Experimental quantum simulations of many-body physics with trapped ions

Ch Schneider1,2, Diego Porras3 and Tobias Schäetz1,2

1 Max-Planck-Institut für Quantenoptik, Hans-Kopfermann-Straße 1, 85748 Garching, Germany
2 Albert-Ludwigs-Universität Freiburg, Physikalisches Institut, Hermann-Herder-Str. 3, 79104 Freiburg, Germany
3 Departamento de Física Teórica I, Universidad Complutense, 28040 Madrid, Spain
E-mail: tobias.schaetz@mpq.mpg.de

Received 24 August 2009, in final form 10 August 2011
Published 17 January 2012
Online at stacks.iop.org/RoPP/75/024401

Abstract
Direct experimental access to some of the most intriguing quantum phenomena is not granted due to the lack of precise control of the relevant parameters in their naturally intricate environment. Their simulation on conventional computers is impossible, since quantum behaviour arising with superposition states or entanglement is not efficiently translatable into the classical language. However, one could gain deeper insight into complex quantum dynamics by experimentally simulating the quantum behaviour of interest in another quantum system, where the relevant parameters and interactions can be controlled and robust effects detected sufficiently well. Systems of trapped ions provide unique control of both the internal (electronic) and external (motional) degrees of freedom. The mutual Coulomb interaction between the ions allows for large interaction strengths at comparatively large mutual ion distances enabling individual control and readout. Systems of trapped ions therefore exhibit a prominent system in several physical disciplines, for example, quantum information processing or metrology. Here, we will give an overview of different trapping techniques of ions as well as implementations for coherent manipulation of their quantum states and discuss the related theoretical basics. We then report on the experimental and theoretical progress in simulating quantum many-body physics with trapped ions and present current approaches for scaling up to more ions and more-dimensional systems.

(Some figures may appear in colour only in the online journal)
1. Introduction

Simulations and deeper understanding of the dynamics of some tens of interacting spins are already intractable with the most powerful classical computers. For instance, the generic state of 50 spin-1/2 particles is defined by $2^{50}$ numbers and to describe its evolution a $2^{50} \times 2^{50}$ matrix has to be exponentiated [1]. Recently, one of the ten most powerful supercomputers, JUGENE in Jülich, was exploring this regime. The current record was set by simulating a system of 42 quantum bits (qubits), equivalent to 42 spin-1/2 particles [2, 3].

In any case, it will not help to increase the impressive classical calculation capabilities to simulate only slightly larger quantum systems. Each doubling of the computational power will just allow the addition of one spin/qubit to the system (approximately after two years, according to Moore’s law [4]). Furthermore, simply pursuing this path of exponential growth in the computer’s classical capabilities would require an exponential shrinking of its electronic components [5]. The structure size currently amounts to approximately 30 nm, a distance spanned by roughly 100 atoms. If gifted engineers further miniaturize the sizes of their structures, ‘currents’ of a few electrons will ‘flow’ on ‘wires’ spanned by a few atoms only. As a consequence, quantum effects will have to be considered for future classical computers, leading to serious consequences. Electrons charging a capacitor, for example, currently realize a storage of logical information: a charged capacitor represents a ‘one’, a discharged capacitor a ‘zero’. What, if the few electrons, classically well caught within the potential of the capacitor, follow their natural quantum mechanical paths and simply escape through the walls by tunnelling?

However, allowing for quantum effects in a controlled way might also be exploited as a feature. Richard Feynman originally proposed [5] using a well controlled quantum system to efficiently track problems that are very hard to address on classical computers and named the device a ‘quantum computer’ (QC). Nowadays his proposal can be seen closer to the description of a quantum simulator (QS) [6]. In any case, his idea has been theoretically investigated and further developed to the concept of a universal QC. Fulfilling a well-defined set of prerequisites, known as diVincenzo’s criteria [6, 7], should make possible running any classical and quantum algorithm by a stroboscopic sequence of operations. These have to act on single qubits, for example, changing their state, and on pairs of qubits performing changes on one qubit, conditional on the state of its mate.

Hundreds of groups worldwide work on many approaches in different fields of atomic, molecular and solid-state systems to realize their version of the envisioned QC. For a concise review see, for example, [8].

However, even assuming an ideal system and perfect operations will require the control of the order of $10^3$ logical qubits as a basis for translating any algorithm or the quantum dynamics of a complex system into a sequence of stroboscopic gate operations on a potential universal QC [9]. Residual decoherence will cause computational errors and must be minimized to allow for high operational fidelities (∼99.999%–99.999%) [10]. At present, only then the errors could be overcome by quantum error correction, at the price of a reasonable but still tremendous overhead of ancilla qubits, approximately another 100 per logical qubit. In total, of the order of $10^5$ qubits are required. Even though there appear to be no fundamental obstacles for enhancing the fidelities of the operations and for scaling the size of the systems [11], there is still challenging technological development ahead. The realization of a universal QC is not expected within the next decades.

A shortcut via analogue QS has been taken into consideration [5] to allow deeper insight into the dynamics of quantum systems. ‘Analogue’ emphasizes that the dynamics of the system are not translated into an algorithm of gate operations on subsets of qubits. In contrast, a system of quantum particles is required, where (1) the initial state and its dynamics can be precisely controlled, (2) as many relevant parameters as possible manipulated and (3) the readout of the important characteristics of the final state performed in an efficient way. If the system’s evolution was governed by a Hamiltonian suspected to account for the quantum effects of interest, we would be able to experimentally investigate the physics of interest isolated from disturbances, close to Feynman’s original proposal. The requirements on the number of quantum particles and fidelities of operations for analogue QS are predicted to be substantially relaxed compared with QC [12]. However, it remains to be investigated which

---

4 The electronic components are arranged in two dimensions and, only recently, the third dimension is exploited. However, sufficient cooling has to be provided.

5 Depending on the context, the abbreviations ‘QC’ and ‘QS’ may also stand for quantum computation and quantum simulation, respectively.
realistic assumption on different sources of decoherence in the particular system will lead to a sufficiently small impact on the dedicated QS [13]. It is predicted that QS are less prone to decoherence, for example, in simulating robust effects such as quantum phase transitions (QPT). Therefore, they do not require any precautions in contrast to QC, which suffer from the costly overhead due to quantum error correction. It was even proposed to establish decoherence as an asset [1]. In this context, decoherence is not to be seen as a source of errors, as in the field of universal QC, but as a resource to simulate its natural counterpart. For example, decoherence is suspected to be responsible and required for enhanced efficiencies of (quantum) processes in biological systems at $T \sim 300$ K [14,15].

To discuss the different requirements for different analogue QS, we can distinguish between two categories of simulations. One category deals with problems where QS provide a simulated counterpart that allows intriguing questions that are not directly tractable in the laboratory to be experimentally addressed. Examples are highly relativistic effects such as Hawking and Unruh radiation or the zitterbewegung of a freely moving particle predicted by Dirac’s equation (see also [12]). The second category of simulations deals with objectives that are (probably fundamentally) not accessible with classical computation, for example, the complex quantum dynamics of spins in solid-state systems, as mentioned above. A promising strategy is to initialize an analogue QS in a state that can be prepared easily in the system of choice according to step (1) introduced above. Evolving the system adiabatically by changing its parameters according to (2) allows a new state to be reached that is hard or impossible to reach otherwise, for example, via a QPT. The aim here is not to simulate the effects including all disturbances and peculiarities, because the analogue QS would then become as complex as the system to be simulated. The aim can be to investigate whether the simplified model still yields the effects observable in nature and, thereby, to gain a concise deeper understanding of their relevant ingredients. However, there remains room for the important discussion as to whether the specific dynamics emulate nature or simulate the implemented model (Hamiltonian) and whether the results allow the drawing of further conclusions.

In any case, it has to be emphasized that analogue QS are intrinsically not universal. That is, different realizations of a QS will allow the simulation of different systems. Even more important, different approaches for the identical models (Hamiltonians) might allow cross-checking of the validity of the QSs [16].

There are several systems proposed to implement analogue QS, offering different advantages [12] to address the physics in many-body systems. One of them consists of neutral atoms within optical lattices [17–20]. Another promising candidate is based on trapped ions [21–23], originally suggested by Cirac and Zoller in 1995 [24] in the context of QC. Trapped ions already compete at the forefront of many fields, where ultimate accuracy and precision is required, such as metrology (see, for example, [25]). Trapped ions offer unique operational fidelities, individual addressability and short- as well as long-range interactions due to Coulomb forces.

Many models of both categories of QS are promising candidates or already addressed by trapped ions. Examples for the first category are emanating from the fields of cosmology [26–28], relativistic dynamics [29–34], quantum field theory [35], quantum optics [36] including quantum walks as a potential tool for QSs [15,37–40], chemistry [41], and biology [42,43]. For the second category, quantum spin Hamiltonians [21], Bose–Hubbard [22] and spin–boson [44] models were proposed to describe solid-state systems and their simulation would allow the observation and investigation of a rich variety of QPTs [45]. A summary and concise description of theoretical proposals on QS of both categories based on trapped ions and first experimental results up to the year 2008 can be found in [46].

This report aims to describe the current status of the field of experimental, analogue QS addressing many-body physics, its challenges and possible ways to address them. The first proof-of-principle experiment was achieved [47] and extended recently [48,49] on a few trapped ions in linear radio-frequency (RF) traps. The main challenge for QS remains to scale up towards 50–100 ions or even beyond. A simulated system of this size would reach far beyond the regime accessible via classical computation and, even more importantly, allow open scientific questions to be addressed.

The report is organized as follows. In section 2 we introduce the tools available for QS by briefly summarizing the types of traps, different ions species and different technical implementations of the control of the electronic and motional degrees of freedom. In section 3 we derive the mathematical description based on [21,50,51]. We aim at extending the existing formalism to be directly applicable to more dimensions and individual trapping conditions envisioned in arrays of ions. We apply this formalism to a basic building block of QC, a two qubit phase gate on the radial modes measured in our group, and emphasize similarities and differences between the application of similar operations for analogue QS. This section is supplemented by a detailed appendix. In section 4, we interpret the interactions in the context of analogue QS, which should be sufficient for understanding the subsequent discussion of the experimental implementations without going through the details of section 3. In section 5, we first depict the proof-of-principle experiments on a few trapped ions in linear RF traps. Based on the state-of-the-art capabilities we present in the second part of this section a summary of proposals to study many-body physics in a variety of solid-state systems. The two following sections are dedicated to two proposals aiming for scaling up the systems. In section 6, we discuss potential realizations of a two-dimensional array of RF surface-electrode traps. They are conceptually similar to promising approaches in Penning traps [50,52,53]. We will also introduce an alternative approach based on ions in optical traps in section 7, thus, trying to combine the advantages of trapped ions and optical lattices. Finally, we conclude in section 8.
to oscillate at frequencies directly. The time averaged confining pseudopotential allows the ion quadrupole electrodes (yellow circles) interacts with a charged atom magnitude smaller than $\omega_1/\Omega_1$ shift of the electronic levels of the atom or ion. The frequency applied via laser beams that provide an intensity dependent ac Stark of $k_B$, the minimum in the centre of the quadrupole (field lines indicated by black arrows). Typical depths of the pseudopotential are of the order of $k_B \times 10^4 \text{K}$. ($\omega_1/\Omega_1$) In optical traps, the optical field is typically applied via laser beams that provide an intensity dependent Stark shift of the electronic levels of the atom or ion. The frequency $\Omega_{\text{opt}}$ of the laser can be detuned blue (red) with respect to the relevant electronic resonance frequency $\omega_{\text{electron}}$ and therefore forces the atom/ion to seek low (high) fields. Typical depths of the pseudopotential are of the order of $k_B \times 10^{-4} \text{K}$.

2. Tools required for experimental quantum simulations

In this section, we describe the requirements to implement analogue QS based on trapped ions. Most of these tools have been developed over the last decades, many for the purpose of quantum information processing (QIP) with the main focus on QC.

2.1. Ion traps and Coulomb crystals

Isolating and trapping of individual particles as well as the precise control of their motional (external) degrees of freedom is key for many high precision measurements. Several trapping concepts have been developed for and implemented with ions, such as RF traps [54], Penning traps [55] and optical traps [56]. The physics of these devices, for example, of RF traps and optical dipole traps, is closely related. Electromagnetic multipole fields act on the charge or induce electric dipole moments. The resulting forces on the particles lead in time average to a confining pseudopotential. The two concepts are compared in figure 1.

However, there was a delay of more than a decade between trapping charged atoms in RF fields [57, 58] and trapping neutral particles with optical fields [59]. One explanation is that RF traps provide potential depths of the order of several eV $\approx k_B \times 10^4 \text{K}$, while optical traps typically store particles up to $k_B \times 10^{-3} \text{K}$ only. This discrepancy is mainly due to the comparatively large Coulomb force that RF fields can exert on charges. The RF field at typical frequencies $\Omega_{\text{RF}}/(2\pi) = 10 \text{MHz–100 MHz}$ directly acts on the massive ion. The related motional frequencies within the deep pseudopotential amount to a few MHz. Optical fields, in contrast, oscillate more than six orders of magnitude faster: too fast for the massive atomic core to follow. In a simplified picture, the optical field has to induce a dipole moment of the electron and the atomic core to follow. In a simplified picture, the optical field has to follow. In a simplified picture, the optical field has to follow.

Here we focus first on ions in linear RF traps. The concept for the radial confinement is depicted in figures 1(a) and 2. The RF field applied to two opposing electrodes of the quadrupole can provide a radially confining pseudopotential. Similar to a quadrupole mass filter, one can find voltages for given parameters (electrode geometry and mass/charge ratio of the ion species) that allow for stable confinement in two dimensions. Additional dc voltages add a static harmonic potential to complete the three-dimensional confinement that can be assumed to be harmonic. Dependent on the application, these dc voltages can be applied to electrodes realized as rings or needles along the axis or by a segmentation of the quadrupole electrodes (see figure 2(b)). A confined ion will oscillate with frequency $\omega_Z/(2\pi)$ along the trap axis and with frequencies $\omega_X/Y/(2\pi)$ in the radial directions. The radial oscillation is superimposed by a fast oscillation at frequency $\Omega_{\text{RF}}/(2\pi)$ (so-called micromotion), which increases with increasing distance of the ion from the trap centre, such that the RF field does not vanish anymore.

Typical parameters for conventional setups are a minimal ion–electrode distance $h \sim 100 \mu \text{m}$–$1000 \mu \text{m}$ allowing for RF voltages of the order of 1000 V.

Different laser cooling schemes can be applied to reduce the total energy of motion of the ion [51]. Doppler cooling [61–64] of several ions already allows a regime to be entered, where the kinetic energy $(k_B T \sim \text{mK})$ of the ions becomes significantly smaller than the energy related to the mutual Coulomb repulsion. Hence, the ions cannot exchange their position anymore. A phase transition from the gaseous (liquid)
plasma to a crystalline structure occurs [65, 66]. On the one
hand, the resulting Coulomb crystals (see figure 3) provide
many similarities with solid state crystals already partially
explaining why Coulomb crystals appear naturally suited to
simulate many-body physics. (1) The ions reside on individual
lattice sites. (2) The motion of the ions (external degree
of freedom) can be described easiest in terms of common
motional modes with the related quanta being phonons. The
phonons in Coulomb crystals allow long-range interactions to
be mediated between the spins associated with the ions. In
a different context, the phonons can also be interpreted as
bosonic particles, for example, capable of tunnelling between
lattice sites simulated by the ions (see also section 5.2). On
the other hand, there are advantageous differences compared
with solid-state crystals. (3) Coulomb crystals typically build
up in ultra-high vacuum ((10^{-9} – 10^{-11}) mbar) and are very
well shielded against disturbances from the environment, thus
providing long coherency times. (4) Coulomb crystals feature
lattice constants of a few micrometres (see figure 3), dependent
on the trapping potential counteracting the mutual Coulomb
repulsion. Compared with a solid, where distances are of the
order of Ångstroms (10^{-10} m) the density of the structure in
one dimension is reduced by five, in three dimensions by fifteen
orders of magnitude. This allows for individual addressing of
the ions and for individual preparation, control and readout
of their electronic and motional states. (5) The Coulomb
interaction between the charged ions is not shielded within the
crystal, as in Coulomb crystals the charge of all ions has the
same sign in contrast to ionic crystals in solid-state systems.
However, it should be mentioned that, as opposed to quantum
solids, the quantum statistics of the ions is not relevant due to
the suppression of the mutual exchange processes.

It has to be pointed out that it is possible to
deterministically achieve phase transitions between different
structures of Coulomb crystals for large numbers of ions
[65, 66]. When the ratio of radial to axial confinement is
reduced or the number of confined ions is increased, we
observe the transition from a linear chain of ions via a two-
dimensional zigzag structure to a three-dimensional structure
(see figures 3(b)–(d)).

Despite the unique conditions in Coulomb crystals in
linear RF traps and the high fidelities of operations, current
experimental approaches to QS (and QC) are still limited
to a small number of ions. The approaches include of the
order of ten ions arranged in a linear chain [49, 67]. This
is accomplished by choosing the radial confinement much
stronger than the axial one. The linear chain orients along
the weakest (Z) direction, where tiny oscillations of the cooled
ions around the minimum of the pseudopotential (X and Y) and
thus micromotion still remains negligible.

For the purposes of a QC and QS, scaling to a larger
number of spins and more dimensions while keeping sufficient
control over all required degrees of freedom remains the
challenge of the research field. Using longer linear chains
confined in anharmonic axial potentials [68] might provide a
way to reach a number of ions in the system that in principle
already exceeds capabilities of a classical supercomputer.
Another way might be the use of RF ring traps offering periodic
boundary conditions for static Coulomb crystals [65, 66] and
even (more-dimensional) crystalline beams of ions [69–71].
A microfabricated ring trap is currently being developed and
fabricated at Sandia National Laboratories [72].

The two main limitations for further scaling of the number
of ions in a common potential, from a practical point of view,
are (1) the emergence of 3N normal modes for N ions plus their
sum and difference frequencies that lead to an increasingly
crowded phonon spectrum (already for each spatial dimension
separately). Individual spectral components become difficult
to identify and off-resonant couplings to ‘spectator’ transitions
[50] are hard to avoid. However, under certain conditions, QSs
are predicted to allow for coupling to all modes simultaneously,
see for example [21]. (2) QSs based on ions in large,
more-dimensional Coulomb crystals suffer from additional
challenges, for example, intrinsic micromotion (due to the
displacement from the minimum of the pseudopotential), an
inhomogeneous ion spacing (due to space charge effects) and
the coupling between modes of all three spatial dimensions.

One approach for scalability might be to generate a spin-
off from the QIP community based on their new concept
of a surface-electrode geometry for RF traps [73, 74] (see
figure 14). Currently, this design is tested with the aim to allow
for networks of interconnected linear traps. This constitutes
a promising possibility to realize the multiplex architecture of

Figure 3. Fluorescence images of laser-cooled ions in a common
confining potential of a linear RF trap (see figure 2), forming
differently structured Coulomb crystals. (a) A single ion (Mg\textsuperscript{+}).
(b) A linear chain of 40 ions at \(\omega_{X/Y} \gg \omega_Z\). The axis of the chain
coincides with the trap Z-axis, which is identically orientated in the
rest of the images. (c) A linear chain embedding a two-dimensional
zigzag structure of 60 ions for \(\omega_{X/Y} > \omega_Z\). (d) A three-dimensional
structure of more than 40 ions at \(\omega_{X/Y} \gtrapprox \omega_Z\). The enhanced
signal-to-noise ratio in (d) is achieved by extended exposure.
Structural phase transitions can be induced between one-, two-
and three-dimensional structures, for example by reducing the ratio
of radial to axial trapping frequencies.
memory and processor traps for universal QC [75]. However, for QS we need a miniaturized array of traps allowing for more-dimensional interactions, as discussed in section 6.

It has to be emphasized that there are other concepts for trapping ions, a prominent one being Penning traps. Penning traps provide trapping potentials of similar parameters as RF traps. A strong, static magnetic field and a dc electric field yield a stable confinement of large, rotating Coulomb crystals. Storing many cool ions in a Penning trap, naturally provides, for example, a large triangular lattice of ions [76–78] that is also predicted to be well suited for QS [79]. Promising results are on their way [52, 53]. Another challenging proposal for QS involves trapping ions optionally simultaneously with atoms in optical lattices [80] (see section 7).

2.2. Ions

A large variety of different atomic ions have already been used for the purpose of QIP. Every ion or, more specifically, every isotope has different properties, for example, regarding the level scheme or the charge–mass ratio, and thus can meet different requirements of a QS. However, they all have a single valence electron leading to an alkali-like level scheme. Most prominent are the earth alkali ions Be\(^+\), Mg\(^+\), Ca\(^+\), Sr\(^+\) and Ba\(^+\). A similar electronic structure has Zn\(^+\), Cd\(^+\) and Hg\(^+\), followed by Yb\(^+\) [81].

Typically, two electronic levels with sufficiently long coherence times are chosen as qubit or spin states \(|\downarrow\rangle\) and \(|\uparrow\rangle\), respectively. (In principle, however, the restriction to two states is not required and the use of up to 60 states has been proposed for (neutral) holmium [82].) The types of qubits can be divided into two classes: in optical qubits, the states are encoded in two states with a dipole-forbidden transition at an optical frequency. An example is \(^{40}\text{Ca}^+\) with \(|\downarrow\rangle := |S_{1/2}\rangle\) and \(|\uparrow\rangle := |D_{5/2}\rangle\). The lifetime of \(|\uparrow\rangle\) is on the order of 1 s, which defines the upper bound for its coherence time. In hyperfine/Zeeman qubits, two sublevels from the ground state manifold are chosen as \(|\downarrow\rangle\) and \(|\uparrow\rangle\). An applied magnetic field lifts the degeneracy within the manifolds of electronic levels to allow for spectrally resolving the dedicated states. The states of hyperfine/Zeeman qubits have extremely long lifetimes and coherence times on the order of minutes have been observed [83, 84]. As an example for a hyperfine/Zeeman qubit, an excerpt of the level scheme of \(^{25}\text{Mg}^+\) is shown in figure 4. The transition frequencies in hyperfine/Zeeman qubits are in the microwave regime.

2.3. Basic operations

The quantized oscillation of the ions in the harmonically approximated potential of the trap gives rise to motional states, which are typically expressed in terms of Fock states \(|n\rangle\). Independent of the choice of qubit we will require three different types of couplings to electronic states and/or motional states to assemble the toolbox for QC and QS based on trapped ions (for details see section 3).

(a) Coupling of the electronic states only \(|\downarrow\rangle|n\rangle \Leftrightarrow |\uparrow\rangle|n\rangle\).

This operation can be used to implement Rabi flops between the electronic states and serves as a one-qubit gate of a potential QC. In the context of QS it can be interpreted as simulated magnetic field (see also section 3.2).

(b) Coupling of the electronic and motional states \(|\downarrow\rangle|n\rangle \Leftrightarrow |\uparrow\rangle|n\rangle\).

This operation can drive Rabi flops between electronic states and different motional states, for example on the first red state \((n' = n - 1)\) or blue sideband \((n' = n + 1)\) (see also section 3.2). It can be used to create entanglement between the electronic and motional states and is an important ingredient for both sideband cooling and the readout of the motional state (see below).

(c) State-dependent forces (for example, \(|\downarrow\rangle|n\rangle \rightarrow |\downarrow\rangle|n + 1\rangle\)). These forces lead to state-dependent displacements. They can be used for conditional interactions between multiple ions, which are exploited for quantum gates (see sections 3.3 and 3.4) or effective spin–spin interactions in the simulation of quantum spin Hamiltonians (see section 3.5).

Operations (a)–(c) can be realized for both classes of qubits in the optical regime and for hyperfine/Zeeman qubits additionally via microwave fields [50, 51].
spin–spin interaction [21, 47].

Simulation of the quantum Ising Hamiltonian to create the effective Hamiltonians. (interactions can be used for single-qubit gates in QC and to simulate transitions driven by a pair of laser beams (shown without motional implemented, for example, by two-photon stimulated-Raman geometric phase gate for the displacement pulse [110, 123] orientation the frequency of a motional mode. This interaction is used in the form of an interaction with a single beam of frequency \( \omega \) can be substituted by two beams with frequencies \( \omega_1 \) and \( \omega_2 \), sections 3.3).

For operation (a), the frequency \( \omega_I \) has to (approximately) meet the transition frequency between \( |↓⟩ \) and \( |↑⟩ \) becomes site-dependent due to position-dependent Zeeman shifts. The ions can be individually addressed by applying microwave fields with these site-dependent frequencies \( \omega_l \). In addition, this causes state-dependent forces as in the Stern–Gerlach experiment and allows for coupling to the motional modes. The main challenge here is to provide sufficiently large magnetic field gradients and to cope with state-dependent transition frequencies, if high operational fidelities are required (see, for example, [90]).

As an alternative to the static magnetic field gradients, alternating magnetic fields due to microwave currents in electrodes of surface-electrode traps (see section 6) have been proposed [91] and first promising results have been achieved [92, 93]. Due to the small height of the ion above the electrode in this type of trap, a sufficiently large ac Zeeman shift can be generated, which can be treated analogously to the ac Stark shift created by laser beams in two-photon stimulated-Raman transitions discussed above. However, the small height above the electrodes leads to further challenges (compare section 6) and high microwave powers are required.

Coupling via optical fields. Optical qubit states can be linked by a single, nearly resonant laser beam with frequency \( \omega_l \) and wave vector \( \vec{k}_l \). Due to the lifetime of the \( |↑⟩ \) state of \( 1/\Gamma \sim 1 \text{ s} \), the linewidth of the laser has to be very narrow (\( \sim 1 \text{ Hz} \)). Operations (a) and (b) can be implemented directly (see section 3.2).

State-dependent forces (see type (c) in the text) can be provided by a bichromatic light field (see, for example, [85–87] and also section 3.3).

In hyperfine/Zeeman qubits, the single laser beam can be substituted by two beams with frequencies \( \omega_1 \), \( \omega_2 \) and wavevectors \( \vec{k}_1 \), \( \vec{k}_2 \) driving two-photon stimulated-Raman transitions. The beams are detuned by \( \Delta \gg \Gamma \) from a third level, for example, a P level (compare figure 4) with a typical lifetime \( 1/\Gamma \) of the order of few nanoseconds. In the mathematical treatment, this third level can be adiabatically eliminated for large detunings and the interaction gains the form of an interaction with a single beam of frequency \( \omega_l = |\omega_1 - \omega_2| \) and wavevector \( \vec{k}_l = \vec{k}_1 - \vec{k}_2 \). The requirement of a narrow linewidth holds only for the difference frequency \( \omega_l \), which can be fulfilled comparatively easily: the two beams can be generated from the same laser using acousto-optical modulators driven by a stable microwave source, while the requirements on the frequency stability of the laser are relaxed. For operation (a), the frequency \( \omega_l \) has to (approximately) meet the transition frequency of the qubit states (see figure 5(a)). For operation (b), \( \vec{k}_l \) in addition must not vanish to achieve a sufficient momentum transfer to the ions, see figure 4 and section 3.2. Therefore, the two beams are typically orthogonal (\( |\vec{k}_l| \approx \sqrt{2} |\vec{k}_1| \)) or counter-propagating (\( |\vec{k}_l| \approx 2 |\vec{k}_1| \)). The state-dependent forces (c) can be implemented by nearly resonant beams (\( \omega_l \approx 0 \)) and beam geometries as for operation (b) (see figure 5(b), sections 3.3–3.5 and section 4).

The main technical drawback of using two-photon stimulated-Raman transitions is decoherence due to spontaneous emission after off-resonantly populating the third level. This limitation can be mitigated by increasing the detuning \( \Delta \) and the intensities \( I_{1/2} \) of the beams, since the interaction strength scales with \( I_{1/2}/\Delta \), while the spontaneous emission rate scales with \( I_{1/2}/\Delta^2 \).

Coupling via microwave fields. Alternatively, transitions between the electronic states in hyperfine/Zeeman qubits can be driven laser-less by microwave fields. This allows the direct realization of operation (a). However, due to the comparatively long wavelength and the related small momentum transfer (\(|\vec{h}\vec{k}_l| \rightarrow 0 \)), only negligible coupling to the motional modes can be achieved directly and additional efforts are required to provide operations (b) and (c) [88, 89]: by applying a static magnetic field gradient along the axis of an ion chain, the transition frequency between \( |↓⟩ \) and \( |↑⟩ \) becomes site-dependent due to position-dependent Zeeman shifts. The ions can be individually addressed by applying microwave fields with these site-dependent frequencies \( \omega_l \).

In this type of trap, a sufficiently large ac Zeeman shift can be generated, which can be treated analogously to the ac Stark shift created by laser beams in two-photon stimulated-Raman transitions discussed above. However, the small height above the electrodes leads to further challenges (compare section 6) and high microwave powers are required.

2.4. Initialization and readout

Initialization of motional and electronic states. The initialization into one of the qubit states, for example \( |↓⟩ \), can be achieved with near-unity efficiency by optical pumping [94]. Regarding the motional modes, the initialization includes Doppler cooling in all three dimensions leading to a thermal state with an average phonon number \( \bar{n} \) of typically a few to ten quanta. This pre-cooling is required to reach the Lamb–Dicke regime (see section 3.2), where subsequent resolved sideband cooling [95–97] or cooling utilizing electromagnetically induced transparency can be applied [98, 99]. These cooling schemes lead close to the motional ground state \( |0⟩ \) (\( n = 0 \) with probability of 98% in [97]) of the dedicated modes.

Readout of electronic and motional states. We distinguish the two electronic states by observing state-dependent laser fluorescence. The dipole allowed transition to an excited state starting in the state \( |↓⟩ \) is driven resonantly (see the transition labelled ‘BD’ in figure 4) in a closed cycle completed by spontaneous emission back to the state \( |↓⟩ \) due to selection rules. For state \( |↑⟩ \) the detection laser is off-resonant. The ion

Figure 5. Implementations of different interaction types for hyperfine/Zeeman qubits. (a) An operation of type (a) can be implemented, for example, by two-photon stimulated-Raman transitions driven by a pair of laser beams (shown without motional dependence) or directly by a microwave field. These types of interactions can be used for single-qubit gates in QC and to simulate effective Hamiltonians in the simulation of the quantum Ising Hamiltonian. (b) State-dependent forces (see type (c) in the text) can be created by two beams detuned by approximately the frequency of a motional mode. This interaction is used in the geometric phase gate for the displacement pulse [110, 123] or in the simulation of the quantum Ising Hamiltonian to create the effective spin–spin interaction [21, 47].

\[ |↓⟩ \rightarrow |↑⟩ \]

\[ |↑⟩ \rightarrow |↓⟩ \]
The following calculations (section 3.1 and 3.2) are the theoretical excursion. Therefore, it appears 'bright' for quantum computing and derived from the result of the spin state detection described above [109].

3. Theoretical excursion

The following calculations (section 3.1 and 3.2) are the mathematical description of the toolbox that is available for both QCs and QSSs. Detailed discussions can be found, for example, in [30] or [51]. We summarize important equations in the following and extend the mathematical description to be applicable to scaled approaches of QSSs, for example, two-dimensional arrays of ions in individual traps. We will continue with the description of an implementation of the effective spin–spin interaction for ions appearing in quantum spin Hamiltonians. In order to investigate it isolated from other interactions, we will first discuss it from the point of view of quantum gates [110] in sections 3.3 and 3.4. Finally, in section 3.5, we use all tools to derive and discuss the quantum Ising Hamiltonian based on [21] as an example. The mathematical descriptions will be required to pursue proposals described in section 5.2.

This section aims at deriving the mathematical treatment of the simulation of a quantum Ising Hamiltonian. It underlines the approximations and transformations applied in these calculations and discusses the related corrections compared with the ideal quantum Ising model, which grow in importance in regard to scaled systems described in section 6. Readers who are more interested in a pictorial description of the simulation of a quantum Ising Hamiltonian may skip this section and continue to section 4.

3.1. Theoretical basics

In the following we consider two-level systems only. The Hamiltonian describing the energy of the electronic states of \( N \) such systems is given by

\[
\hat{H}_e = \sum_{i=1}^{N} \frac{\hbar \omega_1}{2} \hat{a}_i^\dagger \hat{a}_i + \frac{N \hbar \omega_{\mathrm{I}} + \omega_{\mathrm{J}}}{2},
\]

(3.1)

where \( \hbar \omega_{1/4} \) denote the energies of the states \( |\downarrow\rangle \) and \( |\uparrow\rangle \), respectively, \( \omega_{\mathrm{I}} := \omega_{\mathrm{I}} - \omega_{\mathrm{J}} \), and the operator \( \hat{a}_{i}^{\dagger} \hat{a}_{i} \) the Pauli operator (compare appendix B, equation (B.1)) acting on the \( i \)th ion.

The ions are considered to be trapped in a common harmonic potential or several individual potentials, which can be approximated to harmonic order. The corresponding Hamiltonian in terms of the normal modes of the oscillation reads

\[
\hat{H}_m = \sum_{m=1}^{3N} \hbar \omega_m \left( \hat{a}_m^\dagger \hat{a}_m + \frac{1}{2} \right).
\]

(3.2)

Here, \( \hat{a}_m^\dagger \) and \( \hat{a}_m \) are the annihilation and creation operators of the \( m \)th mode, respectively, and \( \omega_m \) the corresponding frequency. In the following, the constant terms \( \hbar \omega_{m/2} \) will also be omitted and the abbreviation \( \bar{H}_0 := \bar{H}_e + \bar{H}_m \) will be used.

An interaction of an ion with the electric field \( \vec{E} \) of a laser beam is described by \( -\vec{\mu} \cdot \vec{E}(\vec{r}, t) \), where \( \vec{\mu} \) denotes the electric dipole operator for the transition \( |\downarrow\rangle \leftrightarrow |\uparrow\rangle \) and \( \vec{E}(\vec{r}, t) \) the field at the site of the ion. The Hamiltonian describing the interaction of the field with \( N \) ions becomes

\[
\hat{H}_{\mathrm{I}} = \sum_{i=1}^{N} \hbar \Omega^{(i)}_\mathrm{I} \left( e^{i(\bar{K}_1^{(i)} x^{(i)} - \omega_{\mathrm{I}} t + \phi^{(i)})} + \text{h.c.} \right) \cdot \vec{\mu}^{(i)}.
\]

(3.3)

Here, \( \Omega^{(i)}_\mathrm{I} = -\mu E^{(i)}/2 \in \mathbb{R} \) is the interaction strength at site \( i \), \( \bar{K}_1^{(i)} \) the wavevector at site \( i \), \( x^{(i)} \) the position of the \( i \)th ion, \( \omega_{\mathrm{I}} \) the frequency of the field and \( \phi^{(i)} \) an additional phase.

In the most general form, the operator \( \vec{\mu}^{(i)} \) can be expressed as a linear combination of Pauli operators \( \vec{\sigma}^{(i)/z} \) and the identity operator \( 1^{(i)} \) (see appendix B):

\[
\vec{\mu}^{(i)} := 0 \cdot 1^{(i)} + \alpha_x \sigma_x^{(i)} + \alpha_y \sigma_y^{(i)} + \alpha_z \sigma_z^{(i)},
\]

(3.4)

with the prefactors \( \alpha_x, \alpha_y, \alpha_z \in \mathbb{R} \), which are determined by the polarization of the electric field and angular momenta of the states encoding \( |\downarrow\rangle \) and \( |\uparrow\rangle \) (see also section 3.4 for some examples).

The position operator \( \vec{r}^{(i)} \) in equation (3.3) is decomposed into the equilibrium position \( \vec{x}_0 \) and the displacement \( \vec{x}^{(i)} = \vec{r}^{(i)} - \vec{x}_0 \). The terms \( \vec{k}_1^{(i)} \cdot \vec{x}_0 \) give rise to a constant phase, which we absorb into \( \phi^{(i)} \) and \( \vec{k}_1^{(i)} \cdot \vec{x}^{(i)} \rightarrow \phi^{(i)} \).

The displacement of the ion from its equilibrium position \( \vec{x}^{(i)} \) is expressed in terms of the normal modes of motion

\[
\vec{\xi}^{(i)} := \sum_{m=1}^{3N} \left( b_{m,i} \vec{q}_m \cdot \vec{c}_X + b_{m,i+N} \vec{q}_m \cdot \vec{c}_Y + b_{m,i+2N} \vec{q}_m \cdot \vec{c}_Z \right),
\]

(3.5)

where \( b_{m,i} \) are the elements of an (orthogonal) transformation matrix (compare appendix A, equation (A.8)). Expressing the operators \( \vec{q}_m \) of the normal modes in terms of the creation and annihilation operators yields

\[
\vec{q}_m = q_{m0} \left( \hat{a}_m + \hat{a}_m^\dagger \right)
\]

(3.6)

with \( q_{m0} \) denoting the scale factor. Hence, the scalar product appearing in the Hamiltonian yields

\[
\vec{k}_1^{(i)} \cdot \vec{\xi}^{(i)} = \sum_{m=1}^{3N} \eta_{m}^{(i)} (\hat{a}_m + \hat{a}_m^\dagger),
\]

(3.7)

where the Lamb–Dicke parameter of the \( m \)th mode and \( i \)th site has been introduced:

\[
\eta_{m}^{(i)} := q_{m0} \left( b_{m,i} \vec{k}_1^{(i)} \cdot \vec{c}_X + b_{m,i+N} \vec{k}_1^{(i)} \cdot \vec{c}_Y + b_{m,i+2N} \vec{k}_1^{(i)} \cdot \vec{c}_Z \right).
\]

(3.8)
To summarize, the interaction term of the Hamiltonian gains the form
\[ \hat{\mathcal{H}}_I = \sum_{i=1}^{N} \hbar \Omega_i^{(i)} \left( \exp \left[ i \sum_{n=1}^{N} \eta_n^{(i)} (\hat{a}_n + \hat{a}_n^\dagger) - \omega t + \phi_n^{(i)} \right] + \text{h.c.} \right) \hat{\sigma}^{(i)}. \]
(3.9)

The transformation into the interaction picture
\[ \hat{\mathcal{H}}'_I = \hat{U}_0^\dagger \hat{\mathcal{H}}_I \hat{U}_0 \quad \text{with} \quad \hat{U}_0 = e^{-i\hat{\sigma}_0 \hat{\sigma}_z / \hbar}, \]
(3.10)
can be carried out for each site separately, hence
\[ \hat{\mathcal{H}}^{(i)}'_I = \hat{U}^{(i)}_I \hat{H}^{(i)}_I \hat{U}^{(i)\dagger}_I \quad \text{with} \quad \hat{U}^{(i)}_0 = e^{-i\hat{\sigma}_0 \hat{\sigma}_z / \hbar}, \]
(3.11)
where \( \hat{\mathcal{H}}^{(i)}_I \) is the Hamiltonian corresponding to the \( i \)th ion and \( \hat{H}^{(i)}_I = \sum_{n=1}^{N} \hat{H}^{(i)}_n \).

The operator \( \hat{\kappa}^{(i)} \) (equation (3.4)) related to the electronic states reads as follows in the interaction picture (compare equation (B.15)):
\[ \hat{\kappa}^{(i)} = e^{i\beta t}\hat{\sigma}^{(i)}_z / 2 \hat{\kappa}^{(i)} e^{-i\beta t}\hat{\sigma}^{(i)}_z / 2 \]
(3.12)
\[ = \frac{1}{2} \left( \alpha s^{(i)} I + (\alpha_1 + \alpha_2 / i) e^{i\omega t} \hat{\sigma}^{(i)} + \alpha_3 \hat{\sigma}^{(i)}_z + \text{h.c.} \right). \]
(3.13)
Here, we introduced \( \hat{\sigma}^{(i)} = \hat{\sigma}^{(i)}_x + i \hat{\sigma}^{(i)}_y \) and \( \hat{\sigma}^{(i)} = \hat{\sigma}^{(i)}_x - i \hat{\sigma}^{(i)}_y \).

The terms in equation (3.9) containing the motional operators transform as follows (compare equation (C.10)):
\[ e^{i\beta t}\hat{\sigma}^{(i)}_z \hat{a}_n e^{-i\beta t}\hat{\sigma}^{(i)}_z \hat{a}_n = e^{i(\beta t + \phi_1^{(i)})} \left( \hat{a}_n e^{i\beta t} \hat{\sigma}^{(i)}_z + \hat{a}_n^\dagger e^{-i\beta t} \hat{\sigma}^{(i)}_z + \text{h.c.} \right). \]
(3.14)
Hence, it yields the following expression for the complete Hamiltonian in the interaction picture:
\[ \hat{\mathcal{H}}^{(i)}'_I = \hbar \Omega_i^{(i)} \left( \exp \left[ i \sum_{m=1}^{3N} \eta_m^{(i)} (\hat{a}_m e^{-i\omega t} + \hat{a}_m^\dagger e^{i\omega t}) - \omega t + \phi_1^{(i)} \right] + \text{h.c.} \right) \hat{\kappa}^{(i)} \hat{\sigma}^{(i)} + \text{h.c.} \]
(3.15)
At this point fast rotating terms which average out on short timescales are neglected (rotating wave approximation, RWA). For \( \Omega_i^{(i)} \ll \omega, \phi \), we distinguish between two cases: In the first case, \( \omega \ll \omega \); terms containing \( e^{i\omega t} \) are neglected (see figure 5(b) for an example of an implementation). (If \( \omega_1 = \omega_2 = 0 \), nothing will change and the Hamiltonian will still be exact.) In the second case, \( |\omega - \omega_1| \ll \omega_1 \), all terms but \( e^{i(\omega t - \omega n)\hat{\sigma}} \) are neglected (see figure 5(a)):
\[ \hat{\mathcal{H}}^{(i)(RWA)}_I = \hbar \Omega_i^{(i)} \exp \left[ i \sum_{m=1}^{3N} \eta_m^{(i)} (\hat{a}_m e^{-i\omega t} + \hat{a}_m^\dagger e^{i\omega t}) - \omega t + \phi_1^{(i)} \right] + \text{h.c.} \]
(3.16)
for \( \omega \ll \omega_1 \).
\[ \hat{\mathcal{H}}^{(i)(RWA)}_I = \frac{\hbar}{2} \Omega_i^{(i)} \exp \left[ i \sum_{m=1}^{3N} \eta_m^{(i)} (\hat{a}_m e^{-i\omega t} + \hat{a}_m^\dagger e^{i\omega t}) - (\omega - \omega_1) t + \phi_1^{(i)} \right] \left( \alpha_1 + \frac{\alpha_2}{i} \right) \hat{\sigma}^{(i)} + \text{h.c.} \]
(3.17)
for \( |\omega - \omega_1| \ll \omega_1 \).

### 3.2. \( \hat{\sigma}_z/\hat{\sigma}_y \) interaction

The time evolution corresponding to \( \hat{\mathcal{H}}^{(i)(RWA)}_I \) in equation (3.17) is involved. The time evolution is calculated for a single ion \( i \) and a single motional mode \( m \), for example, in [50, 51]. As some simplifications (Lamb–Dicke regime, see below) are not always justified for experiments, we will summarize this calculation here. (The index \( m \) is skipped in this section.)

In this case the Hamiltonian simplifies to
\[ \hat{\mathcal{H}}^{(RWA)}_I = \frac{\hbar}{2} \Omega_i \exp \left[ i(\eta_\alpha (\hat{a}_\alpha e^{-i\omega_1 t} + \hat{a}_\alpha^\dagger e^{i\omega_1 t}) - (\omega_1 - \omega_1 t + \phi_1) \right] \times \left( \alpha_1 + \frac{\alpha_2}{i} \right) \hat{\sigma}_z + \text{h.c.} \]
(3.18)

Writing the state vector in the basis of electronic states \( |s\rangle \) and motional Fock states \( |n\rangle \),
\[ |\psi(t)\rangle = \sum_{s \in \{\uparrow, \downarrow\}} \sum_n c_{s,n}(t) |s, n\rangle, \]
(3.19)
the Schrödinger equation yields
\[ i\hbar \dot{c}_{s,n}(t) = \sum_{s' \in \{\uparrow, \downarrow\}} \sum_n \langle s', n' | \hat{\mathcal{H}}^{(RWA)}_I | s, n \rangle c_{s', n}(t). \]
(3.20)

Matrix elements of the Hamiltonian vanish for \( s' = s \). We obtain for the non-vanishing matrix elements [95, 111] (compare appendix D)
\[ \langle \uparrow, n' | \hat{\mathcal{H}}^{(RWA)}_I | \downarrow, n \rangle = \frac{\hbar}{2} \Omega_i \exp \left[ -i(\omega_1 - \omega_1 t + \phi_1) \right] \left( \alpha_1 + \frac{\alpha_2}{i} \right) \]
(3.21)
\[ \times \langle n' | \hat{D} (i\eta_\alpha e^{i\omega_1 t}) | n \rangle \langle \uparrow | \hat{\sigma}_z | \downarrow \rangle \]
\[ = \hbar \Omega_{\alpha} | n' \rangle \langle n' | \exp \left[ i(\omega_1 - \omega_1 t + \phi_1) \right] \]
where \( \hat{D}(\lambda) := \exp e^{i\lambda \hat{\sigma}_z} \) denotes the displacement operator and
\[ \Omega_{\alpha} := \Omega_i e^{-\eta_\alpha^2 / 2} \sum_n \left( \frac{1}{n-\eta_\alpha^2} \right) L_{n-\eta_\alpha^2}(\eta_\alpha^2). \]
(3.22)
Here, \( L_n^{\{a\}}(x) \) are the associated Laguerre polynomials, \( n := \min(n', n) \), and \( n' := \max(n', n) \). Analogously, we obtain \( \langle \downarrow, n | \hat{\mathcal{H}}^{(RWA)}_I | \uparrow, n' \rangle = \langle \downarrow, n' | \hat{\mathcal{H}}^{(RWA)}_I | \downarrow, n' \rangle \).

We define \( \delta := (\omega_1 - \omega_1 t) - (n-n') \omega \). For small detunings \( |\delta| \ll \omega \) and interaction strengths \( |\Omega_{\alpha}| \ll \omega \) (resolved sideband regime), we apply an RWA neglecting terms rotating faster than \( e^{i\omega t} \). Equation (3.20) can then be solved for each subset \( [\uparrow, n'] \) and \( [\downarrow, n'] \) separately:
\[ c_{\uparrow,n}(t) = -i\Omega_{\alpha} | n' \rangle \langle n' | \left( \alpha_1 + \frac{\alpha_2}{i} \right) e^{i(\omega_1 n' - \omega_1 t + \phi_1)} c_{\downarrow,n}(t). \]
(3.23)
\[ c_{\downarrow,n}(t) = -i\Omega_{\alpha} | n' \rangle \langle n' | \left( \alpha_1 + \frac{\alpha_2}{i} \right)^* e^{-i(\omega_1 n' - \omega_1 t + \phi_1)} c_{\uparrow,n}(t). \]
(3.24)

The solution of the system of differential equations yields Rabi oscillations between the states \( |\uparrow, n'\rangle \leftrightarrow |\downarrow, n'\rangle \) (compare appendix E):
\[
\left( \begin{array}{c}
 c_{\uparrow,n}(t) \\
 c_{\downarrow,n}(t)
\end{array} \right) = \left[ \begin{array}{c}
 \cos(X_{\uparrow,n}t) + \frac{\delta}{2} X_{\uparrow,n} \sin(X_{\uparrow,n}t) e^{-i\delta t/2} \\
 -\frac{Y_{\uparrow,n}}{X_{\uparrow,n}} \sin(X_{\uparrow,n}t) e^{i\delta t/2}
\end{array} \right] \left( \begin{array}{c}
 \frac{Y_{\uparrow,n}}{X_{\uparrow,n}} \sin(X_{\uparrow,n}t) e^{-i\delta t/2} \\
 \cos(X_{\uparrow,n}t) - \frac{\delta}{2} X_{\uparrow,n} \sin(X_{\uparrow,n}t) e^{i\delta t/2}
\end{array} \right) \left( \begin{array}{c}
 c_{\uparrow,n}(0) \\
 c_{\downarrow,n}(0)
\end{array} \right)
\]

with \( Y_{\uparrow,n} := -i\Omega_{\uparrow,n}(\alpha_1 + \alpha_2/|n|)\text{Re}^{|n|-n}^\text{Im}^\text{Im}^\text{Re}^{|n|} \) and \( X_{\uparrow,n} := \sqrt{(\delta^2/4) + |Y_{\uparrow,n}|^2}. \)

In the Lamb–Dicke regime, \( \eta(|\bar{\alpha} + \bar{\beta}^2|^{1/2} \ll 1, \) equation (3.22) can be expanded to first order in \( \eta: \)
\[
\begin{align*}
\Omega_{\uparrow,n-1,n} = & \Omega_l \sqrt{n} \\
\Omega_{\downarrow,n} = & \Omega_1 \text{ (carrier)}, \\
\Omega_{\uparrow,n+1,n} = & \Omega_l \eta \sqrt{n + 1} \text{ (first blue sideband).}
\end{align*}
\]

Successive red sideband transitions \( |\downarrow\rangle|n\rangle \to |\uparrow\rangle|n-1\rangle \) followed by dissipative repumping to \( |\downarrow\rangle|n\rangle \) with high probability are routinely used for sideband cooling close to the motional ground state \( |n = 0\rangle. \)

If the Lamb–Dicke parameter becomes effectively zero, the motional dependence will vanish (see equations (3.26) and (3.28)). The only remaining transition is the carrier transition equation (3.27) affecting the electronic states only. This is the case, for example, for two-photon stimulated-Raman transitions with co-propagating beams or for microwave driven transitions in hyperfine qubits, where \( k_1 \approx 0. \) In systems with more than one ion, the ions will not be motionally coupled. That is why equation (3.25) also holds for each site separately in such systems.

\[
\hat{U}_l^{(\text{LDR}(\text{RWA}))} (t_0) = \exp \left[ i \sum_{m=1}^{N} \sum_{i=1}^{N} \frac{\Omega_1^{(i)} \eta_{m}^{(i)} (e^{-i\omega_m t_0} - 1) e^{-i\delta m t_0} e^{i\delta m t_0} \hat{a}_m^{\dagger} \hat{a}_m^{\dagger} (\alpha_0 \hat{1}^{(i)} + \alpha_3 \hat{\sigma}_z^{(i)} + \text{h.c.})} {\delta_m} \right] \\
\times \exp \left[ -i \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{m=1}^{3N} \frac{\Omega_1^{(i)} \eta_{m}^{(j)} \eta_{m}^{(j)}} {\delta_m^{(i)}} (\alpha_0 \hat{1}^{(i)} + \alpha_3 \hat{\sigma}_z^{(i)}) \otimes (\alpha_0 \hat{1}^{(j)} + \alpha_3 \hat{\sigma}_z^{(j)}) \right] \\
\times \delta_m (t - t_0) \cos(\varphi_1 - \varphi_1) - \sin(\delta_m (t - t_0) - (\varphi_1 - \varphi_1)) \right].
\]

Equation (3.25) simplifies for resonant carrier transitions \( (\delta = 0) \) and a pure \( \hat{\sigma}_z \) interaction \( (\alpha_1 = 1 \text{ and } \alpha_2 = 0) : \)
\[
\left( \begin{array}{c}
 c_{\uparrow,n}(t) \\
 c_{\downarrow,n}(t)
\end{array} \right) = \hat{R}(\vartheta, \varphi) \left( \begin{array}{c}
 c_{\uparrow,n}(0) \\
 c_{\downarrow,n}(0)
\end{array} \right),
\]
where
\[
\hat{R}(\vartheta, \varphi) := \begin{pmatrix}
\cos(\vartheta/2) & -ie^{i\varphi} \sin(\vartheta/2) \\
-ie^{i\varphi} \sin(\vartheta/2) & \cos(\vartheta/2)
\end{pmatrix},
\]
\( \vartheta := 2\Omega_{\uparrow,n} t \) and \( \varphi := \varphi_1. \) The rotation matrix \( \hat{R}(\pi/2, \varphi) \) describes a \( \pi/2 \)-pulse and \( \hat{R}(\pi, \varphi) \) a \( \pi \)-pulse with phase \( \varphi. \)

3.3. Effective \( \hat{\sigma}_z \otimes \hat{\sigma}_z \) interaction

We will now discuss the case of equation (3.16) with \( \alpha_1 = \alpha_2 = 0. \) Hence, we omit the superscript of the Hamiltonian indicating an RWA. In the Lamb–Dicke regime, \( \eta_{m}^{(i)} (\hat{a}_m + \hat{a}_m^{\dagger})^{1/2} \ll 1, \) the Hamiltonian can be expanded to first order in the Lamb-Dicke parameters \( \eta_{m}^{(i)}. \) A subsequent RWA neglecting terms rotating faster than \( e^{i\omega_m t} \) with \( \delta_m := \omega_1 - \omega_m \) yields
\[
\hat{H}_l^{(i)(\text{LDR}(\text{RWA}))} = \hbar \Omega_l^{(i)} e^{i(\omega_1 t + \omega_1 t)}
\]
\[
\times \left( (\alpha_0 \hat{1}^{(i)} + \alpha_3 \hat{\sigma}_z^{(i)}) + \text{h.c.} \right)
\]
\[= \hbar \Omega_l^{(i)} e^{i(\omega_1 t + \omega_1 t)} \left( (\alpha_0 \hat{1}^{(i)} + \alpha_3 \hat{\sigma}_z^{(i)}) + \text{h.c.} \right)
\]
Note that equation (3.32) breaks up into a sum over terms that depend on only one mode \( m \) and one site \( i \) each.

With the excursion in appendix F the total time evolution operator in the interaction picture reads

The interaction described by equation (3.33) can be interpreted as follows: the first exponential function has the form of a displacement operator \( \hat{D}(\lambda) = e^{i\lambda \hat{a} - \lambda^* \hat{a}^\dagger} \), which leads to a displacement of a coherent state by \( \lambda \) in phase space. Due to the \( e^{i(\omega_m t - \omega_m t)} \) proportionality of the exponent, the trajectory for a coherent state of each mode describes a circle in phase space (or a straight line in the limit \( \delta_m = 0 \)). The coherent state returns to its initial position at times \( T_m = 2\pi/\delta_m \) with \( l \in \mathbb{N} \), where the exponent vanishes. The second exponential can be expanded into a \( \hat{\sigma}_z \otimes \hat{\sigma}_z \) interaction, a \( \hat{\sigma}_z \) interaction and a global phase. The \( \hat{\sigma}_z \otimes \hat{\sigma}_z \) terms give rise to a geometric phase, which increases in time \( t \), and the \( \hat{\sigma}_z \) terms lead to a dynamic phase [112, 113]. The area in phase space enclosed by the trajectory is proportional to these phases.

3.4. Geometric phase gates

The collective interaction of multiple ions with the same laser(s) has been proposed for the implementation of quantum
gates [85, 114–117]. These gates are described in the $z$-basis by equation (3.33) and have been first implemented in [110, 118]. Mølmer–Sørensen gates can be mathematically treated analogously in a rotated basis and are described in detail in [86, 87]. Implementations are reported in [119–122].

We will exemplarily discuss geometric phase gates based on the $\hat{\sigma}_z \otimes \hat{\sigma}_z$ terms in equation (3.33) in the following. They offer excellent tools to investigate a pure $\hat{\sigma}_z \otimes \hat{\sigma}_z$ interaction required for the simulation of more involved Hamiltonians such as quantum spin Hamiltonians. The geometric phase gates also allow the discussion of the $\hat{\sigma}_x \otimes \hat{\sigma}_z$ interaction in a familiar frame, while for the quantum Ising Hamiltonian a canonical transformation is introduced, which leads to a more involved dressed-state picture (see section 3.5).

The interaction according to the Hamiltonian can be implemented [47, 110, 123] by stimulated-Raman transitions driven by two beams with wavevectors $\vec{k}_1$, $\vec{k}_2$ and difference frequency close to a (several) motional mode(s) (see figure 5(b)). On average the differential ac Stark shift between the levels $|\downarrow\rangle$ and $|\uparrow\rangle$ caused by the two beams can be compensated by choosing appropriate polarizations of the beams. Still, on short timescales $\sim 2\pi/\delta_m$ the ions experience a state-dependent force that leads to the above displacement in the phase spaces of the corresponding modes.

In the original implementation of the geometric phase gate [110] two $^9\text{Be}^+$ ions are used. The state-dependent forces amount to $\vec{F}_1 = -2\vec{F}_2$. This means that the operators $(\alpha_0 \hat{I}^{(i)} \otimes \alpha_3 \hat{a}^{(j)} \hat{b}^{(j)})$ have diagonal elements $1$ and $-2$, which is fulfilled for $\alpha_0 = -1/2$ and $\alpha_3 = 3/2$. The effective wavevectors $\vec{k}_1^{(i)} = \vec{k}_1 - \vec{k}_2$ point along the axis of the linear trap and the laser beams are detuned by $\delta_{\text{STR}} = 2\pi \times 266 \text{kHz}$ from the stretch (STR) mode. The effect of the centre-of-mass (COM) mode can be neglected ($\delta_{\text{COM}} \approx 1000\delta_{\text{STR}}$). The ions are placed at the same phase of the stimulated-Raman interaction ($\phi_1^{(i)} = \phi_2^{(j)} = 0$).

For $t = T_\text{g} = 2\pi/\delta_{\text{STR}}$ the time evolution operator equation (3.33) simplifies to

\[
\hat{U}_1^{(\text{LDR|RWA})}(T_\text{g}, 0) \approx \exp \left( -2\pi i \sum_{i=1}^{2} \sum_{j=1}^{2} (-1)^{i-j} \Omega_i^2 \eta_{\text{STR}}^2 \delta_{\text{STR}} \right) \times \left[ \alpha_3^2 \hat{a}^{(i)} \otimes \hat{a}^{(j)} + \alpha_0 \alpha_3 \left( \hat{a}^{(i)} + \hat{a}^{(j)} \right) \right],
\]

where we have used $\eta_{\text{STR}} := \eta_{\text{STR}}^{(1)} = -\eta_{\text{STR}}^{(2)}$ and neglected the global phase arising from the $\hat{I}^{(i)} \otimes \hat{I}^{(j)}$ terms. The sequence of the gate is similar to the one in figure 7, but without the second displacement pulse $\hat{D}_2$. Ideally, the initial state $|\Psi_i\rangle = |\downarrow\downarrow\rangle \eta_{\text{COM}} = 0, \eta_{\text{STR}} = 0\rangle$ is rotated to $1/2(|\downarrow\downarrow\rangle + |\downarrow\uparrow\rangle + |\uparrow\downarrow\rangle + |\uparrow\uparrow\rangle) \eta_{\text{COM}} = 0, \eta_{\text{STR}} = 0\rangle$ by the first $\hat{R}(\pi/2, \pi/2)$ pulse (the phase $\psi$ of the first pulse can be chosen arbitrarily). The only non-vanishing contributions arise from the $\hat{\sigma}_z \otimes \hat{\sigma}_z$ terms for the $|\downarrow\downarrow\rangle$ and $|\uparrow\uparrow\rangle$ states, which gain a geometric phase

\[
\Phi_{\text{STR}}^{\downarrow\downarrow\uparrow\uparrow} = -2\pi \times 4 \frac{\Omega_i^2 \eta_{\text{STR}}^2}{\delta_{\text{STR}}} \alpha_3^2.
\]

By choosing appropriate beam intensities and thus $\Omega_i$, these phases equal $\Phi_{\text{STR}}^{\downarrow\downarrow\uparrow\uparrow} = -\pi/2$. The subsequent $\hat{R}(\pi, \pi/2)$ and $\hat{R}(\pi/2, \pi/2)$ pulses lead to the final Bell state $|\tilde{\Psi}\rangle = 1/\sqrt{2} (|\downarrow\downarrow\rangle + i |\uparrow\uparrow\rangle)$, which is achieved experimentally with a fidelity of $F = 97\%$ [110].

A similar implementation of the geometric phase gate is reported in [123] based on two $^{25}\text{Mg}^+$ ions. The state-dependent forces amount to $\vec{F}_1 = -3/2\vec{F}_2$ ($\alpha_0 = -1/4$ and $\alpha_3 = 5/4$). Furthermore, the detuning from the STR mode amounts to $\delta_{\text{STR}} = -2\pi \times 266 \text{kHz}$ and simultaneously the detuning from the COM mode $\delta_{\text{COM}} = -2\pi \times 1330 \text{kHz}$ (compare figure 6(a)). Hence, the effect of the COM mode is also exploited for the gate. As the detuning from the COM mode is chosen to be an integer multiple of the detuning from the STR mode ($\delta_{\text{COM}} = -5\delta_{\text{STR}}$), hence, there is no entanglement left between the electronic and motional modes at the gate duration $T_\text{g} = (2\pi/\delta_{\text{COM}}) = 3.75 \mu\text{s}$. (Note that the spin-echo sequence is not included in $T_\text{g}$.) (b) The parameters correspond to a phase gate on two of the radial motional modes. The radial centre-of-mass (COM) and rocking (ROC) mode have a comparatively small frequency difference of only $2\pi \times 130\text{kHz}$. The detunings from both modes are chosen to have the same absolute values resulting in (approximately) equal contributions to the acquired geometric phase from both modes. The gate duration according to the original implementation would amount to $T_\text{g} = (2\pi/\delta_{\text{COM}}) = 15.4 \mu\text{s}$. As the displacement pulse is repeated in the second gap of the spin-echo sequence (compare figure 7) to cancel dynamic phases (compare [118] and see text), the duration increases by an additional factor of two.

![Figure 6. Comparison between parameters of geometric phase gates [110] with two ions using the axial motional modes and radial motional modes. (a) The parameters correspond to the gate from [123] (b) The parameters correspond to a phase gate on two of the radial motional modes. The radial centre-of-mass (COM) and rocking (ROC) mode have a comparatively small frequency difference of only $2\pi \times 130\text{kHz}$. The detunings from both modes are chosen to have the same absolute values resulting in (approximately) equal contributions to the acquired geometric phase from both modes. The gate duration according to the original implementation would amount to $T_\text{g} = (2\pi/\delta_{\text{COM}}) = 15.4 \mu\text{s}$. As the displacement pulse is repeated in the second gap of the spin-echo sequence (compare figure 7) to cancel dynamic phases, the duration increases by an additional factor of two.

By choosing appropriate beam intensities and thus $\Omega_i$, these phases equal $\Phi_{\text{STR}}^{\downarrow\downarrow\uparrow\uparrow} = -\pi/2$. The subsequent $\hat{R}(\pi, \pi/2)$ and $\hat{R}(\pi/2, \pi/2)$ pulses lead to the final Bell state $|\tilde{\Psi}\rangle = 1/\sqrt{2} (|\downarrow\downarrow\rangle + i |\uparrow\uparrow\rangle)$, which is achieved experimentally with a fidelity of $F = 97\%$ [110].

A similar implementation of the geometric phase gate is reported in [123] based on two $^{25}\text{Mg}^+$ ions. The state-dependent forces amount to $\vec{F}_1 = -3/2\vec{F}_2$ ($\alpha_0 = -1/4$ and $\alpha_3 = 5/4$). Furthermore, the detuning from the STR mode amounts to $\delta_{\text{STR}} = -2\pi \times 266 \text{kHz}$ and simultaneously the detuning from the COM mode $\delta_{\text{COM}} = -2\pi \times 1330 \text{kHz}$ (compare figure 6(a)). Hence, the effect of the COM mode is also exploited for the gate. As the detuning from the COM mode is chosen to be an integer multiple of the detuning from the STR mode ($\delta_{\text{COM}} = -5\delta_{\text{STR}}$), the first exponential in equation (3.33) still becomes unity for the gate duration of $T_\text{g} = (2\pi/\delta_{\text{STR}})$. (In other words, all circular trajectories in
all phase spaces return to their initial position for $T_g$.) As a result, there is no entanglement left between the electronic and motional states.

Analogous to equation (3.35), but considering $\delta_{\text{STR}} < 0$ and $\delta_{\text{COM}} > 0$ for the detunings and $\eta_{\text{COM}} = \eta_{\text{COM}}$ for the Lamb-Dicke parameters of the COM mode, the geometric phases yield

$$\Phi_{\text{STR}1/1/1} = 2\pi \times 4 \frac{\Omega_{\text{COM}}^2 \eta_{\text{COM}}^2}{\delta_{\text{STR}}^2} \alpha_3^2,$$

$$\Phi_{\text{COM}1/1/1} = -2\pi \left| \frac{\delta_{\text{COM}}}{\delta_{\text{STR}}} \right| \times 4 \frac{\Omega_{\text{COM}}^2 \eta_{\text{COM}}^2}{\delta_{\text{COM}}^2} \alpha_0 \alpha_3^2.$$  \hfill (3.36)

By adjusting the beam intensities appropriately the differential phase between $\downarrow\downarrow/\uparrow\uparrow\uparrow$ and $\downarrow\uparrow/\uparrow\downarrow\downarrow$ can be adjusted to fulfill $\Phi_{\text{STR}1/1/1} - \Phi_{\text{COM}1/1/1} = \pi/2$. As $\Phi_{\text{STR}1/1/1}$ has the opposite sign compared with $\Phi_{\text{COM}1/1/1}$, the geometric phase gate makes use of two motional modes simultaneously.

However, some of the dynamic phases from the COM mode do not vanish:

$$\Phi_{\text{COM}1/1/1} = \pm 2\pi \left| \frac{\delta_{\text{COM}}}{\delta_{\text{STR}}} \right| \times 8 \frac{\Omega_{\text{COM}}^2 \eta_{\text{COM}}^2}{\delta_{\text{COM}}^2} \alpha_0 \alpha_3^2.$$  \hfill (3.37)

These phases have an absolute value of $2\alpha_0 / \alpha_3$ of the geometric phase from the COM mode and lead to a small deviation from the ideal state at the end of the gate.

Compared with the original implementation in [110], the gate is speeded up by approximately a factor of 10 and the fidelity $F$ for the Bell state exceeds 95%. (Note that the duration of the spin-echo sequence is not included in $T_g$, because its rotations could be much faster and empty gaps can in principle be removed.)

The radial motional modes are interesting, because they are similar to the normal modes in systems of individual traps for each ion (compare section 6), which are promising candidates for scalable systems in quantum simulations. To investigate the differences between the axial and radial modes of motion the geometric phase gate with $^{25}\text{Mg}^+$ is performed on a pair of radial modes (see also [122] for a Mølmer–Sørensen gate performed on the radial modes).

The detunings from the COM and ROC mode (abbreviation for ‘rocking’ mode, the equivalent to the STR mode in terms of the axial motional modes) are chosen to have the same absolute values $\delta_{\text{COM}} = -\delta_{\text{ROC}} = 2\pi \times 65 \text{ kHz}$ (see figure 6). The geometric phases acquired on each motional mode are basically the same as in equation (3.36), where ‘STR’ has to be replaced by ‘ROC’, and equation (3.37). (However, the signs change due to a change of the signs of the detunings.) The contributions to the total differential geometric phase between $\downarrow\downarrow/\uparrow\uparrow\uparrow$ and $\downarrow\uparrow/\uparrow\downarrow\downarrow$ due to the COM and ROC mode are (approximately) equal now. However, the dynamic phase (analogous to equation (3.38)) arising from the COM mode can no longer be neglected.

The pulse scheme of the geometric phase gate is modified by adding a second displacement pulse in the second gap of the spin-echo sequence (see figure 7 and compare [118]). The intensities of the beams are now adjusted for differential geometric phases due to each displacement pulse of $\pi/2$. While the geometric phases of both displacement pulses add up to $\pi/2$, the dynamic phases cancel each other, as the $\pi$ pulse of the spin-echo sequence interchanges $\downarrow\downarrow \leftrightarrow \uparrow\uparrow\uparrow$ (and $\downarrow\uparrow \leftrightarrow \uparrow\downarrow\downarrow$). Additionally, the more symmetric pulse scheme enhances the robustness.
of the gate against uncompensated differential ac Stark shifts between $|\downarrow\rangle$ and $|\uparrow\rangle$).

The total fluorescence from the two ions as a function of the total duration of the displacements $2\tau_D$ is shown in figure 8. The gate duration due to the smaller detunings and the second displacement pulse is more than a factor of eight longer than for the gate in [123]. Still, the fidelity exceeds 95% (see figure 9).

3.5. Quantum Ising Hamiltonian

Above we have introduced $\hat{\sigma}_z \otimes \hat{\sigma}_z$ interactions that are used in quantum gates. In the following we will present a slightly different approach, in which Ising spin–spin interactions are continuously induced by means of optical forces.

The spin–spin interaction as proposed in [21] and experimentally realized in the simulation of a quantum Ising Hamiltonian in [47] is identical to the interaction $H_I$ described in section 3.3. (Note that a similar proposal involving the spin–spin interaction $\hat{H}_I$ has to be applied to equation (3.15) (and subsequent equations) and $e^{\pm i\omega t}$ to 1 to equation (3.32). Hence, $H_I$ reads in the new interaction picture (including the expansion to first order in the Lamb–Dicke parameters and the RWA):

$$\hat{H}_I^{\text{LDR}(\text{RWA})} := \sum_{i=1}^{N} \sum_{m=1}^{3N} \hbar \Omega_m^{(i)} \eta_m^{(i)} e^{i\psi_m^{(i)}} \hat{a}_m^\dagger \hat{a}_m + \text{h.c.}$$

(3.41)

However, the full Hamiltonian in the interaction picture now also involves

$$\hat{H}_I = \hat{U}_0^\dagger \hat{H}_I \hat{U}_0 = \hat{H}_I \hat{U}_c.$$

(3.42)

where the transformation is the identity, because trivially $[\hat{H}_\theta, \hat{H}_I] = 0$.

To gain the form of a spin–spin interaction, we apply a canonical transformation (compare [21]) to the Hamiltonian. We want to note that an adiabatic elimination of phonons would yield the same effective description of the system; however, the canonical transformation provides a systematic method to also calculate the corrections to the ideal quantum Ising model:

$$\hat{H}_I^{\text{LDR}(\text{RWA})} + \hat{H}_I^{\text{c}} := \hat{U}_c \left( \hat{H}_I^{\text{LDR}(\text{RWA})} + \hat{H}_M \right) \hat{U}_c^\dagger$$

(3.43)

with

$$\hat{U}_c := \exp \left( -\frac{N}{\hbar} \sum_{i=1}^{N} \sum_{m=1}^{3N} \frac{1}{\hbar \delta_m} \left[ \hat{\xi}_m^{(i)} \hat{\sigma}_z^m - \hat{\xi}_m^{(i)\dagger} \hat{\sigma}_z^m \right] \right) \quad (3.44)$$

and

$$\hat{\xi}_m^{(i)} := i \hbar \Omega_m^{(i)} \eta_m^{(i)} e^{i\psi_m^{(i)}} \left( \alpha_0 \hat{\mathbf{l}}_m^{(i)} + \alpha_3 \hat{\sigma}_z^{(i)} \right).$$

(3.45)

Using the calculations from appendix G, the transformed Hamiltonian reads

$$\hat{H}_I^{\text{LDR}(\text{RWA})} + \hat{H}_I^{\text{c}} = \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{m=1}^{3N} \frac{1}{\hbar \delta_m} \hat{\xi}_m^{(i)} \otimes \hat{\xi}_m^{(j)\dagger} + \hat{H}_\phi$$

$$= \hbar \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{m=1}^{3N} \frac{\Omega_m^{(i)}}{\delta_m} \frac{\eta_m^{(i)} \eta_m^{(j)\dagger} e^{i\psi_m^{(i)} - i\psi_m^{(j)}}}{\delta_m}$$

$$\times \left( \alpha_0 \hat{\mathbf{l}}_m^{(i)} + \alpha_3 \hat{\sigma}_z^{(i)} \right) \otimes \left( \alpha_0 \hat{\mathbf{l}}_m^{(j)} + \alpha_3 \hat{\sigma}_z^{(j)} \right) + \hat{H}_\phi.$$  

(3.46)
The Hamiltonian can be expanded into a pure $\hat{\sigma}_z \otimes \hat{\sigma}_z$ interaction, a 'bias' term with $\hat{\sigma}_z$ interaction, and a constant term that can be neglected. The 'bias' term acts as a longitudinal magnetic field and leads to a deviation from the quantum Ising model. At first glance, this is not desired and it will be treated as an error in the following discussion. However, by including a 'bias' term in a controlled way we could also explore an extended phase diagram with the longitudinal field as an additional parameter.

We want to stress the similarity between the spin-spin interaction according to equation (3.46) and the $\hat{\sigma}_z \otimes \hat{\sigma}_z$ interaction discussed in section 3.3. The canonical transformation has the form of a displacement operator and looks very similar to the first exponential function in equation (3.33) (except for the time dependence of the latter). The similarity to the geometric phase term in equation (3.33) can be best seen comparing the time evolution operators. As the Hamiltonian in equation (3.46) is time-independent, the time evolution simply reads

$$\hat{U}_S^{\text{RWA}}(t, t_0) = \exp \left( -i \sum_{j=1}^N \sum_{m=1}^{3N} \frac{\Omega S(j) m j}{\delta m} (\sigma_0^j \hat{\sigma}_z^j + \sigma_3^j \hat{\sigma}_z^j) \right) \times \exp \left( i \sum_{m=1}^{3N} \delta_m (t - t_0) \hat{a}_m^\dagger \hat{a}_m \right).$$

(3.47)

Before we can apply the easier time evolution of equation (3.47), in which electronic states are decoupled from motional states, the state vector $|\psi\rangle$ has to be transformed from the original picture to $|\psi\rangle^{\text{syn}} := \hat{U}_c |\psi\rangle$. As $\hat{U}_c$ depends on the electronic state, the transformation will in general lead to an entangled state and the canonical transformation can be interpreted as dressed-state picture (electronic states 'dressed' with motional states). As the states for the simulation of the quantum Ising Hamiltonian are prepared in the original (undressed) picture, but the Hamiltonian acts in the dressed picture, an error is introduced into the simulation (see, for example, [125]). However, as long as the effect due to $\hat{U}_c$ is small ($\Omega S(j) m j / \delta m \ll 1$), we can use the approximation $|\psi\rangle^{\text{syn}} \approx |\psi\rangle$. In terms of the geometric phase gate this corresponds to the case when the circles in phase space are small and the entanglement between electronic and motional states can be neglected at any time.

The same holds for the measurements of observables: they are performed in the original (undressed) picture, in which electronic states are entangled with the motional states, and in general a further error is introduced in the simulation. However, the measurement of the states is typically insensitive to the motional states and involves a projection to one of the electronic states, for example, the $|\downarrow\rangle$ state: $\hat{P}^{(i)} := |\downarrow\rangle^{(i)} \langle \downarrow|$. As $[\hat{U}_c, \hat{P}^{(i)}] = 0$, the projector does not change under the canonical transformation and the readout of $\hat{\sigma}_z$ eigenstates (without any rotations of the bases applied beforehand) does not introduce further errors.

The form of the 'bias' term proportional to $\hat{\sigma}_z$ can be simplified in the case of a linear Paul trap with equal Rabi frequencies $\Omega S(j)$ and equal phases $\phi_S(j)$ for all ions: The sum over $j$ extends over the Lamb-Dicke parameters $\eta_m(j)$ only. This sum is non-zero only for centre-of-mass modes, for which the $\eta_m(j)$ are additionally independent of the site $j$. Hence, the three sums simplify to a sum over $\hat{\sigma}_z^{(i)}$ with constant prefactor [21]:

$$2\Omega S(j) |\gamma_0(i)|^2 \sum_{m=0}^{M} \frac{\eta_m(j)}{\delta_m} \left( \sum_{i=1}^N \hat{\sigma}_z^{(i)} \right).$$

(3.48)

However, this simplification does not necessarily hold for two-dimensional arrays of individual traps for each ion.

In the following, we will discuss the magnetic field term, which originates from a $\hat{\sigma}_z$ interaction described by equation (3.17) (with $a_1 = 1$ and $a_2 = 0$). In principle, we have to apply the substitution $\hat{e}^{\text{syn} \sigma j} \rightarrow \hat{e}^{\text{syn} \sigma j}$ due to the new interaction picture here, too. However, we consider a magnetic field term without motional dependence in the following ($\eta_m(j) = 0$, compare section 3.2) and thus the terms containing the motional creation/annihilation operators vanish:

$$\hat{\sigma}_z^{(i)} = \hat{h}(i) + \hat{\sigma}_z^{(i)} + \hat{a}_m^\dagger \hat{a}_m + \hat{h}. \quad (3.49)$$

The canonical transformation can be rewritten as

$$\hat{U}_c = \exp \left( \sum_{i=1}^N \hat{h}(i) \left( \hat{a}_m^\dagger \hat{a}_m + \hat{\sigma}_z^{(i)} \right) \right)$$

(3.50)

with the Hermitian operator

$$\hat{h}(i) := \left( \sum_{m=1}^N [\xi_m(i) \hat{a}_m^\dagger + \zeta_m(i) \hat{a}_m] \right) \quad (3.51)$$

and

$$\xi_m(i) := -\frac{\hbar \Omega_S(i) \eta_m(i) e^{\phi_S(i)}}{\hbar \delta_m(i)}. \quad (3.52)$$

Trivially, the commutator $[\hat{h}(i), \hat{\sigma}_z^{(i)}] = 0$. The canonical transformation of $\hat{H}_M^{\text{RWA}}$ (see equation (3.49)) is thus equivalent to a transformation of the $\hat{\sigma}_z^{(i)}$ operator as in equations (B.16) and (B.17):

$$\hat{H}_M^{\text{RWA}} = \hat{U}_c \hat{H}_M^{\text{RWA}} \hat{U}_c^\dagger \quad (3.53)$$

The expansion to first order in $\xi_m(i)$ (and thus to first order in $\hat{h}(i)$) yields

$$\hat{H}_M^{\text{RWA}} \approx \sum_{i=1}^N \frac{\hbar \Omega_S(i)}{2} \left( \xi_m(i) \hat{a}_m^\dagger + \xi_m(i) \hat{a}_m \right) \quad \left( 1 + 2i\alpha \sum_{m=1}^N \xi_m(i) \hat{a}_m^\dagger + \xi_m(i) \hat{a}_m \right) \hat{\sigma}_z^{(i)} + \text{h.c.}$$

(3.54)
The magnetic field term after the canonical transformation deviates to order $\hat{\Omega}^{(i)}_m \eta^{(i)}_m \alpha_i / \delta_{m}$ due to the pure $\hat{\delta}$ interaction $\hat{\Omega}^{(RWA)}_m$. This introduces an additional error in the simulation. If the condition $|\Omega^{(i)}_m \eta^{(i)}_m \alpha_i / \delta_{m}| \ll 1$ is met, it can be small or even negligible and we, effectively, will obtain the desired magnetic field term.

To summarize, the complete Hamiltonian is obtained by adding equation (3.46) and equation (3.54). It consists of a spin–spin interaction term and a simulated magnetic field pointing in the $x$-direction, which add up to the ideal quantum Ising Hamiltonian. Assuming a resonant interaction for the simulated magnetic field ($\omega_M = \omega_M^i = 0$) and neglecting the phases ($\varphi^{(i)} = \varphi^{(i)}_M = 0$), the quantum Ising part can be written as

$$
\hat{\mathcal{H}}^{\text{Ising}} := H_{\text{M}} + J \sum_{i=1}^N \sum_{j \neq i}^N J(i,j) \hat{\sigma}^{(i)}_z \otimes \hat{\sigma}^{(j)}_z,
$$

(3.55)

where

$$
B^{(i)} := \hbar \Omega^{(i)}_m \text{ and } J^{(i,j)} := \hbar \sum_{m=1}^N \frac{\Omega^{(i)}_m \eta^{(i)}_m \eta^{(j)}_m}{\delta_m} \sigma^2_z.
$$

(3.56)

( Note that the superscripts of $B^{(i)}$ and $J^{(i,j)}$ indicating the site will be omitted in the following sections, if the interaction strengths for all ions are equal.)

Depending on the sign of the detunings $\delta_m$, the effective spin–spin interaction can be of ferromagnetic ($J^{(i,j)} < 0$) or antiferromagnetic ($J^{(i,j)} > 0$) nature. The range and spatial structure of the spin–spin interactions can be partially controlled by choosing the absolute values of the detunings $\delta_m$. There are two different limits in which the interaction can be shaped in a very controlled way. On the one hand, if the laser is tuned close to the frequency of a given mode $m$ ($|\delta_m| \gg |\delta_{m'}| \forall m'$), then $J^{(i,j)}$ is a long-range interaction, whose spatial dependence is governed by the phonon wavefunction corresponding to $m$, as can be seen directly in equation (3.56). On the other hand, if the motional modes contribute to the spin–spin coupling ($|\delta_m - \delta_{m'}| \ll |\delta_{m'}| \forall m, m'$), the interaction shows a power-law decay $J^{(i,j)} \propto 1/|i - j|^{3}$ [21, 124]. This can be understood by the partial interference between the contributions from each mode’s wavefunction.

Furthermore, the freedom in the choice of the coupling strengths $\Omega^{(i)}_m$ for the individual ions as well as the trapping geometry and laser direction going into $\eta^{(i)}_m$ allows us to shape the individual coupling strengths $J^{(i,j)}$. This opens up the possibility of simulating a rich variety of models beyond the quantum Ising Hamiltonian, such as spin-frustrated systems [48] (see section 5.1). An illustration of the normal modes of quantum Ising Hamiltonian, such as spin-frustrated systems, is a long-range interaction, whose spatial dependence is governed by the phonon wavefunction corresponding to $m$, as can be seen directly in equation (3.56).

The first 'bias' term is further discussed in the context of the experimental realization, see [47] and section 5.1. The second term $\hat{\mathcal{H}}_{\delta}$ can be interpreted as an energy offset, which cancels by applying an appropriate redefinition of the energy scale. As mentioned above, the last term leads to only a small or even negligible error for $|\Omega^{(i)}_m \eta^{(i)}_m \alpha_i / \delta_{m}| \ll 1$. For a more detailed discussion of the errors in the simulation of quantum spin Hamiltonians we refer the reader to [125].

2003280 Hz 2000411 Hz 2000411 Hz
1100000 Hz 1097011 Hz 1097011 Hz
900911 Hz 900911 Hz 896345 Hz

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure10}
\caption{Example for normal modes in a surface-electrode trap geometry similar to the one discussed in section 6. It consists of three trapping zones (red spheres) arranged in a triangle at mutual distances of 40 μm. Each trapping zone has a (bare) frequency of $2\pi \times 2$ MHz corresponding to the vibration towards the centre of the triangle, $2\pi \times 1.1$ MHz for the out-of-plane vibration and $2\pi \times 0.9$ MHz for the third, perpendicular direction. The range and spatial structure of the spin–spin interactions can be partially controlled by choosing the absolute values of the detunings $\delta_m$. There are two different limits in which the interaction can be shaped in a very controlled way. On the one hand, if the laser is tuned close to the frequency of a given mode $m$ ($|\delta_m| \gg |\delta_{m'}| \forall m'$), then $J^{(i,j)}$ is a long-range interaction, whose spatial dependence is governed by the phonon wavefunction corresponding to $m$, as can be seen directly in equation (3.56). On the other hand, if the motional modes contribute to the spin–spin coupling ($|\delta_m - \delta_{m'}| \ll |\delta_{m'}| \forall m, m'$), the interaction shows a power-law decay $J^{(i,j)} \propto 1/|i - j|^{3}$ [21, 124]. This can be understood by the partial interference between the contributions from each mode’s wavefunction.}
\end{figure}
We want to emphasize that $[\hat{H}_B, \hat{H}_J] \neq 0$, such that the time evolution of the total quantum Ising Hamiltonian $\hat{H}_\text{QIsing}$ cannot be simply described by the time evolutions of $\hat{H}_B$ and $\hat{H}_J$ separately.

4. Operations interpreted for experimental quantum simulations

To realize a QS for a quantum spin Hamiltonian, we have to (1) simulate the ‘spin’, provide (2) its initialization and (3) the interaction of this ‘spin’ with a simulated magnetic field, (4) realize an interaction between several spins (spin–spin interaction) and (5) allow for efficient detection of the final spin state. Additional diversity for QS arises by the capability of precise initialization, control and readout of the motional states.

The mathematical derivation and description of the individual operations have been described in section 3. In this section, we explain in a simplified pictorial way the related generic building blocks in terms of an adiabatic QS of a quantum spin Hamiltonian within a linear chain of ions. No specific ion species or trapping concept are required. A whole family of quantum spin Hamiltonians related generic building blocks in terms of an adiabatic QS this section, we explain in a simplified pictorial way the individual operations have been described in section 3.1. (1) Phonons do not have to be restricted to states. The states are coupled via electro-magnetic radiation (see section 2.2). (2) Topological defects in the zigzag structure of two-dimensional Coulomb crystals (see figure 13) are proposed to simulate solitons [127].

4.1. Simulating the spin

The mutual distance between the ions/spins in linear RF traps is typically of the order of several micrometres (see figure 3). Therefore, the direct interaction between their electronic states remains negligible, which is advantageous, because the related interaction strength could hardly be tuned or even switched off. Therefore, the spin-1/2 states are implemented like qubit states (see section 2.2).

4.2. Simulating the magnetic field

Implementing an artificial spin allows us to shape artificial fields to implement a precisely controllable interaction and related dynamics between the ‘spin’ and the ‘field’. To simulate an effective magnetic field, the two electronic states $|\downarrow\rangle$ and $|\uparrow\rangle$ are coupled via electro-magnetic radiation (see section 2.3, operation (a)). The related coherent oscillation of the state population between the two levels can be described in terms of Rabi flopping. In the Bloch sphere picture, the tip of the electronic state vector rotates during one flop continuously from state $|\downarrow\rangle$ to $|\uparrow\rangle$ and vice versa. For continuous coupling this can be interpreted as the precession of a spin exposed to a perpendicular magnetic field.

The rotation matrix in equation (3.30) exactly describes this interaction with a single spin (see also sections 3.2 and 3.4). For example, if we start with $|\downarrow\rangle$ and apply a pulsed rotation $\hat{R}(\pi/2, \pi/2)$, we will obtain an eigenstate of $\hat{J}_z$, which is abbreviated by $|\rightarrow\rangle := \sqrt{2}/2(|\downarrow\rangle + |\uparrow\rangle)$. In the Bloch picture, this corresponds to a 90° rotation of the Bloch vector around the y-axis, such that it will point in the direction of the x-axis. Continuing with a second identical rotation we just flip the spin to $|\uparrow\rangle$ as if we applied $\hat{R}(\pi, \pi/2)$ or a 180° rotation around the y-axis, respectively. However, we can replace the second operation by $\hat{R}(\pi/2, 0)$, which corresponds to a rotation around the x-axis. As the state $|\rightarrow\rangle$ is an eigenstate of $\hat{J}_x$, it will not be affected.

Stroboscopic rotations have been introduced in section 3.4 to implement single-qubit gates for a QC. Continuous versions of these single-qubit operations can be interpreted in the context of analogue QS as simulated magnetic field (first term of equation (3.55)).

4.3. Simulating the spin–spin interaction

Let us first discuss the implementation of a basic spin–spin interaction close to the original proposal in [21]: two ions are confined in a linear RF trap and a standing wave provides state-dependent dipole forces. The ions are located at the same phases ($\varphi^{(i)} = 0$), such that ions in different spin states are pulled/pushed in opposite directions.

If both ions are in the same spin state, they will be pulled in the same direction. Hence, their mutual distance and mutual Coulomb energy, respectively, remain unchanged. However, if the two spins are in different states, one ion will be pulled and the other one pushed. Their mutual distance and as a result their mutual Coulomb energy will change. This is exactly the essence of a spin–spin interaction, where the energy corresponding to a spin state depends on the states of its neighbours. To interpret interactions as ferromagnetic or antiferromagnetic it is advantageous to consider the mutual Coulomb energy in longer chains of spins (see figure 11).

The technical realization in [47] avoids the difficulties arising from standing waves and resonantly enhances the interaction strengths by implementing the spin–spin interactions with stimulated-Raman transitions as in the case of quantum gates [124] (see section 3.4). In a pictorial interpretation, the standing waves are replaced by ‘walking’ waves and instead of static displacements we obtain driven oscillations of the ions. However, the mathematical description yields exactly the same spin model in an appropriately chosen frame (see section 3.5). The sign of $J$ can additionally be changed by choosing a different sign for the detunings $\delta_m$ from the modes (see equation (3.56)).

4.4. Geometric phase gate versus adiabatic quantum simulations

It might be helpful to emphasize the differences and similarities of the interactions in QC and analogue QS: to realize a
energetically preferred, such that the mutual Coulomb energy will be decreased. (Note that the mutual Coulomb energy will not change. For alternating chain of ions with the same spin states will again only be displaced and the mutual Coulomb energy is increased and decreased and the mutual Coulomb energy is due to its $1/d$ dependence. Here $d$ denotes the distance between neighbouring spins. The ferromagnetic order is energetically preferred, such that $J < 0$ for this interaction. (b) A chain of ions with the same spin states will again only be displaced and the mutual Coulomb energy will not change. For alternating spin states the distances between neighbouring ions will increase, such that the mutual Coulomb energy will be decreased. (Note that this should not to be confused with the structural zigzag phase-transition (see figure 3), where the radial displacements are typically orders of magnitude larger and spin-independent.) The antiferromagnetic order is energetically preferred, such that $J > 0$.

In contrast, we consider an adiabatic evolution according to the quantum Ising Hamiltonian in the case of analogue QS (see section 5.1 for the experimental protocol). We have to make sure that the entanglement between electronic and motional states remains small at any time during the simulation (see discussion of errors in section 3.5). Additionally, running the simulation on many spins simultaneously will result in contributions from many motional modes simultaneously. As a result, a large detuning from all modes has to be chosen, such that the difference of the radial frequencies can be neglected and a net effect from all modes remains. Choosing the right parameters allows the simulation of spin–spin interactions of different strength, different signs and even range of interaction [21].

Furthermore, a scan of the duration of the displacement pulses $T_D$ in geometric phase gates leads to a periodic evolution from $|↓↓⟩$ to $|↑↑⟩$ and vice versa (see figure 8).

In contrast, the distinct contributions ($\tilde{\mathcal{H}}_B$ and $\tilde{\mathcal{H}}_I$) of the quantum Ising Hamiltonian are not stroboscopically alternated but applied simultaneously. As mentioned in section 3.5, the time evolution according to the quantum Ising Hamiltonian is not simply the time evolution according to $\tilde{\mathcal{H}}_B$ and $\tilde{\mathcal{H}}_I$ separately. As a consequence, applying the spin–spin interaction for a longer duration and/or increasing its strength does not alter the state anymore.

5. Towards simulating many-body physics

In the first part of this section we want to assemble the building blocks described above to illustrate how an analogue QS of a quantum spin Hamiltonian can be implemented. For this purpose, we will describe the realization of first proof-of-principle experiments on the quantum Ising Hamiltonian (see equation (3.55)). In the second part we aim to summarize, to the best of our knowledge, the existing proposals addressing many-body physics with the described and available toolbox.

5.1. Proof-of-principle experiments on quantum spin Hamiltonians

First, we will describe the basic implementation of the experimental protocol on the axial modes for the case of two spins [47], as illustrated in figure 12. Subsequently, we will emphasize the differences and additional information explored in [48, 49]. For the details on the individual experimental parameters we refer to these references.

For the case of two spins, the protocol has been realized following five steps. (1) The two ions are initialized by Doppler cooling, sideband cooling and optical pumping (see section 2.4) in the state $|↓↓⟩$ ($n_{\text{STR}} = 0$). (2) Both spins are prepared by a common $\hat{R}(\pi/2, \pi/2)$ rotation in the $\hat{\sigma}_z$ eigenstate $|\rightarrow\rangle$ ($n_{\text{STR}} = 0$). (3) An effective magnetic field of amplitude $B_x$ is applied equivalent to a continuous $\hat{R}(2\Omega_M t, 0)$ rotation (see equations (3.55) and (3.56)). At this step, the state $|\rightarrow\rangle$ ($n_{\text{STR}} = 0$) represents the ground state of the first term of the quantum Ising Hamiltonian in equation (3.55) that can be 'easily' prepared. Note that the rotation is slightly off-resonant to mimic an additional $\hat{\sigma}_x$ interaction counteracting the 'bias' field (see also [47]). (4) The effective spin–spin interaction $J$ is ramped up adiabatically with respect to the timescale $1/\Omega_M$ defined by the simulated magnetic field, until $|J| \gg B_x$. The system adiabatically evolves into its new ground state, which is an equal superposition of the two energetically preferred states of the ferromagnetic order: $|\sqrt{2}(|↓↓⟩ + |↑↑⟩)$ (5) Finally, both interactions are switched off. The readout of the final spin state is performed by state-dependent detection. This
projects the spin state to one out of the four eigenstates of the measurement basis (|↓↓⟩, |↓↑⟩, |↑↓⟩, |↑↑⟩). Steps (1)–(5) are repeated many times to obtain the populations related to these states.

To investigate the degree of entanglement of the final spin state, an additional parity measurement is performed as in the case of the geometric phase gates (see section 3.4). The populations of |↓↓⟩ and |↑↑⟩ as a function of |J|/Bx and the entanglement fidelity are summarized in figure 12.

The experimentally observed entanglement of the final states confirms that the transition from paramagnetic to ferromagnetic order is not caused by thermal fluctuations that drive thermal phase transitions, but by the so-called quantum fluctuations [45, 128] driving QPTs in the thermodynamic limit at zero temperature. In this picture tunnelling processes induced by Bx coherently couple the degenerate states |↑⟩ and |↓⟩ with an amplitude \( \propto B_x/|J| \). For \( N \) spins the amplitude for the tunnelling process between \( |\Psi_{N+1}⟩ = |↑↑\ldots↑⟩ \) and \( |\Psi_{N}⟩ = |↓↓\ldots↓⟩ \) is proportional to \( (B_x/|J|)^N \), since all \( N \) spins must be flipped. In the thermodynamic limit (\( N \to \infty \)) the system is predicted to undergo a QPT at \( |J| = B_x \). At values \( |J| > B_x \) the tunnelling between \( |\Psi_{\infty+1}⟩ \) and \( |\Psi_{\infty}⟩ \) is completely suppressed. In our case of a finite system \( |\Psi_{2+1}⟩ \) and \( |\Psi_{2}⟩ \) remain coupled and the sharp QPT is smoothed into a gradual change from paramagnetic to ferromagnetic order (see figure 12).

It has to be noted that the performance of such a simulation on a large number of spins in a one-dimensional chain requires several technical improvements. Recently, a group at the University of Maryland pioneered a substantial step for scaling by investigating the emergence of magnetism in the quantum Ising model up to nine ions [49]. To achieve these results they mediated the interactions via the radial modes of motion [21, 122] (see also section 3.4). Furthermore, they implemented the effective spin–spin interactions in a rotated frame using Mølmer–Sørensen interactions [85, 87] on robust hyperfine clock states. Thereby, they do not depend on the phases \( \phi(i) \) of the laser beams at the sites of the ions. To perform an adiabatic transition, the simulated magnetic field had been adiabatically turned off, while the effective spin–spin interaction remains constant.

Their results allow much more than simply increasing the number of spins: they enter a new regime of intriguing questions. The crossover of the quantum magnetization [47] from paramagnetic to ferromagnetic order is sharpening as the number of ions is increased from two to nine, ‘prefacing the expected QPT in the thermodynamic limit’ [49]. Even though the results can still be calculated on a classical computer, they provide a possibility to critically benchmark QS aiming for only slightly larger systems.

Increasing the number of ions to three and adapting the individual spin–spin interactions including their signs allows spin frustration to be addressed in the smallest possible magnetic network [48]. Spin frustration of the ground state can be pictorially understood in a two-dimensional triangular spin lattice featuring antiferromagnetic spin–spin interactions. Here, it becomes impossible for neighbouring ions to have pairwise opposite states. Classically, two ions will adopt different states, while the state of the third one is undetermined. During an adiabatic evolution of the quantum mechanical system (starting from the paramagnetic order) nature will choose a superposition of all degenerate states, leading to massive entanglement in a spin-frustrated system. In the realization of the experiment, the three ions are still trapped in a one-dimensional chain. However, almost complete control over the amplitudes and signs of \( J^{(1,2)}, J^{(2,3)}, J^{(3,1)} \) is gained by coupling to particular collective modes of motion and choosing appropriate detunings [48].

It has to be mentioned that for an increased number of spins the energetic gap between ground and excited states further shrinks and the requirement on adiabaticity enforces longer simulation durations related to a longer exposure to decohering disturbances. Still, as mentioned in section 1, the influence of decoherence might destroy the entanglement within the system, but this might not be relevant for the observable of interest. Here it will be crucial to investigate the role of the decoherence effects with respect to the specific analogue QS.

With respect to digital (stroboscopic) QS it should be emphasized that no quantum error correction is required for proof-of-principle experiments on a few ions. Promising results of a stroboscopic version of the simulation of the quantum Ising Hamiltonian with two spins have been shown recently [129].
Condensed matter met atomic, molecular and optical physics not so long ago, when trapping techniques for ultracold neutral atoms and ions allowed experimentalists to generate lattices and crystals, where models from solid-state physics may be implemented. Combining the fields has led to a very rich interdisciplinary research activity, as well as to several misunderstandings between scientists looking at the same system from different points of view. In the particular case of trapped ion experiments, the outlook for quantum simulation of many-body models is very exciting, but some knowledge on the details of this physical system is required to understand both the limitations and the amazing possibilities of this setup.

In the following we review many-body models that have the potential to be simulated with trapped ions. There have been several contributions both from theory and experiments to this research line. Most of them share the common feature that they are inspired by known models from condensed matter physics, but their implementation with trapped ions turns out to lead to a rich variety of new physical phenomena, which may even require new theoretical paradigms that go beyond the conventional ones in the solid state. The three main reasons for this are (1) trapped ion experiments are naturally performed in a non-equilibrium regime, whereas solid-state physics typically deals with thermal equilibrium, (2) trapped ion systems may in principle be controlled and measured at the single particle level, (3) ion crystals are typically mesoscopic systems, in the sense that they may reach a number of particles (spins, phonons, etc) large enough to show emergent many-body physics, but still finite size effects are important. All those peculiarities have to be kept in mind, since they provide us with unique features for analogue QS.

5.2.1. Quantum spin models. Following the experimental advances in QIP, the most natural degree of freedom to be used for QS seems to be the electronic states for spins and the phonons to mediate their mutual interactions. However, one has to identify conditions where interesting phenomena arise, such as, for example, quantum critical phases. This has already lead to the promising proof-of-principle experiments discussed above.

A unique feature that we can exploit with trapped ions is the fact that the effective spin–spin interactions can be implemented showing a dipolar decay, \( J^{(i,j)} \propto \frac{1}{|i - j|^{3}} \). In the case of the Ising interaction, the cubic dependence does not change the critical universality class of the model, as shown, for example, by the numerical calculations in [125]. However, even in this case, long-range entanglement is induced by the long-range interaction, which is absent in conventional nearest-neighbour quantum Ising chains. On the other hand, when considering other interacting schemes, such as the XXZ-Hamiltonian, the dipolar interaction may lead to the formation of quasi-crystalline phases of spin excitations [130]. Furthermore, the simulation of the hexagonal Kitaev model with ions in an optimized, two-dimensional surface-electrode trap has been proposed [131].

Moreover, intrinsic properties of the trapped ion crystal, such as the linear–zigzag transition, have been demonstrated to be a QPT of the universality class of the Ising model in a transverse field [132, 133].

Several pieces have been added to the toolbox of quantum simulation, which definitely allow us to explore physics beyond conventional solid-state paradigms. For example, a theoretical proposal has been presented to implement models, whose ground states show topological features [134]. Also, methods to implement three-body spin–spin interactions have been designed, see [135]. Finally, dissipation in trapped ion systems has been proved to be useful to engineer quantum phases that arise as steady-state of dissipative processes [136].

The many-body physics of dissipative systems is a much more unexplored area than equilibrium properties, even for theorists. For that reason, adding dissipation to quantum magnetism opens an exciting perspective for trapped ions.

5.2.2. Interacting boson models. A variety of exciting quantum many-body systems may also be simulated using the collective motional degrees of freedom (phonons) to realize models of interacting bosons. In particular, whenever the motional coupling between ions is small compared with the trapping frequency, the phonon number is conserved and becomes a good quantum number to characterize the quantum state of the system. This principle was introduced and exploited in [22] to show that the physics of radial modes in Coulomb chains is effectively described by a Bose–Hubbard model. Vibrational couplings between two ions, say 1 and 2, induced by the Coulomb interaction, have a typical form \( \propto \hat{x}^{(1)} \hat{x}^{(2)} \), where \( \hat{x}^{(i)} \) is the ion displacement operator. Under the approximation of phonon number conservation, those terms become tunnelling couplings of the form \((\hat{a}_1^\dagger \hat{a}_2 + \text{h.c.})\). The same idea applies to quartic anharmonicities of the trap, which yield Hubbard interactions, \((\hat{a}_m^\dagger \hat{a}_n)^2\). Anharmonicities may be induced and controlled with optical forces, as shown in [22]. This analogy between phonons and interacting bosons opens an exciting avenue of research, where experiments might be relevant even with a single ion, realizing a single anharmonic quantum oscillator.

The ground state of those phonon-Hubbard models in Coulomb chains was extensively studied in [137], where it was shown that phonon Luttinger liquid phases may arise. Very recent experiments indeed show the tunnelling of phonons between ions trapped by different potentials, realizing thus an important step towards the use of phonons for quantum simulation [138, 139]. Exploiting phonon tight-binding models has also been shown to allow the implementation of models with disorder showing Anderson localization [140], as well as synthetic gauge potentials using periodic driving of the trap frequencies, see [126]. Using dipole forces acting on ions confined in a microtrap array (see section 6), motional couplings can be controlled such that phonons simulating charged particles experience synthetic gauge fields.

5.2.3. Spin–boson models. The natural convergence of the proposals presented above leads to the quantum simulation of
spin–boson models. This is a paradigmatic model for quantum impurities in solids, which typically describes a single spin coupled to a continuous bath of harmonic oscillators with a power-law spectral density. Surprisingly, the coupling of the electronic levels of a single ion to the axial phonons of a Coulomb chain yields a spin–boson model with a quasi-ohmic spectral density [44]. The physics to be simulated here is equivalent to some celebrated models in condensed matter physics, such as the Kondo effect. The finite size effects that are intrinsic to trapped ion systems turn out to yield features beyond the conventional physics of these models, in particular quantum revivals associated with the reflection of vibrational waves along the chain. Quite recently, it has been proposed to study a situation in which spins and phonons are coupled, in such a way that a Jaynes–Cummings–Hubbard model is simulated [141]. In this model phonons follow a tight-binding Hamiltonian and, in addition, they are locally coupled to spins. The system has been shown to undergo a superfluid–Mott insulator QPT.

5.2.4. Inhomogeneous many-body models: impurities and topological defects. The tools for QS in ion traps are not restricted to electronic and motional degrees of freedom only. It has been proposed to exploit impurities in the Coulomb crystal. On the one hand, for example, by embedding ion(s) of a different species (different mass) into the crystal and taking advantage of the altered spectrum of the modes and scattering of phonons [142] and the option to include larger simulated spins ($S > 1/2$) [143]. On the other hand, by creating localized topological defects within the more dimensional structure of the Coulomb crystal (see figure 13). In [127] it was suggested to induce a structural phase transition from a linear chain of ions (see figure 3(b)) to a zigzag structure (see figure 3(c)), for example, by lowering the radial confinement. Changing the parameters in a non-adiabatic way (fast compared with the phonons mediating information within the crystal) should cause independent domains of ‘zigzag’ and ‘zagzag’ structure, respectively. At their clash, topological defects were predicted and have recently been observed (see figure 13). The number of the created defects should scale according to the Kibble–Zurek prediction [144–146]. The defects themselves can be interpreted as solitons [127]. Solitons are defined as localized solutions of nonlinear systems, which depend essentially on nonlinearity. Such solitons have a unique spectrum of frequencies with modes which are localized to the soliton and whose frequency is separated by a gap from the other phonons. A quantum mechanical time evolution of these modes was calculated numerically and it is expected to remain coherent for hundreds of oscillations [147]. QS could allow us to explore their potential applications for QIP [127] as well as the quantum behaviour of these ‘objects’ themselves. Solitons appear in all branches of the natural sciences and have been extensively investigated in solid-state systems [148]. Among others, classical solitons were observed in waveguide arrays [149, 150] and Bose–Einstein condensates (BECs) [151], where they are mean field solutions. Discrete solitons were investigated in the Frenkel–Kontorova (FK) model [152, 153], which describes chains of coupled particles interacting with a local nonlinear potential. In a different realization, a variant of the FK model can also be realized in the ion trap by adding an optical lattice to a linear chain [154–156].

The important requirement to address any of these intriguing models will be to increase the number of ions and the dimensionality of the system. Trapping ions in two-dimensional arrays would allow the study of hard-core boson phases, showing the effect of frustration, quantum spin liquid phases and quantum states with chiral ordering [157]. Two approaches for scaling will be described in more detail in the following sections.

6. Scaling analogue quantum simulations in arrays of radio-frequency surface-electrode traps

One possible way to overcome the limitations on scalability of trapped ions in a common potential well (see section 2.1) is to store them in an array of individual RF traps.

6.1. One-dimensional radio-frequency surface-electrode traps

Conventional RF traps with their three-dimensional geometry of electrodes (see figure 2) individually fabricated with conventional machining were unique ‘masterpieces’ with unique characteristics.

In 2005 and 2006, a group at NIST pioneered the miniaturization of RF traps by projecting the electrodes onto a surface [73, 74] (see figure 14(a)), very similar to chip traps for neutral atoms [158]. Introducing photolithographic techniques for the trap fabrication opened up exceptional precision and the production of small series of identical traps, see for example [159, 160]. Within these linear RF surface-electrode traps, motional ground state cooling was achieved at a height of the ion over the electrode surface of $h = 40 \mu m$ and with a comparatively small motional heating rate of the order of 1 quantum ms$^{-1}$ [74].

Motional heating rates scale with $\sim h^{-4}$ [161]. The exact heating mechanisms are not yet fully understood and the groups
at NIST, in Berkeley, at MIT and others are currently putting a lot of effort into further investigations. However, the groups at MIT [162], NIST and the University of Maryland demonstrated a significant reduction of the heating rates in cryogenic (surface electrode) traps for QC purposes (see also [50]). The inverse of these heating rates has long been compared with typical operational durations of a QC of tens of microseconds (see also section 3.4).

For scaling towards a universal QC it might suffice to interconnect linear ion traps via junctions on a two-dimensional surface to a network of one-dimensional traps [163], realizing the ‘multiplex ion trap architecture’ [75]. That is, ions are proposed to be shuttled between processor and memory traps only interacting in the processor traps. This would allow the subdivision of the large total number of ions into small groups in many individual traps and to reduce the local requirements to a technically manageable effort. One-dimensional RF surface-electrode traps with more than 150 individual dc electrodes and several junctions have been realized [163], allowing ions at moderate heating rates to be shuttled.

In addition, the opportunity arose to deliver identical traps to different groups. One example is the linear RF surface-electrode trap (denoted by ‘Sandia Linear Trap’ in the following) [159], which was designed by the groups in Oxford, Innsbruck and Sandia National Laboratories. The latter fabricated a small series of identical replicas. The traps have been tested in several laboratories and the individually measured trapping parameters are in good agreement with the design values. Publications are in preparation by the groups at Oxford and Sandia (see also [159]).

It has to be emphasized that pursuing the multiplex approach for scaling universal QC is not applicable to the proposed analogue QS, where the ensemble of spins is supposed to evolve uniformly as a whole.

Figure 14. Schematic to illustrate the projection of the electrodes of the RF (yellow) and dc (shaded) electrodes on a surface as a way to scale towards two-dimensional arrays of ions. The black crosses indicate the positions of the minima of the pseudopotentials.

(a) Cross section of the electrodes of a conventional linear RF trap with three-dimensional geometry and the electrode structures projected onto a surface. The dashed arrows point at the new location of the electrodes, the white areas represent isolating gaps.

(b) Cross section (upper part) and top view (lower part) of the stripe electrodes. It has been proposed to concatenate several linear RF surface-electrode traps as depicted in (a) as a basic unit to span a two-dimensional array of ions [164] (red and blue disks representing ions in opposite spin states). For sufficiently small mutual ion distances and decoherence rates of the ions, this is an approach to scale analogue QS.

Figure 15. Illustration of the optimization results for the electrode structure for a basic triangular lattice with respect to the height of the ions above the traps at constant mutual ion distance. The white gap isolates RF and dc patches. The three red disks symbolize three ions at a constant distance of \( d = 40 \mu m \), hovering above the surface at a height of (a) \( h = 30 \mu m \), (b) \( h = 40 \mu m \) and (c) \( h = 50 \mu m \). (Courtesy of Roman Schmied.)

6.2. Optimized two-dimensional arrays of radio-frequency surface-electrode traps

Shortly after the invention of RF surface-electrode traps it was proposed to concatenate linear traps sufficiently close, such that the ions experience mutual Coulomb interaction in two dimensions [164] (see figure 14(b)). However, for a real two-dimensional lattice at sufficiently small and uniform ion distances of \( d \leq 40 \mu m \) in two dimensions, this proposal requires the ions to approach the disturbing surface to \( h \leq d/2 = 20 \mu m \) [73].

Schmied et al implemented a method to calculate the global optimum of the electrode shapes for arbitrary trap locations and curvatures (originally only for periodic boundary conditions) [165]. The gaps between neighbouring electrodes were neglected. The authors exemplarily optimized a trap array with comparatively stiff horizontal confinement.

The idea of optimizing electrode structures can also be used for designing traps for analogue QSs with partially converse requirements. In a collaboration of R. Schmied, NIST, Sandia National Laboratories, and us, such a surface-electrode trap has been designed and is currently in fabrication. The trap will provide three trapping zones arranged in a triangle (similar to figure 15) and is intended as a first step towards larger arrays of ions. For this purpose, the optimization method was extended to finite-sized traps.

It has to be emphasized that there are currently several proposals and approaches for arrays of surface-electrode traps mainly for QC. Groups in Berkeley and Innsbruck aim at trap arrays with individually controlled RF electrodes. They have the advantage of selectively lowerable trap frequencies for individual traps and thus increasable interaction strengths between ions in different traps, while especially the height of the ions above the surface can be larger and the trap depth of other traps can be sustained [166] (see also the discussion in the following subsections). This approach can in principle be extended to quasi-micromotion-free shuttling of ions in arrays of RF traps [167] at the expense of precise control of the RF voltage for each RF electrode. Another proposal suggests individual coils to be included for each trap to allow for laser-less interactions mainly for QS [168]. Different trap geometries specifically for QSs have been designed by a group in Sussex [169]. Arrays of Penning traps with surface electrodes are advanced by the groups at Imperial College [170] and the University of Mainz [171].
In the following subsections we discuss the optimization goals for a surface-electrode trap for an analogue QS, their implications and the perspectives for scalability of this approach.

6.2.1. Maximization of interaction strengths. The crucial prerequisite for QSs is to maximize the interaction strength (see section 3). As opposed to the QC case with multiple ions in a common potential, this has to be achieved in QS for ions in individual potentials. The increased mutual ion distances in arrays of individual traps substantially reduces the strength of the effective spin–spin interaction. It has be taken into account that the conditional forces have a limited strength, for example, because the laser power is limited or the assumptions in the theoretical model impose constraints as for the quantum Ising Hamiltonian (see section 3.5). However, a reduced stiffness of the individual potentials compared with the example [165] (trap frequencies on the order of $2\pi \times 20$ MHz) results in larger displacements of the ions by the same forces. This is related to an increased mutual Coulomb energy and thus larger interaction strengths. Still, a lower bound for the trap frequencies (on the order of $2\pi \times 1$ MHz for Mg$^+$) is imposed by the constraints for efficient ground state cooling.

6.2.2. Minimization of decoherence. The ions will inevitably approach the disturbing electrode surfaces, if the distance between the individual traps is reduced. We now reinvest the reduced requirements on the stiffness of the horizontal confinement to increase the height of the ions above the surface $h$ keeping the mutual ion distances $d$ constant. Some results for the scenario of a basic triangular lattice are depicted in figure 15, which demonstrates the adapted shape of the electrodes due to different optimization goals. Note that the influence from electrodes of neighbouring traps increases for an increased height $h$. The optimization allows for an increase in the height by more then a factor of 2, still maintaining realistic trapping parameters (see below). Hence, the related motional heating rates (in units of energy per time) are expected to be reduced by more than an order of magnitude. In addition, the increased $h$ should help to protect the electrodes from the high intensity of the laser beams parallel to the electrode surfaces.

We additionally include required isolating gaps between electrodes into subsequent simulations to deduce deviations in the resulting trapping potential [172] (see figure 15). The influence of the gaps turned out to be negligible for the example shown in figure 15, however, for further miniaturized traps these influences will grow in importance due to technically limited gap sizes.

6.2.3. Maximization of the lifetime of trapped ions. The reduced frequencies and increased height above the surface come at the price of a reduced trap depth. First, sufficiently deep potentials have to be provided to assure adequate loading rates out of thermal atomic beams, preferably via efficient photoionization [173, 174]. Second, sufficient lifetimes for many ions within the potentials of scaled traps have to be achieved. Currently, the average lifetime in a room temperature surface-electrode trap exceeds one hour (for the Sandia Linear Trap operated in our laboratory).

Deeper trapping potentials for surface traps were already achieved by a conductive mesh with controlled voltage (85% light transmittance) a few millimetres above the electrode surface [175]. It has also been successfully tested for the Sandia Linear Trap. The mesh shields the ions from charges on the camera viewport and provides a wavelength-independent alternative to a conductive coating (see, for example [176]).

6.2.4. Control of the symmetry of interaction. We additionally gain control over the individual orientations of trap axes or the relative orientations of axes of different traps, respectively (see figure 16). This allows the interaction to be shaped for a given direction of motional excitation between ions in different traps (see section 3). It also allows cooling of all spatial degrees of freedom with laser beams, which have to propagate parallel to the trap surface to minimize scattering off the surface. We can rotate the individual trap axes from pointing towards the centre of the structure (see figure 16(a)) into a parallel alignment and additionally include the required tilt of the vertical (Z) axes, which will result in a different symmetry of the electrodes (see figure 16(b)).

6.2.5. Control of the potential in individual traps. First, splitting dc electrodes into several separately controllable segments allows for the individual compensation of displacements of the ions from the minima of the pseudopotential due to stray fields and space charge effects (compare section 2.1 and see [176, 177] for schemes of micromotion compensation). Second, for further scaling, these electrodes can be used to compensate boundary effects. Due to the larger number of inner ions, outer ones would be shifted to larger mutual distances. The further increased density of electrodes on the surface requires their connections in a multilayer structure with vertical wiring (vias) [159, 160].

Figure 16. Electrode structures for basic triangular lattices with different orientation and tilt of the principal axes. Red disks symbolize ions trapped in the potential minima for parameters comparable to those in figure 15(b). (a) One principal axis points in the vertical direction with the $X$- and $Y$-axis lying in the horizontal plane of the electrodes. The $X$-principal axis of each trap points towards the centre of the triangle. (b) The respective principal axes of all traps point in the same direction and additionally the $Z$-axis is tilted with respect to the surface by more than $10^\circ$, which results in a different symmetry of the electrodes. The tilt of the $Z$-axis is essential to reach all spatial degrees of freedom with laser beams restricted to a plane parallel to the electrodes. (Courtesy of Roman Schmied.)
6.2.6. Estimation of parameters. We estimate the strength of simulated spin–spin interactions for the case of Mg$^+$ ions in such devices with currently available laser equipment. We assume a typical laser power of 400 mW (max. 600 mW are available) at 280 nm from an all solid-state laser source [178]. We further assume the beam to have a cylindrical profile with waists of 10 µm × 100 µm and an electrode structure as depicted in figure 16(b) (d = h = 40 µm). For a trap depth of 100 meV and a minimal oscillation frequency of the ions of $2\pi \times 2$ MHz, the interaction strengths by far exceed $2\pi h \times 1$ kHz.

In a different approach, we could think of using the motional degrees of freedom for QSs. This scheme would have the advantage that bare motional couplings are already in the $2\pi \times 5$ kHz regime. In that sense, they are stronger than effective spin–spin interactions, since the latter are slowed down with respect to the original motional couplings by the requirement of adiabaticity. A recent theoretical proposal by some of us has shown that using periodic modulations of the trapping frequencies, some phenomena from solid-state physics may be simulated, such as photon assisted tunnelling [126] (see section 5.2).

6.3. Perspectives of our approach

As depicted in figure 16(b), in a first step three ions will reside on the vertices of a triangle and the interaction between the spins can be simulated as in [47, 48] (compare section 3.5) or [126] (compare section 5.2). The above parameter estimates should already suffice for proof-of-principle experiments and mesoscopically scaled QS. Motional modes in two-dimensional trap arrays will behave similarly to radial modes in linear RF traps for all three dimensions [21, 124] and the effective spin–spin interaction will prefer antiferromagnetic order for far, red detuning from all modes. Thus, the systems should give us the possibility to study spin frustrations in a spatial, triangular configuration (see also section 5.1).

Based on the results of these investigations further scaling of the surface-trap architecture to large-scale (triangular) lattices of tens or even hundreds of spins might be pursued (see figure 17). In addition to the optimization of the trapping parameters, further technical difficulties have to be considered.

Decoherence due to motional heating as a result of the vicinity to the electrode surfaces could be mitigated within a cryogenic setup [50, 162]. The reduced vacuum pressure could additionally help to increase the lifetime of Mg$^+$, which is currently limited by photochemical reactions with hydrogen (mostly $\text{H}_2 + \text{Mg}^{2+} \rightarrow \text{MgH}^+ + \text{H}^+$) and collisions with heavy components of the rest gas. The reaction can also be inverted by pulsed laser beams [179]. However, scaling the system to tens or hundreds of ions will still require frequent and efficient reloading. Increasing the loading efficiency and preserving the vacuum conditions could be achieved by photoinizing cold atoms from a magneto-optical trap (MOT) [180].

Currently, the available laser power should not impose any restrictions on the realization of systems of few tens of ions (see [178] currently providing up to 600 mW). Higher laser powers for magnesium are in reach [181–183] and could allow for even larger arrays of simultaneously coupled ions. In addition to that, efforts in optics, for example arrays of lenses [184, 185], fibres integrated into the trap [186] or integrated mirrors [187–189], could provide individual addressing and high light intensities at the position of the ions. To further mitigate the problem of scattered light from surfaces, one could think of realizing surface traps on partially transparent substrates [74]. Alternatively, laser-less coupling could be used as mentioned in section 2.3 [88, 89, 91–93, 168].

Last but not least, it still has to be identified how to measure observables that permit the verification of frustration effects without the need for full (exponentially complex) state tomography.

7. Scaling quantum simulations based on ions in optical lattices

Some groups aim to merge the two fields of QS based on ions in RF traps and atoms confined in optical lattices. It has already been proposed to combine Coulomb crystals in a harmonic confinement of a common RF trap of three-dimensional geometry with (commensurate) optical lattices to shape anharmonic trapping potentials providing new possibilities to simulate interactions [157]. Another proposal deals with the
simulation of the Frenkel–Kontorova model using a standing wave aligned with the trap axis [156]. Kollath et al suggested exploiting a trapped ion to coherently couple (such as a scanning microscope) to the atoms confined in an optical lattice [190].

Optical ion trapping was realized with a single Mg$^+$ ion trapped in a dipole trap [80]. We can now dream of spanning an array of ions (even simultaneously with neutral atoms) within an optical lattice. It has to be emphasized that the smaller trap depth of optical traps (see figure 1) renders it highly unlikely that optically trapping charged atoms will allow us to outperform the achievable trapping parameters or coherence times of both, ions in RF traps and of optically trapped neutral atoms. However, in our opinion, this is not required. The advantage of equally and closely spaced traps might be combined with individual addressability and, most importantly, the long-range interaction provided by Coulomb forces between the ions.

In the following, we will first describe how trapping of an ion in a dipole trap was achieved. Still facing a huge variety of challenges, the new possibilities will be discussed afterwards.

7.1. Trapping of an ion in a dipole trap

The procedure used in [80] to load a magnesium ion ($^{24}$Mg$^+$) into a dipole trap consists of the following steps: an atom is photoionized out of a thermal beam and trapped and Doppler cooled in a conventional RF trap. Next, stray electric fields are minimized at the site of the ion using the ion as a sensor. Then a Gaussian laser beam providing the dipole trap is focused onto the ion and the RF drive of the RF trap is switched off. From that time on, the ion is confined in the dipole trap in the directions perpendicular to the beam propagation. The depth of the dipole trap potential amounts to $U_0 \approx 2\pi\hbar \times 800$ MHz or $U_0 \approx k_B \times 38$ mK, respectively, and the detuning of the dipole trap beam from the relevant transition ($S_{1/2}$ $\leftrightarrow$ $P_{3/2}$) to $\Delta \approx -6600\Gamma$, where $1/\Gamma$ determines the lifetime of the $|P_{3/2}\rangle$ state. Static electric fields provide the confinement in the direction of beam propagation. After a few milliseconds the RF drive is switched on again and the presence of the ion can be verified via its fluorescence during Doppler cooling.

For the given parameters a half-life of approximately 2.5 ms is achieved. This value is in very good agreement with the theoretical predictions, assuming exclusively the heating process related to off-resonant scattering of the trapping light by the ion, the so-called recoil heating. It can be concluded that the heating and subsequent loss of ions from the optical potential is not dominated by heating effects related to the charge of the ion, for example, due to the vicinity of electrodes or fluctuating stray electric fields. Thus, state-of-the-art techniques for neutral atoms should allow effective enhancement of the lifetime and coherence times [191].

7.2. Lifetime and coherence times of optically trapped ions

We aim to increase the lifetime by cooling the ion in the dipole trap. Due to the large ac Stark shift and its large position dependence, simple Doppler cooling within the existing setup is challenging. Possibilities of cooling the ions directly towards the ground state of motion within the dipole potential are currently being investigated theoretically and experimentally.

An alternative approach suggests to use cold atoms or even a BEC to sympathetically cool ions [192]. On longer timescales the approach of cavity assisted cooling of ions in conventional RF traps reported in [193] might also provide long lifetimes without affecting the electronic state of the ion.

Currently, the coherence time of the electronic state of the ion is limited to a few microseconds due to the high spontaneous emission rate. If longer coherence times are required (which is not necessarily the case for every scenario), they can be achieved in two ways. (1) As for two-photon stimulated-Raman transitions the spontaneous emission rate can be reduced by increasing the detuning. A larger beam intensity could sustain the potential depth. (2) Another option would be to work with blue detuned light, where the potential depths can remain identical, however, the ions seek low intensity and exhibit less spontaneous emission.

7.3. Towards ions and atoms in a common optical lattice

It has still to be demonstrated that one or several ions can be confined within one- or more-dimensional optical lattices. With currently available laser sources a mutual ion distance within each dimension of the order of tens of micrometres could be achieved, which corresponds to one ion at approximately every 40th to 50th lattice site (see methods in [80]). Therefore the mutual ion distance could be smaller than the currently envisioned distances between neighbouring traps in the RF surface-electrode trap approach (see section 6).

Since the photoionization scheme applied so far ionizes out of a thermal beam of magnesium atoms, the average kinetic energy of the atoms is much larger than the depth of the optical potential and, in addition, the local vacuum is severely affected. The loading efficiency for RF traps could be largely enhanced by ionizing Mg atoms from a MOT [194], which would also allow direct loading of atoms into an optical trap. In addition, after loading neutral atoms into the lattice, some of them could be photoionized on site.

Ions and atoms confined in a common optical lattice could offer an approach to exploit the physics of charge transfer reactions. This might allow for a complete new class of QS, for example, of solid-state systems, where atoms in a completely occupied lattice (at an initially small density of ions) share electrons by tunnelling causing highly entangled states of the compound system. The resulting quantum many-body dynamics should be governed by the interplay of the quantum state of the trapped neutral atoms and the electron tunnelling from neutrals to ions.

8. Conclusions

In the last few years the basic building blocks for a scalable architecture of a quantum information processor (QC) with trapped ions have been demonstrated for a few qubits. Additionally, a large variety of new techniques have already been tested that might considerably extend the available
toolbox. For example, interactions based on magnetic field gradients and RF fields, fibre-coupled optical support on chip or economically and technologically facilitated cryogenic environments. Despite the fact that it will be a non-trivial challenge to scale the system to approximately $10^5$ qubits, no fundamental limitations have been identified so far.

On a shorter timescale, intriguing problems might be studied by realizing analogue quantum simulators (QS), by far exceeding the capabilities of classical computers. They can be based on similar techniques as a potential QC, but with less severe constraints on the fidelity of operations and the number of required ions.

Currently available operational fidelities are predicted to allow for studying many-body physics, for example in systems described by quantum spin Hamiltonians, the Bose–Hubbard and the spin–boson models. First proof-of-principle experiments simulating Ising type interactions with a few ions have already been successfully demonstrated.

The required increase of the number of ions and the accessible dimensions is proposed within two-dimensional arrays of RF surface-electrode traps. However, the approach is still at the level of proof-of-principle experiments and further challenges might arise during its development. Alternative approaches include Penning traps or optical lattices.

Even though the enthusiasm within this quickly growing field seems to be justified, it has to be emphasized that efficient analogue QS still require more than simply scaling. Examples of other important challenges are (1) to investigate carefully the influence of different sources of decoherence on the fidelity of the simulation. Thus, it must be distinguished for the dedicated application, which decoherence the simulation will be robust against, which decoherence can be considered in the simulation and which decoherence is even essential to be included. (2) To identify possibilities to cross-check the validity of the output or to benchmark it against other QS approaches, as soon as the achieved output is not accessible with a classical computer anymore.

In the future it might be beneficial to combine advantages of several systems for hybrid QS. On longer timescales, the experiences gained by developing an analogue QS based on trapped ions might culminate in approaches incorporating solid-state devices that might allow for ‘easier’ scaling. With the realization of a universal QC, universal QS will also become accessible.

Acknowledgments

CS and TS acknowledge support by the Max-Planck-Institut für Quantenoptik (MPQ), Max-Planck-Gesellschaft (MPG), Deutsche Forschungsgemeinschaft (DFG) (SCHA 973/1-6), the European Commission (The Physics of Ion Coulomb Crystals: FP7 2007–2013. grant no. 249958) and the DFG Cluster of Excellence ‘Munich Centre for Advanced Photonics’. DP acknowledges support from C.A.M. Project QUTEMAD, RyC Contract Y200200074, and MICINN FIS2009-10061. Martin Enderlein, Thomas Huber, and Hector Schmitz have participated in the measurements of the geometric phase gate utilizing the radial modes of motion presented in section 3.4. Günther Leschhorn and Steffen Kahra have observed and taken the picture of the structural defect in a Coulomb crystal of Mg$^+$ ions (see figure 13(b)). Benni Reznik and Haggai Landa provided the corresponding simulation (see figure 13(a)). Roman Schmied kindly provided several simulation results and illustrations of optimized electrode structures of surface-electrode traps (see section 6). We want to thank Geza Giedke and Alex Retzker for discussions and Martin Enderlein, Thomas Huber, and Dietrich Leibfried for comments on the manuscript. We also thank Ignacio Cirac and Gerhard Rempe for their intellectual and financial support.

Appendix A. Normal modes and frequencies

This section describes a more general derivation of the normal modes and frequencies compared with the one-dimensional treatment for the linear Paul trap as in [195]. The equations are extended to three dimensions and an arbitrary trap potential, as long as the potential at the equilibrium position $\vec{x}_i^0$ of each ion can be well approximated by a harmononic potential.

The position of the $i$th ion is expressed in the Cartesian coordinates of the laboratory frame

$$\vec{r}^{(i)} = r_i \vec{e}_x + r_{i+N} \vec{e}_y + r_{i+2N} \vec{e}_z.$$  \hfill (A.1)

The decomposition into the equilibrium position $\vec{x}_i^0$ and displacements $\vec{z}^{(i)}$ yields

$$\vec{r}^{(i)} = \vec{x}_i^0 + \vec{z}^{(i)}$$  \hfill (A.2)

$$= (x_{0,i} + x_i) \vec{e}_x + (x_{0,i+N} + x_{i+N}) \vec{e}_y + (x_{0,i+2N} + x_{i+2N}) \vec{e}_z.$$  \hfill (A.3)

The Lagrangian for $N$ ions takes the form

$$\mathcal{L} = \frac{1}{2} M \left[ \sum_{k=1}^{3N} \dot{x}_k^2 - \sum_{k=1}^{3N} \sum_{j=1}^{3N} \frac{\partial^2 V}{\partial r_k \partial r_j} (x_k = x_j = 0) \right]_{\omega = 0},$$  \hfill (A.4)

where $M$ denotes the mass of an ion, the index of the partial derivatives signifies its evaluation at the equilibrium positions and $V$ denotes the potential consisting of the trap potential $V_0$ and the Coulomb potentials of the ions:

$$V = V_0 + \frac{Q^2}{8\pi \epsilon_0} \sum_{i=1}^{N} \sum_{j \neq i}^{N} \frac{1}{|\vec{r}^{(i)} - \vec{r}^{(j)}|}.$$  \hfill (A.5)

Here, $Q$ denotes the charge and $\epsilon_0$ the electric constant.

For practical purposes the trap potential can be expressed by the harmonic terms corresponding to each ion:

$$V_0 = \frac{1}{2} M \sum_{i=1}^{N} \sum_{j=1}^{3} \omega_j^{(i)} \left( \vec{z}^{(i)} - \vec{p}^{(i)} \right) \hat{a}_j^{(i)} \otimes \hat{a}_j^{(i)} (\vec{r}^{(i)} - \vec{p}^{(i)}).$$  \hfill (A.6)

Here, $\omega_j^{(i)}$ denotes the $j$th frequency of the harmonically approximated potential of the $i$th ion, $\hat{a}_j^{(i)}$ the unity vector of the principle axis corresponding to $\omega_j^{(i)}$ and $\vec{p}^{(i)}$ the position of
the local minimum of the potential for the \(i\)th ion. Note that the frequencies, the vectors of the principle axes, and the minima of the potentials become equal for all ions in the special case of a linear Paul trap.

The eigenvalues of the Hessian \(A := (a_{ij})\) (see equation (A.4)) yield the squares of the frequencies \(\omega_m\) of the normal modes and its eigenvectors \(\vec{b}_m\) determine the ions’ motion of the \(m\)th mode:

\[
q_m = \vec{b}_m \cdot \vec{x} \quad \text{with} \quad \vec{x} := (x_1, \ldots, x_{3N}). \tag{A.7}
\]

With the abbreviations \(\vec{q} := (q_1, \ldots, q_{3N})\) and \(B := (\vec{b}_1, \ldots, \vec{b}_{3N})^T\), where the \(\vec{b}_m\) shall be understood as rows of \(B\), we can express the relation in a more compact way:

\[
\vec{q} = B \vec{x} \iff \vec{x} = B^T \vec{q}. \tag{A.8}
\]

Typically, the eigenvalues and eigenvectors of \(A\) have to be determined numerically.

### Appendix B. Transformations of Pauli operators

The definitions of the Pauli operators are repeated here to avoid confusions concerning their normalization:

\[
\hat{\sigma}_x := \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}, \quad \hat{\sigma}_y := \begin{pmatrix} 0 & i \\ i & 0 \end{pmatrix}, \quad \hat{\sigma}_z := \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}. \tag{B.1}
\]

The Pauli operators obey the relations

\[
[\hat{\sigma}_i, \hat{\sigma}_j] = 2i\epsilon_{ijk}\hat{\sigma}_k, \tag{B.2}
\]

\[
[\hat{\sigma}_i, \hat{\sigma}_j] = 2\delta_{ij}, \tag{B.3}
\]

\[
\hat{\sigma}_i^2 = \hat{1}. \tag{B.4}
\]

A more convenient notation in some contexts is

\[
\hat{\sigma}_+ := \hat{\sigma}_x + i\hat{\sigma}_y = \begin{pmatrix} 0 & 2 \\ 2 & 0 \end{pmatrix}, \tag{B.5}
\]

\[
\hat{\sigma}_- := \hat{\sigma}_x - i\hat{\sigma}_y = \begin{pmatrix} 0 & 0 \\ 0 & 2 \end{pmatrix} \tag{B.6}
\]

with the normalization as in [50]. They fulfill the following relations:

\[
[\hat{\sigma}_+, \hat{\sigma}_-] = \pm 4\hat{\sigma}_z, \tag{B.7}
\]

\[
[\hat{\sigma}_z, \hat{\sigma}_\pm] = \pm 2\hat{\sigma}_\pm. \tag{B.8}
\]

The transformations of the Pauli operators into the interaction picture involve terms of the form

\[
\hat{\sigma}_i' := e^{ik\hat{\sigma}_i} \hat{\sigma}_i e^{-ik\hat{\sigma}_i}. \tag{B.9}
\]

The transformation leaves \(\hat{1}\) and \(\hat{\sigma}_z\) unchanged. The non-trivial cases \(i = x\) and \(i = y\) can be calculated using the Baker–Campbell–Hausdorff formula

\[
e^{-\hat{\theta} \hat{A}} \hat{e}^\hat{B} = \sum_n \frac{1}{n!} [\hat{A}, \hat{B}]_n^{(n)} \\
= \hat{A} + [\hat{A}, \hat{B}] + \frac{1}{2} [[\hat{A}, \hat{B}], \hat{B}] + \cdots \tag{B.10}
\]

with \(\hat{B} = -i\hbar \dot{\hat{A}}, \hat{\dot{A}} = \dot{\hat{A}}_{\perp} = \hat{\sigma}_z \hat{\dot{\hat{A}}}_z - m_+ := 2\) and \(m_- := 2i\).

The commutators are given by

\[
[\hat{\sigma}_+ \pm \hat{\sigma}_-, \dot{\hat{\sigma}}_z] = \pm 2\hbar \left( \hat{\sigma}_z \mp \hat{\sigma}_- \right), \tag{B.11}
\]

\[
[\hat{\sigma}_+ \pm \hat{\sigma}_-, \dot{\hat{\sigma}}_z]^{(2)} = 2\hbar \left( \hat{\sigma}_z \mp \hat{\sigma}_- \right), \tag{B.12}
\]

\[
[\hat{\sigma}_+ \pm \hat{\sigma}_-, \dot{\hat{\sigma}}_z]^{(2n-1)} = (2\hbar)^{n-1} \left( \hat{\sigma}_z \mp \hat{\sigma}_- \right), \tag{B.13}
\]

\[
[\hat{\sigma}_+ \pm \hat{\sigma}_-, \dot{\hat{\sigma}}_z]^{(2n)} = (2\hbar)^n \left( \hat{\sigma}_z \pm \hat{\sigma}_- \right). \tag{B.14}
\]

Hence, the Pauli operators in the interaction picture read

\[
e^{i\hbar \dot{\hat{\sigma}}_z / \hbar} \hat{\sigma}_\pm e^{-i\hbar \dot{\hat{\sigma}}_z / \hbar} = e^{i\hbar \dot{\hat{\sigma}}_z / \hbar} \hat{\sigma}_\pm e^{-i\hbar \dot{\hat{\sigma}}_z / \hbar} = \frac{1}{m_\pm} \sum_n (2\hbar)^n \left( \hat{\sigma}_z \mp \hat{\sigma}_- \right). \tag{B.15}
\]

The operators \(\hat{\sigma}_\pm\) transform as:

\[
e^{i\hbar \dot{\hat{\sigma}}_z / \hbar} \hat{\sigma}_+ e^{-i\hbar \dot{\hat{\sigma}}_z / \hbar} = e^{i\hbar \dot{\hat{\sigma}}_z / \hbar} \hat{\sigma}_+ e^{-i\hbar \dot{\hat{\sigma}}_z / \hbar} = e^{-i\hbar \dot{\hat{\sigma}}_z / \hbar}. \tag{B.16}
\]

### Appendix C. Transformations of motional operators

The creation operator \(\hat{a}\) and the annihilation operator \(\hat{a}^\dagger\) fulfil the relations

\[
\hat{a} |n\rangle = \sqrt{n} |n - 1\rangle, \tag{C.1}
\]

\[
\hat{a}^\dagger |n\rangle = \sqrt{n + 1} |n + 1\rangle, \tag{C.2}
\]

\[
[\hat{a}, \hat{a}^\dagger] = \hat{1}. \tag{C.3}
\]

The transformation of the Hamiltonians into the interaction picture requires the knowledge of the transformation of \(e^{i\hbar \hat{a}^\dagger \hat{a}}\).

It can be performed using the special case of the Baker–Campbell–Hausdorff formula from appendix B again. The commutators appearing in the formula are

\[
[\hat{a} + \hat{a}^\dagger, -i\hbar \hat{a}^\dagger \hat{a}] = -i\hbar (\hat{a} - \hat{a}^\dagger), \tag{C.4}
\]

\[
[\hat{a} + \hat{a}^\dagger, -i\hbar \hat{a}^\dagger \hat{a}]^{(2)} = -i\hbar [\hat{a} - \hat{a}^\dagger, -i\hbar \hat{a}^\dagger \hat{a}] = -(\hbar)^2 (\hat{a} + \hat{a}^\dagger), \tag{C.5}
\]

\[
[\hat{a} + \hat{a}^\dagger, -i\hbar \hat{a}^\dagger \hat{a}]^{(n)} = -(\hbar)^n (\hat{a} + (-1)^n \hat{a}^\dagger). \tag{C.6}
\]

Hence, the full transformation reads

\[
e^{i\hbar \hat{a}^\dagger \hat{a}} \hat{a} e^{-i\hbar \hat{a}^\dagger \hat{a}} = \sum_n (\hbar)^n (\hat{a} + (-1)^n \hat{a}^\dagger) \tag{C.8}
\]

\[
= \hat{a} e^{-i\hbar} + \hat{a}^\dagger e^{i\hbar}. \tag{C.8}
\]
From this relation we can immediately derive
\[
\begin{align*}
\hat{a} e^{i\lambda \hat{a}} \hat{a} e^{-i\lambda \hat{a}} &= \left[ e^{i\lambda \hat{a}} (\hat{a} + \hat{a}^\dagger) e^{-i\lambda \hat{a}} \right]^n \\
&= \left[ \hat{a} e^{-i\lambda} + \hat{a}^\dagger e^{i\lambda} \right]^n
\end{align*}
\] (C.9)
by making use of the unitarity of \(e^{i\lambda \hat{a}}\). We obtain for the transformation
\[
\begin{align*}
e^{i\lambda \hat{a}} e^{i\xi} (\hat{a} + \hat{a}^\dagger) e^{-i\lambda \hat{a}} &= e^{i\lambda \hat{a}} \sum_n \binom{\text{tr}}{n} \frac{i^\dagger}{n!} (\hat{a} + \hat{a}^\dagger)^n e^{-i\lambda \hat{a}} \\
&= \sum_n \binom{\text{tr}}{n} \left[ \hat{a} e^{-i\lambda} + \hat{a}^\dagger e^{i\lambda} \right]^n \\
&= \exp \left( i \xi \left[ \hat{a} e^{-i\lambda} + \hat{a}^\dagger e^{i\lambda} \right] \right)
\end{align*}
\] (C.10)

**Appendix D. Matrix elements of displacement operator**

The following derivation is based on [95]. A difference is that we do not restrict the displacement to purely imaginary \(\lambda\) in the following. A similar derivation can be found in appendix B of [111].

The simple Baker–Campbell–Hausdorff formula
\[
e^{A+B} = e^A e^B e^{-[A,B]/2}
\] (D.1)
holds for \([A, [A, B]] = [B, [A, B]] = 0\).

We obtain for the annihilation operator
\[
\hat{a}^m |n\rangle = \begin{cases} \sqrt{\frac{n!}{(n-m)!}} |n-m\rangle & \text{for } m \leq n, \\ 0 & \text{else}. \end{cases}
\] (D.2)

Using the above form of the Baker–Campbell–Hausdorff formula, we can rewrite the displacement operator as
\[
\hat{D}(\lambda) = e^{i\lambda \hat{a}^\dagger - \lambda^* \hat{a}} = e^{-|\lambda|^2/2} e^{i\lambda \hat{a}^\dagger} e^{-\lambda \hat{a}}
\] (D.3)

With
\[
e^{-\lambda \hat{a}} |n\rangle = \sum_m \frac{(-\lambda)^m}{m!} \hat{a}^m |n\rangle
\] (D.4)
this yields for \(n' \geq n\)
\[
|n'\rangle \langle n' | \hat{D}(\lambda) |n\rangle = e^{-|\lambda|^2/2} \sum_{m=0}^{n-n'} |m\rangle \langle m | \frac{(-\lambda)^m}{m!}
\times \left( \frac{n!}{(n'-m)!} \right)^{1/2} \left( \frac{n!}{(n-m)!} \right)^{1/2}
= e^{-|\lambda|^2/2 |n'-n|} \sum_{m=0}^{n-n'} \frac{(-1)^m |\lambda|^2 m}{m! \sqrt{(n'-n)!} \sqrt{(n-m)!}} \\
\times |n'\rangle \langle n' | L_m^{n-n'} (|\lambda|^2)
\] (D.5)
where \(L_m^{n-n'} (|\lambda|^2)\) denotes the generalized Laguerre polynomials [196]. Analogously, we obtain for \(n' \leq n\)
\[
|n\rangle \langle n | \hat{D}(\lambda) |n'\rangle = \langle n' | \hat{D}(\lambda) |n\rangle = e^{-|\lambda|^2/2} \left( -\lambda^\dagger \right)^{n-n'} \frac{n!}{n'} L_m^{n-n'} (|\lambda|^2). \tag{D.6}
\]

For values \(\lambda = i\eta e^{i \theta_0}\) with \(\eta \in \mathbb{R}\), we can write equations (D.5) and (D.6) as
\[
|n\rangle \langle n | \hat{D}(i \eta e^{i \theta_0}) |n'\rangle = e^{-\eta^2/2} \langle n' | e^{i \theta_0} |n\rangle |\eta'|^{n-n'} | e^{i \theta_0} |n'-n| \times \sqrt{\frac{n!}{n_{\perp}}} L_m^{n-n'} (\eta^2),
\] (D.7)
where \(n_{\perp} := \min(n', n)\) and \(n_{\parallel} := \max(n', n)\).

**Appendix E. System of differential equations of the Rabi problem**

The Rabi problem consists of the following system of differential equations:
\[
\begin{align*}
\dot{c}_1 &= \frac{\lambda}{2} c_2 \\
\dot{c}_2 &= -\frac{\lambda}{2} c_1 - i \omega \eta c_2
\end{align*}
\] (E.1)

It can be solved by differentiating with respect to \(t\)
\[
\begin{align*}
\dot{c}_2 &= \lambda e^{-i \omega t} c_1 - i \omega \eta c_2 \\
\dot{c}_1 &= -\lambda e^{-i \omega t} c_2 - i \omega \eta c_1
\end{align*}
\] (E.2)

and inserting equation (E.1):
\[
\begin{align*}
\dot{c}_2 &= -i \omega \dot{c}_2 - |\lambda|^2 c_2 \\
\dot{c}_1 &= i \omega \dot{c}_1 - |\lambda|^2 c_1
\end{align*}
\] (E.3)

Using the ansatz \(c_1 = a_1 e^{i \theta_0}\) we obtain the characteristic equations
\[
-\kappa_2^2 = \omega \kappa_2 - |\lambda|^2, \quad (E.5)
-\kappa_2^2 = -\omega \kappa_1 - |\lambda|^2, \quad (E.6)
\]
which have the solutions
\[
\kappa_2 = -\frac{\omega}{2} \pm \frac{\sqrt{\omega^2 + |\lambda|^2}}{2}, \quad (E.7)
\kappa_1 = \frac{\omega}{2} \pm \frac{\sqrt{\omega^2 + |\lambda|^2}}{2}, \quad (E.8)
\]
Here, we introduced the abbreviation \(\kappa_1 := \sqrt{(|\omega|^2/4) + |\lambda|^2}\).

The solutions of equations (E.3) and (E.4) read
\[
\begin{align*}
c_2 &= a_2 e^{i \theta_2 e^{-\kappa_2 t}} + a_{-2} e^{-i \theta_2 e^{-\kappa_2 t}} \\
&= \left( a_2 e^{i \theta_2} + a_{-2} e^{-i \theta_2} \right) e^{-i \kappa_2 t/2}, \quad (E.9)
c_1 &= a_1 e^{i \theta_1 e^{-\kappa_1 t}} + a_{-1} e^{-i \theta_1 e^{-\kappa_1 t}} \\
&= \left( a_1 e^{i \theta_1} + a_{-1} e^{-i \theta_1} \right) e^{i \kappa_1 t/2}. \quad (E.10)
\end{align*}
\]

Inserting them into the original system of differential equations equation (E.1), we obtain the following relations for the constants \(a_{1,2,\pm}\):
\[
\begin{align*}
\kappa_{2,\pm} a_{2,\pm} &= \lambda a_{1,\pm}, \quad (E.11)
\kappa_{1,\pm} a_{1,\pm} &= -\lambda a_{2,\pm}. \quad (E.12)
\end{align*}
\]
We replace $a_{1,\pm}$ by $a_{2,\pm}$ using equation (E.11) and obtain

$$c_2 = (a_{2,\pm} e^{i\omega t} + a_{-2,\pm} e^{-i\omega t}) e^{-i\omega t/2}, \quad (E.13)$$

$$c_1 = (\mu a_{2,\pm} e^{i\omega t} + \mu a_{-2,\pm} e^{-i\omega t}) e^{i\omega t/2}, \quad (E.14)$$

where we introduced the (temporary) abbreviation $\mu_{\pm} := i\kappa_{\pm}/\lambda$.

The constants $a_{2,\pm}$ can now be expressed in terms of the initial values $c_{20} := c_{2}(t = 0)$ and $c_{10} := c_{1}(t = 0)$. Setting $t = 0$ in equations (E.13) and (E.14), we obtain a system of linear equations with the solutions

$$a_{2,\pm} = \frac{\mu_c c_{20} - c_{10}}{\mu_c - \mu_+}, \quad (E.15)$$

$$a_{-2,\pm} = -\frac{\mu_c c_{20} - c_{10}}{\mu_+ - \mu_c}, \quad (E.16)$$

Hence,

$$c_2 = \frac{(\mu_c c_{20} - c_{10}) e^{i\omega t} - (\mu_c c_{20} - c_{10}) e^{-i\omega t}}{\mu_+ - \mu_-} e^{-i\omega t/2}, \quad (E.17)$$

$$c_1 = -\frac{\mu_c (\mu c_{20} - c_{10}) e^{i\omega t} - \mu_+ (\mu c_{20} - c_{10}) e^{-i\omega t}}{\mu_+ - \mu_-} e^{i\omega t/2}, \quad (E.18)$$

and by expressing $\mu_{\pm}$ in terms of $\omega$, $\kappa_{\pm}$, and $\lambda$, we obtain

$$c_2(t) = \left(\cos(\kappa t) + i \frac{\omega}{2 \kappa} \sin(\kappa t)\right) e^{-i\omega t/2} c_2(0) + \frac{\lambda \kappa}{\kappa} \sin(\kappa t) e^{-i\omega t/2} c_1(0), \quad (E.19)$$

$$c_1(t) = -\frac{\lambda \kappa}{2} \sin(\kappa t) e^{-i\omega t/2} c_2(0) + \left(\cos(\kappa t) - i \frac{\omega}{2 \kappa} \sin(\kappa t)\right) e^{i\omega t/2} c_1(0). \quad (E.20)$$

### Appendix F. Time evolution operator

The calculation of the time evolution operator involves terms of the form

$$\hat{T}^{(i)}(t) = i \xi_m^{(i)} e^{i\delta_m(t-t')} \hat{a}_m^\dagger + \text{h.c.}, \quad (F.1)$$

where $\xi_m^{(i)} \in \mathbb{R}$ and the total Hamiltonian reads $\hat{H}(t) := \sum_{i=1}^{N} \sum_{m=1}^{2N} \hat{H}_m^{(i)}(t)$. (More generally, the constants $\xi_m^{(i)}$ represent Hermitian operators $\hat{\xi}_m^{(i)}$ with $[\hat{\xi}_m^{(i)}, \hat{\xi}_n^{(j)}] = 0\forall i, j, m, n$.)

The commutator of two of these terms will trivially vanish for all $i, j$ and all times $t', t''$, if both terms belong to different modes $m \neq n$:

$$[\hat{T}^{(i)}(t'), \hat{T}^{(j)}(t'')] = 0 \quad \text{for } m \neq n. \quad (F.2)$$

However, for $m = n$, the commutators do not vanish. Using the relation

$$[e^{\lambda \hat{a}^\dagger} e^{-i\kappa \hat{a}} - e^{i\kappa \hat{a}^\dagger} e^{-\lambda \hat{a}}] = e^{i(\lambda - \kappa)} (\hat{a} \hat{a}^\dagger - \hat{a}^\dagger \hat{a}) - e^{i(\lambda - \kappa)} (\hat{a} \hat{a}^\dagger - \hat{a}^\dagger \hat{a})$$

$$= 2i \sin(\lambda - \kappa) \hat{I} \quad \text{(F.3)}$$

yields

$$[\hat{T}_m^{(i)}(t'), \hat{T}_n^{(j)}(t'')] = 2i \sin(\delta_m(t' - t'') - (\psi^{(i)} - \psi^{(j)})).$$

The time evolution operator can be calculated using a Magnus expansion [197, 198]. As commutators with higher ‘nesting level’ trivially vanish, the expansion simplifies to

$$\hat{U}(t, t_0) = \exp \left( -\frac{i}{\hbar} \int_{t_0}^{t} dt' \hat{H}(t') \right) - \frac{1}{2\hbar^2} \int_{t_0}^{t} dt' \int_{t_0}^{t} dt'' [\hat{H}(t'), \hat{H}(t'')] \quad \text{(F.5)}$$

The single integrals yield

$$\int_{t_0}^{t} dt' \hat{V}_m^{(i)}(t') = -\frac{\xi_m^{(i)}}{\delta_m} \left( e^{-i\kappa_n(t-t_0)} - 1 \right) e^{-i\kappa_n t} e^{i\psi^{(i)}} \hat{a}_m^\dagger + \text{h.c.} \quad \text{(F.6)}$$

and the double integrals of the commutators yield

$$\int_{t_0}^{t} dt' \int_{t_0}^{t} dt'' [\hat{V}_m^{(i)}(t'), \hat{V}_n^{(j)}(t'')] = 2i \xi_m^{(i)} s_{m,n}^{(i,j)} \quad \text{(F.7)}$$

Note that in the time evolution operator corresponding to $\hat{H}(t)$ the terms $\sin(\psi^{(i)} - \psi^{(j)}) = -\sin(\psi^{(i)} - \psi^{(j)})$ cancel each other.

### Appendix G. Canonical transformation

The unitary operator of the canonical transformation has the form

$$\hat{U}_c := e^{-\frac{i}{\hbar}(\lambda \hat{a}^\dagger - \kappa \hat{a})} \quad \text{with } \lambda := \frac{\xi}{\kappa} \quad \text{(G.1)}$$

and is applied to

$$\hat{H} := (\xi \hat{a}^\dagger + \xi^* \hat{a}) + (-\kappa \hat{a}^\dagger \hat{a}) \rightarrow \hat{H}' = \hat{U}_c \hat{H} \hat{U}_c^\dagger. \quad \text{(G.2)}$$

Here, the constants $\xi \in \mathbb{C}$ and $\kappa \in \mathbb{R}$.

We use the Baker–Campbell–Hausdorff formula from appendix B to do the transformation. The commutators for $\hat{H}_1$ yield

$$[\xi \hat{a}^\dagger + \xi^* \hat{a}, \lambda \hat{a}^\dagger - \kappa \hat{a}] = \xi \lambda (\hat{a}^\dagger \hat{a} + \hat{a} \hat{a}^\dagger) - \xi \kappa (\hat{a}^\dagger \hat{a} + \hat{a} \hat{a}^\dagger)$$

$$= \left( \xi \lambda^* + \xi^* \kappa \right) \hat{I} \quad \text{(G.3)}$$

$$= \frac{2 \xi \kappa}{\kappa} \hat{I}.$$
Commutators with higher ‘nesting levels’ trivially vanish. Hence, the complete transformation of $\hat{H}_1$ reads

$$\hat{H}'_1 := \hat{U} \hat{H}_1 \hat{U}^\dagger = (\xi \hat{a}^\dagger + \xi^* \hat{a}) + \frac{2\xi \xi^*}{\kappa} \hat{1}. \quad (G.4)$$

The commutators for $\hat{H}_2$ read

$$[-\kappa \hat{a}^\dagger \hat{a}, \lambda \hat{a}^\dagger - \lambda^* \hat{a}] = -\kappa (\hat{a}^\dagger [\hat{a}, \lambda \hat{a}^\dagger - \lambda^* \hat{a}]) + [\hat{a}^\dagger, \lambda \hat{a}^\dagger - \lambda^* \hat{a}]$$

$$= -\kappa (\lambda \hat{a}^\dagger + \lambda^* \hat{a})$$

$$= -(\xi \hat{a}^\dagger + \xi^* \hat{a})$$

and

$$[-\kappa \hat{a}^\dagger \hat{a}, \lambda \hat{a}^\dagger - \lambda^* \hat{a}]^{[2]} = -(\xi \hat{a}^\dagger + \xi^* \hat{a}), \lambda \hat{a}^\dagger - \lambda^* \hat{a}$$

$$= -\frac{2\xi \xi^*}{\kappa} \hat{1}, \quad (G.5)$$

where we used equation (G.3). Higher order terms in the expansion trivially vanish again. The complete transformation of $\hat{H}_2$ reads

$$\hat{H}'_2 := \hat{U} \hat{H}_2 \hat{U}^\dagger = -\kappa \hat{a}^\dagger \hat{a} - (\xi \hat{a}^\dagger + \xi^* \hat{a}) - \frac{2\xi \xi^*}{\kappa} \hat{1}. \quad (G.7)$$

Hence, the transformation of the full Hamiltonian reads

$$\hat{H}' = \hat{H}'_1 + \hat{H}'_2 = \frac{\xi \xi^*}{\kappa} \hat{1} - \kappa \hat{a}^\dagger \hat{a}. \quad (G.8)$$

We want to stress that the canonical transformation is exact in this case.

References

[29] Lamata L, León J, Schütz T and Solano E 2007 Dirac equation and quantum relativistic effects in a single trapped ion Phys. Rev. Lett. 98 253005


[183] Toptica Photonics AG 2010 private communication