new Hamiltonian \(K\) is a function of \(I\) alone, independent of \(\theta\), yielding the solution of Hamilton's equations \(I=\text{const}\) and \(\theta=t+\theta_0\), where \(\omega=\partial K/\partial I\) is the frequency of oscillation or rotation. The level curves of \(K\) are the straight lines shown in Fig. 1b.

For \(N\) degrees of freedom, the motion can also be described using the action-angle variables of the unperturbed system, for which the Hamiltonian takes the form

\[
K = K_0(I) + \epsilon K_1(I, \theta)
\]  

(3)

with

\[
\omega = \frac{\partial K_0}{\partial I}
\]  

(4)

the \(N\)-vector of unperturbed frequencies.

It is convenient to introduce a Poincaré surface of section \(\Sigma_R\) in the phase space (Fig. 2a). First recall that if \(z=(p,q)\) represents \(2N\) canonical variables, then Hamilton's equations with Hamiltonian \(H(z)\) cause an initial phase point \(z\) to evolve in time to a new point \(z(t)\). The transformation from \(z\) to \(z(t)\) is canonical; i.e., it preserves the Poisson bracket structure
\[
[p_i, p_j] = [q_i, q_j] = 0, \quad (5)
\]

\[
[q_i, p_j] = \delta_{ij},
\]

where

\[
[q_i, p_j] = \sum_{k=1}^{N} \left( \frac{\partial q_i}{\partial q_k} \frac{\partial p_j}{\partial p_k} - \frac{\partial q_i}{\partial p_k} \frac{\partial p_j}{\partial q_k} \right). \quad (6)
\]

One consequence of (5) is that a Hamiltonian flow preserves the phase space volume (Liouville's Theorem).

Since the motion lies on a constant energy surface \( H(z) = \text{const} \), one of the variables, say \( p_N \), can always be expressed in terms of the others:

\[
p_N = f(x, q_N) \quad (7)
\]

where

\[
x = (p_1 \ldots p_{N-1}, q_1 \ldots q_{N-1}) \quad (8)
\]

has dimension \( M = 2N - 2 \). We define a surface of section \( \Sigma_R \) by the equation

\[
q_N = g(x, p_N), \quad (9)
\]

where the function \( g \) is arbitrary, subject to the restriction that \( g \) be nowhere tangent to the unperturbed flow. As the phase point \( z \) evolves with time, it repeatedly pierces (in the same sense) the surface of section.
As shown in Fig. 2a, the successive intersections $n, n+1, n+2$ etc of the trajectory with the surface of section generate a map of dimension $M$,

$$x_{n+1} = T(x_n).$$ (10)

Since different $x_n$'s usually return to $\Sigma_R$ at different times, yielding the $x_{n+1}$'s, the transformation $T$ is not generally canonical in $x$. This implies, for example, that the map is not volume preserving (but measure preserving instead). However, it can be shown that a particular $g$ can be chosen to preserve the Poisson bracket structure (5) for $x$. In this case, the map $T$ is canonical and volume preserving. We assume in the succeeding discussion that this choice for $\Sigma_R$ has been made.

For two degrees of freedom, as shown in Fig. 2b, the phase space is four dimensional, but the motion lies on the three dimensional energy surface $H(p_1, q_1, p_2, q_2) = \text{const}$. Choosing a surface of section $q_2 = g$ yields the two dimensional surface of section $(p_1, q_1)$ on which the map is defined. For three degrees of freedom, the phase space is six dimensional, with the motion on a five dimensional energy surface. Choosing a surface of section $q_3 = g$ yields a four dimensional surface of section, Fig. 2c(2). The successive intersections of the motion with this surface of section can be projected onto the $(p_1, q_1)$ and $(p_2, q_2)$ planes, as shown in Fig. 2c(3).

The generic behavior of near integrable systems with two degrees of freedom is now reasonably well known (Lichtenberg and Lieberman, 1983, Sec. 3.2). The successive intersections of various trajectories with the
(p₁, q₁) and (I₁, θ₁) surfaces of section are illustrated in Fig. 3a and Fig. 3b respectively for a driven pendulum with Hamiltonian

\[ H = \frac{p_1^2}{2} - \cos q_1 - \epsilon \cos(q_1 - \Omega t), \]

where \( \Omega \) is the external driving frequency. First, there is a set of measure zero of singly periodic (closed) orbits that are dense in the phase space. If an orbit pierces the surface of section exactly \( k \) times, then a set of fixed points of period \( k \) is generated in the surface of section. These sets of fixed points are dense in the surface. Their stability is determined by two characteristic exponents. We will see that these exponents are either purely imaginary, corresponding to stable elliptical orbits encircling each fixed point, or they are purely real, corresponding to unstable, hyperbolic orbits. As a system parameter is varied, the stable orbits can undergo transitions such as period doubling bifurcations that can alter orbit stability, create new periodic orbits, and generate local chaotic behavior.

Almost all trajectories, however, are not singly periodic. A finite fraction of the trajectories are the quasiperiodic ("regular") trajectories of KAM theory, and the remaining fraction are nonperiodic ("stochastic"). The regular trajectories depend discontinuously on initial conditions. Stochastic and regular trajectories are intimately comingled, with some stochastic trajectory lying arbitrarily close to every point both in the four dimensional phase space and in the two dimensional surface of section. The stochastic trajectories form in the neighborhood of resonances of the motion between the two degrees of freedom,
\[ \mathbf{m} \cdot \omega = m_1 \omega_1 + m_2 \omega_2 = 0, \]  

(12)

where \( \omega \) is given by (4) and \( \mathbf{m} \) is a vector of integers. The stochastic motion appears in thin layers surrounding the separatrices associated with these resonances. The thickness of the layers increases with increasing perturbation strength. For weak perturbation, stochastic layers associated with different resonances are isolated from each other by KAM surfaces. The motion is stable, lying either in a KAM surface or within a thin stochastic layer bounded by nearby KAM surfaces. As the perturbation increases, the thickness of the layers expands, leading to resonance overlap (Chirikov, 1979; Lichtenberg and Lieberman, 1983, Sec 4.2), the destruction of the last KAM surface separating the layers. This signals the sudden appearance of strong stochasticity in the motion, in which the previously separated layers merge, and the trajectory freely moves across the layers.

The nature of the motion in systems with three or more degrees of freedom is similar to the above in most respects, but there are some major differences, probably not all discovered. For singly periodic orbits, a new type of instability appears that is described by complex characteristic exponents. Furthermore, the supposedly "universal" period doubling bifurcation sequences that appear generically in all dissipative, as well as in two degree of freedom Hamiltonian systems, are not generic to Hamiltonian systems having three or more degrees of freedom. Turning to the structure of the nonperiodic trajectories, we find the same intermixing of stochastic and regular (KAM) orbits in the 2N-dimensional
phase space. Stochastic layers form near the separatrices associated with resonances of the motion among the degrees of freedom. For strong perturbation, resonance overlap leads to motion across the layers and the presence of strong stochasticity, as for two degrees of freedom. In the limit of weak perturbation, however, resonance overlap does not occur. A new physical behavior of the motion then makes its appearance: stochastic motion along the resonance layers—the so-called weak instability or Arnold diffusion. This motion is the consequence of two fundamental properties of systems having three or more degrees of freedom:

1. Resonance layers are no longer isolated by KAM surfaces. Generically, the layers intersect, forming a connected web dense in the phase space.

2. Conservation of energy no longer prevents large chaotic motions of the actions along the layers over long times. As a result, large long-time excursions of the actions along resonance layers are generic in systems with three or more degrees of freedom. Furthermore, the interconnection of the dense set of layers ensures that the chaotic motion, stepping from layer to layer, can carry the system arbitrarily close to any region of the phase space consistent with energy conservation.

In this manuscript, we first examine the linear stability of singly periodic orbits, showing how complex instability arises. We then summarize our present understanding regarding period doubling bifurcation sequences in Hamiltonian systems. Finally, we discuss the phenomenon of Arnold diffusion.
II. LINEAR STABILITY

We consider for simplicity a period 1 fixed point $x_0 = Tx_0$. Linearizing (10) about $x_0$, we obtain the equation

$$\Delta x_{n+1} = S \Delta x_n,$$  \hspace{1cm} (13)

where, as a consequence of (5), $S$ is an $M \times M$ symplectic matrix; i.e., $S$ has the symmetry property

$$S^T \Gamma \cdot S = \Gamma,$$  \hspace{1cm} (14)

where

$$\Gamma = \begin{bmatrix} 0 & -U_1 \\ U_1 & 0 \end{bmatrix},$$  \hspace{1cm} (15)

and $U_1$ is the unit $(M/2) \times (M/2)$ matrix. Letting $\Delta x_n = \Delta x_0 \lambda^n$ in (13), we obtain the eigenvalues $\lambda$ from the equation

$$P(\lambda) = \det(S - \lambda U) = 0,$$  \hspace{1cm} (16)

where $U$ is the $M \times M$ unit matrix. Using (14) and (15), it is easy to show that the polynomial $P$ is reflexive:

$$P(\lambda) = \lambda^{2M} P(1/\lambda).$$  \hspace{1cm} (17)

Thus the eigenvalues occur in reciprocal pairs $(\lambda, 1/\lambda)$ [See Lichtenberg and Lieberman, 1983, Sec. 3.3]. The characteristic exponents $s$ are
defined by the equation \( \lambda = e^S \).

For two degrees of freedom, \( M=2 \) and

\[
P(\lambda) = \lambda^2 - A\lambda + 1 = 0 \tag{18}
\]
yields two roots. The roots must either be complex conjugates that lie on the unit circle in the complex \( \lambda \) plane, as shown in Fig. 4a, or they must lie on the real axis, as shown in Fig. 4b. The former case, obtained for \( |A| < 2 \), yields stable orbits encircling the fixed point in the surface of section; the latter case yields unstable behavior, since there is always an eigenvalue with magnitude greater than unity. As a system parameter is varied, the two roots on the unit circle may collide at either \( \lambda = 1 \) or \( \lambda = -1 \), yielding a transition to unstable behavior via either a tangent or a period doubling bifurcation.

For three degrees of freedom, \( M=4 \), and the eigenvalue equation

\[
P(\lambda) = \lambda^4 - A\lambda^3 + B\lambda^2 - A\lambda + 1 = 0 \tag{19}
\]
yields four roots. Pairs of roots may each be stable or unstable, as shown in Figs. 4a and 4b. However a new case of "complex instability" arises in which a 4-tuple of complex roots \((\lambda, \lambda^*, 1/\lambda, 1/\lambda^*)\) lies off of the unit circle(Fig. 4c). Since two of these roots have magnitude greater than unity, the motion is unstable.

The transition from stability to complex instability as a system parameter is varied occurs as follows: Initially, two pairs of complex conjugate roots lie on the unit circle. The roots collide, producing one of
the two cases shown in Fig. 4d. Either the roots "pass through" each other (the motion remains stable), or they move off of the unit circle into the complex plane, yielding complex instability. Which case occurs depends on the sign of a quadratic form,

\[ \sigma_i = \text{sgn} \Omega_i, \]  

(20)

where

\[ \Omega_i = \mathbf{x}_i \cdot \mathbf{S} \cdot \mathbf{r} \cdot \mathbf{x}_i \]  

(21)

and \( \mathbf{x}_i \) is the eigenvector having eigenvalue \( \lambda_i \) (or \( \lambda_i^* \)). The quantity \( \sigma_i \) is called the Krein signature (Moser, 1958; Howard and MacKay, 1985). For \( \sigma_1 = \sigma_2 \), the roots pass through each other; for \( \sigma_1 = -\sigma_2 \), the roots move off of the unit circle, and the system becomes unstable. Figure 5 (Howard and MacKay, 1985) shows the pattern of the eigenvalues in various regions of the \((B,A)\) parameter plane; the region of stability is shown crosshatched. Detailed studies of the motion in the region of complex instability have not been made.

III. PERIOD DOUBLING BIFURCATIONS

We consider the behavior of a singly periodic orbit in a two degree of freedom Hamiltonian flow as a parameter \( C \) is varied. We assume that the orbit pierces the surface of section once, yielding a stable period 1 fixed point \( x_1 \) for \( C > C_1 \). A generic behavior of such orbits can be as follows (Lichtenberg and Lieberman, 1983, Sec. 7.3a and Appen. B): As \( C \rightarrow C_1 \), the two eigenvalues on the unit circle collide at \( \lambda = -1 \) and move out along the
real axis (Fig. 4e). Thus $x_1$ is unstable for $C < C_1$. However, two stable fixed points having period 2 bifurcate from $x_1$ at $C = C_1$. This pair in turn destabilizes by the same mechanism at $C = C_2$, giving birth to a stable period 4 set of points, and so on, with a period $2^k$ stable orbit appearing at $C_k$. The sequence of parameters $C_1, C_2, C_3, \ldots$ converges to a finite limiting value $C_\infty$, beyond which the orbit is nonperiodic (stochastic).

A remarkable feature of this behavior is that $C_k$ converges geometrically to $C_\infty$ as

$$C_k - C_\infty \sim \delta^{-k},$$

(22)

where, for two degree of freedom Hamiltonian systems, $\delta$ is a universal constant: $\delta \approx 8.72$.

An example of this behavior (van Zeyts, 1981; Helleman, 1980) is shown in Fig. 6 for the quadratic map

$$x_{n+1} = 2Cx_n + 2x_n^2 - y_n$$

(23)

$$y_{n+1} = x_n.$$

In this numerically generated figure, successive images of the surface of section $(x_{n+1}, x_n)$ are shown magnified by the factor $A$; the first four bifurcations (period 16 stable orbit) can be clearly seen.
The same generic behavior is found for all dissipative (volume contracting) maps, no matter what their dimensionality! For these maps, (22) holds, but with a different universal constant: \( \delta \approx 4.66 \).

A final surprise is that (22) does not hold for Hamiltonian systems having three or more degrees of freedom. The period doubling bifurcation sequence appears not to exist generically. Apparently (Contopoulos, 1983), two mechanisms terminate the bifurcation sequence before it passes to completion: inverse pitchfork (period doubling) bifurcations or complex instability. We described complex instability in Sec. II. Figure 7 illustrates the two types of pitchfork bifurcations. A component of \( x \) is plotted against a parameter \( \eta \) of the map. In Fig. 7a, a stable periodic orbit (solid line) at \( x_1 \) destabilizes at \( \eta_1 \), giving birth to a period 2 stable orbit for \( \eta > \eta_1 \). This is the normal period doubling bifurcation mechanism. In Fig. 7b, the stable orbit at \( x_1 \) destabilizes at \( \eta_1 \) and gives birth to two unstable orbits that restabilize at \( x^* \). Between \( \eta_1 \) and \( \eta_1^* \), a stable period 1 and period two orbit coexist. The discontinuous jump from \( x_1 \) to \( x^* \) as \( \eta \) is varied can terminate the sequence of period doubling bifurcations.

IV. ARNOLD DIFFUSION

For autonomous Hamiltonian systems having \( N > 2 \) degrees of freedom, the resonance layers near separatrices are not isolated by KAM surfaces. This can be seen from the following table, which compares the dimension of the energy and KAM surfaces in the \( 2N \) dimensional phase space.
We recall that the energy surface (K=const) has dimension 2N-1. The KAM surfaces have dimension N, being perturbed forms of the integrable n-tori (I=const) of the unperturbed Hamiltonian K₀.

To divide a 2N-1 dimensional surface into distinct pieces, the dividing surface must have dimensionality 2N-2. The situation is analogous to that shown in Fig. 8, where one dimensional "lines" can divide a two dimensional "plane" into distinct parts, but cannot so divide a three dimensional "volume". For two degrees of freedom, we see that the two dimensional KAM tori do divide the three dimensional energy surface into distinct pieces. For three degrees of freedom, the three dimensional KAM tori cannot divide the five dimensional energy surface into distinct pieces. Thus, all (five dimensional) stochastic layers are connected together to form the "Arnold web". The web is dense in the phase space, and its measure varies wildly, but is never zero. For an initial condition within the web, the chaotic motion on the web can carry the system point arbitrarily close to any point on the energy surface. For an initial condition not on the web, the motion is regular, and the unperturbed actions I are conserved to within of order ε¹/².

An essential feature of Arnold diffusion is the existence of long-time chaotic motion along the resonance layers. If we look at a resonance layer (I₁, θ₁) in a three dimensional projection, adding an additional action
variable $I_2$, we obtain the structure shown in Fig. 9. The resonance layer has thickness of order $\epsilon^{1/2}$ and extends along $I_2$. It is easy to see that driving a large chaotic excursion along $I_2$ requires a system with at least three degrees of freedom. First, let's consider the dynamics with just two degrees of freedom. The change in the Hamiltonian is

$$\Delta K = \Delta K_0 + \epsilon \Delta K_1 = 0$$

(24)

since energy is conserved. Thus

$$\Delta K_0 = \omega_1 \Delta I_1 + \omega_2 \Delta I_2 = O(\epsilon),$$

(25)

where the $\omega$'s are given by (4). Since $I_1$ is confined to the stochastic layer, $\Delta I_1 = O(\epsilon^{1/2})$. From (25), it follows that $\Delta I_2 = O(\epsilon^{1/2})$, and large excursions of the actions are forbidden.

For three degrees of freedom, (25) is replaced by

$$\Delta K_0 = \omega_1 \Delta I_1 + \omega_2 \Delta I_2 + \omega_3 \Delta I_3 = O(\epsilon).$$

(26)

Even if $\Delta I_1 = O(\epsilon^{1/2})$, large excursions of the two actions $I_2$ and $I_3$ along the resonance are permitted by energy conservation, provided

$$\omega_2 \Delta I_2 + \omega_3 \Delta I_3 = O(\epsilon^{1/2}).$$

(27)

To understand Arnold diffusion geometrically, we consider the motion in the $N$ dimensional, unperturbed action space $I$ in which all the angle variables $\Theta$ have been projected out. For the unperturbed system, the
actions are conserved, and each trajectory is a stationary point. For example, for free particle motion in three dimensions,

\[ K_0 = I_1^2 + I_2^2 + I_3^2. \]  

(28)

The energy surface is a sphere, and the resonance surfaces (12) are flat planes intersecting the origin, as shown in Fig. 10. Another example is the two degree of freedom Hamiltonian

\[ K_0 = I_1^2 + 36I_2^2 \]  

(29)

shown in Fig. 11, where the energy surface is an ellipse, and the resonance surfaces are lines passing through the origin of the action space.

Let us consider a perturbed system

\[ K = K_0(I) + \epsilon \sum V_k(I) \exp(i m_k \cdot \theta), \]  

(30)

where the sum over \( k \) is for all sets of integer vectors \( m_k \), each having driving amplitude \( V_k \). The motion in action space is given by

\[ \dot{I} = -(\partial K/\partial \theta) = -i \epsilon \sum m_k V_k \exp(i m_k \cdot \theta), \]  

(31)

and we see that each component \( k \) drives an oscillation in \( I \) in the direction \( m_k \). For a given \( I \), there is some value \( k_R \) that drives a
resonant motion

\[ m_R \cdot \omega(I) = m_R \cdot (\partial K_0 / \partial I) \approx 0, \tag{32} \]

yielding the chaotic motion across the resonance layer shown in Fig. 9. Equation (32) implies that a resonance vector lies in an energy surface, as shown in Fig. 11. In general, the direction \( m_R \) of the resonance action excursion is not perpendicular to the resonance surface.

It can be seen from Fig. 11 that for arbitrary \( m \), the resonance surfaces do not intersect on a constant energy surface. This property is generic for systems having two degrees of freedom. For three or more degrees of freedom, the resonance surfaces generically intersect, as shown in Fig. 12a for the free particle Hamiltonian. Two resonance planes intersect at nonzero actions along a line. The resonance surfaces also intersect the spherical energy surface in great circle meridians. An energy conserving motion from one resonance to another is possible. The motion may proceed along a meridian of one resonance to an intersection, turn sharply, and move along a new meridian. This type of motion is generic to systems with three or more degrees of freedom. The intersection of resonances in the constant energy surface generates a dense, interconnected network, the Arnold web. The web for this example is illustrated in Fig. 12b, with all resonances shown for which \( |m_j| < 3 \).

A number of systems exhibiting Arnold diffusion have been studied, most of them by numerical techniques. The merging of stochastic trajectories into a single web was proven by Arnold (1964) for the specific
Hamiltonian

$$K = \frac{(I_1^2 + I_2^2)}{2} + (\epsilon \cos \theta_1 - 1)(1 + \mu \sin \theta_2 + \mu \cos t). \quad (33)$$

Another system studied analytically and numerically is the driven, two dimensional nonlinear oscillator (Chirikov et al, 1979)

$$H = \frac{(p_{1}^2 + p_{2}^2)}{2} + \frac{(q_{1}^4 + q_{2}^4)}{4} - \mu q_1 q_2 - \epsilon q_1 f(\Omega t). \quad (34)$$

Other examples more amenable to extensive numerical computations that have been studied are symplectic maps such as coupled sets of "standard" maps (Froeschlé, 1971, 1972; Froeschlé and Scheidecker, 1973), or the three dimensional billiards problem (Tennyson et al, 1979; Lieberman, 1980; Lieberman and Tennyson, 1982). We describe this latter problem as representative of systems exhibiting Arnold diffusion.

The billiards system, shown schematically in Fig. 13a, is that of a ball bouncing back and forth between a smooth wall at $z=h$ and a fixed wall at $z=0$ that is rippled in two dimensions $x$ and $y$. The surface of section is given in terms of the ball positions in the $x_n$ and $y_n$ directions and the trajectory angles

$$\alpha_n = \tan^{-1}(v_x/v_z), \quad \beta_n = \tan^{-1}(v_y/v_z),$$

evaluated just before the $n$th collision with the rippled wall. The definition of the variables in the $x$, $z$ plane is shown in Fig. 13b.
Assuming that the ripple is small and the trajectories are not at grazing angles, the rippled wall may be replaced by a flat wall at \( z=0 \) whose normal vector is a function of \( x \) and \( y \), analogous to the idea of a Fresnel mirror. The simplified map exhibits the general features of the exact equations and may be written in explicit form

\[
\begin{align*}
\alpha_{n+1} &= \alpha_n - 2a_x k_x \sin k_x x_n + \mu k_x \gamma_c, \\
x_{n+1} &= x_n + 2h \tan \alpha_{n+1}, \\
\beta_{n+1} &= \beta_n - 2a_y k_y \sin k_y y_n + \mu k_y \gamma_c, \\
y_{n+1} &= y_n + 2h \tan \beta_{n+1},
\end{align*}
\]

where

\[\gamma_c = \sin (k_x x_n + k_y y_n),\]

\( a_x \) and \( a_y \) are the amplitudes of the ripple in the \( x \) and \( y \) directions, respectively, and \( \mu \) is the amplitude of the diagonal ripple and represents the coupling between the \( x \) and \( y \) motions.

If \( \mu=0 \), the system breaks into two uncoupled parts describing motion in \( x-z \) and \( y-z \) separately. Figure 14 shows the motion in the \( \alpha-x \) surface of section for the uncoupled case. Several different orbits are shown, each run for 1000 iterations. We see the usual features of a system having two degrees of freedom: (a) regular (KAM) orbits, (b) resonance island orbits, and (c) stochastic orbits. The island orbits are examples of "higher order" KAM curves. The central resonance at \( \alpha=0, x=0 \) corresponds to a stable motion for which the ball bounces up and down along \( z \) in the valley of the rippled wall. The island orbits correspond to "adiabatic"
motion in the valley with a small oscillation back and forth in x occurring over many bounce times in z. There are two major stochastic regions visible. The thick layers for $\alpha$ near $\pm \pi/2$ are regions produced by all overlapping resonances having one bounce period in $z$ equal to one or more periods along $x$, corresponding to grazing angle trajectories. A thin layer has also formed near the separatrix of the central resonance, corresponding to motion in $x$ for which the ball is either just reflected or just transmitted over a hill. These regions of stochasticity are separated by KAM curves.

A numerical calculation of Arnold diffusion in the coupled system is given in Fig. 15. The surface of section $(\alpha, x, \beta, y)$ is represented in the form of two plots $(\alpha, x)$ and $(\beta, y)$. Thus two points, one in $(\alpha, x)$ and one in $(\beta, y)$, are required to specify a point in the four dimensional section. In the figure, the two plots have been superimposed for convenience, and $x$ and $y$ have been normalized to their respective wavelengths $2\pi/k_x$ and $2\pi/k_y$, respectively. The initial condition has been chosen on an island orbit in $x$ and within the thin separatrix layer in $y$. This corresponds to an initial regular motion in $x$, well confined in the valley, while the stochastic $y$ motion just reaches or passes over a hill. The successive stages of the Arnold diffusion of the $x$ motion are shown in Figs. 15b-d for $1.5 \times 10^5$, $3.5 \times 10^6$ and $10^7$ iterations of the map, respectively. In the absence of coupling ($\mu=0$), the $x$ motion would be confined to a smooth closed curve. For finite coupling, $x$ diffuses slowly due to the randomizing influence of the stochastic $y$ motion.

An analytical calculation of the Arnold diffusion rate was first
performed by Chirikov (1971,1979) and his collaborators. For the billiards problem, the diffusion rate has been calculated by Tennyson et al (1979) [see also Lieberman (1980) and Lichtenberg and Lieberman (1983, Sec. 6.2)]. The basic procedure is to break the original three degree of freedom system into two systems that each have two degrees of freedom, which are successively solved. The simplest decomposition considers only three resonances. The first (guiding) resonance can be chosen arbitrarily and defines the local region within the Arnold web where the diffusion is to be determined. The strongest remaining resonance drives the chaotic motion across the separatrix layer of the guiding resonance. The remaining strongest resonance drives the Arnold diffusion.

A form of diffusion related to Arnold diffusion is modulational diffusion, in which chaotic motion is driven along a band of overlapping resonances by a slow modulation of the driving perturbation. Both Arnold and modulational diffusion only occur in (autonomous) systems having three or more degrees of freedom. Whereas Arnold diffusion is universal and generally weak, modulational diffusion requires that an overlap condition be satisfied for a band of primary resonances, but the diffusion can be strong. An example of a system exhibiting modulational diffusion is the Hamiltonian

\[ K = \frac{(I_1^2 + I_2^2)}{2} - \epsilon \cos(\theta_1 + \lambda \sin \Omega t) - \mu \cos (\theta_1 - \theta_2). \]  

(36)

Here, the \( \theta_1 \) and \( \theta_2 \) motions might represent vertical and radial betatron oscillations that are coupled by the parameter \( \mu \), and the \( \lambda \sin \Omega t \) term might represent the coupling of a slow synchrotron oscillation to the \( \theta_1 \).

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V. REFERENCES


VI. FIGURE CAPTIONS

Figure 1. Phase space for one degree of freedom; (a) in \((p,q)\) coordinates; (b) in action angle coordinates \((I,\theta)\).

Figure 2. Motion in phase space and definition of the Poincaré surface of section. (a) Intersections of a trajectory with the surface of section. (b) Two degrees of freedom showing: (1) four dimensional phase space with the trajectory on a three dimensional energy surface; (2) projection of the trajectory onto the \((p_1,q_1,q_2)\) volume; and (3) successive intersections of the trajectory with the two dimensional surface of section \(q_2 = g\). (c) Three degrees of freedom showing: (1) six dimensional phase space with a trajectory on a five dimensional energy surface; (2) three successive intersections of the trajectory with the four dimensional surface of section \(q_3 = g\); and (3) projections of these intersections of the surface of section onto the \((p_1,q_1)\) and \((p_2,q_2)\) planes.

Figure 3. Near integrable motion in (a) the \((p_1,q_1)\) and (b) the \((I_1,\theta_1)\) surface of section for two degrees of freedom.

Figure 4. Eigenvalues \(\lambda_i\) of a fixed point in the complex \(\lambda\) plane. The eigenvalues must occur in reciprocal pairs \((\lambda,1/\lambda)\). (a) stability; (b) instability; (c) complex instability for three or more degrees of freedom; (d) a Krein collision; and (e) a period doubling bifurcation.

Figure 5. Pattern of eigenvalues in the \((B,A)\) parameter plane for three degrees of freedom. The dots are the eigenvalues \(\lambda_i\), and the circle is a unit circle in the complex \(\lambda\) plane. The stable region is hatched.
Figure 6. Surface of section plots of the period doubling sequence for the map (23). Here \( x_{n+1} \) is plotted versus \( x_n \) for various values of the parameter \( C \). The parameter \( A \) indicates the magnification of the plot.

Figure 7. Two types of bifurcations in \( x \) as a parameter \( \eta \) is varied: (a) period doubling and (b) inverse pitchfork. The solid(dotted) lines are the stable(unstable) branches.

Figure 8. Dividing a surface into pieces. In (a) the two dimensional plane is divided by lines into a set of distinct areas; in (b) the three dimensional volume is not divided by lines into distinct volumes.

Figure 9. Illustrating the directions of the fast diffusion across a resonance layer and the slow diffusion along the layer.

Figure 10. The three dimensional action space showing energy surfaces (spheres) and resonance surfaces (planes) for the unperturbed free particle Hamiltonian.

Figure 11. Resonance curves (lines) and energy contours (ellipses) in two dimensional action space for the Hamiltonian (29). The resonance labels are the values of \( m_1 \), where \( \omega_1 m_1+\omega_2=0 \). The direction of a resonance vector \( m_R \) is shown.

Figure 12. Illustrating Arnold Diffusion. (a) Intersection of two resonance surfaces in an action space having three degrees of freedom. An energy conserving motion (wiggly line) from one resonance surface to another is possible. (b) The Arnold web for the free particle Hamiltonian; only some of the intersecting resonances are shown.
Figure 13. The three dimensional billiards system. (a) A point particle bounces back and forth between a smooth and a periodically rippled wall. (b) Motion in two degrees of freedom, illustrating the definition of the trajectory angle $\alpha_n$ and the bounce position $x_n$ just before the $n$th collision with the rippled wall.

Figure 14. Motion in the $(\alpha, x)$ surface of section for the uncoupled billiards system. The parameters are $\mu=0; \lambda_x:\hbar; a_x$ as 100:10:2; $\lambda_x=2\pi/k_x$. Fifteen initial conditions at $x=0$ are each iterated for 1000 collisions with the rippled wall.

Figure 15. A numerical computation showing Arnold diffusion. The initial condition is close to the central resonance in the $(\alpha, x)$ plane and within the separatrix stochastic layer in the $(\beta, y)$ plane. The parameters are $\mu/\hbar=0.004; \lambda_x:\hbar; a_x$ and $\lambda_y:\hbar; a_y$ as 100:10:2.
Stochastic Layers / Isolating KAM Surfaces

Resonance Fixed Points

(a)

(b)

I_1

\theta_1
Unit Circle

(a) $|A| < 2$

(b) Tangent Bifurcation $A > 2$

(c) Period-doubling Bifurcation $A < -2$

Complex Instability

Stable

Collision

Unstable
$A = 1$
$C = -0.9$

$2^0$

$A = 1$
$C = -1.02$

$2^1$

$A = 1$
$C = -1.05$

$3^1$

$A = 5$
$C = -1.10$

$3^2$

$A = 5$
$C = -1.24$

$3^3$

$A = 5$
$C = -1.253$

$3^4$

$A = 25$
$C = -1.253$

$3^5$

$A = 100$
$C = -1.26$
(b)

\[ V = 100 \]

\[ C = -1.2632 \]

(c)

\[ V = 100 \]

\[ C = -1.2634 \]

\[ V = 100 \]

\[ C = -1.2636 \]

\[ V = 200 \]

\[ C = -1.2636 \]

\[ V = 200 \]

\[ C = -1.2636 \]

\[ V = 200 \]

\[ C = -1.264 \]

\[ V = 200 \]

\[ C = -1.265 \]

\[ V = 200 \]

\[ C = -1.266 \]

\[ V = 800 \]

\[ C = -1.266 \]

\[ V = 800 \]

\[ C = -1.266 \]
Resonance surfaces

Energy surface

Energy-conserving motion along resonances

(a)

(b)
Surface of section

(a) 

(b) 

\[ \alpha, \beta \]

\[ \theta = k_x x \]

\[ \phi = k_y y \]

\[ N = 1.5 \times 10^5 \]

(c) 

\[ \alpha, \beta \]

\[ \theta = k_x x, \phi = k_y y \]

\[ N = 3.5 \times 10^6 \]

(d) 

\[ \alpha, \beta \]

\[ \theta = k_x x, \phi = k_y y \]

\[ N = 10^7 \]