We present a comprehensive phase-space treatment of the motion of charged particles in electrodynamic traps. Focusing on five-wire surface-electrode Paul traps, we study the details of integrable and chaotic motion of a single ion. We introduce appropriate phase-space measures and give a universal characterization of the trap effectiveness as a function of the parameters. We rigorously derive the commonly used (time-independent) pseudopotential approximation, quantify its regime of validity, and analyze the mechanism of its breakdown within the time-dependent potential. The phase-space approach that we develop gives a general framework for describing ion dynamics in a broad variety of surface Paul traps. To probe this framework experimentally, we propose and analyze, using numerical simulations, an experiment that can be realized with an existing four-wire trap. We predict a robust experimental signature of the existence of trapping pockets within a mixed regular and chaotic phase-space structure. Intricately rich escape dynamics suggest that surface traps give access to exploring microscopic Hamiltonian transport phenomena in phase space.

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I. INTRODUCTION AND MAIN RESULTS

Surface-electrode Paul traps are becoming increasingly widespread, in large part due to their potential for scalable microfabrication [1,2]. Introduced only recently [3,4], they differ from traditional linear and hyperbolic Paul traps by the strong nonlinearity of the potential, leading to complex and partly chaotic motion even for a single ion. Furthermore, the small scale of surface traps implies a limited trapping volume, and the importance of perturbations, intrinsic or extrinsic to the trap, grows significantly. Therefore, on the practical side, a comprehensive understanding of the different dynamical regimes in these traps is important for optimizing ion loading, cooling, storage, and manipulation. From a more fundamental perspective, surface traps offer a unique setup for studying the interplay of nonlinearity, chaos, and microscopic stochastic forces.

Nonlinear and stochastic ion dynamics have attracted fundamental and practical interest since the first crystallization of charged particles in electrodynamic fields [5]. The first systematic studies of chaos in Paul traps followed the demonstrations of a cloud-crystal phase transition with small clusters [6,7] and clouds [8]. Since Paul traps are based on (periodic) time-dependent electric fields, the system does not conserve energy and so particles can eventually heat up and escape the trap. Deterministic chaos was found to be a predominant source of heating that balances the laser cooling and stabilizes the cloud phase. The smallest system with appreciable nonlinearity (and hence, possibly, chaotic motion) in the earlier Paul traps was that of two ions, and it was studied in detail [9–15], including an analysis of all integrable cases [18–21]. Because of their importance in high-accuracy quantum applications, the effects of the periodic driving (known as the “micromotion”) are extensively studied in various regimes [22–30]. The nonlinear (and time-dependent) regimes of motion in a surface trap remain largely unexplored, however.

In this paper, we develop a phase-space framework for studying classical dynamics of charged particles in surface Paul traps. We introduce appropriate phase-space measures that allow us to quantify and compare the trap’s effectiveness at different parameter values. Our quantitative results apply directly to any five-wire trap of any size (for the model that we consider). The general conclusions of the study apply to a broad range of surface traps and present a systematic approach to the investigation of the nonlinear and time-dependent motion.

We consider an ion trapped in vacuum above a set of planar electrodes in the five-wire configuration, carrying a combination of direct current (DC) and radio-frequency (rf) periodically modulated voltages (see Sec. II and Fig. 1). The axial $x$ motion (parallel to the electrodes’ symmetry axis) is that of a decoupled, time-independent harmonic oscillator with frequency $\omega_x$, the axial harmonic trapping frequency. The $yz$ motion is coupled, nonlinear, and driven periodically in time, and hence the phase space $\{y, z, p_y, p_z\}$ whose dimension is 4 (with $p_y$ and $p_z$ the canonical momenta) has to be in practice increased to include the time $t$, making it effectively five-dimensional (5D).

A 5D phase space is a big place [31]. Moreover, the number of independent parameters of the full model can become quite high [32]. However, we show that an essential understanding of the dynamics can be obtained by a systematic study starting from a two-dimensional (2D) time-independent reduction, and continuing through more elaborate three-dimensional (3D), which is the effective dimension of a 2D phase space with

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a time-dependent Hamiltonian, and four-dimensional (4D) phase-space analysis. This procedure, which we develop in this work, requires also that we limit the parameters at each step. Hence, as a starting point, we assume a trap configuration with equal y and z harmonic DC antitrapping coming from the axial electrode and no further harmonic DC terms. In this case, an ion starting with the initial conditions \( y = 0 \) (in the coordinate parallel to the electrodes’ plane), remains at \( y = 0 \), and the \( z \) direction (normal to the electrode surface) decouples exactly. We study this motion in detail in Sec. III, also assuming at first that there is no further “bias” DC voltage on the rf electrodes. Then the dynamics depend on two parameters—the rf voltage \( U_{rf} \) and the axial DC trapping voltage \( U_{dc} \times \propto \omega_z^2 \). Studying the rf-driven \( z \) motion begins with the commonly used approximation that replaces the time-dependent potential with an effective, time-independent pseudopotential. The latter depends in this setup on a single nondimensional parameter that we name the “pseudopotential parameter,”

\[
\lambda = (m \omega_z w^2 / \sqrt{2e U_{rf}})^2 \propto U_{dc}/U_{rf}^2, \tag{1}
\]

where \( m \) and \( e \) are the ion’s mass and charge, \( \Omega \) is the rf frequency, and \( w \) is the electrodes’ width and separation (see Sec. II for details). Since the pseudopotential is time independent, the phase space is 2D, and the motion is integrable without any chaos. Within the pseudopotential approximation, a maximization of the trap depth is misleadingly simple—it is a monotonously decreasing function of \( \lambda \), so it decreases with \( \omega_z \) and increases with \( U_{rf} \).

This simple dependence is completely altered within the time-dependent potential. The latter can be thought of as composed of the pseudopotential (characterized by \( \lambda \)) and a time-dependent perturbation scaled by \( U_{rf} \). Since the equations of motion depend explicitly on time, the phase space is now effectively 3D. With \( U_{rf} \) below a threshold value, the entire 3D phase space is very close to regular and the pseudopotential approximation is admissible for any \( \lambda \). Increasing \( U_{rf} \) gradually makes the phase space “mixed”—chaotic motion develops inside bounded strips within the regular phase space, and most notably within a large connected “chaotic sea” that reaches from the top of the trap and penetrates the phase space toward the trap center. Once inside the chaotic sea, the ion will quickly escape past the trap’s barrier, and therefore increasing \( U_{rf} \) beyond an optimal value leads to a degradation of the trap’s effectiveness. We find that the maximal trapping can be obtained by taking \( U_{rf} \) at the threshold of validity of the pseudopotential approximation, together with \( \lambda \to 0 \) (requiring the reduction of the axial trapping \( \omega_z \)). If \( \omega_z \) is constrained to a certain value or range of values, optimal trapping can be obtained by tuning \( U_{rf} \) to a \( \lambda \)-dependent value.

With this characterization of the 3D phase space of \( z \) motion, we turn in Sec. IV to the coupled \( yz \) motion, starting with the 4D phase space of the pseudopotential approximation. Remarkably, we find that the motion is very close to being completely integrable, i.e., with almost no chaotic signatures. This is quite surprising since the pseudopotential is highly nonlinear and \( a \ priori \) there is no reason to expect its integrability. We continue by first studying in detail the effect of a bias DC voltage \( (U_b) \) applied to the rf electrodes, still within the pseudopotential’s 4D phase space. The bias voltage leads first to an increase of the trap depth, but beyond a threshold magnitude causes the gradual destruction of the regular phase space. As a function of the energy, the phase space may split into a few (nearly regular) trapping “islands” separated by chaotic regions.

Reintroducing the time dependence of the trap, we study how the integrability-breaking perturbation that scales with \( U_{rf} \) leads the ion to escape from the trap, due to the combined effect of the bias and time dependence. In the 5D phase space, the regions of regular and chaotic motion can be intertwined in a complicated way. This introduces new possibilities for transport mechanisms in phase space, as we briefly discuss below. At the same time, it presents a challenge for visualizing and exhaustively analyzing the motion. For a practical, first step study of the ion’s escape from the trap, we present a simple measure of phase-space volume that enables quantification of these competing processes and their dependence on the three parameters \( \lambda, U_{rf}, \) and \( U_b \).

We find (in Sec. IV B) that the border of validity of the pseudopotential (for approximately regular motion) found for the \( z \) motion holds also for the coupled \( yz \) motion, restricted to intermediate values of \( \lambda \) and \( U_b \). Above this threshold, all chaotic trajectories escape, and the regular region gradually shrinks as well. We also find an optimal regime of the parameters maximizing the (mostly regular) trapping phase space volume of \( yz \) motion in the model five-wire trap. We find that this regime is obtained for \( U_{rf} \) just below the threshold of validity of the pseudopotential, with \( \lambda \to 0 \) (requiring to reduce the axial trapping \( \omega_z \)), and an optimal intermediate value of \( U_b \). These parameters are distinctly different from a naïve pseudopotential approach that would lead to increasing \( U_b \) and \( U_{rf} \) as much as possible.

The near integrability of the pseudopotential for the symmetric, unbiased five-wire configuration makes it a natural starting point to analyze dynamics in surface traps, since the
motion remains close to integrable for any perturbation that is not too large. This includes the full rf-driven motion in 5D phase space, and indeed we confirm that this perturbation can be considered small within the 5D phase space, in the same regime where it is small in the 3D phase space of z motion. A systematic treatment based on the integrable pseudopotential limit of the symmetric five-wire trap gives a framework for studying further coupling or asymmetry, or the application to different geometries. The consideration of circular traps [33,34], and even of electrons guided on a chip [35], could be another interesting direction, immediately applicable (the latter is essentially a surface trap, operating at much higher frequencies since the electrons are much lighter).

Motivated by practical questions on the motion of trapped ions, this work is focused mostly on the ion’s escape from the trap. However, from a broader point of view, one can study how transport between different parts of the phase space proceeds within the Hamiltonian motion. As described in the following, for a phase space which is effectively 3D or 4D, the motion is amenable to classification using 2D Poincaré sections, and the transport properties can be scrutinized. With more degrees of freedom, there are richer options for transport and there is a fundamental interest in unveiling the involved mechanisms. The ion trap offers a system with excellent prospects for experimenting with this physics, at the level of elementary particles. The trap parameters are highly controllable, the dimensionality of the effective phase space can be altered, and as we find in the following, further control parameters can be devised. There are some limitations with respect to directly measuring microscopic properties of the motion that nonetheless can be circumvented.

In Sec. V, we propose an experiment probing the nonlinear and chaotic motion of an ion in a surface trap. Our analysis is based on a four-wire trap described in Ref. [36], with an electrode configuration that is more complicated than the model of the five-wire trap discussed above. We consider adding a controlled “tickle” perturbation (a small-amplitude voltage modulation) on one of the electrodes, at a frequency above resonance with the secular motion. It gives a robust experimental procedure resulting in a clear signal with a nonmonotonous and sensitive dependence on the parameters.

Running numerical simulations of this trap configuration with real experiment parameters, we find clear evidence (within the limitations of an initial study) of trapping “pockets” induced in the trap’s phase space. We predict a variety of distinct functional forms for the ion’s escape probability from these pockets, which present a challenge for further theory and experiments on Hamiltonian transport in a mixed phase space [37–39]. We also find quantitatively very similar escape probabilities with the pseudopotential approximation for this trap, indicating that it is being operated not far from its integrability limit.

The theory presented in this work gives a general framework that can be applied to various surface-electrode traps. With an understanding of the mechanism that determines the nature of the dynamics, the scope of the conclusions that we draw (Sec. VI) extends beyond the particular models here studied. In the appendixes, we collect a set of phase-space tools that allow one to treat arbitrary trap potentials which are nonlinear and periodically driven. In particular, we rigorously derive the pseudopotential approximation for a generic periodic potential, as a canonical transformation from the original coordinates, that results in a time-independent pseudo-Hamiltonian (which may a priori depend on the canonical momenta [40,41]). The derivation of the pseudopotential from a canonical transformation allows us to keep the Hamiltonian phase-space structure, and in particular the structure of the invariant tori of the motion (in the near-integrable regime). As briefly discussed in the outlook of Sec. VI, this Hamiltonian structure turns out to be invaluable in a treatment of stochastic motion in phase space, using a Fokker-Planck approach [42].

II. THE FIVE-WIRE SURFACE TRAP

In this section, we present the details of the trap model and set the notation for the rest of this work. The total trap potential is the sum of two parts: the time-dependent potential produced by two rf surface-electrodes and a time-independent harmonic potential contribution (quadratic in the coordinates). As shown in Fig. 1, the surface electrodes lie in the z = 0 plane and are assumed to have an infinite extent along the x axis. The y axis is perpendicular to the infinite direction of the electrodes in the electrode plane, and the z axis is orthogonal to the electrode plane. The electric potential from the surface electrodes is given by [32,43–45]

$$V_{5w}(y, z) = [V_w(y, z; w) - V_w(y, z; -w)],$$  (2)

with the potential due to one planar electrode of width w centered at +w and held at unit voltage being

$$V_w(y, z; w) = \frac{1}{\pi} \left[ \arctan \left( \frac{y - w/2}{z} \right) - \arctan \left( \frac{y - 3w/2}{z} \right) \right].$$  (3)

The field due to the potential of Eq. (2) vanishes at a saddle point $\vec{R}_0 = \{0, 0, z_0\}$ at a height $z_0 = \sqrt{3}w/2$ above the electrodes. Since this potential does not trap along the x axis, additional (DC) electrodes must be included. Here we approximate the potential confining the ion along x by an ideal static quadrupole (harmonic) potential along x that leads to antitrapping in the radial directions. The total dimensional potential energy of an ion in the trap is

$$\hat{V}_{\text{trap}} = e \sum_{a} \frac{U_{\text{DC}}(R_a - R_{0,a})^2}{2c_a} + e(U_b - U_{\text{rf}} \cos \Omega t)V_{5w}(y, z),$$  (4)

where $a \in \{x, y, z\}$, $\vec{R}$ is the vector coordinate of the ion, $\Omega$ is the rf frequency, and $e$ is the charge of the ion. $U_{\text{rf}}$ and $U_b$ are the rf and the DC “bias” voltages (respectively) applied to the electrodes, and $U_{\text{DC}}$ characterizes the strength of the static harmonic potential. The geometric properties of the latter are defined by $c_a$ (which are of squared-length dimension), and by the origin of the static quadrupole that we choose to coincide with the saddle point of $V_{5w}$, that is $R_{0,a} = \{0, 0, z_0\}$.

Measuring distances in units of the width w and rescaling time by half the rf frequency, $\Omega/2$,

$$\vec{R}_a \rightarrow \vec{R}_a/w \quad t \rightarrow \Omega t/2,$$  (5)
we find the nondimensional potential

$$V_{\text{trap}} = V_h + (a_5 - 2q_5 \cos 2t)V_{5,0}(y,z),$$

(6)

where

$$V_h = \frac{1}{2}[a_x x^2 + a_y y^2 + a_z(z - z_0)^2],$$

(7)

and the nondimensional parameters are

$$a_x = \frac{4 e U_{\text{DC}}}{m\Omega^2 c_a}, \quad q_x = \frac{2 e U_{\text{rf}}}{m w^2 \Omega^2}, \quad a_z = \frac{4 e U_b}{m w^2 \Omega^2}.$$  

(8)

In this setup, the motion in the $x$ direction is that of a simple harmonic oscillator (with frequency $\omega_x$), decoupled from the motion in the radial $yz$ plane. As a consequence of the Laplace motion, the harmonic axial DC voltage gives rise to radial antitrapping, which we take to be symmetric:

$$a_z = a_y = -\frac{1}{2} a_x, \quad a_x > 0.$$  

(9)

The linearized secular frequencies of the ion near the center of the trap are given by [46]

$$\omega_a = \frac{\Omega}{2},$$

(10)

where $v_a(a_x, q_a)$ is the nondimensional characteristic exponent of the corresponding Mathieu equation, with parameters $a_x, q_a$. The latter ($q_a$) can be obtained by a linearization of the trap potential $V_{\text{trap}}$ of Eq. (6) around $[y = 0, z = z_0]$, giving, for $a_5 = 0$,

$$q_x = 0, \quad q_c = -q_y = \frac{2}{c_3 \pi q_s}.$$  

(11)

In the limit $a_x, q_a^2 \ll 1$, we can approximate

$$v_a \approx \sqrt{a_x + q_a^2/2},$$

(12)

which results in

$$\omega_x = \frac{c_3 \pi q_s}{2}, \quad \omega_c \approx -\frac{a_x + q_a^2}{2} \frac{\Omega}{2}.$$  

(13)

For the figures and numerical analysis in this paper, we consider both nondimensional parameters and physical parameters that give a specific example of real-world values. All the dimensional values are given for $^9$Be$^+$ ions in a trap with electrode width and rf frequency [that appear in Eq. (5)], given by

$$w = 50 \mu m, \quad \Omega = 2\pi \times 100 \text{ MHz}.$$  

(14)

### III. HAMILTONIAN MOTION IN ONE SPATIAL DIMENSION

In this section, we consider the unbiased configuration $U_b = 0$ (i.e., $a_5 = 0$) and characterize the motion of the ion in the rf potential, and the corresponding pseudopotential, of the $z$ spatial coordinate orthogonal to the electrode plane. In Sec. IIIA, we introduce the notation, and in Sec. IIIB, we present the possible types of phase space structures and discuss the qualitative implications. Then in Sec. IIIC, we define quantitative measures that enable characterization of the trap’s dependence on its parameters. The effect of the bias voltage $U_b$ will be studied in Sec. IV, for the coupled $yz$ motion.

### A. Reduction of the equation of motion

Setting $y = \dot{y} = 0$ in the equations of motion derived from Eq. (6) results in $y(t) = 0$ for all times, and thus the $z$ motion can be studied independently. Since $a_5 = 0$, the potential depends on two parameters, $a_x \propto \omega_x$ and $q_5$, that are proportional to the static quadrupole and the rf voltages respectively [Eq. (8)]. With $y = 0$ in Eq. (6), the rf potential can be simplified to

$$V_{\text{rf}} = a_z(z - z_0)^2 - \frac{4}{2\pi} q_5 \cos 2t \left[ \arctan \left( -\frac{1}{2} \right) - \arctan \left( -\frac{3/2}{z} \right) \right].$$

(15)

In the rest of the paper, we refer to this potential as the 1D “rf potential.” Since the rf frequency sets the fastest frequency scale, the method of averaging is a natural approach, allowing one to simplify the description of the motion and reduce the equations of motion to an autonomous system. Using this method, it is possible to derive an effective time-independent pseudopotential approximation to the rf potential that is well known [47,48]. In Appendix A, we present a derivation of the pseudopotential as a time-dependent canonical transformation, from the original phase-space coordinates of motion within the rf potential, to new coordinates. This transformation eliminates the $\pi$-periodic component of the position and momentum, while preserving the Hamiltonian structure (with a new time-independent Hamiltonian). Thus, although the new Hamiltonian is time independent, the new variables are actually time-dependent functions of the original ones (and vice versa). In the following subsections, we will consider in detail the regime where the pseudopotential gives a simpler and accurate approximation and its limits of applicability.

The 1D pseudopotential approximation to Eq. (15), derived in Appendix A, is

$$V_{\text{pseudo}}^{1D} = a_z(z - z_0)^2 - \frac{4}{2\pi} q_5 \cos 2t \left[ \frac{16(3 - 4z^2)}{2} + \frac{16(3 - 4z^2)}{2} \right].$$

(16)

By a second rescaling of time using

$$t \rightarrow q_5 t,$$

(17)

we get the rescaled pseudopotential,

$$V_{\lambda}^{1D} = V_{\text{pseudo}}^{1D} = -a_z(z - z_0)^2 - \frac{16(3 - 4z^2)}{2} + \frac{16(3 - 4z^2)}{2} \pi^2(9 + 40z^2 + 16z^4).$$

(18)

governed by a single pseudopotential parameter defined equivalently to Eq. (1) by

$$\lambda = \frac{a_5}{q_5^2} = \frac{a_x}{2q_5^2} > 0,$$

(19)

where we have used the fact that in the symmetric case that we consider, $a_x = -a_z/2 < 0$, as in Eq. (9).

The pseudopotential has two fixed points, defined as the solutions of

$$dV_{\text{pseudo}}^{1D}/dz = 0.$$

(20)
A close-up of the region around the last unbroken torus of panel (c). A chaotic "sea" emerges from the region near the unstable fixed point.

The first point, \( z_s \), is the stable fixed point at the center of the trap (that is, the pseudopotential minimum). The second point, \( z_u \), is an unstable fixed point, a local maximum of the pseudopotential, beyond which the ion escapes the trap. As discussed above, we assume a configuration such that the \( z \) origin of the harmonic term in Eq. (18) coincides with the point where the force from \( V_{sw} \) is 0. In that case, the stable fixed point is independent of the parameters \( a_z \) and \( q_z \). Within the pseudopotential as well as the rf potential, \( z_0 = z_s \) for all \( \lambda \). For \( \lambda = 0 \), \( z_u \) is maximal, while for \( \lambda > 0 \), \( z_u \) moves toward \( z_s \). The numerical values of the fixed points are given by

\[
\begin{align*}
  z_s &= \frac{\sqrt{3}}{2} \approx 0.866, \\
  z_u |_{\lambda=0} &= \sqrt{3/4 + \sqrt{3}} \approx 1.575. 
\end{align*}
\]

The linearization near \( z_s \) of \( V_{\text{pseudo}}^{1D} \) of Eq. (16) gives the harmonic oscillator potential

\[
V_{\text{pseudo}}^{1D} \rightarrow V_{\text{h.o.}}^{1D} = \frac{1}{2} \nu_z^2 z^2, \tag{22}
\]

and the linearization of \( V_{rf}^{1D} \) of Eq. (15) gives the Mathieu oscillator potential

\[
V_{rf}^{1D} \rightarrow V_{\text{M.o.}}^{1D} = \frac{1}{2} (a_z - 2 q_z \cos 2 t) z^2. \tag{23}
\]

The rf potential is, in contrast, time dependent and \( \pi \) periodic, so the motion can be thought of as occurring within a 3D phase space defined by \( \{ z, p_z, t \} \), with the \( t \) dimension periodic. The ion trajectories can be studied by constructing a 2D cut of phase space taken at fixed value of \( t \) (mod \( \pi \)); see Fig. 2(b). The continuous Hamiltonian motion as a function of time is then "stroboscopically" projected to a discrete map of phase space onto itself at \( \pi \) time intervals. The ion does not follow continuously as a function of time any curve in this planar surface of section, but rather revisits this plane every \( \pi \) period of the external drive.

In contrast to the pseudopotential, within the time-dependent potential two different types of motion can arise—regular and chaotic. For bounded regular motion, the ion’s return points to the 2D surface of section lie on a closed, 1D curve, or a finite set of such curves in the case of a chain of resonance “islands.” In the latter case, in each period the ion revisits a different island from the chain in a fixed order. For chaotic motion, the ion’s return points will in the limit of infinite time fill a region with nonzero area. The structure of unbroken tori (invariant curves), chaotic strips, and resonance island chains is fractal [repeating itself in smaller and smaller scales—see Figs. 2(c) and 2(d)]. Each closed curve and every
connected region of chaotic trajectories, are invariant under the
time evolution. Therefore, a stroboscopic map taken at any
value of $t \pmod{\pi}$ will contain the corresponding structures
with the same enclosed area, distorted in time with periodicity
$\pi$; see Fig. 3. This important property of the time evolution
will be used below.

Figures 2(a) and 2(b) show the similarity between the pseudopotential
phase space and that of the rf potential for small $q_5$. This can be understood by decomposing the rf potential
as a sum of its time-independent-averaged pseudopotential
(whose phase-space structure depends on the single parameter
$\lambda$) and a time-dependent perturbation, whose amplitude is
controlled by $q_5$. Therefore, we expect the rf potential to be
well approximated by the pseudopotential in the limit of small
$q_5$ for any $\lambda$. The pseudopotential is integrable, and the measure
of chaotic trajectories in the phase space increases with $q_5$.
When $q_5$ is small, the phase space is composed of mostly
regular motion and as it is increased, an increasing area of
phase space becomes chaotic. Figures 2(c) and 2(d) show the
stroboscopic map with a moderate value of both $\lambda$ and $q_5$, for
which there is significant deviation from integrability with the
aforementioned mixed structure of regular tori, chaotic strips,
resonance islands, and a large chaotic sea leading through the
unstable fixed point to escape from the trap. Figure 3 shows the
phase space with only a small central regular island remaining,
for a larger $q_5$. In this figure also the micromotion dressing of
the pseudopotential motion is visualized.

To gain more understanding of the motion within the rf
potential, we can use a correspondence between the phase
space of the time-dependent rf potential and a higher dimen-
sional phase space of a time-independent potential (see, e.g.,
Ref. [49] and references therein). It is a general method that
relates also to approaches for treating dynamical modes of
periodic Hamiltonians [24,50,51]. The motion generated by the
Hamiltonian

$$H(z, p_z, t) = p_z^2/2 + \psi(z, t)$$

(24)
can be embedded into a larger, 4D time-independent phase
space, using the correspondence

$$H(z, p_z, t) \rightarrow H(q, p, Q, P) = H(q, p, Q) + P.$$  

(25)

In Appendix B, we show that this embedding reproduces the
original motion of $\{z(t), p_z(t)\}$ for a fixed energy shell
of $H(q, p, Q, P) = 0$. Since the conserved energy of a 4D
time-independent Hamiltonian defines a 3D manifold, this
justifies our statement that the original motion takes place in
an effectively 3D phase space.

Using this embedding, we can deduce that the phase space
of the rf potential has a further important property. Since this
phase space is effectively 3D, any 2D invariant torus within
it divides the space into two disconnected regions—“inside”
and “outside.” This means that it is possible to define the “last
unbroken torus,” that is, the torus with maximal volume. In the
stroboscopic map, it is simply the “last” continuous invariant
curve, having maximal area. Since phase-space trajectories
cannot cross, any motion started with an initial condition within
the last unbroken torus will remain bounded. Even chaotic
motion, within this torus, is disconnected from the chaotic
sea outside. The existence of a torus that divides the phase
space into two disconnected components facilitates a rigorous
characterization of the trap.

C. Trap characterization

As a quantitative measure of the trap’s ability to trap
particles, we take the phase space area within the last unbroken
torus. This measure is in fact proportional to the action of the
canonical action-angle coordinates for integrable motion. It
is a phase-space invariant conserved in time, which is well
defined both for the pseudopotential and the rf potential (for
which the energy, in contrast, is not conserved) and allows an
accurate comparison of the capability of the trap to store ions.
The action is also monotoneous with the ion’s maximal kinetic
energy (Fig. 2). Beyond the last unbroken torus, the chaotic
region is connected to the unstable fixed point and the ion is
likely to quickly escape the trap.

For the pseudopotential, the phase-space area within a given
closed invariant curve can be easily calculated by integrating
along the curve,

$$J = \int p_z dz = 2 \sqrt{2[E - \psi(z)]} dz,$$  

(26)

where the integration on the right-hand side is to be performed
from the minimal $z$ to the maximal $z$ of the closed curve, and $E$

is the energy of the particle. The phase-space area of the
rescaled pseudopotential of Eq. (18) is given by $J_\lambda = J/q_5$

coming from Eq. (17). The trapping phase space area, $J_{\text{max}}$, is
defined for each set of parameters as the maximal value of $J$

for the bounded motion. In fact, it is the area bounded by the
separatrix—the closed curve that passes through the unstable
fixed point.

Within the rf potential, the area within the last unbroken
torus can be calculated directly from the stroboscopic map.
However, this calculation is cumbersome when $p_z$ (restricted
to positive values, say), is not a function of $z$, as can happen
near resonance islands; see Fig. 2(d). We can use the fact that
the stroboscopic map of $H$ coincides with a Poincaré surface of
section of $\mathcal{H}$ defined in Eq. (25), as discussed in Appendix B,
and calculate the phase-space area within a closed curve of the
stroboscopic map using a method developed for calculating the
action of 2D tori surfaces in the 4D phase space, described in
Appendix C. With this method, we calculate $J_{\text{max}}$, the phase-
space trapping area, which is the area bounded by the last
unbroken torus of the rf potential phase space.

To map the phase space of both the rf potential and the pseudopotential, we simulate the dynamics with initial
conditions of $p_z(0) = 0$ and small increments in $z(0)$ between
$z_\text{i}$ and $z_\text{u}$. This allows us to cover the entire phase space (except
some resonance islands) and also to construct numerically the
transformation to action-angle variables throughout the region
of regular motion. The simulation is repeated for different trap
parameters, giving a complete characterization of the depend-
ence on the parameters. An invariant curve is numerically
identified if the curve obtained from a single trajectory is
continuous and has no width (to some accuracy). For the
verification of the numerics, the pseudopotential simulations
are compared with direct quadrature, numerically minimizing
Eq. (20) and integrating Eq. (26). The phase-space trapping
area is also calculated directly for convex curves from the
FIG. 4. Rescaled (nondimensional) trapping phase-space area, plotted as function of $\sqrt{\lambda} \propto \omega_x/U_{rf}$, at increasing values of $q_5$ [see text for details and also Eqs. (5) and (14)]. The points are obtained by numerical simulation of the equations of motion in the rf potential, and the dotted curve is obtained for the pseudopotential from Eq. (26). Up to $q_5 \approx 0.5$ ($U_{rf} \approx 25$ V), the pseudopotential gives a very good approximation of the global trap characteristics in the rf potential, with all simulated trap parameters falling very close to the pseudopotential curve, a function only of $\lambda$. The sharp jumps for larger $q_5$ values are due to the last unbroken torus breaking and being replaced by a nested one.

FIG. 5. Trapping phase space area $J_{\text{max}}$ (nondimensional), as a function of the axial trapping frequency $\omega_x$, for increasing values of $U_{rf}$ (equivalently, $q_5$; see text for details). For $U_{rf} = 20$ V, the results of the rf potential simulations overlap the pseudopotential curve. With $U_{rf} = 30$ V, $J_{\text{max}}$ increases for both the pseudopotential and the rf potential, but is lower within the rf potential for $\omega_x/2\pi < 3$ MHz as compared with the pseudopotential curve. For higher values of $U_{rf}$, the deviation becomes even more pronounced, since the pseudopotential trapping phase-space area (not shown) only increases, while within the rf potential $J_{\text{max}}$ decreases due to chaos. While for the pseudopotential $J_{\text{max}}$ depends monotonously on $\omega_x$ and $U_{rf}$, within the rf potential this dependence is nonmonotonous and is generally more complicated due to the role played by chaotic motion. The maximal trapping is obtained with $U_{rf} \approx 25$ V and $\lambda \to 0$.

FIG. 6. (a) Trapping phase-space area and (b) escape height [both in nondimensional units; see Eq. (5)], as a function of the linearized axial trapping frequency $\omega_x$ at the stable fixed point, for increasing values of the axial trapping frequency $\omega_x$. The global trapping characteristics are nonmonotonic with $\omega_x$ and tend to become independent of $\omega_x$. In addition, it can be seen that the escape height and phase-space trapping area do not reach their maximum together as a function of the parameters.
To maximize the total phase-space trapping area, taking $U_{rf} \approx 25$ V at the border of the validity of the pseudopotential is optimal, and $\lambda \to 0$ (equivalently, $\omega_z \to 0$) is required. Otherwise, if $\omega_z$ is constrained, the maximal trapping will be obtained at different values of $U_{rf}$. In experiments with trapped ions, it is often desirable to have the linearized trapping frequencies $\omega_z$ as high as possible. However, increasing $\omega_z$ causes antitrapping in the radial coordinates and increases $\lambda$ (at fixed $\omega_z$). Increasing $\omega_z$ (at fixed $\omega_z$) requires increasing $U_{rf}$ and hence the nonintegrable perturbation. Both effects eventually degrade the trap.

Finally, in Fig. 6(b) we plot the escape height from the trap, defined as the minimal position $z > z_s$, at $p_z = 0$ and $t = 0 \ (mod \ \pi)$, for which the ion escapes the trap. The escape height and the phase-space trapping area do not attain their maximum together, which emphasizes the importance of the trap’s capability to trap high-momentum particles, measured by the phase-space area.

**IV. HAMILTONIAN MOTION IN TWO SPATIAL DIMENSIONS**

The pseudopotential for the coupled $yz$ motion, including the bias voltage, can be decomposed as

$$V_{\text{pseudo}}^{2D} = \frac{1}{2} a_z [y^2 + (z - z_0)^2] + \frac{q_5^2}{4} [V_{avg}^{2D}] + a_5 [V_{sw}(y, z) - \frac{1}{2}],$$

(27)

with $V_{sw}$ defined in Eq. (2), the factor of $1/3$ makes the potential vanish at its minimum, and $V_{avg}^{2D}$ the time-independent approximation to the time-dependent part of Eq. (6), can be derived as outlined in Appendix A and reads

$$V_{avg}^{2D} = \frac{16[16(y^2 + z^2)^2 + 24(y - z)(y + z) + 9]}{2^2[(1 - 2y)^2 + 4z^2][3 - 2y]^2 + 4z^2][3 - 2y + 2z^2][2y + 2z + 2z^2].$$

(28)

With the second rescaling of time ($t \to q_5 t$) given in Eq. (17), we get the rescaled pseudopotential

$$V_{\lambda}^{2D} = -\frac{1}{2} \lambda [y^2 + (z - z_0)^2] + V_{avg}^{2D} - \lambda_b [V_{sw}(y, z) - \frac{1}{2}],$$

(29)

where we have defined the pseudopotential bias parameter,

$$\lambda_b = -\frac{a_5}{q_5} \propto -\frac{U_b}{(U_{rf})^2}.$$  

(30)

Let us first consider the unbiased potential ($\lambda_b = 0$). By running careful numerical simulations of the dynamics over a fine grid of initial conditions, we have found almost no signatures of chaos throughout the phase space up to the unstable fixed point. Pinning down the reason for this near-integrability and identifying the implied additional constant of motion is beyond the scope of the current work. However, this integrability implies that any added perturbation, if not too large, introduces only a limited structural change of the phase space. In this section, we demonstrate this statement for two separate perturbations of the integrable pseudopotential and then their combined effect. First, we consider (in Sec. IV A) the effect of a negative bias voltage on the structure of the pseudopotential phase space. Then, in Sec. IV B, we study the full rf potential, which can be considered as a second perturbation scaled by $q_5$ that is added to the pseudopotential, including the bias.

**A. The 4D pseudopotential phase space**

In the 4D phase space of the $yz$ pseudopotential, increasing $\lambda_b$ increases the trap depth by pushing the unstable fixed point $z_u$ up, and eventually completely eliminating it, allowing the ion to be trapped without escaping for any energy. At the same time, this perturbation gradually deforms the phase space, breaks the main island into several smaller islands, and introduces a chaotic region where the ion is expected to be more susceptible to heating and can escape from the trap (within the rf potential, to be studied in Sec. IV B). Hence, the bias voltage induces a competition between two mechanisms of opposite nature that we investigate in the following.

We have simulated the equations of motion derived from Eq. (27), varying the parameters. We vary $\lambda$ between $\sqrt{\lambda} \approx 0.016$ and $\sqrt{\lambda} \approx 0.065$ [$\omega_z = 2\pi \times 0.5$ MHz to $\omega_z = 2\pi \times 2$ MHz at $U_{rf} = 20$ V], corresponding to the first quarter of the horizontal axis of Fig. 4. We vary $\lambda_b$ between $\sqrt{\lambda_b} = 0.21$ [$U_b = -0.2$ V at $U_{rf} = 20$ V] and $\sqrt{\lambda_b} = 0.43$ [$U_b = -0.8$ V at $U_{rf} = 20$ V], in addition to the unbiased potential $\lambda_b = 0$.

For each pair ($\lambda$,$\lambda_b$), we run the simulation on a grid of energies between 0 and up to a maximal energy, which we take to be the energy of a particle at rest at a certain height $z_u^{max}$. For $\lambda_b = 0$, this height is just $z_u^{max} = z_u$. For $\lambda_b > 0$, $z_u$ is pushed up significantly, and we truncate the simulations at an arbitrary energy shell corresponding to a particle at rest at a height of

$$z_u^0 \approx 2.3 \times z_u \approx 2.$$  

(31)

As we discuss in detail in the conclusions of the following subsection, trajectories that reach very high above the electrode are expected to be less relevant from an experimental point of view, and hence the condition above is a reasonable one.

We begin by examining the structure of the phase space as a function of $\lambda$ and $\lambda_b$. For a small value of the bias potential, we find that hardly any parts of phase space become chaotic as $z_u$ is pushed up. Figure 7 demonstrates this for different values of $\lambda$. Each panel shows a Poincaré surface of section, constructed by taking simulations with different initial conditions at a fixed energy, and plotting a point in the plane of $\{z, p_z\}$, every time a simulated trajectory goes through $y = 0$, with $p_y > 0$. Except for the largest $\lambda$ value, the phase space contains a large central regular island. For the two highest $\lambda$ values shown, the cusp in the trajectories indicates the proximity of the unstable fixed point in a higher energy shell. For the maximal value of $\lambda$, the phase space is completely broken into smaller resonance islands, although the motion still stays almost regular.
We find that for this relatively small value of \( \lambda \) the ion dose not escape for any initial condition and the energy shell is close to the maximal simulated volume of interest, to be defined below. Since we can reduce the 4D indicator functions will be significantly up, while the phase space remains mostly regular (albeit with large resonance islands). We note that in contrast to Figs. 2 and 3, here the island centers do not correspond to any fixed point. In addition, each island is seen to be a section of a different, disconnected family of tori.

Fixing the parameters, with \( \lambda \) at the maximal value we have used and \( \lambda_b \) at an intermediate value, Fig. 8 shows Poincaré sections at different, increasing energies. The phase space splits into two central islands with an increasing chaotic sea between them. In Fig. 9, for the largest value of \( \lambda_b \), a similar phase-space splitting occurs (with the main island centered at lower \( z \) values), together with island destruction and an emergence of a new island. However, since in Fig. 9 the value of \( \lambda \) is small, even in the high-energy shells there can be a large fraction of (nearly) regular motion—and we note that the scale of the \( p_z \) axis is \( \sim 1.6 \) times that of Fig. 8 and twice that of Fig. 7.

In order to quantify the trap’s effectiveness within the pseudopotential approximation, we single out a 4D phase-space volume with a given property, to be defined below. Since such a volume is a function of the conserved energy \( E \) up to which it is measured, it can be obtained by integration of a 3D volume of interest \( \Lambda^E(3D) \) over the energy shells,

\[
\Lambda^E(3D) = \int_0^E dE \left[ \Lambda^E(3D)' \right],
\]

with \( \tilde{q} = (y, z) \), \( \tilde{p} = (p_y, p_z) \), \( H \) the pseudopotential Hamiltonian, and \( \chi(\tilde{q}, \tilde{p}) \) an indicator function (taking therefore either the value 0 or 1) that selects some region of phase space with a specific property (different \( \chi \) functions will be introduced below). We note that by its definition, \( \Lambda^E(3D) \) is independent of the rescaling of time [in Eq. (17)], leading to the pseudopotential of Eq. (29), but \( \Lambda^E(4D) \) is not.

Within each energy shell, \( \Lambda^E(3D) \) can be calculated in a straightforward manner from the Poincaré section, as outlined in Appendix D. We stress here that by its definition, the Poincaré section contains the full information on the motion (at each energy). Within the pseudopotential approximation, the Poincaré section is 2D, which simplifies the analysis significantly, since we can reduce the 4D indicator functions \( \chi(\tilde{q}, \tilde{p}) \) to their 2D counterparts \( \chi(z, p_z) \) within the Poincaré section.

We introduce two indicators, \( \chi(z, p_z) \) and \( \chi(z, p_x) \), and assign their value within each Poincaré section (gridded appropriately). For the first, we set \( \chi(z, p_z) = 1 \) on every point of the Poincaré section such that the corresponding motion is...
bounded within the pseudopotential approximation. Hence, \( \chi_t \) measures the total pseudopotential phase-space volume. We further define \( \chi_t(z,p_r) \) to equal 1 on points of the Poincaré section that (in the pseudopotential approximation) are within the last unbroken torus of the center islands (around \( p_z = 0 \)), and 0 otherwise. With this definition, it is relatively fast to compute \( \chi_t \) numerically, and it gives a good approximation of the regular motion volume. A more accurate quantification where regular regions outside the last unbroken torus are included and chaotic regions within the last unbroken torus are excluded would be significantly more complicated to implement for very extensive simulations, and not necessarily more relevant.

Figure 10 compares these two measures of the pseudopotential phase-space volume, as a function of \( \lambda, \lambda_b \). The limit of integrability appears to be near \( \lambda \rightarrow 0 \) and \( \lambda_b \rightarrow 0 \). For the smallest \( \lambda \) value, even a large \( \lambda_b \) (up to the largest one) allows a large gain of (mostly) regular phase space (Fig. 9). Likewise, for the smallest value of \( \lambda_b \), the phase space can change its structure but maintain primarily regular motion up to a large value of \( \lambda \) (Fig. 7). However, as both \( \lambda \) and \( \lambda_b \) increase beyond their minimal values, the increased useful phase-space volume due to the bias may be saturated at a certain (\( \lambda \)-dependent) \( \lambda_b \) value, by the destruction of the regular motion regions that come with it (Fig. 8). Within the pseudopotential, the dynamics are independent of the value of \( U_{rf} \), which only rescales the \( \lambda \) and \( \lambda_b \) values for the smallest value of \( \lambda \), the 4D phase-space volume. A more accurate quantification could be obtained by using the 2D Poincaré sections constructed for the 4D phase space of the pseudopotential.

Repeating the same set of simulations as described in the previous subsection (for the pseudopotential), within the full \( yz \) rf potential of Eq. (6), we take \( U_{rf} \) between 20 and 35 V (corresponding to \( q_S \in [0.43, 0.76] \)), as suggested by the analysis of Sec. III. At each value of \( U_{rf} \), we simulate exactly the same initial conditions as simulated with the pseudopotential. We run the simulation for 20 000 rf periods, a duration longer than the typical lifetime of the trajectories that were observed to escape. In addition to \( \chi_t \) and \( \chi_r \), that characterize the motion solely within the pseudopotential, we can now define \( \chi_q(z,p_r) = 1 \) on every point of the pseudopotential Poincaré section if it belongs to a trajectory that does not escape from the trap within the rf potential simulation, when initiated at time \( t = 0 \) with the same initial condition (and 0 otherwise).

In Figs. 11 and 12 we compare, as a function of \( \lambda, \lambda_b \), and \( U_{rf} \), the total pseudopotential phase-space volume nonescaping within the rf potential \( (\chi_q \chi_r) \), with the pseudopotential volume bounded within the last unbroken torus of center islands, and nonescaping within the rf potential \( (\chi_q \chi_r) \). Although within the pseudopotential, the dynamics are determined by \( \lambda \) and \( \lambda_b \) are identical for any value of \( U_{rf} \) at fixed \( \lambda \), the 4D phase-space volume \( \lambda^{(4D)} \) in fact scales \( \propto (U_{rf})^2 \). This scaling would imply an increase of the volume with \( (U_{rf})^2 \) for fixed \( \lambda \) and \( \lambda_b \). However, the competition with the effect of trajectories gradually escaping from the trap due to the increase of \( U_{rf} \), in particular in the higher energy shells and larger values of \( \lambda \) and \( \lambda_b \), determines the effective gain due to the increase of \( U_{rf} \).

By comparing Fig. 10 with Fig. 11(a), we find that for intermediate parameter values, \( \sqrt{\lambda} \lesssim 0.043 \) and \( \sqrt{\lambda_b} \lesssim 0.30 \), almost the entire pseudopotential phase-space volume is bounded within unbroken tori, and hardly any of that escapes
within the rf simulation at $U_{\text{rf}} = 20$ V. For larger $\lambda$, $\lambda_0$ values, there could be a large chaotic volume that remains trapped. Comparing with Fig. 11(b), we see that the trapped volume increases (and is mostly regular) if increasing $U_{\text{rf}}$ to 25 V for $\sqrt{\lambda_0} \lesssim 0.3$, while for larger $\lambda_0$, we see a crossover from an increase to a decrease of the volume (depending on $\lambda$), due to trajectories escaping within the rf potential. Figure 12(a) shows that for $U_{\text{rf}} = 30$ V and $\sqrt{\lambda_0} \gtrsim 0.3$, trajectories continue to escape from the rf potential and lead to a decrease of trapped phase-space volume as compared with $U_{\text{rf}} = 25$ V, since even the regular parts shrink. For $U_{\text{rf}} = 35$ V [Fig. 12(b)], even for $\lambda_0 \gtrsim 0.2$, the volume decreases (as compared with $U_{\text{rf}} = 30$ V).

Hence, we see again how the near-integrable pseudopotential behaves under the addition of combined perturbations. The pseudopotential volume bounded by the last unbroken torus of center islands appears to give a good measure of the trajectories that are robust to escape under the rf potential. It indicates the trap’s capability to trap particles of a high kinetic energy, and it is a relatively simple quantity to calculate numerically. The border of validity of the pseudopotential (for approximately regular motion) was found in Sec. III to be $q_x \approx 0.5 (U_{\text{rf}} \approx 25$ V), for any value of $\lambda$. We can conclude that this threshold holds also in the coupled $y_z$ motion, restricted to intermediate values of $\lambda$ and $\lambda_0$. Above this $U_{\text{rf}}$ threshold, all chaotic trajectories escape, and the regular region shrinks as well. For lower values of $U_{\text{rf}}$, even parts of the chaotic motion could remain trapped (for very long times), but for $\sqrt{\lambda} \lesssim 0.043$ and $\sqrt{\lambda_0} \lesssim 0.30$, the phase space is primarily regular. For higher values of $\lambda$ or $\lambda_0$, the picture is more complicated and sensitive to an interplay of the parameters, as described above.

Figure 11 also allows us to deduce how to maximize a trapped phase-space volume that is (mostly) regular. A value of $U_{\text{rf}}$ that is just below the threshold of pseudopotential validity is apparently close to optimal. At lower $U_{\text{rf}}$, the phase-space volume shrinks because of the energy scaling, and at higher $U_{\text{rf}}$, significantly more trajectories escape. For the trap parameters of Eq. (14), it is $U_{\text{rf}} \approx 20$ V. Taking $\lambda \to 0$ is required, and $\sqrt{\lambda_0} \approx 0.37$ can be considered to give the maximal volume which is robust to small integrability-breaking perturbations. Although increasing $\lambda_0$ allows [according to Fig. 11(a)] to continue and increase the trapped volume, by comparing Fig. 11(a) with Fig. 11(b) we see that this added volume is expected to be unstable to small perturbations (in particular the trap’s rf potential itself). At higher values of $U_{\text{rf}}$, as in Fig. 12, restricting the parameters to lower values of $\lambda_0$ is favorable.

The above simulations were based on constraining the maximal height in the calculation of the phase-space volume as a function of the different trap parameters. Beyond being useful for numerical reasons, it is relevant experimentally for a few reasons. Motion far from the trap center in any direction would be in the edges of the laser-cooling beam waist and would damp slowly, possibly slower than typical heating rates. In particular, motion at high energy that approaches very closely the electrodes is potentially more susceptible to heating effects which scale with $1/e^a$, where $a \approx 4$ [53]. Finally, while being cooled toward lower energy, the ion could possibly hit the chaotic sea just because of the topology of the regular trapping islands, shown, e.g., in the higher energy shells of Fig. 9.

We have truncated the simulations at the height given in Eq. (31), which [with Eq. (14)] corresponds to a pseudopotential trap depth of at least $E/k_B \approx 70$ K [with $U_{\text{rf}} = 20$ V and $U_{\text{dc}} = -0.2$ V]. Since atomic ions are trapped in vacuum and routinely cooled by Doppler cooling to milli-Kelvin temperatures, with the entire setup possibly refrigerated to a few Kelvin, this is a very high energy scale. It is only smaller than the initial ion temperatures after their evaporation from an oven and ionization. We have not directly explored the motion at the scale of oven temperatures in our simulations, and the operation of laser cooling during the loading process remains an interesting open question.
V. EXPERIMENTAL PROBE OF NONEQUILIBRIUM ION DYNAMICS

Experimentally probing the phase-space structure of a trap is a challenging task. To avoid any non-Hamiltonian perturbations, the experiment has to be done on a short time scale (so that heating is negligible), and “in the dark” (with the laser cooling beam turned off after the initial cooling step). An ideal experimental sequence would consist of exciting the ion into various well-resolved initial conditions, and then measuring microscopic properties of the trajectory. Both aspects, however, are nontrivial to achieve with large-amplitude motion in 3D. In this section, we present an initial proposal of an experiment procedure that allows one to probe Hamiltonian transport in a mixed phase space with a trapped ion, while circumventing these difficulties.

We consider an indirect method of exciting the ion to a large amplitude motion by an additional low-amplitude periodic modulation [54] of one electrode’s potential (known as a “tickle”) that produces an oscillatory electric field at the equilibrium position of the ion. The tickle perturbation gradually (as a function of its amplitude) destroys the regular motion parts of phase space, turning them into mixed regions, with islands (that vanish and emerge), and chaotic strips that grow into contiguous chaotic regions. The latter may become connected to the regions already chaotic in the unperturbed Hamiltonian, with a structure and geometry which are very sensitive to the tickle parameters. Numerical simulations allow one to construct the ion’s survival probability in the trap as a function of the tickle amplitude and its duration, which shows distinct features that can be averaged over the initial conditions. Thus this approach uses the properties of chaotic motion—sensitive dependence on initial conditions, whose memory is, however, quickly erased (assuming it is mixing), to avoid the necessity of initializing the ion to precise initial conditions. It further replaces the difficulty of experimentally measuring the details of the trajectory with the more accessible reconstruction of the ion’s survival probability as a function of the initial ion energy, the electric field amplitude of the tickle, and the duration of the tickle. As we show below, the survival probability shows strong features that are distinguishable from simple thermalisation over a barrier, or a simple oscillator excitation process.

We must now depart from the five-wire trap in the symmetric configuration, modeled so far in this work. Hitherto, it was chosen because it enabled us to build the theory in steps of increasing complexity of the phase space and parameter dependence. In hindsight, it allowed us to uncover a near-integrable limit of surface traps, which plausibly stands at the radial yz plane. Nevertheless, the yz motion corresponds to a 5D phase space as discussed in Sec. IV. For the parameters of the simulation, the rf frequency is \(\Omega = 2\pi \times 163\) MHz, the stable fixed point is approximately \(40\) \(\mu\)m above the electrode surface, and the linearized secular oscillation frequencies around it are \(\omega_{x,y,z} \approx 2\pi \times \{1.49, 9.16, 10.36\}\) MHz. (34)

Numerical simulations are performed by solving the equations of motion of the ion in the 3D rf potential, starting with initial conditions picked at random from a thermal distribution at the Doppler cooling limit. A periodically modulated voltage (the tickle) is added to an electrode potential [55], with the form

\[
U_d(t) = U_0 \cos(\omega_d t + \phi_0),
\]

where \(\phi_0\), the initial phase of the tickle modulation, is a (uniformly distributed) random variable, and \(\omega_d = 1.1 \times \omega_z\). (36)

Simulating 200 initial conditions at each value of tickle modulation for a fixed maximal duration of 0.4 ms, we obtain the probability that the ion remains trapped as a function of time \(t\) and tickle amplitude \(U_0\). The ion’s survival probability at fixed values of \(U_0\) is shown in Fig. 13, for \(U_0\) in the range 192–244 mV, in steps of 4 mV. We define the survival probability function as

\[
P_s(t; U_0) = 1 - P_c(t < t; U_0),
\]

FIG. 13. The simulated survival probability [Eq. (37)] as a function of time for different tickle amplitudes \((U_0)\), in a four-wire trap (see the text for details of the model and parameters). The simulations show a sharp threshold for the ion’s escape out of the trap, followed by a nonmonotonous dependence on \(U_0\). We interpret the saturation of (some) curves as the coexistence of long-lived trapping pockets that are destroyed and re-emerge as a function of the perturbation amplitude. These features together with the different functional forms of \(P_s(t; U_0)\) present a potentially strong experimental signature that could be used to indirectly probe the trap’s phase-space structure, and a promising direction for studying Hamiltonian transport in a mixed-phase space.
where \( P_s(\tau \leq t; U_0) \) is the probability that the escape time \( \tau \) (which is a random variable in each realization) is between 0 and \( t \), for a fixed value of \( U_0 \).

With the frequency of the tickle modulation slightly higher than \( \omega_z \), the center of the main island (close to resonant with the tickle), is gradually destroyed as \( U_0 \) is increased, but can remain as a pocket of long-lived chaotic motion (longer than the time reached in our simulation), protected from escaping by a broad region of (mostly) regular tori. Indeed for tickle amplitude \( U_0 \approx 192 \text{ mV} \) we observe no escape of the ion. Increasing the tickle amplitude beyond this threshold value, we find a sharp onset of ion escape. After a short-time flat part of the survival probability, the ion escapes at a slow rate (almost linearly with time for \( U_0 = 200 \text{ mV} \)). Surprisingly, the escape rate is not monotonic with the tickle amplitude, and \( P_s(t; U_0) \) returns to nearly unity for \( U_0 = 208–216 \text{ mV} \), before dropping sharply again.

With \( U_0 \approx 220 \text{ mV} \), the functional shape of \( P_s(t; U_0) \) changes strongly and approximates an exponential distribution with a saturation. This is possibly the result of a branching process where the ion, depending on the exact (random) initial conditions, may get trapped in a long-lived “pocket,” or otherwise escape the trap. The escape plausibly occurs from a (mixing) chaotic region through a “bottleneck,” leading to the unstable fixed point, typically a memoryless process with an exponential distribution. The trapping is likely to occur in a large region in phase space that is connected to the region of initial conditions by chaotic motion, but weakly connected to the unstable fixed point, so the probability to escape the pocket is low on the studied timescale. On a longer timescale, the trapping pockets could eventually decay. For \( U_0 \approx 240 \text{ mV} \), there are no observable pockets, but the ion lifetime increases with the perturbation amplitude, and new pockets are formed. It should be stressed that the escape probability shown in Fig. 13 is strictly different from the exponential decay of a simple thermal activation mechanism above a barrier and also cannot be explained as a linear or parametric resonance of a harmonic oscillator, relevant in the low amplitude regime \([56,57]\). This rich nonmonotonic behavior remains to be explored with a more detailed analysis of the phase space, the pockets, and the statistical properties of the motion, but the salient features could be observed with a relatively simple experiment.

Figure 13 indicates that the escape probability beyond \( t \gtrsim 0.1 \text{ ms} \) is weakly dependent on time. In Fig. 14, we plot the value of the survival probability at fixed \( t = 0.3 \text{ ms} \) as a function of the tickle voltage (with data from more simulations). The nonmonotonic dependence of the survival probability on \( U_0 \) is a clear experimental signal that may be obtained by just cooling the ion to the Doppler limit and varying the tickle amplitude. We note that due to the signal being based on ensuing ion escape, a full set of experiments would require a large number of lost and reloaded ions. Beyond a certain number of repetitions, this might degrade the trap stability due to uncontrolled stray charges of the trap electrodes produced by the loading, a technical weakness of the proposed scheme. An extension of the proposed experiment to a multizone trap, separating the loading region from the escape region, may avoid these shortcomings. Another possibility would be observing transfer between multiple wells without going over the trap’s barrier, thus avoiding the ion’s escape altogether.

**VI. SUMMARY AND OUTLOOK**

The main goal of this paper has been to provide a description of Hamiltonian dynamics within surface-electrode Paul traps. We have aimed at developing a general framework that enables us, put simply, to understand both what is going on in the phase space and why it is like that. This understanding of the nature of the dynamics has furthermore allowed us to answer practical questions such as the optimal choice of experimental parameters to obtain the largest possible trapping regions.

To achieve that goal, we have increased the complexity of the treatment gradually, starting from a time-independent one-dimensional model describing the motion along the \( z \) direction perpendicular to the trap surface and including successively the effects of the time dependence and of coupling with the \( y \) direction (in plane but orthogonal to the direction of the rf wires). In this process, we find that although many regimes, each of them having their own idiosyncrasy, exist, the
global picture that emerges is characterized by a few simple observations. The first is that in the regime where presumably most traps are operated in practice, chaotic trajectories usually do not remain trapped. Thus, we can identify the trapped volume with the volume of regular (near-integrable) dynamics. As a consequence, there is in general a competition between increasing the strength of the applied trapping forces, which we would intuitively expect to increase trapped phase-space volumes, and the necessity to avoid entering a regime which is too chaotic. The optimal choice of parameters is determined by that competition, controlled by a few parameters that play multiple roles, and for the five-wire configuration that we have considered in detail, we provide explicit values (while, of course, real-life traps would always contain elements omitted from our model).

More broadly, the theory presented in this work gives a general framework that can be applied to various surface-electrode traps. It can be a starting point for asking more refined questions on the dynamics, for example, including dissipative processes like cooling, or heating. Of the general conclusions that we can draw, an important first step in a future analysis would be to identify an integrable or near-integrable limit of a trap. Then, the parameters should be studied according to their role in increasing the phase space volume on the one hand, and introducing chaotic motion on the other hand. As we saw, typically each parameter introduces both effects, with some value forming a threshold between the two. A further general conclusion is that a classification of the phase-space motion within the pseudopotential approximation could be sufficient for answering many questions for practical parameters and regimes of motion. A natural and robust “boundary” is formed by those parameter values for which the motion is largely integrable within the pseudopotential, up to the point at which a change of the parameters introduces a lot of chaotic motion. The same holds for the rf potential, which has a threshold for the drive amplitude, beyond which a lot of chaos is quickly introduced. Thus below this threshold, for any new parameter that has to be considered, we expect that a study of the pseudopotential is sufficient (up to the point where a significant volume of chaotic motion is introduced), which implies a tremendous simplification. Above this rf threshold, at least for the examples analyzed here, we find that there is little benefit because the trajectories quickly escape.

This work represents also a first and necessary step to study in detail Hamiltonian and non-Hamiltonian phase-space transport effects. We have in this work merely touched on the complex motion possible in the 5D phase space. We have presented a preliminary study indicating that rich Hamiltonian transport mechanisms can be explored in the ion trap, and we hope to provoke further endeavours in this direction. The phase-space framework is also a natural starting point for a semiclassical consideration of quantum mechanical effects. The strength of a charged particle’s interaction with the environment also implies that stringent conditions on the system’s isolation are required in order to observe such effects. The manifestations of the phase-space structures that we studied in this work, when the motion becomes sufficiently coherent [37], could be an interesting future direction.

Finally, a quantitative study of non-Hamiltonian effects, such as heating by electric noise, or laser cooling dynamics within the Paul trap, is also of importance and interest. In particular, the combination of micromotion together with stochastic processes presents a challenge both for experimental control and theoretical description. We will study these effects in future publications. In Ref. [42], we will study in detail laser cooling in the nonlinear and time-dependent potential, and its interplay with position-dependent noise, where the phase-space framework will again prove essential to our approach of the problem. Strong position dependence is one characteristic of a dominant source of noise in Paul traps, the so-called anomalous heating [53], which is also characterized by a “colored” (frequency-dependent) noise spectrum. The treatment of colored noise, again using a phase-space approach, will be studied separately [58].

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APPENDIX A: THE PSEUDOPOTENTIAL

The pseudopotential [48] is a time-independent approximation [41] of the time-dependent rf potential. We will now derive the pseudopotential approximation using canonical perturbation theory [59]. We introduce a formal perturbation parameter $\epsilon$, and we consider a particle subject to the Hamiltonian

$$H(q, p, t) = H_0(p) + \epsilon H_1(q, t) + \epsilon^2 H_2(q) = p^2/2 + \epsilon f(t)V_2(q) + \epsilon^2 V_0(q), \quad (A1)$$

where $f(t)$ is periodic with a period $T = 2\pi/\Omega$. At the end of the calculation, we can set $\epsilon = 1$, but we will see that $\epsilon$ can be assigned a physical meaning, and that the ordering of the terms in Eq. (A1) is meaningful as well. We avoid an explicit vector notation here, but the dimension of phase space is arbitrary. We are searching for a canonical transformation to the new set of conjugate coordinates $(Q, P)$, derived from a generating function $F(q, P, t)$. The transformation will depend explicitly on time and is required to reduce to the identity at $t = 0$. Starting with $q(t = 0)$ and $p(t = 0)$, we will construct the canonical transformation to describe the stroboscopic map of $H(q, p, t)$, such that $(Q(t = T), P(t = T))$ approximate $[q(t = T), p(t = T)]$ within the Poincaré surface of section after one iteration. We will see that the evolution of $(Q, P)$ can be derived from a time-independent Hamiltonian expanded in the perturbation parameter

$$K(Q, P) = K_0(Q, P) + \epsilon K_1(Q, P) + \epsilon^2 K_2(Q, P) + \cdots. \quad (A2)$$
Hence, with the canonical transformation defined by \( q = q(Q, P, t) \) and \( p = p(Q, P, t) \), we require that \( F \) obeys

\[
F(q, P, t) = qP + \epsilon F_1 + \epsilon^2 F_2 + \ldots, \quad p = \frac{\partial F}{\partial q}, \quad Q = \frac{\partial F}{\partial P},
\]

and we expand similarly the transformation functions,

\[
q(Q, P, t) = Q + \epsilon q_1 + \epsilon^2 q_2 + \cdots, \quad p(Q, P, t) = P + \epsilon p_1 + \epsilon^2 p_2 + \cdots.
\]

Now by plugging \( q = q(Q, P, t) \) using Eq. (A4) into the definition of \( F \) from Eq. (A3), Taylor expanding the terms of \( F \) with respect to \( q \), and collecting the terms up to order \( \epsilon^2 \), we have

\[
p_1 = \left[ \frac{\partial F_1(q, P, t)}{\partial q} \right]_{q=Q}, \quad p_2 = \left[ \frac{\partial^2 F_1(q, P, t) q_1(Q, P, t) + \frac{\partial F_2(q, P, t)}{\partial P}}{\partial q} \right]_{q=Q},
\]

where the partial derivatives are to be taken with respect to argument \( Q \), and then \( q \) is simply to be replaced by the variable \( Q \) (and not by the function \( q(Q, P, t) \), which has already been used in the expansion). Similarly,

\[
q_1 = -\left[ \frac{\partial F_1(q, P, t)}{\partial P} \right]_{q=Q}, \quad q_2 = -\left[ \frac{\partial^2 F_1(q, P, t) q_1(Q, P, t) + \frac{\partial F_2(q, P, t)}{\partial P}}{\partial q} \right]_{q=Q}.
\]

Finally, the transformed Hamiltonian, expanded to the same order, is given by

\[
K(Q, P) = H(q, p, t) + \frac{\partial F(q, P, t)}{\partial t} = H_0(P + \epsilon p_1 + \epsilon^2 p_2) + \epsilon H_1(Q + \epsilon q_1, t) + \epsilon^2 H_2(Q) + \epsilon \left[ \frac{\partial F_1}{\partial t} \right]_{q=Q+\epsilon q_1} + \epsilon^2 \left[ \frac{\partial F_2}{\partial t} \right]_{q=Q},
\]

where the first step of the expansion has been indicated, and the next steps will be performed in the following.

Equating Eq. (A2) with Eq. (A7), we get at order \( \epsilon^0 \) simply

\[
K_0(P) = H_0(p = P) = P^2/2.
\]

We now recall that if \( G(Q, P, t) \) is a function of phase-space variables with an explicit dependence on time, its total time evolution (on an orbit in phase space, for motion generated by the Hamiltonian \( K_0 \)) has the derivative (using Poisson brackets),

\[
\frac{dG(Q(t), P(t), t)}{dt} = \frac{\partial G(Q, P, t)}{\partial t} + [K_0, G] = \frac{\partial G}{\partial t} + \frac{\partial K_0}{\partial P} \frac{\partial G}{\partial Q} - \frac{\partial K_0}{\partial Q} \frac{\partial G}{\partial P} = \frac{\partial G}{\partial t} + F_1 \frac{\partial G}{\partial Q},
\]

At order \( \epsilon^1 \) we get the equation

\[
K_1(Q, P) = p_1(Q, P, t)P + H_1(q = Q, t) + \frac{\partial F_1}{\partial t} \bigg|_{q=Q},
\]

which, substituting \( p_1 \) from Eq. (A5) and using Eq. (A9), can be seen to have the form

\[
\frac{dG_1(Q, P, t)}{dt} = K_1(Q, P) - H_1(Q, t),
\]

where \( Q \) evolves along the orbits of \( K_0(P) \) [a free particle], and

\[
G_1 = [F_1]_{q=Q}.
\]

Equation (A11) can be solved by the method of integrating the right-hand side along these orbits [59], and \( F_1 \) would be that integral, with \( q \) replacing \( Q \) and the initial condition \( F_1(t = 0) = 0 \). Skipping the details of the derivation, it is easy to verify that \( G_1 \) is given by

\[
G_1(Q, P, t) = \int_0^t \left[ K_1(Q + P(t' - t), P) - H_1(Q + P(t' - t), t) \right] dt'.
\]

and hence, \( F_1(Q, P, t) = G_1(Q, P, t) \big|_{q=Q} \).

So far the treatment has been quite general, and there is still a lot of freedom in the choice of \( K_1 \). The natural choice that defines the pseudopotential is to take \( K_1 \) to be the average of \( H_1 \) along the orbit of \( K_0 \), over a period \( T \). This will make \( \{Q, P\} \) the mapping of \( \{q, p\} \) under one iteration of the stroboscopic map, and also \( F_1 \) will remain well behaved—without terms that are unbounded in time. We find

\[
K_1(Q, P) = \frac{1}{T} \int_0^T H_1(Q + P(t' - T), t') dt' \approx \frac{1}{T} \int_0^T f(t') [V_2(Q) + \nabla V_2(Q) P(t' - T)] dt',
\]

where the potentials have been Taylor expanded to the required order (as will be justified below), and we use \( \nabla = \partial / \partial Q \) to emphasize that the results can be generalized to any dimension as noted above. We will now further simplify the algebra in the following by assuming that \( f(t) \) obeys the two conditions,

\[
\int_0^T f(t') dt' = 0, \quad \int_0^T f(t') t' dt' = 0,
\]

which hold for any \( T \)-periodic and time-reversal-invariant function \( f(t) = f(-t) \), and in particular for \( f(t) = \cos \Omega t \). In this case, we get simply

\[
K_1(Q, P) = 0.
\]

We can now obtain the canonical transformation functions (at order \( \epsilon \) explicitly,

\[
p_1(Q, P, t) = \frac{\partial G_1}{\partial Q} = -\int_0^t dt' f(t') \nabla V_2(Q + P(t' - t)), \quad q_1(Q, P, t) = -\frac{\partial G_1}{\partial P} = \int_0^t dt' f(t')(t' - t) \nabla V_2(Q + P(t' - t)).
\]
At order \( \epsilon^2 \) we get the equation
\[
K_2(Q, P) = p_2(Q, P, t)P + \frac{1}{2}[p_1(Q, P, t)]^2 \\
+ \left[ H_2(q) + \frac{\partial H_1(q, t)}{\partial q} q_1(Q, P, t) \\
+ \frac{\partial^2 F_1}{\partial q \partial t} q_1(Q, P, t) + \frac{\partial F_1}{\partial t} \right]_{q=Q},
\]
which, as before, substituting \( p_2 \) from Eq. (A5) and using Eq. (A9), can be rearranged to have the form
\[
\frac{dG_2(Q, P, t)}{dt} = K_2(Q, P) - H_2(Q) - \frac{1}{2}[p_1(Q, P, t)]^2 \\
+ \frac{\partial}{\partial Q} \left[ p_1(Q, P, t)P + H_1(Q, t) \\
+ \frac{\partial F_1}{\partial Q} Q, P, t \right] q_1(Q, P, t),
\]
with \( G_2 = [F_2]_{q=Q} \). From Eq. (A10), the term in square brackets is just \( K_1(Q, P) \) and by Eq. (A15) it is 0. We will not need the explicit solution for \( G_2 \) [that can be obtained as in Eq. (A12)]; however, we are interested in the pseudopotential \( K_2(Q, P) \), that is chosen to make the mean of \( G_2 \) over a period \( T \) vanish. Hence, we have
\[
K_2(Q, P) = \frac{1}{T} \int_0^T \left[ H_2(Q + P(T' - T)) \\
+ \frac{1}{2}[p_1(Q + P(T' - T), P, T')] \right] dt'.
\]
which gives, using Eq. (A16),
\[
K_2(Q, P) \approx V_0(q) + \frac{1}{2T} \int_0^T dt' \left[ \int_0^{t'} dt_2 f(t_2) \right]^2 [\nabla V_2(Q)]^2.
\]
For \( f(t) = \cos \Omega t \), we have thus derived the familiar pseudopotential expression (setting \( \epsilon = 1 \)),
\[
K(Q, P) = \frac{P^2}{2} + V_0(q) + \frac{1}{4\Omega^2} [\nabla V_2(Q)]^2.
\]
We note that \( K \) is independent of the momentum \( P \) due to the conditions in Eq. (A14). We are left only with justifying the Taylor expansion of the potentials. The motion within the pseudopotential \( K(Q, P) \) is characterized by a frequency \( \omega \), and the nondimensional frequency \( \nu = 2\omega/\Omega \) can serve as the perturbation parameter. The scale of the variables and functions above is given by
\[
Q \sim 1, \quad P \sim \omega Q \sim \frac{\nu}{\Omega} Q, \\
K \sim V_0 \sim K_2 \sim \omega^2 Q^2 \sim \nu^2 \Omega^2 Q^2,
\]
which is consistent with \( V_2 \sim \nu \Omega^2 Q^2 \) and \( \nabla V_2 \sim \nu \Omega^2 Q \) when Eq. (A21) is used. Looking at the order of the terms in Eqs. (A13) and (A18) [for short times of order \( \Omega^{-1} \)] justifies \textit{a posteriori} the approximation in these equations, showing that pseudopotential of Eq. (A21) is an approximation to second order in \( \nu \). This is consistent with the perturbation parameter that was used in the derivation, \( \epsilon = \nu \) [24], and the ordering of the terms in Eq. (A1).

**APPENDIX B: EMBEDDING IN 4D PHASE SPACE**

In Eq. (25), we have written the embedding of the motion generated by a general 1D time-dependent Hamiltonian, \( H(z, p, t) \rightarrow H(q, p, Q, P) = H(q, p, Q) + P \). To see that this embedding reproduces the original motion as solutions of Hamilton’s equations, we can directly calculate the latter. The equations for \( q \) and \( p \) coincide with those of \( z \) and \( p_z \), and in addition we have \( Q = 1 \) (and hence \( f = 1 \) which is required by consistency), and \( P = -\partial H(q, p, Q)/\partial Q \). Since \( H \) is time independent, each of its trajectories lies on a 3D manifold of constant “energy” \( \mathcal{H}(q, p, Q, P) \equiv \epsilon \), in the 4D phase space. The original motion is reproduced for \( \mathcal{H} = \epsilon = 0 \), since this gives using Eq. (25), \( 0 = H(t) + P(t) \), compatible with the equation for \( P \), as can be verified directly.

A Poincaré surface of section is constructed by taking simulations with different initial conditions at a fixed energy, and plotting a point in the plane of the two variables (e.g.) \( q \) and \( p \), every time a simulated trajectory goes through a fixed value of (e.g.) \( Q \), in a definite direction. Here since \( H \) depends on \( Q \) only through \( \cos 2Q \), we can understand that the stroboscopic map of \( H \), taken at a fixed value of \( t \) (mod \( \pi \)), is equivalent to a Poincaré surface of section of \( H \), taken within the fixed energy subspace \( \epsilon = 0 \), with \( \cos 2Q = \text{const} \). Therefore, the phase-space area within a closed curve of the stroboscopic map can be calculated using a method developed for calculating the action of 2D tori surfaces in the 4D phase space, described in Appendix C. This method can also be applied to the calculation of the action for the \( yz \) motion within the pseudopotential approximation, studied in Sec. IV.

**APPENDIX C: IN Variant TORI IN 4D PHASE SPACE**

Here we describe (based on Ref. [37]) the calculation of action invariants in the 4D phase space of the canonical variables \( \{q, p, Q, P\} \), that can correspond to the time-independent pseudopotential in two spatial dimensions, or to the time-dependent potential in one spatial dimension using the correspondence in Eq. (25). On each invariant torus one can draw two independent paths \( C_1 \) and \( C_2 \), and define the corresponding area \( J_a = \int_{C_a} [pdq + QdP] \), and action \( I_a = I_{1a}/2\pi \), with \( a = 1, 2 \). We are working within the integrable approximation (ignoring any chaotic structure in the phase space), inside the main island up to the last unbroken torus, where there exists a local canonical action-angle transformation \( \{q, p, Q, P\} \rightarrow (I_1, I_2, \theta_1, \theta_2) \). The energy is conserved, i.e.,
\[
\mathcal{H}(J_1, J_2) = \epsilon,
\]
and we can choose \( C_1 \) to be in the Poincaré surface of section defined by the intersection of the torus with \( Q = 0 \). The action integral over \( C_1 \) is then \( J_1 = \int_{C_1} pdq \). To obtain both the value of \( J_1 \) and \( J_2 \) (the latter cannot be calculated directly), we use the fact that Eq. (C1) determines an implicit function \( J_2(J_1) \). Taking its differential,
\[
\frac{\partial \mathcal{H}}{\partial J_1} dJ_1 + \frac{\partial \mathcal{H}}{\partial J_2} dJ_2 = 0 \Rightarrow \frac{dJ_2}{dJ_1} = -\frac{\nu_1}{\nu_2} \equiv -\alpha,
\]
where $\alpha$ is the winding number of the torus, allows us to define the Legendre transform of $J_2(J_1)$,

$$J(\alpha) = J_2(J_1(\alpha)) + \alpha J_1(\alpha). \quad (C3)$$

The total action, $J$, is given by the integration of the Lagrangian over the time,

$$J = \int \{ p\dot{q} + P \dot{Q} \} dt.$$ 

The winding number can be approximated by $r/s$, where $r$ is the number of turns around the center of the Poincaré section and $s$ is the number of times, starting from $q(0), p(0)$ in the $Q = 0$ plane, that the trajectory crosses the Poincaré section before returning (within some accuracy) to the initial point. By the definition of the Legendre transform, $dJ/d\alpha = J_1$. Therefore, simulating the equations of motion on a narrow grid covering the phase space, and calculating the corresponding values of $J$ and $\alpha$, allow one to obtain $J_1$ and $J_2$.

For the particular case of Eqs. (24) and (25), the action is given by

$$J = \int \{ p\dot{q} + P \dot{Q} \} dt = \int \{ p\dot{z} - H(z, p_z, t) \} dt$$

$$= \int \{ \dot{z}^2 - H(z, p_z, t) \} dt = \int \left[ \frac{1}{2} \dot{z}^2 - V^{(1D)}(z, t) \right] dt. \quad (C5)$$

**APPENDIX D: VOLUME IN 4D PHASE SPACE**

To calculate any volume of a 4D phase space with a specific property, let us define an indicator function of phase space $\chi_{\vec{q}, \vec{p}}$, taking the value 1 on phase-space structures to be counted, and 0 otherwise. Here, $\vec{q} = (q, Q)$, and $\vec{p} = (p, P)$.

Within a fixed energy shell $E$, the restricted 3D volume of interest can be defined by

$$\Lambda_{\chi}^{(3D)}(E) = \int dqdp \chi_{\vec{q}, \vec{p}}(E - H(\vec{q}, \vec{p})),$$

with the Hamiltonian $H(\vec{q}, \vec{p})$. We now employ a canonical transformation,

$$\vec{q}, \vec{p} \rightarrow \vec{Z} \equiv \{ q, p, t, -E \},$$

where $\{ q, p \}$ are the coordinates within the Poincaré section that has to be constructed. The integral in Eq. (D1) then transforms according to

$$\Lambda_{\chi}^{(3D)} = \int d\vec{Z} \delta(E - H(\vec{Z})) \chi(\vec{Z}),$$

$$= \int dqdp T(q, p) \chi(q, p).$$

where $T(q, p)$ is the return time of the particle to the Poincaré section—the time along the trajectory from a point starting on the Poincaré surface, to its next return to the section. Then a total 4D phase space volume of interest up to energy $E$ can be obtained by integrating over the energy shells,

$$\Lambda_{\chi}^{(4D)}(E) = \int_0^E \Lambda_{\chi}^{(3D)}(E') dE'.$$

**References**


[31] To paraphrase the quote by Carlton Cave on Hilbert spaces, the complete characterization of the motion in 5D phase space can be very challenging.
[55] The tickle is applied in the simulation on electrode C1 in Fig. 1 of Ref. [36]. It is a very elongated electrode along the x axis and consequently produces an electric field with a negligible
component in this direction at the equilibrium position of the ion. The field components in the $y$ and $z$ directions are approximately equal and correspond to approximately 4 kV/m for a tickle amplitude of 1 V on the elongated electrode.


