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<tr>
<td><strong>Publication date</strong></td>
<td>2013</td>
</tr>
<tr>
<td><strong>Type of publication</strong></td>
<td>Article (peer-reviewed)</td>
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</table>
[http://dx.doi.org/10.1103/PhysRevA.87.050304](http://dx.doi.org/10.1103/PhysRevA.87.050304) |
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Entangling two defects via a surrounding crystal

T. Fogarty,1,2 E. Kajari,3 B. G. Taketani,3 A. Wolf,3 Th. Busch,1,2 and Giovanna Morigi3,4
1Physics Department, University College Cork, Cork, Ireland
2Quantum Systems Unit, Okinawa Institute of Science and Technology, Okinawa 904-0495, Japan
3Theoretische Physik, Universität des Saarlandes, D-66123 Saarbrücken, Germany
4Departament de Física, Universitat Autònoma de Barcelona, E-08193 Bellaterra, Spain
(Received 6 August 2012; published 28 May 2013)

We theoretically show how two impurity defects in a crystalline structure can be entangled through coupling with the crystal. We demonstrate this with a harmonic chain of trapped ions in which two ions of a different species are embedded. Entanglement is found for sufficiently cold chains and for a certain class of initial, separable states of the defects. It results from the interplay between localized modes which involve the defects and the interposed ions, it is independent of the chain size, and it decays slowly with the distance between the impurities. These dynamics can be observed in systems exhibiting spatial order; viable realizations are optical lattices, optomechanical systems, or cavity arrays in circuit QED.

DOI: 10.1103/PhysRevA.87.050304 PACS number(s): 03.67.Bg, 03.65.Yz, 03.67.Mn, 42.50.Dv

Entanglement is a quantum mechanical property with no classical counterpart. One of its peculiar features are the correlations between the measurement outcomes for two entangled objects, even when they are at large distances. This makes entanglement a precious resource for quantum communication protocols and quantum metrology. On the other hand, its quantum mechanical nature makes it fragile against external perturbations. The issue of identifying open-system dynamics in which entanglement is robust recently motivated a series of studies, which showed that a bath can mediate entanglement between two objects interacting with it. Studies which analyzed these dynamics in abstract terms assumed that the physical objects coupling with the bath are characterized by the same coupling strength to the bath degrees of freedom. In realistic models, this corresponds to assuming that both objects couple to the same particle of the bath. Further works addressed the question of how bath-mediated entanglement scales with the distance between the objects. In a system composed by a linear chain of oscillators it was shown that entanglement between any pair of oscillators decays quickly on a length scale of the order of the interparticle distance. The authors argued on the basis of a phenomenological Markovian model that the bath cutoff wavelength, which is the inverse of the cutoff frequency, is the characteristic length over which bath-mediated entanglement vanishes. In a linear chain of oscillators this coincides with the interparticle distance.

In Ref. [16], however, it was shown that Markovian models cannot capture all dynamical processes which can lead to entanglement between two objects. Moreover, in Ref. [16] a microscopic model was proposed in which two defect oscillators embedded in the chain become entangled via the coupling with the chain instead of thermalizing at the chain’s temperature. This is possible since the impurities break the discrete translational symmetry of the lattice, giving rise to a set of normal modes involving the defects and the ions between them. Such modes are essential for the generation of entanglement at steady state. We remark that a linear chain of oscillators in a thermal state and coupled to a single impurity is a realization of Rubin’s model; it leads to thermalization of the impurity and is an example of Brownian motion in a solid-state environment. The addition of a second impurity introduces a new symmetry modifying this behavior. In the context of recent studies on non-Markovianity, one can say that correlations between the two defects are stored and mediated by a bath pseudo-mode, which is composed by oscillations of the bath particles spatially interposed between the two defects.

The findings of Ref. [16] imply that two impurity defects can be entangled by the surrounding bulk when the latter exhibits some sort of spatial order. In addition, entanglement could be found between two defects at any distance, independent of the size of the bulk. The latter statement thus goes beyond existing studies, which analyze entanglement generation between two systems at the edges of a chain. It further addresses the question of whether it is possible to generate entanglement between two macroscopically distant objects through interaction with a large common bulk. Understanding this issue is relevant at the fundamental level, for example, in the context of recent studies of thermalization in quantum systems. Moreover, it is important for studies on biological systems and for implementing protocols based on dissipative dynamics in quantum networks. In this Rapid Communication we analyze the dependence of bath-mediated entanglement on the distance by considering the specific physical system of two ions in a linear Paul trap, which are embedded in a linear chain of ions of different species, as shown in Fig. 1. Let $N$ and $Q$ be the number of ions and their charge. The ions are aligned along the $z$ axis inside a linear Paul trap with position (canonically conjugated momentum) $\vec{r}_j = (x_j, y_j, z_j)$ $[p_j = (p_{jx}, p_{jy}, p_{jz})]$, where $j = 1, \ldots, N$ labels the ions. The impurity defects are two ions of different species far away from the chain edges, whose mass $M$ is larger than the mass $m$ of the other ions composing the chain. The trap potential for the ion $j$ reads $V_{\text{trap}}(\vec{r}_j) = \left[U_j z_j^2 + U_{\perp,j}(x_j^2 + y_j^2)\right]/2$, where $U_j$ is determined by the static quadrupole potential and $U_{\perp,j} = (U_0/m_j - U_j)/2$ by the radio-frequency potential creating the transverse confinement. We note the dependence of the transverse potential strength on the mass.
of the ion: This implies that the defect’s transverse motion is a localized oscillation in the chain for sufficiently large ratios $M/m$. The ions are at a sufficiently low temperature $T$ such that they perform harmonic vibrations about their respective equilibrium positions $z_j^{(0)} = (0,0,0)$, at which the trap force and the Coulomb repulsion mutually balance. In this limit the dynamics is governed by the quadratic Hamiltonian $H_0 = H_\parallel + H_\perp + H_\perp^{(y)}$, with

$$H_\parallel = \sum_{j=1}^{N} \left( \frac{p_j^2}{2m_j} + \frac{1}{2} U_j q_j^2 + \frac{1}{4} \sum_{q\neq j} \mathcal{K}_{j,q}(q_j - q_q)^2 \right),$$

$$H_\perp^{(y)} = \sum_{j=1}^{N} \left( \frac{p_{j,x}^2}{2m_j} + \frac{1}{2} U_{j,x} x_j^2 - \frac{1}{8} \sum_{q\neq j} \mathcal{K}_{j,q}(x_j - x_q)^2 \right),$$

and $H_\perp^{(y)} = H_\perp^{(x-y)}$. Here, $q_j = z_j - z_j^{(0)}$, while $\mathcal{K}_{j,q} = 2Q^2/|z_j^{(0)} - z_q^{(0)}|^3$ is the coupling due to the Coulomb repulsion.

We first analyze the spectrum of Hamiltonian $H_0$. For simplicity, we assume equally spaced axial equilibrium positions with interparticle distance $d = z_j^{(0)} - z_{j+1}^{(0)}$, which can be found in the central region of long ion chains [35] and in anharmonic potentials [36,37]. Figure 2(a) displays the spectra of the axial and transverse modes for $M \approx 2.87m$, which corresponds to $\text{In}^+$ ions embedded in a $\text{Ca}^+$ chain [38]. Two degenerate normal mode frequencies for each transverse spectrum are separated from the respective transverse branch by a gap. If this gap is sufficiently large, these modes approximately coincide with the defects transverse vibrations. In this limit they are strongly coupled with each other via the Coulomb interaction and weakly coupled to the rest of the chain. Their dynamics are governed by a beam-splitter type of interaction: The two defects can become entangled by the unitary evolution after preparing each transverse mode in a squeezed state [39]. We verify this behavior assuming that initially the chain is in a thermal state at temperature $T$ while the transverse modes are prepared in identical squeezed vacuum states along the $x$ direction with variances $\Delta x_j^2 = x_j^2 e^{\Delta^2}/2$ and $\Delta x_j^2 = x_j^2 e^{\Delta^2}/2$. Here $x_0 = \sqrt{h/(M\Omega_{\perp})}$ is the size of the ground state of the defect oscillator with frequency $\Omega_{\perp} = \sqrt{U_{\perp}(M)/M}$, $\rho_0 = h/\sqrt{\Delta}$, and $s$ is the real-valued squeezing parameter [39]. The defect’s state remains Gaussian under the evolution given by the quadratic Hamiltonian, Eq. (2), and is hence described by the first moments and the covariance matrix [40]. Entanglement between the defect modes is quantified by means of the logarithmic negativity $E_N$ obtained from the symplectic eigenvalues of the partially transposed covariance matrix [41]. The grey curves in Fig. 2(b) display $E_N$ as a function of the time $t$ for a fixed distance $d = 7a$: $E_N$ grows.

FIG. 1. (Color online) A setup where the dynamics we predict could be tested is an ion chain in a Paul trap, which embeds two ions of larger mass. A standing-wave laser illuminates the defect ions and couples their transverse and axial displacements via the mechanical effects of light. For a certain class of initial separable states, the defect’s transverse motion becomes entangled by the coupling with the axial phonons of the chain.

FIG. 2. (Color online) (a) Spectrum of excitations when a chain of $\text{Ca}^+$ ions contains two $\text{In}^+$ ions with mutual distance $d = 7a$. The mass ratio is $M/m \approx 2.87$ [38]. The eigenfrequencies $\omega_k$ of $H_\parallel$ (solid line) and $H_\perp$ (dashed line) are displayed as a function of the quasimomentum $k$ (in units of $\pi/a$). The isolated transverse branches, indicated by the circle, almost coincide with the frequency of the defect transverse oscillator. The parameters are $U_{j,x}(m)/U_j = 2000$ and $\omega_j = 2\pi \times 26.1\text{kHz}$, while the frequencies are in units of $\omega_0 = \sqrt{U_{\perp}/m}$. (b) Logarithmic negativity $E_N$ as a function of time $t$ (in units of $\sqrt{\omega_j}/N$) for $N = 50$, initial squeezing $s = 0.5$, while the rest of the chain is initially prepared at $T = 8\hbar \omega_j/\hbar k_b \approx 10\mu\text{K}$. The black (grey) curves represent the case with laser coupling switched on (off), with coupling strength $\gamma = 18.2m\omega_0^2$. (c) Logarithmic negativity as a function of time for a chain with $N = 800$ (solid line), 1000 (dashed), and 1200 (dotted) ions, for $d = 17a$, $T = 0$, $s = 0.5$, and $\gamma = 0.036 m\omega_0^2$. For each curve, the time is in units of $\omega_j(N)^{-1}$, with $\omega_j(N) = \omega_{0}\sigma\sqrt{\log N}/N$ and $\omega_{0\sigma} = 2\pi \times 659.6\text{kHz}$. Insets: corresponding spectral densities (in units of $m\omega_0^2$) as a function of the frequency. The solid (dashed) lines correspond to $J_{\perp}(\omega)$ [$J_{\parallel}(\omega)$]. The arrows indicate the defect frequencies for the solid curves in (b) and (c) [44].
with \( t \) and increases with \( s \). The defects become entangled also when they are initially not squeezed (\( s = 0 \)), which is due to the fact that the normal modes of the isolated frequencies include some small displacements of the other ions of the chain. Numerical simulations show that \( E_N \) at a given time \( t \) decreases with the strength of the direct Coulomb coupling, which scales with the distance as \( 1/d^2 \).

We now demonstrate that entanglement between the defects can be generated and substantially enhanced by coupling with the axial phonons of the chain. For this purpose we consider an interaction which couples the defects’ transverse and axial displacements. The dynamics are now given by the Hamiltonian \( H = H_0 + H_\perp(t) \), where

\[
H_\perp(t) = \frac{\gamma(t)}{2} \left[(x_j - q_j)^2 + (x_j - q_j)^2\right],
\]

with \( \gamma(t) = \gamma \Theta(t) \) being an effective coupling strength and \( \Theta(t) \) the Heaviside function. This coupling could be realized, for instance, using a standing-wave laser in the \( x\)-\( z \) plane illuminating the defects in the Lamb-Dicke regime and with modes at their equilibrium positions [42]. For this geometry the motion along \( y \) is decoupled and will be ignored from now on. At \( t > 0 \) a displacement of each defect along \( x \) excites a wave packet of axial phononic modes.

We first note that these dynamics lead to thermalization of a single defect [17,43], but are significantly modified when both defects are coupled. In order to understand why, let us consider that the defects are symmetrically placed with both defects are coupled. In order to understand why, let us consider that the defects are symmetrically placed with both defects are coupled. In order to understand why, let us consider that the defects are symmetrically placed with both defects are coupled. In order to understand why, let us consider that the defects are symmetrically placed with both defects are coupled. In order to understand why, let us consider that the defects are symmetrically placed with both defects are coupled. In order to understand why, let us consider that the defects are symmetrically placed with both defects are coupled. In order to understand why, let us consider that the defects are symmetrically placed with both defects are coupled.

The logarithmic negativity is determined after numerically evaluating the defect dynamics, starting from the formal solution of the coupled Heisenberg equations of transverse, axial, and defect modes of the Hamiltonian \( H \) [43]. The solid curves in Fig. 2(b) display \( E_N \) in a chain of 50 ions and for times over which finite-size effects are negligible. Entanglement builds up after a transient time and reaches values which are an order of magnitude larger than the values found in the absence of the coupling laser. It increases with the strength of the initial squeezing \( s \), and we checked that it also increases when decreasing the temperature of the chain, which is consistent with the results of previous works [10,16,43]. We note that although cooling a large chain to ultralow temperatures is a challenging task, the basic requirement for observing the dynamics predicted here is that the axial mode of the chain resonant with the defect frequency is prepared in the ground state. In a chain of trapped ions this could be realized, for example, by cooling the motion of the defect ions while the laser coupling transverse and axial displacement is switched on [47]. Noise will not significantly affect the predicted dynamics, provided that the dark mode is cold and protected from external noise sources [48].

We now demonstrate that the created entanglement is not a finite-size effect, and is indeed independent of the chain size. Figure 2(c) displays the logarithmic negativity as a function of time for various chain sizes and a mutual distance of \( d = 17a \) between the defects. Here the c.m. motion is decoupled by tuning its frequency to a value at which the corresponding spectral density vanishes (see inset). The curves are displayed for times that are shorter than the revival time and have been rescaled by the size-dependent frequency \( \omega_1 = \omega_{\text{ref}} \sqrt{\log N/N} \), where \( \omega_{\text{ref}} \) is a constant and the size-dependent factor warrants that the interparticle distance at the chain center is independent of \( N \) (therefore keeping constant the cutoff frequency) [35]. One observes that the logarithmic negativity oscillates about a (quasi) stationary state, whose value is independent of the chain size. For increasing chain sizes the revival time, and thus the time window over which this entanglement is found, correspondingly increases.

We finally show that the entanglement generated by the interaction with the chain slowly decays with the mutual distance. Figure 3 displays the mean value of the logarithmic negativity, \( \bar{E}_N \), averaged over the time in which it regularly oscillates, as a function of the mutual distance when the c.m. is decoupled. The different points correspond to different zeros of the spectral density: optimal entanglement is thus achieved by suitably tuning the frequency of the impurity, so that it matches the frequency of an optimal dark normal mode. Steady-state entanglement is found on time scales \( t_0 \) of the order of \( d/c_s \), where \( c_s \approx \omega_0/a \) is the sound velocity [35]. For the parameters and the chains here considered, \( t_0 \sim 100 \mu s \). At large \( d \), \( E_N \) decays linearly with the distance, which is confirmed by a systematic analysis performed on a model with nearest-neighbor interactions [49].

The resulting entanglement can be measured by extending the method developed in Ref. [50] to two Gaussian modes. The procedure consists of coupling the transverse oscillation of each defect with an ancillary qubit, such as an electronic transition of the defect ion. This can be done using the
Hamiltonian $H_{\text{int}}^{\text{def}} = \hbar \gamma (\sigma_j^2 + \frac{1}{4} \sigma_j^4)$, where $\sigma_j^2 = (x_j^2 + y_j^2 + z_j^2)$. Parameters are set to $N = 800$, $T = 0$, $s = 0.5$, and coupling $\gamma = 3230 m \omega_0^2$. Inset: Spectral densities for $d = 15 \sigma$.

The authors acknowledge discussions with C. Cormick, J. Brito, G. De Chiara, C. Kurz, and E. Lutz, and support by the European Commission (IP AQUITE, STREP PICC), the BMBF (QuOREp), the German Research Foundation, and the Irish Research Council through the Embark Initiative.

The study sheds light on the role of the reservoir in establishing quantum correlations. It further identifies the basic requirements for entangling two distant objects which interact with a bulk exhibiting long-range order. Here, nonlocal, decoherence-free subspaces appear which are associated with normal modes involving the defect vibrations. In principle, it is thus possible to entangle arbitrarily distant objects via a common bath. An intrinsic limitation to such dynamics is the fact that the required time linearly scales up with the distance, while the zeros of the dark normal modes become denser and denser. A further challenge is the realization of arbitrary large regular lattices. Such limitations could be overcome by applying local time-dependent operations.

Tuning the defect frequency is the key for entanglement generation in other physical platforms. These dynamics can also be observed, for instance, in dipolar quantum gases in the Mott insulator phase in an optical lattice, where the optical traps can be engineered in order to tune the frequency of the transverse oscillations of different species. Other examples are cavity arrays in circuit QED and optomechanical systems as in [55], where the frequencies can be controlled by means of proper designs and the use of refractive media.

The authors acknowledge discussions with C. Cormick, J. Brito, G. De Chiara, C. Kurz, and E. Lutz, and support by the European Commission (IP AQUITE, STREP PICC), the BMBF (QuOREp), the German Research Foundation, and the Irish Research Council through the Embark Initiative.