Surface traps for freely rotating ion ring crystals

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Received 14 May 2015, revised 5 August 2015
Accepted for publication 18 August 2015
Published 16 September 2015

Abstract
Rings of trapped ions are an attractive system to study quantum-many body dynamics with closed boundary conditions as well as symmetry breaking. One of the biggest challenges towards such experiments is to sufficiently suppress rotational imperfections of the trapping potential and to allow for instance freely rotating ring structures. We show how to overcome this challenge with a surface trap design and perform numerical calculations to analyze the consequences of various imperfections in detail. We conclude that trap electrode imperfections, external stray electric fields, and local charging of the trap electrodes can be controlled sufficiently well to allow ion rings to rotate freely even near their rotational ground state.

Keywords: ion trap, ring traps, quantum simulations
(Some figures may appear in colour only in the online journal)

1. Introduction

The electronic and motional degrees-of-freedom of ions trapped with electromagnetic fields are extremely well decoupled from their environment. In addition, lasers and electromagnetic fields allow for excellent control of both degrees-of-freedom on the single quantum level [1, 2]. Both of those properties make ion crystals nearly perfect systems to study many-body physics in closed systems [3–6]. While most experiments are carried out with linear ion strings, a particularly interesting structure is a ring of trapped ions. Proposals include mini-accelerators [7, 8], dynamics of kinks [9], quantum emulation of ring molecules, and the acoustic analog of Hawking radiation [10]. Recently, also rings of trapped ions have been suggested to realize the concept of so-called time crystals [11, 12].

However, starting from experiments on how to implement rings of trapped ions by Schütz et al [8, 13], it has become clear that imperfections and charging will make it very hard to implement such experiments. Thus, a few design improvements have been proposed [14–16]. Furthermore, the Sandia group has implemented a ring trap on surface trap technology [17]. Common to those designs and experiments is that the resulting ring potential has relatively large diameters, making it difficult to compensate for imperfections.

Inspired by [18, 19], we study a novel design, deviating from the idea of bending a conventional linear trap into a ring. In addition, with a planar electrode design amenable to microfabrication, we seek to reduce inevitable imperfections from the geometry as well as local charging of trap electrodes. The main feature of our geometry is to trap the ion ring far away from the trapping electrodes as compared to the typical ion–ion distance and the ring diameter itself. Thus local imperfections from stray charges affect the rotational symmetry of the ion ring much less as if the ions were trapped close to the trap electrodes.

Our design is composed of concentric planar ring electrodes (see figure 1). Trapping is accomplished by applying suitable radio frequency (rf) voltage to those rings. Work by Clark suggests that the multipoles of such a trapping potential can be adjusted over a wide range by changing the rf voltage to the various rings and the size of the rings [19]. In view of the fact that it is experimentally difficult to keep several rf high-voltage sources in phase, we design a trap requiring only one rf high-voltage source. Fixing this parameter, we aim to
find a trap geometry yielding a rotationally symmetric potential minimum at the desired ring diameter and height. For the study of the physics of symmetrization of the wavefunction of bosonic and fermionic ions discussed in the context of time crystals [11, 12], it will be important that the ion ring can freely rotate at the level of single rotational quanta. Keeping that in mind, we consider ion ring can freely rotate at the level of single rotational quanta. We simulated a gap size of 400 μm above the surface, which alter the potential only very little when the ion is less than 100 μm above the trap surface. figure 1. Schematic of the trap showing ring-shaped surface electrodes in the center. A ring of ions (floating balls in the figure) are trapped about 400 μm above the surface. The fan-shaped electrodes are compensation electrodes.We simulated a gap size of 15 μm, which alter the potential only very little when the ion is less than 100 μm above the trap surface.

For identical fermions, the energy scale is the same, although its dependence on α can be different. For an ion ring of 100 \(^{40}\)Ca\(^{+}\) ions with a diameter of 100 μm, the energy scale of the collective rotation is \(E_{\text{gap}}/k_{B} = 0.2\) nK. We seek to create a ring potential with sufficiently small imperfections such that a classical ion ring with rotational energy corresponding to the ground state energy \(E_{\text{gap}}/2\) would not be pinned by the imperfections.

In view of these considerations, it is important to trap the ion ring far away from trapping electrodes, while at the same time keeping the ion ring as compact as possible. Advantages of this approach are that effects of trap imperfections on the ion crystal’s rotational symmetry are reduced and that heating of the ion motion should be manageable. Additional design constraints are ease of symmetric trap fabrication as well as reasonable trapping voltages while maintaining an appreciable trap depth. To this end we target an ion ring with a diameter of 100 μm trapped 400 μm above the trap electrodes.

The remainder of the paper is organized as follows. In section 2, we briefly summarize the methods outlined in [19] on how to efficiently calculate rotationally symmetric potentials. Armed with the potential, we study the structure of ion crystals forming in such ring shaped potentials in section 3. We then analyze various imperfections breaking the rotational symmetry in section 4, most notably external stray fields, electrode edge irregularities, and local charging of the trap electrodes. Section 5 addresses the process of cooling and pinning such a ring of ions.

2. Calculation of the trapping potential

We start with the trap design proposed in [19] composed of planar ring electrodes of different radii and applied voltages. Given this cylindrically symmetric boundary condition, the analytic solution to Laplace’s equation is given by [21]

\[
\Phi(z, r) = \int_{0}^{\infty} J_0(kr) e^{-kz} A_0(k) dk.
\]

Where \(J_i\) is the Bessel function of ith order. \(A_0(k)\) can be expressed as \(A_0(k) = \sum_{n=1}^{N} A_n(k)\), and \(A_n(k)\) is given by

\[
A_n(k) = V_i(b_i J_i(kb) - a_i J_i(ka)).
\]

Where \(b_i\) and \(a_i\) are the outer and inner radius of each ring electrode and \(V_i\) is the amplitude of the rf voltage applied to each electrode. In order to study ions in this oscillating trapping potential, we approximate the potential by the time-averaged pseudopotential

\[
\Psi(z, r) = \frac{Q^2}{4 M \Omega_{\text{rf}}^2} |\bar{E}(z, r)|^2.
\]

This approximation is valid when the oscillation frequency of the trapped ion is much smaller than the rf frequency.

 Trap design. Reference [19] showed that multipole surface traps can be built from concentric rings with particular sets of ring diameters and applied voltages. In view of the fact that it is experimentally difficult to keep several rf high-voltage sources in phase, we design a trap targeting only one rf high-voltage source. Our design, shown in figure 2, is composed of three concentric ring electrodes with outer radius \(b_j = \{126, 600, 1100 \mu m\}\), with the second ring grounded and the other two connected to a fixed rf driving source of amplitude \(V_{\text{rf}} = 1000\) V. As the rf driving frequency, we choose \(\Omega = 2 \pi \times 6\) MHz. In what follows, we also assume Calcium ions with mass \(M = 40\) amu. The design leads to a Mexican-hat-shaped potential in the radial direction and a confining pseudopotential in the axial direction, as shown in figure 3. The trap potential has minimum at radius \(r = 58.0\) μm, height \(h = 385\) μm, leading to Mathieu-q parameters of \(q_r = 0.254\) and \(q_z = 0.255\) in the radial and vertical direction, respectively.

Design variation compensation. In view of fabrication imperfections, we study the effects of small deviation in the size of the center electrode. Our simulation figure 4 shows that the position of the potential minimum is very sensitive to the change in the size of the center electrode. In particular, it
shows that changes of $1 \mu m$ in radius will shift the radius of the minimum by $10 \mu m$. While we expect that microfabrication allows fabrication with tolerances below the micrometer range, we also can tune the potential by adding a small variable rf voltage with the same driving frequency on the second ring, but with the phase exactly opposite to that. Simulations show that the minimum position is shifted radially inward by about $2.5 \mu m V^{-1}$, while the trap depth changes by $0.005 eV V^{-1}$. This small compensation rf voltage provides a powerful tool for fine-tuning the potential in situ.

3. Structure of ring crystals

Of particular interest are the conditions under which ultracold ions form a ring in this potential. For this, we carry out molecular dynamics simulation to analyze the structure of laser-cooled ions in the surface trap [22]. We calculate the trajectories and velocities of the trapped ions by solving Newtonian equations of motion including the Coulomb interaction, the pseudoforce from the rf potential, and a hypothetical damping force term [23]. The extra damping term serves as a friction term that will gradually reduce the energy of the ions, thereby simulating laser cooling. Thus, the ions will eventually reach a steady state, which represents the expected structure of the cold ion crystal. The equations of motion of the $i$th ion can be written

$$m \frac{d^2 \vec{x}_i}{dt^2} = -\gamma \frac{d \vec{x}_i}{dt} + \vec{F}_r + \vec{F}_C.$$  (5)

Where $\gamma$ is the damping coefficient, $\vec{F}_r$ is the pseudoforce and the Coulomb force $\vec{F}_C$ is given by

$$\vec{F}_C = \frac{q^2}{4\pi \varepsilon_0} \sum_{i=1}^{N} \frac{\vec{r}_i - \vec{r}_j}{R_{ij}^3},$$  (6)

where $N$ is the number of ions in the trap.
Subsequently, the equation of motion is numerically solved by fourth-order Runge–Kutta method with a time step of 20 ns. To ensure that the numerical simulation finds the stable crystal configuration, we linearly ramp down the friction term during the simulation. We start with a damping coefficient $\gamma = 2 \times 10^{-18} \text{ kg s}^{-1}$, and turn it off linearly with time. As a result, our simulation shows that a ring crystal can be formed with up to 92 ions with the parameters and geometry discussed above (outer radius of the inner electrode 126 $\mu$m). As shown in figure 5(a), the 92-ion ring has diameter 116 $\mu$m and height 385 $\mu$m.

Keeping all parameters fixed, but adding one more ion yields a 93-ion ring crystal of two layers with about 1 $\mu$m separation in the plane perpendicular to the trap surface, and has a kink structure as a stable ground state, as shown in figure 5(b). Adding one more ion, we yield a similar 94-ion ring crystal with no kink, as shown in figure 5(c). We can study this phase transition from single-layer ion rings to double-layer ion rings by fine-tuning the trapping potential. This can be done by adjusting the compensation rf voltage on the second ring, as we have discussed in section 2. The kink dynamics in a ring might be an interesting subject in its own right [9]. Contrary to studies in linear traps [24–26], the kinks are in a homogeneous environment and cannot escape by just traveling to the edge of the ion crystal. Furthermore, working with an odd number of ions enforces the presence of an odd number of kinks and thus of at least one, while working with an even number of ions would lead to an even number of kinks.

4. Analysis of imperfections

Of particular interest in our work is to create ion crystals freely rotating even if their rotational energy is comparable to the groundstate energy [11, 12]. With with the criterion established in equation (1), i.e. the energy barrier created by the imperfections should be be smaller than $E_{\text{gap}}/2 = \sim 0.1 \text{ nK}$, we calculate the energy as a function of the angle when rotating the crystal around the symmetry axis. We study three sources of imperfections: a homogeneous electric field, irregularities on the edge of the electrodes, and the effect of a local charging (figure 6).
First we calculate the energy of the ion ring as a function of rotation angle in presence of an homogeneous electric field. The result will be a sinusoidal periodic function whose amplitude represents the classical energy barrier, which we denote $E_B$ (see figure 8). Figure 7 shows the energy barrier $E_B$ as a function of the applied field for 10, 20, 30, 40, and 50 ions. Figure 7 illustrates that the energy barrier is drastically suppressed when the number of ions is increased. Increasing the ion number reduces the ion–ion spacing which is expected to reduce the energy barrier further.

### 5. Laser cooling

The most noticable difference to cooling a ring as compared to linear strings of ions is the rotational degree-of-freedom. Hence, we focus only on the dynamics of the rotation and assume that all other modes are cooled to the standard...
is negative when the detuning $\Delta$ is negative. The $\cos \theta$ arises from the projection of the laser onto the rotational degree-of-freedom of each ion, $\delta_{\text{eff}} = \Delta - k \cdot v$, where $v$ is the velocity of the ion. The decay rate is given as $\Gamma = 1/(7\text{ ns})$ for Calcium, the wavenumber is defined as $k = 2\pi/(397\text{ nm})$, and $s$ is the saturation parameter [1]. In what follows, we assume a saturation parameter of $s = \frac{1}{2}$ and a gaussian beam profile with a beam waist of $200\ \mu m$. We chose a large beam waist to minimize the differences in intensity across the ring of ions and assume a detuning of $\Delta = -10\text{ MHz}$ for optimal cooling.

The Doppler cooling action of the laser on the moving ions will counterbalance the torque from the radiation pressure. If the torque is sufficiently small, the forces will cancel at some finite frequency of rotation. For 30 ions confined to a $R = 58\ \mu m$ ring, we calculate the equilibrium frequency and find a rotation frequency of approximately $1.04\ kHz$ per $1\ \mu m$ displacement of the cooling beam.

To counteract the rotation, we turn to the previous discussion of the effects of electric fields on the ring. By applying a strong electric field, we aim at creating a sufficiently strong energy barrier, thus stopping the ion ring from rotating. The maximum slope of the energy of the ion crystal as function of the rotation angle represents the torque needed to overcome the energy barrier and cause the ions to rotate around the ring

$$\tau_E = \left| \frac{dE}{d\theta} \right|_{\text{max}}.$$  \hfill (10)

Aiming at a static ion crystal, the torque from the cooling laser on a single ion is $\tau_L = R \times F_0$. Next, we look to find an applied electric field that will cancel the torque from the laser for this displacement. This condition can be written as

$$\sum_{i=1}^{N} \tau_L^i < \tau_E.$$  \hfill (11)

Where $\tau_E$ represents the sum over the torque of all ions by a given electric field, with the field orientation chosen to maximize the torque, see figure 10(b).

For ion positions in the trap given an electric field of $75\text{ V m}^{-1}$ optimally aligned against the propagation of the laser, the total sum of the torque from the laser alone is approximately linear for small laser displacements $d\theta$. Analyzing the situation for 30 ions and taking into account their calculated positions in the ion ring, we obtain $\tau = -5.476 \times 10^{-15}\text{ N m}$. The torque from the energy barrier for $75\text{ V m}^{-1}$, given by the maximum slope of the curve in figure 11, is $3.3 \times 10^{-20}\text{ N m}$, allowing for a displacement $d\theta$ of up to $6.0\ \mu m$ to achieve a static ion crystal.

6. Conclusions

We have studied a design of a planar trap providing trapping of a 92 ion-ring of diameter $116\ \mu m$ at height $385\ \mu m$ above its surface. This design can be fabricated using micro-
fabrication methods with high precision. We also studied the rotational motion under three symmetry-breaking imperfections: homogeneous electric field, irregularities of electrode edges from fabrication imperfections, and local charges placed on the trap electrodes. We have shown that the rotational energy barrier induced by these imperfections drastically reduces with an increasing number of ions in the ring. We thereby expect that the energy barrier from the imperfections can be reduced below the rotational ground state energy of large ion crystals. In addition, we have shown that laser alignment and strong homogeneous electric field of 75 V m$^{-1}$ can be utilized to pin and cool the ion ring for trapping and imaging.

Acknowledgments

This work is supported by the WM Keck Foundation. We acknowledge contributions of Anthony Ransford, Hao-kun Li, and Haggai Landa.

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