Quantum simulation of dynamical maps with trapped ions

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This supplementary information provides background material on details of the theory and the experimental implementation of the quantum simulation of open system dynamical maps with trapped ions discussed in the main text. In addition, it covers further experimental simulation results which - due to space restrictions - are not discussed in the main text. Section I provides details on the experimental system and the available set of tools for open-system quantum simulation. In Sec. II we present in more detail the general concept of the engineering of elementary dissipative and coherent Kraus maps, and also provide the specific sequences of operations that have been used in the experimental implementation. In Sec. III we discuss the quantitative experimental characterization of elementary dissipative Kraus maps. Based on these measurements, we describe in more detail a theoretical noise model, which captures the dominant experimental imperfections, and present analytical expressions and information on the numerical simulations for the dynamics of the open many-body spin system under dynamical maps. Sec. IV and V contain background information on the error detection and reduction schemes, as well as experimental details on their implementation. Finally, in Sec. VI we discuss additional experimental simulation results, including the realization of competing dissipative and coherent dynamical maps for varying Hamiltonian interaction strengths and initial states, and the combined implementation of dissipative dynamical maps with the error reduction protocol.

Table I provides an overview of the algorithmic building blocks required for the implementation of elementary dissipative Kraus maps, the realization of Hamiltonian dynamical maps, spectroscopic decoupling of ions, and elements for the detection and reduction of errors in our ion-trap quantum simulator. The table describes the experimental resources needed for the implementation of these elementary sub-routines as well as for the combination of these building blocks in more complex composite dynamical maps. The details of the underlying Kraus map engineering as well as the detailed quantum circuits used in the experiment are presented in the corresponding subsections below.

I. EXPERIMENTAL SYSTEM AND TECHNIQUES

In this section we will describe the available operations in our universal ion-trap quantum simulator.

A. Coherent gates

The qubit is encoded in the $4S_{1/2}(m_j = -1/2)$ and $3D_{5/2}(m_j = -1/2)$ electronic states of the $^{40}$Ca$^+$ ion and is manipulated by precisely timed light pulses on resonance or near resonant with the optical transition.

The laser light can be applied from two different directions as depicted in figure S1, where one beam illuminates the entire ion string homogeneously and the second beam is able to address each ion individually (Schmidt-Kaler et al., 2003).

The set of coherent gates consists of collective single-qubit rotations, addressed single-qubit gates and collective entangling gates. Collective single-qubit rotations are implemented by the globally applied laser beam, resonant with the qubit transition, realizing the unitary

\[ R(\theta, \phi) = \exp \left( - \frac{i \theta \pi}{2} S_{\phi} \right), \]

with $S_{\phi} = \sum_{i=0}^{N} \sigma^{\phi}_i$ a collective spin operator, and

\[ \sigma^{\phi}_i = \sigma^{x}_i \cos(\phi \cdot \pi) + \sigma^{y}_i \sin(\phi \cdot \pi) \]

being a linear combination of the single-qubit Pauli matrices $\sigma^{x}_i$ and $\sigma^{y}_i$, acting on qubit $i$. The rotation angle $\theta$ is determined by $\theta = \Omega \tau/A$, which can be controlled by the Rabi frequency $\Omega$ and the pulse duration $\tau$. In this notation a complete $\pi$-flop inverting the electronic population of the logical states corresponds to a rotation angle $\theta = 1$. On the other hand, tuning the phase $\phi$ of the global laser beam allows one to control the rotation axis lying in the $x$-$y$-plane of the Bloch sphere, around which each of the qubits is rotated. Addressed single-qubit operations

\[ S_z(\theta, i) = \exp \left( - \frac{i \theta \pi}{2} \sigma^{z}_i \right) \]

around the $z$-axis are realized by shining in laser light near resonant with the qubit transition that induces an


### Table I: Summary of the required resources for the elementary and composite dynamical maps and additional tools used in the quantum simulation.

<table>
<thead>
<tr>
<th>Algorithm type</th>
<th>Nr. of MS gates</th>
<th>Nr. of global rotations $R$</th>
<th>Nr. of AC-Stark shifts $S_Z$</th>
<th>Nr. of resets</th>
<th>Total number of operations</th>
<th>Described in section</th>
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<td>V</td>
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<td>V</td>
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<td>V</td>
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<td>44</td>
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Table I: Summary of the required resources for the elementary and composite dynamical maps and additional tools used in the quantum simulation. The required operations for the composite maps do not strictly match the sum of the required elementary operations, since in the implementation of composite maps synergy effects in the resources for the spectroscopic decoupling operations can be exploited.

intensity-dependent AC-Stark shift $\Delta_{AC}$. Again, the rotation angle $\theta$ is determined by the pulse length $\tau$, $\theta = \Delta_{AC} \tau / \pi$, and a $\pi$-pulse corresponds to $\theta = 1$. Finally, collective entangling operations are implemented by a bi-chromatic, globally applied laser field, which effectively realizes two-body Mølmer-Sørensen (MS) type interactions (Mølmer and Sørensen, 1999).

$$MS(\theta, \phi) = \exp \left( -\frac{i \theta \pi}{4} S_Z^2 \right) = \exp \left( -\frac{i \theta \pi}{2} \sum_{i > j} \sigma_i^x \sigma_j^x \right)$$

between all pairs $i$ and $j$ of the ion chain ($i, j = 0, 1, \ldots, N$) (first experimental realization in Ref. (Sackett et al., 2000)). Again, the angle $\phi$ allows one to control whether $\sigma_i^x \sigma_j^x$ (for $\phi = 0$), $\sigma_i^y \sigma_j^y$ (for $\phi = 1/2$) or interactions $\sigma_i^z \sigma_j^z$ corresponding to any other axis $\sigma^\phi$ in the $x$-$y$-plane are realized. In this notation, the angle $\theta = 1/2$ corresponds to a "fully-entangling" MS gate, i.e. a unitary which maps the computational basis states of $N+1$ ions onto multi-particle entangled states, which are (up to local rotations) equivalent to $N+1$-qubit GHZ states. Altogether, operations (1) to (3) form a universal set of gates, enabling the implementation of arbitrary unitaries on any subset of ions (Nebendahl et al., 2009).

### B. Numerical optimization of gate sequences

Any unitary operation required for the implementation of dissipative and coherent maps, as well the error detection and reduction protocols, needs to be decomposed into a sequence of available operations. As discussed in more detail below, such decompositions can be constructed systematically (Müller et al., 2011). However, as such decompositions are in many cases not optimal in terms of the number of required gate operations, it is convenient to resort to a numerical optimal control algorithm (Nebendahl et al., 2009) to search for optimized sequence decompositions involving less gates. Whereas the numerical optimization algorithm becomes inefficient for general unitary operations acting on a large number of qubits, it is well-suited for the optimization of unitaries which act only on a small subset of ions (such as 2+1 ions in the implementation of an elementary dissipative Kraus map), independently of the total system size.

Numerically optimized pulse sequences may include global AC-Stark pulses and MS gates with negative rotation angles $\theta < 0$, which are not directly contained in the available gate set discussed in Sec. I.A. However, as any collective rotation around the $z$-axis of the Bloch sphere can be interpreted as a re-definition of the $x$- and $y$-axes, a global AC-Stark pulse can be omitted if the phases $\phi$ of the following resonant operations are properly adjusted. Regarding MS gates with $\theta < 0$, these can be implemented by MS gates with positive rotation angles, as $MS(-\theta, \phi) \equiv MS(1 - \theta, \phi)$ up to local rotations (see Eq. (9) in Ref. (Müller et al., 2011)).

### C. Spectroscopic decoupling of ions

Despite the globally applied beams for the collective rotations and MS gates, operations on subsets of ions can be realized by spectroscopically decoupling ions not involved in the realization of a certain Kraus map from the dynamics. This is realized by coherently transferring ions to electronic states, where they do not couple to the...
globally applied light fields to a very good approximation. This decoupling technique enables the use of optimized sequences for the realization of Kraus maps on a small number of sites, independently of which subset of ions is currently involved in the map, and independently of the system size, i.e. the total number of sites.

The full decoupling sequence consists of the following parts as also outlined in Fig. 2 in the main-text: (i) First, the population is transferred from the qubit state $|S\rangle = 4S_{1/2}(m_f = -1/2) = |1\rangle = |\uparrow\rangle$ to the $|D\rangle = 3D_{5/2}(m_f = -5/2)$ state. (ii) Then, the population from the other qubit state $|D\rangle = 3D_{5/2}(m_f = -1/2) = |0\rangle = |\downarrow\rangle$ is transferred via $|S\rangle = 4S_{1/2}(m_f = +1/2)$ to the $|D''\rangle = 3D_{5/2}(m_f = -3/2)$ state. The required pulses for decoupling two ions are shown in Table II. Bringing the population from the decoupled states back to the original qubit states is realized by implementing the described sequence in reverse order.

The decoupling technique introduces additional errors that are not included in the theoretical error model. A rigorous treatment of these errors is cumbersome since it cannot be modeled in a qubit system anymore, but involves the full electronic substructure of the ion. However, the effect of the decoupling process in the computational basis can be characterized by quantum process tomography. We found a process fidelity of 94(2)% for decoupling a qubit and bringing it back to the original states. Next, we proved that the decoupled qubit is indeed to a high degree not affected by the manipulation pulses. We checked this in a system of 3+1 ions, where we first decoupled a single qubit, then applied the pulses as required for a single elementary dissipative Kraus map on the remaining two system ions and the ancilla ion, and finally transferred the decoupled qubit back to the original qubit state. The elementary dissipative Kraus maps induce a deterministic AC-Stark shift on the decoupled ion. This Stark shift is measured with a Ramsey-type experiment, and its compensation is performed with the final two pulses in the sequence as shown in Table II. Quantum process tomography on the decoupled qubit, where the systematic Stark shift has been compensated, yields a process fidelity of 93(2)%.

D. Incoherent reinitialization of individual ions

The implementation of an elementary dissipative Kraus map is completed by an incoherent reset of the ancillary ion to its initial state $|S\rangle$, see step (iii) in Fig. 2c in the main text. This reinitialization is realized by an optical pumping technique: First, an addressed pulse is applied to the ancillary ion to transfer the population present in the $|D\rangle$ state to the $|S\rangle$ state. Then $\sigma^-$ polarized light is applied to the entire ion string performing optical pumping from $|S\rangle$ towards $|S\rangle$ via the short-lived $4^2P_{1/2}$ state. This procedure does not affect the information in the system ions encoded in the original qubit states $|S\rangle$ and $|D\rangle$, as the light couples only to the $|S\rangle$ level (Barreiro et al., 2011). The required operations are shown in Table III.

II. ENGINEERING OF DISSIPATIVE AND HAMILTONIAN DYNAMICAL MAPS

In this section we provide details on the engineering and the specifics of the circuit-based experimental implementation of elementary dissipative and Hamiltonian dynamical maps.

A. Action of the dissipative Kraus maps

The elementary dissipative Kraus maps

$$\rho \mapsto E_{i,1}\rho E_{i,1}^\dagger + E_{i,2}\rho E_{i,2}^\dagger$$

(4)

with

$$E_{i,1} = c_i \quad \text{and} \quad E_{i,2} = 1 - c_i^\dagger c_i$$

(5)

<table>
<thead>
<tr>
<th>Transition</th>
<th>Pulse</th>
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<tbody>
<tr>
<td>$</td>
<td>S\rangle \rightarrow</td>
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<td>$</td>
<td>S\rangle \rightarrow</td>
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<td>S\rangle \rightarrow</td>
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<td>D\rangle \rightarrow</td>
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<td>D\rangle \rightarrow</td>
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<tr>
<td>$</td>
<td>D\rangle \rightarrow</td>
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<tr>
<td>$</td>
<td>S\rangle' \rightarrow</td>
</tr>
</tbody>
</table>

TABLE II Pulse sequence for spectroscopic decoupling of ions $i$ and $j$ by coherently transferring their quantum information from the qubit states $|S\rangle$ and $|D\rangle$ to the states $|D''\rangle$ and $|D''\rangle$. The gates listed in the three blocks of the table realize the pulses (i) - (iii) shown in Fig. 2b of the main text.

<table>
<thead>
<tr>
<th>Transition</th>
<th>Pulse</th>
</tr>
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<tbody>
<tr>
<td>$</td>
<td>S\rangle' \rightarrow</td>
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<td>$</td>
<td>S\rangle' \rightarrow</td>
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<tr>
<td>$</td>
<td>S\rangle' \rightarrow</td>
</tr>
<tr>
<td>$</td>
<td>S\rangle' \rightarrow</td>
</tr>
</tbody>
</table>

TABLE III Pulse sequence for the individual reset of qubit $j$. |
are generated by the operators $c_i$ (cf. Eq. (2) in the main text), as given by

$$c_i = (\sigma_i^+ + \sigma_{i+1}^+)(\sigma_i^- - \sigma_{i+1}^-). \quad (6)$$

These operators act bi-locally, i.e. involve two neighboring spins $i$ and $i+1$, whereas the other spins are spectators. It is instructive to examine their action on the basis states of the local Hilbert space of the two qubits $i$ and $i+1$, which is spanned by the singlet and triplet states of the total spin $S_{i,i+1}^2$ of the two spin-$1/2$ particles:

$$S_{i,i+1}^2 \frac{1}{\sqrt{2}}(|01\rangle - |10\rangle) = 0$$

and

$$S_{i,i+1}^2 (|00\rangle, \frac{1}{\sqrt{2}}(|01\rangle + |10\rangle), |11\rangle) = 2 (|00\rangle, \frac{1}{\sqrt{2}}(|01\rangle + |10\rangle), |11\rangle).$$

Here, $S_{i,i+1} = S_i + S_{i+1}$ with $S_i = \frac{1}{2} \sigma_i = \frac{1}{2} (\sigma_i^x, \sigma_i^y, \sigma_i^z \nu)$ and

$$S_{i,i+1}^2 = (S_i + S_{i+1})^2 = 2 + \frac{1}{2} (\sigma_i^x \sigma_{i+1}^x + \sigma_i^y \sigma_{i+1}^y + \sigma_i^z \sigma_{i+1}^z).$$

For simplicity of the notation, we skip the spin indices $i$ and $i+1$ for the states and use the short-hand notation $|00\rangle = |0\rangle_i \otimes |0\rangle_{i+1}$, etc. As illustrated in Fig. S2, the operators $c_i$ induce pumping from the singlet into the triplet $m_S = 0$ state,

$$c_i \frac{1}{\sqrt{2}}(|01\rangle - |10\rangle) = \frac{1}{\sqrt{2}}(|01\rangle + |10\rangle),$$

whereas all triplet states are dark states,

$$c_i |00\rangle = c_i \frac{1}{\sqrt{2}}(|01\rangle + |10\rangle) = c_i |11\rangle = 0.$$

As shown in Fig. 1b of the main text, under this dissipative dynamics a single spin excitation (or hardcore boson) is symmetrically delocalized over the two sites, whereas the states of two ($|11\rangle$) or zero ($|00\rangle$, "vacuum") spin excitations (or hardcore bosons) are left unchanged.

### B. Circuit-based implementation of elementary dissipative Kraus maps

Open-system dynamics according to the elementary dissipative Kraus maps (4) can be realized in a "digital" way, by using quantum simulation tools for open systems, which have been previously developed and demonstrated experimentally in the context of dissipative preparation of Bell and multi-qubit stabilizer states (Barreiro et al., 2011). The key idea in engineering the two-spin dissipative dynamics according to Eq. (4) is to combine the experimentally available gates (see Sec. I.A below for details) with optical pumping on an additional ancillary qubit (see Sec. I.D), which plays the role of a tailored environment.

**General engineering strategy** – The observation that the singlet state is dissipatively converted into the $m_S = 0$ triplet state suggests the following gate-based implementation via a four-step process (shown in Figure S3a), which involves a circuit of unitaries applied to the qubits $i$, $i+1$ (steps (i) to (iii)), followed by the incoherent reset of the ancilla qubit (step (iv)):

(i) First, a unitary $M$, acting on the two system spins $i$ and $i+1$ and the ancillary qubit, coherently maps the binary information whether the two system spins are in the singlet or triplet subspace onto the two logical states $|0\rangle$ and $|1\rangle$ of the ancilla qubit. This is achieved by the unitary

$$M(\theta) = \exp \left( -\frac{i\theta}{2} (S_{i,i+1}^2 - 2) \otimes \sigma_0^x \right) \quad (7)$$

$$= \exp (i\theta P_{i,i+1} \otimes \sigma_0^z)$$

$$= \cos(\theta P_{i,i+1}) \otimes 1_0 - i \sin(\theta P_{i,i+1}) \otimes \sigma_0^z$$

for which we choose $\theta = \pi/2$ so that the unitary reduces to

$$M(\pi/2) = (1 - P_{i,i+1}) \otimes 1_0 - i P_{i,i+1} \otimes \sigma_0^-.$$ 

The unitary $M$ is constructed in a way that the state of the ancilla qubit is rotated conditional on the state of the two system spins, and that the angle of this rotation depends on the operator $(S_{i,i+1}^2 - 2)$ which acts on the two system spins $i$ and $i+1$. Here, $P_{i,i+1} = c_i^\dagger c_i$ is the projector onto the singlet subspace, $(1 - P_{i,i+1})$ its orthogonal complement, and we have used $S_{i,i+1}^2 = 2(1 - P_{i,i+1})$. Under the unitary $M(\pi/2)$, the ancilla qubit is rotated from its initial state $|1\rangle$ into $|0\rangle$ if and only if the two system spins are in the singlet state.

(ii) Next, the transfer of the system qubits from the singlet to the triplet subspace is enabled by a two-qubit

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**FIG. S2** Schematics of the action of the two-spin operator (6) in the local Hilbert space of the two spins $i$ and $i+1$, which is spanned by the singlet state (left sector of the Hilbert space) and the triplet states (right sector). The operator $c_i$ converts the singlet state $\frac{1}{\sqrt{2}}(|01\rangle - |10\rangle)$ into the triplet $m_S = 0$ state $\frac{1}{\sqrt{2}}(|01\rangle + |10\rangle)$. All triplet states are dark states and left invariant.
FIG. S3 Quantum circuits for the realization of elementary dissipative Kraus maps (4) on spins $i$ and $i+1$. a, First, the information whether the two system spin-1/2 particles $i$ and $i+1$ are in the singlet or triplet subspace of the two-spin Hilbert space (see Fig. S2) is coherently mapped onto the logical states $|0\rangle$ and $|1\rangle$ of an ancilla qubit which is initially in $|1\rangle$. (ii) Next, a two-qubit gate is applied, which realizes an effective spin flip on one of the system spins, and thereby conditionally on the state of the ancilla converts the singlet into the triplet $m_S = 0$ state: $\frac{1}{\sqrt{2}}(|01\rangle - |10\rangle) \rightarrow \frac{1}{\sqrt{2}}(|01\rangle + |10\rangle)$. After inverting the mapping (iii), the ancilla ion is optically pumped backed to its initial state $|1\rangle$, so that it can be used for the implementation of subsequent dissipative Kraus maps. (iv) This last steps renders the dynamics irreversible and provides the dissipative ingredient to extract entropy from the two system spins. The probability of the conversion from the singlet to the triplet state is controlled by the angle $\phi$ appearing in the two-qubit gate, and it is given by $p = \sin^2 \phi$. In the limit of $\phi \ll \pi/2$, the conversion from the singlet into the triplet takes place only with a small probability, and the general dissipative Kraus map describing the pumping process reduces to the master equation (11). b, In this work we are interested in deterministic pumping from the singlet into the triplet subspace, i.e. in the case of probability $p = 1$ for $\phi = \pi/2$. In this situation, the circuit simplifies as the inverse mapping to partially disentangle the system spins from the ancilla is not required, and the circuit simplifies to the three-step process, which is also shown in Fig. 2c of the main text.

gate, which acts on the ancilla qubit and the system qubit $i$. It reads

$$C(\phi) = \exp \left( \pm \frac{1}{2} (1 - \sigma_0^i) \otimes i\phi \sigma_1^i \right)$$

$$= |1\rangle\langle 1|_0 \otimes 1_i + |0\rangle\langle 0|_0 \otimes \exp(\pm i\phi \sigma_1^i).$$

This controlled-operation applies $\sigma_1^i$ to the $i$-th system qubit, and thus with probability $p = \sin^2 \phi$ the singlet into the triplet state: $\sigma_1^i \frac{1}{\sqrt{2}}(|01\rangle - |10\rangle) = \frac{1}{\sqrt{2}}(|01\rangle + |10\rangle)$. Here, the ancilla qubit acts as a quantum controller (Baggio et al., 2012; Lloyd and Viola, 2001), which is not observed but controls the coherent feedback which is applied to the system qubits. From Eq. (9) it is clear that if the ancilla is in state $|1\rangle$, corresponding to the case of the two system spins residing in the triplet subspace, no quantum feedback is applied and the state of the system spins remains unchanged. Due to the previous mapping $M$, this assures that the triplet subspace is left invariant.

(iii) The initial mapping (i) is inverted by applying the inverse unitary $M(\pi/2)$. For the system being in the target triplet space from the outset, this re-installs the conditions before step (i). For the special case of deterministic pumping ($p = 1$), as experimentally realized in this work, the inverse unitary can be omitted (see also Figure S3b).

(iv) After these unitary steps, the ancillary qubit remains in general entangled with the two system spins $i$ and $i+1$. It is finally reset to its initial state $|1\rangle$ by optical pumping: this is the physical dissipative mechanism, which renders the dynamics irreversible and allows to extract entropy from the system spins. This refreshes the ancilla qubit and prepares it in the known pure state $|1\rangle$, so that it can be used for the implementation of subsequent dissipative Kraus maps.

For an initially uncorrelated state of the ancilla qubit and the system spins, $|1\rangle\langle 1|_0 \otimes \rho$, the resulting dynamics for the system spins (described by the reduced density matrix $\rho$ of the system spins) is obtained by applying the combined unitary $U = M(\pi/2)C(\phi)M(\pi/2)$ and tracing over the ancilla qubit’s degrees of freedom,

$$\rho \rightarrow tr_0 \{ U(|1\rangle\langle 1|_0 \otimes \rho)U^\dagger \}. \quad (10)$$

Straightforward algebra yields the Kraus map $\rho \rightarrow \sum_{k=1,2} E_{i,k} \rho E_{i,k}^\dagger$ with the operation elements

$$E_{i,1} = \sin \phi c_i \quad \text{and} \quad E_{i,2} = 1 + (\cos \phi - 1) c_i^z c_i.$$ 

One the one hand, for $\phi = \pi/2$ the operation elements reduce to Eq. (5) and correspond to deterministic pumping ($p = 1$) from the singlet into the triplet states, which is the scenario realized in this work. On the other hand, for $\phi \ll \pi/2$ one can expand the operation elements of the Kraus map and recovers the master equation limit

$$\rho \rightarrow \rho + \phi^2 \left( c_i^z \rho c_i - \frac{1}{2} \left( c_i^z c_i \rho c_i^z \right) \right) + O(\phi^3). \quad (11)$$

Additional remarks – We note that our implementation corresponds to an open-loop control scenario, where the ancillary qubit remains unobserved during the simulation. However, we remark that it is possible to measure the state of the ancillary qubit in the computational basis by an addressed fluorescence measurement, before it is reset to its initial state by optical pumping (Barreiro et al., 2011). Such measurement reveals whether under the application a particular dissipative Kraus map the two system spins have undergone a collective two-spin quantum jump from the singlet into the triplet state or not. This information yields ”in-situ” information about the many-body system’s dynamics along a particular trajectory. The temporal statistics of quantum jumps in open, driven many-body quantum spin systems contains valuable information about dynamical phase transitions.
(Ates et al., 2012; Garrahan and Lesanovsky, 2010). In the present work we do not further explore this possibility. We remark that the fluorescence measurement of the ancillary ion, which is associated to the scattering of many photons and non-negligible heating of the vibrational modes, can be combined with subsequent laser recooling of the relevant vibrational modes via the ancillary ion, as demonstrated by (Schindler et al., 2012). This allows one to perform such quantum jump measurements and to afterwards re-initialize the external degrees of freedom, thereby enabling that in the same experimental run further gate operations required for the implementation of subsequent Kraus maps can be still applied with high fidelity.

Specific experimental implementation – For the implementation of the elementary dissipative Kraus maps in the ion-trap simulator, the unitary operations \( M(\pi/2) \) and \( C(\pi/2) \) (see Eqs. (7) and (9)) have to be decomposed into a quantum circuit of available gates: The mapping \( M \) can be written as a product of three unitaries involving 3-body spin interaction terms,

\[
M(\pi/2) = \exp \left( -\frac{i\pi}{4} \sigma_0^x (S_{i,i+1}^z - 2) \right) = \exp \left( -\frac{i\pi}{8} \sigma_0^x \sigma_{i,i+1}^{x} \right) \times \exp \left( -\frac{i\pi}{8} \sigma_0^x \sigma_{i,i+1}^{y} \right),
\]

As discussed in Ref. (Müller et al., 2011) each of the three-body unitaries can be realized by two "fully-entangling" MS gates in combination with single-qubit rotations, such that \( M \) could be implemented by a quantum circuit involving six MS gates and a number of single-qubit gates to rotate the system spins between the \( x, y \) and \( z \)-bases between the different unitaries.

Similarly, the two-qubit controlled operation \( C(\pi/2) \) (see Eq. (9)) is up to local unitaries equivalent to one "fully-entangling" two-ion MS gate (Müller et al., 2011),

\[
C(\pi/2) \sim \exp(-i(\pi/4)\sigma_0^z \sigma_i^z),
\]

acting on the ancilla qubit and the system spin \( i \). In the experiment, we did not use such a systematically constructed circuit decomposition, but instead resorted to the numerical optimization algorithm described in Sec. I.B to further reduce the complexity of the quantum circuit: The gates of the experimentally employed sequence decomposition for the implementation of the unitaries of one elementary dissipative Kraus map are listed in Table IV.

C. Engineering Hamiltonian maps for competing coherent interactions

As discussed in the main text, competing Hamiltonian dynamical maps are realized according to the dimensionless many-body spin Hamiltonian \( H = \sum_i H_i \),

\[
H = \sum_i |\uparrow_i\rangle\langle\uparrow_i| \otimes |\uparrow_i\rangle\langle\uparrow_i| = \sum \frac{(1 + \sigma_i^z)(1 + \sigma_{i+1}^z)}{4}
\]

where the two-body terms correspond to interactions between spin excitations (or hardcore bosons) located on neighboring sites \( i \) and \( i + 1 \). The Hamiltonian dynamical maps \( U_{i,i+1} = \exp(-i\phi H_i) \) acting on spins \( i \) and \( i + 1 \) are up to local rotations equivalent to two-spin MS interactions, \( \exp(-i\phi \sigma_i^z \sigma_{i+1}^z)/4 \). The implementation of the competing Hamiltonian dynamical maps can be realized with two distinct approaches: (i) The elementary two-spin Hamiltonian maps can be implemented sequentially, in analogy to the sequential implementation of the elementary dissipative maps. (ii) Alternatively, since the elementary Hamiltonian maps mutually commute, they can be implemented by a single global unitary operation, acting directly on the entire register of system spins. The unitary according to a sum of pairwise interactions between neighboring spins can for instance be built up from MS gates, which involve pairwise interactions between all pairs of spins, by means of refocusing techniques (Müller et al., 2011). Although one could again try to find numerically optimized sequences of gates for the implementation of the composite Hamiltonian map, we note that such optimization for the unitary acting on the entire register of system spins would have to be done for each system size, and becomes inefficient for increasing system sizes. It is thus advisable to exploit the symmetries of the MS interactions, and to systematically construct sequence decompositions.

Specific experimental implementation – For instance, in the case of 3 system spins (with open boundary conditions), this operation can be decomposed into the following unitaries according to single- and two-body inter-
actions
\[
U_{\text{comp}} = \exp \left( \frac{-i\phi}{4}(\sigma_1^z\sigma_2^z + \sigma_2^z\sigma_3^z) \right) \times \exp \left( -\frac{i\phi}{4}\sigma_2^z \right).
\]

As the ancilla qubit is not required to realize the unitary map, and must not be entangled with the system spins during the operation, it is spectroscopically decoupled during the application of the gate sequence. The experimental sequences used for the implementation of competing Hamiltonian maps for 3 system spins (3+1 ions) and 4 system spins (4+1 ions) are shown in Tables V and VI, respectively.

### Table V
<table>
<thead>
<tr>
<th>Number</th>
<th>Pulse</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(S_z(1.5, 1))</td>
</tr>
<tr>
<td>2</td>
<td>(R(0.5, 0.0))</td>
</tr>
<tr>
<td>3</td>
<td>(MS(1 - 0.25k, 0.5))</td>
</tr>
<tr>
<td>4</td>
<td>(S_z(1, 0.1))</td>
</tr>
<tr>
<td>5</td>
<td>(MS(0.25k, 0.5))</td>
</tr>
<tr>
<td>6</td>
<td>(S_z(1, 0.1))</td>
</tr>
<tr>
<td>7</td>
<td>(R(0.5, 1.0))</td>
</tr>
</tbody>
</table>

**TABLE V** Pulse sequence for the implementation of a composite Hamiltonian map in a 3-spin system, according to Eq. (13). The Hamiltonian strength is controlled by the parameter \(k\), with experimentally implemented values \(k \in \{1, 0.5\}\), corresponding to \(\phi \in \{\pi/2, \pi/4\}\).

### Table VI
<table>
<thead>
<tr>
<th>Number</th>
<th>Pulse</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(R(0.5, 0.5))</td>
</tr>
<tr>
<td>2</td>
<td>(MS(0.5, 1.0))</td>
</tr>
<tr>
<td>3</td>
<td>(R(0.5, 1.5))</td>
</tr>
<tr>
<td>4</td>
<td>(R(0.5, 0.5))</td>
</tr>
<tr>
<td>5</td>
<td>(MS(0.25, 1.0))</td>
</tr>
<tr>
<td>6</td>
<td>(S_z(1, 0.3))</td>
</tr>
<tr>
<td>7</td>
<td>(S_z(1, 0.2))</td>
</tr>
<tr>
<td>8</td>
<td>(MS(0.25, 1.0))</td>
</tr>
<tr>
<td>9</td>
<td>(S_z(1.5, 3))</td>
</tr>
<tr>
<td>10</td>
<td>(S_z(1.5, 2))</td>
</tr>
<tr>
<td>11</td>
<td>(R(0.5, 1.5))</td>
</tr>
</tbody>
</table>

**TABLE VI** Pulse sequence for the implementation of a composite Hamiltonian map in a 4-spin system, for strong competing interactions corresponding to an angle \(\phi = \pi/2\).

### III. EXPERIMENTAL CHARACTERIZATION OF AN ELEMENTARY DISSIPATIVE MAP AND NOISE MODEL

#### A. Modeling an imperfect elementary dissipative map

In the following, we will introduce a theoretical model of the elementary pumping step in the presence of experimental noise. A single elementary dissipative map acting on two system ions can be ideally described by the two-qubit process matrix \(\chi_{id}\) which can be straightforwardly calculated from the Kraus map (4) with generating operators as defined in Eq. (5) (Nielsen and Chuang, 2000). In order to describe the implementation of the elementary dissipative map on the three system qubits (two "active" and one spectroscopically decoupled ion), we will assume a process consisting of the modeled elementary map on the two active system qubits and an identity process on the third qubit. The noise affecting the elementary map is modeled by two independent depolarizing channels acting on each of the two "active" system qubits where the fully depolarizing channel on a single qubit \(i\) can be written as the Kraus map (Nielsen and Chuang, 2000)

\[
\rho \mapsto E^{(i)}_{\text{dep}}(\rho) = \frac{1}{4} (\rho + \sigma_i^x \rho \sigma_i^x + \sigma_i^y \rho \sigma_i^y + \sigma_i^z \rho \sigma_i^z).
\]

The physical noise acting on the register during the individual gates is certainly more complex than this, but the effect of noise on the outcome of any complex algorithm can be approximated by depolarizing noise regardless of the specific characteristics of the noise (Nielsen and Chuang, 2000). We model a noisy elementary dissipative map with the concatenation of two depolarizing channels each acting on one of the system qubits

\[
\rho \mapsto \Pi(\rho) = E^{(i+1)}_{\text{dep}}(E^{(i)}_{\text{dep}}(\rho))
\]

with process matrix \(\chi_{id}\) as

\[
\chi_{\text{idiss}}(\epsilon) = (1 - \epsilon)\chi_{id} + \epsilon\chi_{id}.
\]

In the limit \(\epsilon \to 0\), the imperfect process \(\chi_{\text{idiss}}\) reduces to the ideal two-qubit process \(\chi_{id}\), where in the extreme opposite limit of \(\epsilon \to 1\) depolarizing noise completely dominates and overwrites any effect of the desired engineered dissipative process \(\chi_{id}\). One can now adjust the parameter \(\epsilon\) of this model to the obtained data by a numerical optimization. For this, we maximized the overlap between the expected output state after a single elementary map \(\rho_e\) with noise strength \(\epsilon\) with the actual measured state \(\rho_{\text{exp}}\),

\[
\text{arg}_\epsilon \max \mathcal{F}(\rho_{\text{exp}}, \rho_e).
\]

where the Uhlmann fidelity \(\mathcal{F}(\rho_1, \rho_2)\) for the comparison of two density matrices \(\rho_1, \rho_2\) is used (Gilchrist et al., 2005). We find an optimum for a noise parameter of \(\epsilon = 0.27\).
TABLE VII Results for the numerical simulation of a quantum simulation algorithm, in order to identify the dominant experimental error source. The simulation is performed multiple times with only a single active error source. From the results one can infer that dephasing is the dominant error source.

<table>
<thead>
<tr>
<th>Error source</th>
<th>Overlap with $\Psi_+$</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>77%</td>
</tr>
<tr>
<td>Addressing</td>
<td>95%</td>
</tr>
<tr>
<td>Dephasing</td>
<td>84%</td>
</tr>
<tr>
<td>Intensity fluctuations</td>
<td>99%</td>
</tr>
<tr>
<td>Spectator modes</td>
<td>94%</td>
</tr>
</tbody>
</table>

B. Implementation and analysis of an elementary dissipative map

Here, we will provide a more detailed analysis of the specific implementation of a single dissipative step. During the realization of an elementary dissipative Kraus map gates act on the two ions encoding the system spins $i$ and $i + 1$ and the ancilla ion, while all other ions are spectroscopically decoupled. To quantitatively characterize the implementation of the Kraus map, as realized by the gate sequence shown in Table IV (see also Sec. II.B), we performed a quantum process tomography on the two system qubits. A benchmark for the performance is given by the process fidelity with the ideal two-qubit process. Since the ideal process is not a unitary process, the Choi-Jamiołkowsky process fidelity is a suitable measure (Gilchrist et al., 2005). We find a mean state fidelity of $F = 68(1)\%$.

In order to identify the leading source of imperfections in the implementation, a numerical analysis of the actual physical system is performed. We performed a Monte Carlo simulation of a single elementary map acting on three ions, including noise originating from: laser frequency and intensity fluctuations, magnetic field fluctuations, imperfect state preparation, motional heating, spontaneous decay and crosstalk of the addressed operations. The noise parameters are independently measured and we find an overlap between the numerically predicted and the measured state of $F = 97\%$. In order to determine the dominant error source, we performed the simulation multiple times where only a single noise source affects the evolution. The results are shown in Table VII. From this we find that the main error source is dephasing due to fluctuations in the laser frequency and the magnetic field.

C. Numerical simulation and long-time dynamics under dissipative maps

Based on the study of experimental errors of an elementary dissipative Kraus map in the previous sections, we have realized a numerical study of the dissipative dynamics driving a mesoscopic spin systems ($N = 10$) with initially three spin excitations present ($m_0 = 3$) towards the Dicke state $|D(3,10)\rangle$. The results are shown and discussed in Figure 5a of the main text. To take into account the imperfections in the implementation of the elementary maps, as discussed in the previous section, we assumed a depolarizing noise strength of $\epsilon_{\text{diss}} = 0.02$ for each of the two spins $i$ and $i + 1$ involved in an elementary dissipative map $D_{i,i+1}$. Since the implementation of an elementary Hamiltonian dynamical map $U_{i,i+1}$ on two spins is less complex than the realization of a dissipative map, we assumed a depolarizing noise strength for $U_{i,i+1}$ of $\epsilon_{\text{coh}} = \epsilon_{\text{diss}}/5$. The error reduction protocol to stabilize the system within the desired $m_0 = 3$ excitation subspace can in practice also be only implemented with a finite accuracy. Here we assumed both for the global operation required for one spin extraction procedure and for one injection procedure that the whole spin register is exposed to a global depolarizing noise channel with a probability $\epsilon = 0.02$.

Note that the chosen value of $\epsilon = 0.02$ for the noise strength is roughly one order magnitude smaller than the current experimental value ($\epsilon = 0.27$, see the main text and Sec. III.A above for details). The value of $\epsilon = 0.02$ for the numerical study is chosen in a way to illustrate how the characteristic physical features of the non-equilibrium dynamics, as generated by competing (imperfect) coherent and dissipative dynamical maps, become more and more visible, under a projected future improvement of the quality of the experimental operations by about a factor of ten. To illustrate this, Fig. S4 shows the numerically determined dynamics for a system of 8 spins with initially 3 excitations, as a function of the depolarizing noise strength $\epsilon$. The different curves display the overlap fidelity with the Dicke state $|D(3,8)\rangle$ for different strengths $\phi$ of competing Hamiltonian dynamics. Whereas without active error reduction the competition between engineered coherent and dissipative dynamics is covered by the dominant effect of the noise. However, as anticipated, under the incorporation of the error reduction protocol the crossover for varying competing Hamiltonian strengths becomes clearly visible as the noise level reaches values on the order of $10^{-2}$ and below.

Long-time dynamics and off-diagonal long-range order under imperfect dissipative dynamical maps – As discussed above and in the main text, under experimental imperfections during the pumping into Dicke states, the system initially residing in the $m_0$ excitation subspace suffers population leakage into the subspaces with $m \neq m_0$. In the long time limit, the population of the spin system is homogeneously distributed over these $m$-subspaces, each subspace populated according to its micro-canonical weight, i.e. the number of computational basis states spanning the corresponding subspace.
FIG. S4 Numerical study of competing Hamiltonian and dissipative dynamics in the presence of depolarizing noise. The curves corresponding to varying strengths $\phi$ of competing Hamiltonian dynamics show the overlap with the $|D(3,8)\rangle$ Dicke state for a chain of 8 spins with $m=3$ excitations, a, without and b, with active error reduction, as a function of the strength $\epsilon$ of the depolarizing noise. Under incorporation of the error reduction protocol the dynamical crossover becomes visible for experimental noise levels on the order of $10^{-2}$ and below. Dashed vertical lines serve as a guide to the eye roughly indicating the current experimental noise level ($\epsilon = 0.27$) as well as the smaller value of $\epsilon = 0.02$ used for the numerical simulations in the main text. For details see text.

Within each of the subspaces, not only for $m = m_0$, the dissipative maps are active and pump the population within each subspace towards the corresponding Dicke state $|D(m, N)\rangle$. As a result, for weak noise and in the long time limit, the system is driven into an incoherent mixture of Dicke states of different excitation numbers $m = 0, \ldots, N$.

More quantitatively, a normalized Dicke state with $m$ excitations for $N$ spins is given by

$$|D(m, N)\rangle = (m!)^{-1} \left( \begin{array}{c} N \\ m \end{array} \right)^{-1/2} (S^+)^m |0\rangle^\otimes N$$ (14)

with $S^+ = \sum_{k=1}^N \sigma_k^+$. The number of micro-states of $m$ excitations on $N$ sites is $\left( \begin{array}{c} N \\ m \end{array} \right)$. Thus the incoherent mixture of Dicke states is given by

$$\rho = \frac{1}{2^N} \sum_{m=0}^N \left( \begin{array}{c} N \\ m \end{array} \right) |D(m, N)\rangle\langle D(m, N)|$$ (15)

$$= \frac{1}{2^N} \sum_{m=0}^N \frac{1}{(m!)^2} (S^+)^m |0\rangle^\otimes N |0\rangle^\otimes N (S^-)^m$$

Interestingly, this incoherent mixture of Dicke states is a state with off-diagonal order $\langle \sigma_i^+ \sigma_j^- \rangle \neq 0$,

$$\langle \sigma_i^+ \sigma_j^- \rangle = \frac{1}{N(N-1)} (\langle S^+ S^- \rangle - m),$$ (16)

as we will show in the following. In Eq. (16) we have made use of the symmetry of the Dicke states under permutations of the spin indices. In order to evaluate the global expectation value $\langle S^+ S^- \rangle$, we decompose it as $\langle S^+ S^- \rangle = \langle S^- S^+ \rangle + \langle S^2 \rangle$. Both contributions are easily obtained as $\langle S^2 \rangle = \sum_i \langle \sigma_i^2 \rangle = 2m - N$, and $\langle S^- S^+ \rangle = (m+1)(N-m)$. The latter expectation value can be obtained using the normalization factors for the Dicke states $|D(m, N)\rangle$ and $|D(m+1, N)\rangle$ with $m$ and $m+1$ excitations, respectively. Adding both contributions, we obtain $(S^+ S^-) = m(N+1-m)$. This allows us to evaluate the expectation value $\langle \sigma_i^+ \sigma_j^- \rangle$ with respect to a pure Dicke state:

$$\langle \sigma_i^+ \sigma_j^- |D(m, N)\rangle = \frac{1}{N(N-1)} (\langle S^+ S^- \rangle - m)$$

$$= \frac{m}{N} \left(1 - \frac{m}{N}\right) \cdot \frac{1}{1 - \frac{m}{N}}.$$

In the thermodynamic limit, $N \to \infty$, $m/N = \text{const.}$, we have $\langle \sigma_i^+ \sigma_j^- \rangle \to m/N(1-m/N)$. This expression reflects the "particle-hole" symmetry and shows that at complete filling ($m = N$ spin excitations) or in the "vacuum" state ($m = 0$) there is no off-diagonal order because spin excitations or missing spin excitations cannot be delocalized over the spin chain. At half filling $m = N/2$, where the number of micro-states is maximal, the effect of delocalization and thus the off-diagonal order is maximal. The finite-size factor $1/(1 - 1/N)$ is to be taken into account in small or mesoscopic spin systems, and approaches 1 in the thermodynamic limit.

From this, we can now determine the off-diagonal order of the system in the incoherent mixture of Dicke states as given by the density matrix of Eq. (16). Using the identities

$$\sum_{m=0}^N \left( \begin{array}{c} N \\ m \end{array} \right) m = N 2^{N-1},$$

$$\sum_{m=0}^N \left( \begin{array}{c} N \\ m \end{array} \right) m^2 = N(N+1)2^{N-2},$$
one obtains

\[ \langle \sigma_i^x \sigma_j^x \rangle_{\text{mixture}} = \frac{1}{4}. \]

As expected, this value does not depend on the initial number of excitations anymore, since this information is completely lost, once the system has diffused into the incoherent mixture of Dicke states. The off-diagonal order in the incoherent mixture of Dicke states assumes a value which is independent of the system size \( N \), and remains finite in the thermodynamic limit.

In summary, the experimental imperfections, as described by depolarizing noise and resulting in the non-conservation of the excitation-number during the simulation, lead to a strong decrease of the state overlap between the asymptotically reached many-body body state of Eq. (16) and the ideal "target" dark state \( |D(m_0, N)\rangle \).

Moreover, from a condensed-matter perspective one can state that the imperfections are not too harmful to the off-diagonal long-range order, measured by the two-spin correlation function \( \langle \sigma_i^x \sigma_j^x \rangle \) for \( |i-j| \gg 1 \) as an order parameter. The off-diagonal order is constantly stabilized and re-built by virtue of the repeated application of the engineered (though imperfectly implemented) dissipative dynamical maps.

**IV. QUANTUM ERROR DETECTION METHOD:**

**POST-SELECTIVE QND SCHEME**

**A. General idea**

The post-selective error detection method is based on a quantum non-demolition (QND) measurement of the total number \( m \) of spin excitations present in the system at the end of the sequence of dynamical maps. Simulation outcomes, where due to experimental imperfections the ideally conserved initial spin excitation number \( m_0 \) has changed to a final value \( m \neq m_0 \) are discarded. This leads to an improvement of the simulation accuracy for longer sequences of dynamical maps, at the expense of an increased number of experimental runs to obtain the same measurement statistics. We remark that in a large system, as typical for post-selective techniques, this method becomes inefficient as the probability of remaining within the subspace of initial excitation number \( m_0 \) becomes exponentially small and thus only a vanishingly small number of "successful" runs enter the measurement statistics.

In order to maintain the dissipatively created off-diagonal order in the many-spin system, a crucial property of the excitation number measurement is its QND nature: the spin excitation number \( m \) has to be determined in a way that allows one to only obtain information about the total number of excitations in the system, but no knowledge about the spatial positions of individual excitations along the array. This QND measurement can be realized by a global unitary map, which acts on the entire register of system spins and an ancillary qubit (see Fig. 5b of the main text). Such a unitary is constructed in a way that it maps the binary information whether the register of system spins is in the correct excitation number subspace with \( m = m_0 \) (or not) onto the logical state \( |0\rangle \) (|1\rangle) of the ancillary qubit. It involves the projector \( P_{m_0}^{(N)} \) onto the subspace of \( m_0 \) spin excitations in an array of \( N \) spins, and reads

\[ U_{m_0}^{(N)} = \exp\left( -i \frac{\pi}{2} P_{m_0}^{(N)} \otimes \sigma_0^z \right) \]

\[ = P_{m_0}^{(N)} \otimes (-i \sigma_0^z) + (1 - P_{m_0}^{(N)}) \otimes 1_0. \]

This equation can be understood as follows: The state of the ancilla qubit, initially prepared in \( |1\rangle \), is flipped by the \( \sigma_0^z \) operation if the system spins are in a state with exactly \( m = m_0 \) excitations, whereas the ancilla qubit is left unchanged otherwise. The QND measurement is then completed by a measurement of the ancilla qubit in the computational basis, providing the desired information on whether \( m = m_0 \) or \( m \neq m_0 \), depending on whether the ancilla qubit is measured in \( |0\rangle \) or \( |1\rangle \).

**B. Construction of the projectors onto excitation number subspaces**

The projector \( P_{m_0}^{(N)} \) required for the unitary (17) can be constructed systematically and efficiently for any \( m_0 \) and \( N \) as follows: One starts from the ansatz

\[ P_{m_0}^{(N)} = \sum_{k=0}^{N} \alpha_k S_z^k \]

with \( S_z = \sum_{i=1}^{N} \sigma_i^z \). This ansatz assures that the projector \( P_{m_0}^{(N)} \) (i) is diagonal in the computational basis, (ii) symmetric under any permutation of spin indices, and (iii) does not involve higher powers with \( k > N \) as such terms are already contained in previous terms with \( k \leq N \) due to the property \( \langle \sigma_i^z \rangle^2 = 1 \). Since the computational basis states are eigenstates of \( S_z \),

\[ S_z |0\ldots0\rangle = N |0\ldots0\rangle, \]
\[ S_z |0\ldots0,1\rangle = (N-2) |0\ldots0,1\rangle, \]
\[ \vdots \]
\[ S_z |1\ldots1\rangle = -N |1\ldots1\rangle, \]

**SUPPLEMENTARY INFORMATION**
and the projector per definition fulfills
\[ P_{m_0}^{(N)} |\ldots 0\rangle = 0, \]

\[ P_{m_0}^{(N)} |0\ldots 1\rangle = |0\ldots 01\rangle, \ldots \]

\[ P_{m_0}^{(N)} |1\ldots 1\rangle = 0, \]

its form is uniquely determined by the following coupled system of \( N + 1 \) linear equations,

\[
\begin{pmatrix}
1 & N & N^2 & \ldots & N^N \\
(N - 2) & (N - 2)^2 & \ldots & (N - 2)^N \\
\vdots & \vdots & \vdots & \ddots & \vdots \\
1 & -N & (N - 2)^2 & \ldots & (N - N)^N
\end{pmatrix}
\begin{pmatrix}
\alpha_0 \\
\alpha_1 \\
\vdots \\
\alpha_N
\end{pmatrix} =
\begin{pmatrix}
0 \\
\vdots \\
1 \\
0
\end{pmatrix}
\]

with the only non-zero entry in the \((N - m_0 + 1)\)-th row. This matrix equation is readily solved, yielding for the experimentally relevant case of three system spins with one or two spin excitations the projectors

\[
P_1^{(3)} = |011\rangle\langle 011| + |101\rangle\langle 101| + |110\rangle\langle 110| 
\]

\[
= \frac{1}{16} (9 - 9S_z - S_z^2 + S_z^3) 
\]

\[
= \frac{1}{8} (3 - (\sigma_1^z + \sigma_2^z + \sigma_3^z) \\
- (\sigma_1^z \sigma_2^z + \sigma_1^z \sigma_3^z + \sigma_2^z \sigma_3^z) + 3\sigma_1^z \sigma_2^z \sigma_3^z)
\]

and

\[
P_2^{(3)} = |001\rangle\langle 001| + |010\rangle\langle 010| + |100\rangle\langle 100| 
\]

\[
= \frac{1}{16} (9 + 9S_z - S_z^2 - S_z^3) 
\]

\[
= \frac{1}{8} (3 + (\sigma_1^z + \sigma_2^z + \sigma_3^z) \\
- (\sigma_1^z \sigma_2^z + \sigma_1^z \sigma_3^z + \sigma_2^z \sigma_3^z) - 3\sigma_1^z \sigma_2^z \sigma_3^z)
\]

where \(S_z = \sum_{i=1}^3 \sigma_i^z\). Note that the projectors are closely related and can be transformed into each other by the symmetry operation \(S_z \rightarrow -S_z\), which interchanges the role of up- and down-spins, or occupied and empty sites in the hardcore boson model, respectively.

C. Experimental implementation of the QND measurement

As by the total spin excitation number \(m\) a collective property of the entire spin system is measured, the unitary of Eq. (17) truly is a many-qubit operation: Equations (19) and (20) show that the projectors contains 1, 2 and 3-body spin interaction terms, such that the QND mapping of Eq. (17) involves interaction terms with up to 4-body Pauli operators. A decomposition of the unitary for the QND measurement \(U_{m_0}^{(N)}\) into experimentally available gates, as obtained using the numerical optimization algorithm, is shown in Table VIII. We note that since the QND measurement involves a global unitary, a numerical optimization has to be done separately for any register size and any particular spin excitation number, and furthermore becomes inefficient for increasing register sizes. However, the unitary can be implemented efficiently without resorting to numerical optimization: For a system of \(N\) spins, \(U_{m_0}^{(N)}\) will generally be the product of unitaries corresponding to many-spin interactions, at most \((N + 1)\)-body Pauli operators. These unitaries can be decomposed into sequences of available gates following the recipes outlined in (Müller et al., 2011). Although the implementation of \(U_{m_0}^{(N)}\) becomes experimentally demanding for increasing system sizes \(N\) and in general requires more operations than numerically optimized circuits, the number of required operations for the QND mapping operation still grows polynomially with the number of system spins.

<table>
<thead>
<tr>
<th>Number</th>
<th>Pulse</th>
<th>Number</th>
<th>Pulse</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>R(0.5, -0.5)</td>
<td>11</td>
<td>R(0.146, -0.895)</td>
</tr>
<tr>
<td>2</td>
<td>R(0.5, 0.0)</td>
<td>12</td>
<td>MS(0.375, -1.054)</td>
</tr>
<tr>
<td>3</td>
<td>(S_z(0.5, 3))</td>
<td>13</td>
<td>(S_z(0.364, 3))</td>
</tr>
<tr>
<td>4</td>
<td>MS(0.125, 0.0)</td>
<td>14</td>
<td>MS(0.75, -1.054)</td>
</tr>
<tr>
<td>5</td>
<td>R(0.098, 1.0)</td>
<td>15</td>
<td>R(1.0, 0.0)</td>
</tr>
<tr>
<td>6</td>
<td>(S_z(1.636, 3))</td>
<td>16</td>
<td>(S_z(1.818, 3))</td>
</tr>
<tr>
<td>7</td>
<td>MS(0.25, 0.0)</td>
<td>17</td>
<td>R(0.277, -1.054)</td>
</tr>
<tr>
<td>8</td>
<td>R(0.136, 0.5)</td>
<td>18</td>
<td>(S_z(0.152, 3))</td>
</tr>
<tr>
<td>9</td>
<td>(S_z(0.75, 3))</td>
<td>19</td>
<td>R(0.5, 0.895)</td>
</tr>
<tr>
<td>10</td>
<td>R(0.113, -1.054)</td>
<td></td>
<td></td>
</tr>
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</table>

TABLE VIII Pulse sequence for the QND post-selective measurement of the spin excitation number in a system of 3+1 ions.

Due to the considerable complexity of the mapping operation for the QND measurement, with the optimized circuit involving in total 19 operations (see Table VIII), this method for error detection itself can only be implemented with a certain accuracy and requires a constant resource overhead, independently of the number of dynamical maps applied in the simulation. It can be seen from the data shown in Figs. 3a and b, and Fig. 4a of the main text, that for short sequences such as e.g. only a single elementary dissipative map, where the system without error detection remains with high probability in the desired excitation number subspace, experimental imperfections in the QND measurement itself actually introduce even more errors on the state of the system (red data points) than in the case where it is not applied (blue data points). However, for longer sequences of dynamical
maps, where the population loss out of the initial excitation number subspace becomes more and more significant, the application of the QND post-selective method becomes effective and enables a more accurate simulation of the system dynamics for longer times.

V. QUANTUM ERROR REDUCTION SCHEME - EXCITATION NUMBER STABILIZATION

A. General idea

The second error reduction procedure goes beyond the error detection method described in the previous section: it does not only allow one to detect errors, which have changed the ideally conserved spin excitation number \( m \) during the quantum simulation, but performs an active stabilization of the register of system spins in the wanted subspace of spin excitation number \( m_0 \). In the previous post-selective case the ancilla qubit carried the binary information whether the system is in the correct subspace or not. Here, in contrast, it is necessary to distinguish between at least three cases: (i) The excitation number \( m \) is correct, and thus no error correction process is required; the excitation number is (ii) too small or (iii) too large, as illustrated in figure 5b,c in the main text. This information cannot be stored in a single ancilla qubit with two logical states anymore. Although it is in principle possible to use a higher-dimensional ancillary system, such as multiple ancillary qubits, to store the required information, such an approach would require significantly more complex detection and correction algorithms. Here, we choose an alternative approach, which still allows one to perform the excitation number stabilization with a single ancilla qubit if the stabilization process is implemented as a two-step process: One stabilization step injects a stabilization into the register if there are too few excitations present (single excitation into the register if there are too many excitations present in the system) and can be understood as follows: The state of the ancilla qubit prepared in \( |1\rangle \) is flipped if and only if there are too many spin excitations present in the system; here the system operator \( \sum_{j=m_0+1}^{N} P_j^{(N)} \) denotes the projector onto the subspace of states containing strictly more excitations than the desired value \( m_0 \). It is the sum over all projectors \( P_j^{(N)} \) onto subspaces with \( j > m_0 \) excitations in a system of \( N \) spins, and each of these projectors can be readily constructed following the procedure described in Sec. IV.B.

After the mapping operation between the system spins and the ancilla, the second part of the protocol deals with the actual extraction of a spin excitation from the system. This step faces the difficulty that due to the QND character of the first mapping step, the state of the ancilla qubit only encodes information about whether too many excitations are present in the system, but not on which sites the excitations are located. In order to minimally disturb the state of the system spins it is desirable to devise an extraction scheme with the following properties: (i) An excitation is extracted in a minimally invasive way, i.e. under an extraction of an excitation from a certain site, off-diagonal order among system spins of the rest of the chain is maintained as far as possible. (ii) Second, the scheme should effectively hold and not further alter the state of the system spins once an excitation has been successfully extracted. We have developed and implemented a scheme, which satisfies both criteria, and which exploits a combination of spectroscopic decoupling and optical pumping of the ancillary qubit.

B. Steps of the excitation extraction procedure

Let us now describe in more detail the steps of the excitation extraction procedure, which is illustrated in Fig. S5 for a system of three spins. In step (1) the mapping introduced above is applied, so that the information whether a correctable error (i.e. too many spin excitations in the system) is present (or not), is encoded in the ancilla state \( |0\rangle \) (\( |1\rangle \)). If no correctable error is detected \( (m \leq m_0) \) then the subsequent operations should not affect the system qubits. This can be realized by removing the ancilla qubit incoherently from the computational subspace (step (2)) as will be discussed below. In step (3)
Excitation number stabilization procedure
(spin excitation / hardcore boson extraction protocol)

(1) QND map: m > 1?

(1) if m ≤ 1

m = 1: no error
m = 0: error (not correctable now)

(2) removal of the ancilla by pumping to "parking"

(2) ... no effect

(3) exc. extraction intent from site #1

(3) ... no exc. on site #1

... thus no exc. extracted

(4) removal of the ancilla to "parking" state

(4) ... no effect

(5) exc. extraction from site #2

(5) 1 exc. extracted with certainty

(6) reset of the ancilla to |0⟩

error completely corrected

final state: no further effect of steps (3) - (5)
on the system

final state:

no error or uncorrectable m=0 error

no further effect of step (5)

final state:

error completely corrected or exc. number reduced from m=3 to m=2

FIG. S5 Logical tree illustrating the complete excitation extraction procedure for a 3-spin system with ideally a single excitation present, and for various (unknown) initial states of the system spins. Blue circles correspond to spin excitations (hardcore bosons) localized on particular sites, whereas ellipses indicate situations where excitations (hardcore bosons) can be delocalized over several lattice sites. The error reduction takes place in six steps: By a QND map (1) the information whether or not too many excitations are present in the system, is mapped onto an ancilla qubit. (2) In case of the ideal excitation number or too few excitations, the ancilla is removed from the computational space and the protocol effectively halts. In the opposite case, a swap operation (3) is applied to try to extract an excitation from system site #1. If successful, after the subsequent removal of the ancilla (4) the protocol halts. Otherwise, (5) an extraction by a swap of the ancilla with system site # 2 extracts an excitation. (6) Finally, the ancilla is reset to its initial state |0⟩ for subsequent error reduction steps. See also text for a more detailed explanation.
an extraction attempt on the first site of the spin register is performed by swapping the ancilla, which is now in the state $|0\rangle$ corresponding to an empty site (no spin excitation), with the first qubit of the system. In terms of spins this operation corresponds to a swap (“flip-flop”) process, under which $|0\rangle_0|1\rangle_1 \rightarrow |1\rangle_0|0\rangle_1$, i.e. the spin excitation is coherently exchanged between the ancilla qubit and the first system spin. In the language of hardcore bosons, the process can be interpreted as a tunneling of a boson from the site \#1 onto the previously empty ancilla site (see step (3) in Fig. S5). If after this process the ancilla qubit is in the state corresponding to an occupied site (spin excitation present), a single excitation has been successfully extracted from the system and the process should halt. As will be shown below, this can be performed by removing the population of the ancilla qubit from the computational subspace (step (4)). However, if the ancilla is still in the state $|0\rangle$ corresponding to an empty site after the swap operation, then the first system qubit was in the state corresponding to an unoccupied site and thus no spin excitation (or hardcore boson) could extracted, $|0\rangle_0|0\rangle_1 \rightarrow |0\rangle_0|0\rangle_1$. Thus, this procedure is repeated (step (5)) until the ancilla qubit is found in the state $|1\rangle$ corresponding to an occupied site. In the case of three sites at most two extraction rounds are required (steps (3), (4), and (5)), before the ancilla is reset to $|0\rangle$ (step (6)) for subsequent rounds of error reduction.

**Open vs. closed-loop error reduction scheme** – The described extraction (and injection) protocols are realized in an open-loop fashion, i.e. the ancilla remains unobserved during the whole procedure. This comes at the cost that all pulses (for steps (1) - (6) in the above example) have to be physically applied in every run of the error reduction procedure, even if the ancilla has been already be removed from the computational subspace and in principle no further operations would be required. However, the described protocol could easily be modified into a closed-loop control scenario: In this case one would perform individual measurements of the ancilla qubit after the QND map (1) as well as after the coherent swap operations (steps (3) and (5)). The outcomes of these ancilla measurements would then yield the classical information on whether no error correction is required after step (1) or an excitation has already been extracted successfully in steps (3) or (5). This information could then be used for classical feedback on the quantum system in the sense that one can externally decide whether further steps are required in the error reduction protocol, or whether one can stop the current run of the error reduction protocol and no further operations have to be applied. An advantage of this modified scheme is that dissipative removal operations of the ancilla qubit out of the computational subspace are not required. Experimentally, the price to pay in this modified scheme would be a number of fluorescence measurements, which are slow compared to the application of coherent gates, and the requirement to recool the relevant external degrees of freedom of the ion chain after each measurement as recently demonstrated in (Schindler et al., 2012).

### C. Experimental implementation

In the experiment, the error reduction protocol for a system of three spins, using 3+1 ions was implemented. We considered the case of having ideally $m_0 = 1$ excitation present on the three sites. For the extraction step, the four-qubit unitary $U_{m>1}^{(3)}$ (cf. Eq. (21)) to detect whether too many excitations (i.e. $m = 2, 3$) are present in the system or not, has been obtained by numerical optimization. The resulting circuit consisting of 15 gates is listed in Table IX. On the other hand, for the injection part, the unitary

$$
U_{m<m_0}^{(N)} = \exp \left( -\frac{i\pi}{2} \sum_{j=0}^{m_0-1} P_j^{(N)} \otimes \sigma_0^z \right) 
$$

which interrogates the system whether too few excitations are present, is required. For the case $m_0 = 1$ and $N = 3$, the unitary $U_{m<1}^{(3)}$ realizes a spin flip of the ancilla qubit only if the three system spins are in the state $|000\rangle$. This operation is equivalent to a triple controlled-NOT operation with the three system spins playing the role of the control qubits and the ancilla the target qubit. As we could not obtain a satisfying circuit decomposition for this unitary by means of the usual numerical optimization algorithm, we did not try do directly optimize $U_{m<1}^{(3)}$, but allowed the unitary to add arbitrary phases to states lying outside the desired excitation number subspace $m_0 = 1$. Under these relaxed conditions, the numerical optimizer delivered a circuit decomposition of 19 operations as shown in Table X.

<table>
<thead>
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<th>Number</th>
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<th>Number</th>
<th>Pulse</th>
</tr>
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<tbody>
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<td>1</td>
<td>$R(0.25, 0.5)$</td>
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<td>$S_z(1.0, 3)$</td>
</tr>
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<td>$S_z(1.0, 3)$</td>
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</tr>
<tr>
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<td>$R(0.25, 0.5)$</td>
<td>11</td>
<td>$MS(0.5, 0.5)$</td>
</tr>
<tr>
<td>4</td>
<td>$MS(0.25, 0.5)$</td>
<td>12</td>
<td>$R(1.5, 0.5)$</td>
</tr>
<tr>
<td>5</td>
<td>$R(1.75, 0.0)$</td>
<td>13</td>
<td>$R(0.125, 0.5)$</td>
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<td>$S_z(1.0, 3)$</td>
</tr>
<tr>
<td>7</td>
<td>$R(0.25, 0.0)$</td>
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<td>$R(1.875, 0.5)$</td>
</tr>
<tr>
<td>8</td>
<td>$R(1.75, 0.5)$</td>
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</table>

**TABLE IX** Pulse sequence to implement the QND unitary $U_{m>1}^{(3)}$ (cf. Eq. (21)) as part of the excitation extraction protocol. It maps the information whether there are more than $m = 1$ spin excitations present in a system of three spins, onto the ancilla qubit.

**Implementation of the actual excitation extraction** – We will now discuss the physical mechanism underlying
TABLE X Pulse sequence to implement the QND unitary $U_{m<1}^{(3)}$ (cf. Eq. (22)) as part of the excitation injection protocol. It maps the information whether there are $m = 0$ instead of ideally $m = 1$ spin excitations present in a system of three spins, onto the ancilla qubit.

<table>
<thead>
<tr>
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<td>$R(1.75, 0.5)$</td>
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<td>$S_z(1.0, 3)$</td>
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<td>3</td>
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<td>$R(1.625, 0.0)$</td>
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</tr>
<tr>
<td>5</td>
<td>$R(1.875, 0.0)$</td>
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<td>6</td>
<td>$S_z(1.0, 3)$</td>
<td>17</td>
<td>$R(0.125, 0.0)$</td>
</tr>
<tr>
<td>7</td>
<td>$R(0.125, 0.0)$</td>
<td>18</td>
<td>$S_z(1.0, 3)$</td>
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<td>$R(1.875, 0.5)$</td>
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</table>

FIG. S6 a, Schematic view of the error detection and excitation removal process (cf. Fig. 5c of the main text). b, The excitation removal process on 3 system qubits can be performed by swap operations and effective halting conditions realized by a dissipative decoupling of the ancilla qubit. c, Swap operations (23) between the ancilla qubit and one system spin are implemented by two effective two-qubit MS operations, which are build up from 4 MS operations acting on three ions, interspersed with refocusing operations.

FIG. S7 The dissipative decoupling process of the ancilla qubit used in the injection/removal processes. The goal is to uni-directionally remove population in the computational state $|0\rangle$ (red triangular marker) and add it to electronic population that is possibly already present in the "parking" state $|2\rangle$ (green rectangular marker). Population in the computational state $|1\rangle$ (blue circle) should be left unchanged. The sequence: (i) First, the population in states $|1\rangle$ and $|2\rangle$ is coherently swapped by a π-operation. (ii) Population in $|0\rangle$ is coherently transferred to the state $4S_{1/2}(m_j = +1/2)$. (iii) Optical pumping from this state towards the $|1\rangle$ state. In this step, the populations that were at the beginning of the sequence present in $|2\rangle$ and $|0\rangle$ are added up and temporarily accumulate in state $|1\rangle$. (iv) Finally the populations in states $|1\rangle$ and $|2\rangle$ are again swapped coherently.

the implementation of the removal of a spin excitation (or hardcore boson), as described above and illustrated in Fig. S6. This excitation removal step takes place after the QND mapping (1) and a decoupling operation on the ancilla qubit (2) described below. As mentioned above, the extraction step (3) relies on a swap operation exchanging the excitation of the ancilla qubit with one
of the system spins, say #1. This swap operation is realized by applying a π-pulse, according to the "flip-flop" Hamiltonian

\[ H_{01} = \sigma_0^+ \sigma_1^- + \sigma_0^- \sigma_1^+ = \frac{1}{2}(\sigma_0^x \sigma_1^x + \sigma_0^y \sigma_1^y). \]

The resulting unitary

\[ U_{01} = \exp(-i\frac{\pi}{2}H_{01}) = \exp(-i\frac{\pi}{4}\sigma_0^y \sigma_1^y) \exp(-i\frac{\pi}{4}\sigma_0^y \sigma_1^y) \]

(23)
corresponds to the application of two fully-entangling x- and y-type MS gates (see Eq. (3)) applied to the ancilla ion and the first system ion. This two-ion MS gate can be realized by the global bi-chromatic light field, if all ions but the ancilla qubit and the system qubit #1 are spectroscopically decoupled. However, since step (3) involves the ancilla qubit and system ion #1, and step (5) the ancilla qubit and system ion #2, it is from an experimentally point of view more convenient to spectroscopically decouple only ion #3, but keep both ions #1 and #2 in the "active" qubit states during the extraction procedure. Doing so, each of the x- and y-type MS-gates required for the operation (23) acting on the ancilla (index #0) and system ion #1 can be realized by two MS gates acting on the three ions, interspersed by a π-pulse \( S_z(1, 2) \) applied to the second system ion, as shown in Fig. S6. The application of the "refocusing" pulse in between the MS gates leads to an effective decoupling of ion #2 that is not supposed to participate in the unitary (23) (Müller et al., 2011). This technique is also applied to realize the swap operation required in step (5), which acts on the ancilla qubit and the second system. Using this decoupling technique based on refocusing pulses, no additional spectroscopic decoupling operations are required between the two swap operations corresponding to steps (3) and (5). The experimentally employed sequence of 6 gates is listed in Table XI.

In Sec. V.B we have outlined that the ancilla will be removed from the computational subspace in step (2) if no error is to be corrected in the current error reduction round. In this case, if the ancilla has been removed from the qubit subspace, the two-qubit MS interactions appearing in (23) act only on a single qubit. However, in this "pathological" case where the bi-chromatic laser fields used for the generation of the effective spin-spin interactions of the entangling MS gate of Eq.(3), are applied to a single ion only, these realize up to negligible corrections the identity operation on this single ion (Mølmer and Sørensen, 1999; Roos, 2008). As a consequence, in this case – despite the fact that the MS gate laser pulses of steps (3) and (5) are physically applied to the ions – they do not alter the state of the systems spins, as desired.

**Dissipative decoupling of the ancillary qubit** – As explained above, the ancilla qubit shall be removed from the computational subspace if either (i) no correctable error is detected in the current round of error reduction, or (ii) a superfluous spin excitation has been successfully extracted from the spin system. In these cases, the removal of the ancilla from the qubit subspace guarantees that the error reduction protocol effectively halts (see the logical tree shown in Fig. S5) and the state of the system spins is no further modified, even though after the removal of the ancilla MS gate laser pulses are applied to the ion string. In steps (2) and (4) of the spin extraction protocol, the ancilla qubit is removed from the computational state |1⟩ into the "parking" state |2⟩, which we encode in the \( 3D_{5/2} \left( m_j = -5/2 \right) \) electronic state (see Fig. S7). This removal into |2⟩ takes place if the ancilla ion resides at these instances in |1⟩, if it is in |0⟩ its state remains unaffected.

For this removal it is not possible to use the coherent spectroscopic decoupling technique as used for the implementation of the elementary Kraus maps. The reason is that this would lead to errors in the protocol: for instance, imagine no correctable error is present in the system and thus the ancilla is brought to the decoupling state |2⟩ in step (2). If then another ancilla removal were performed coherently (step (4)), the ancilla would be transferred back from |2⟩ to the computational state |1⟩. This is clearly unwanted as in this case with the ancilla returned to |1⟩, the subsequent swap operation (5) would be performed by mistake. This unwanted behavior can be avoided if the transfer of the ancilla from |1⟩ to |2⟩ is realized dissipatively, by an optical pumping process, which bares similarities with the incoherent reset of the ancilla qubit for the elementary dissipative maps. Such uni-directional, incoherent pumping process from |1⟩ to |2⟩ guarantees that once the ancilla has reached the "parking" state |2⟩, it will in subsequent steps never return to the computational subspace. This optical pumping process for a removal process of the ancilla qubit from one of the computational basis states into |2⟩ is illustrated and described in more detail in Fig. S7.

In the spin excitation protocol the ancilla is removed from |1⟩ (corresponding to an occupied site) to |2⟩, whereas in the spin injection protocol it has to removed

<table>
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<th>Number</th>
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<td>MS(0.25, 0)</td>
</tr>
<tr>
<td>2</td>
<td>( S_z(1.0, j) )</td>
</tr>
<tr>
<td>3</td>
<td>MS(0.25, 0)</td>
</tr>
<tr>
<td>4</td>
<td>MS(0.25, 0.5)</td>
</tr>
<tr>
<td>5</td>
<td>( S_z(1.0, j) )</td>
</tr>
<tr>
<td>6</td>
<td>MS(0.25, 0.5)</td>
</tr>
</tbody>
</table>

**Table XI** Pulse sequence for a swap operation used for excitation injection or removal, see Fig. S6c. Ion \( j \) is the spectator ion, which is not spectroscopically decoupled, but is not intended to participate in the swap operation.
from $|0\rangle$ (empty site) to $|2\rangle$. For both scenarios, the pulse sequence outlined in Fig. S7 can be employed; a $\pi$-pulse resonant with the qubit transition of the ancilla, applied before and after the dissipative decoupling sequence, exchanges the roles of the two computational states $|0\rangle \leftrightarrow |1\rangle$ and thereby allows one to switch between the spin extraction and injection scenario.

D. Experimental results for the stabilization

The active excitation number stabilization procedure can be best tested when applying it to a state that has a considerable amount of the population outside the subspace with the correct excitation number $m_0$. We applied Hadamard operations on the three system qubits initially in $|000\rangle$ to prepare the initial state $|\psi_0\rangle = 1/\sqrt{8}(|001\rangle + |010\rangle + |100\rangle)$, which is an equal-weight superposition of all eight three-qubit computational basis states, each occurring with probability $1/8$. The measured and ideal density matrix of this initial state is shown in Fig. S9a. For $m_0 = 1$ the states $|001\rangle$, $|010\rangle$, $|100\rangle$ span the subspace with the desired spin excitation number. Thus, the initial fraction of population in this subspace is $3/8$ as shown in Fig. 5d of the main text. We then performed the excitation extraction protocol according to the protocol outlined above and summarized in Fig. S5. Ideally this protocol extracts one excitation from the components of the initial state, which contain two or three spin excitations, $m = 2 \rightarrow m = 1$ and $m = 3 \rightarrow m = 2$, thereby pumping the population corresponding to these states into the subspaces with one excitation less. It is a crucial property of the error reduction protocol that the coherences within the subspace of the "correct" excitation number $m = 1$ are ideally preserved, as the dynamics within the desired simulation subspace should be affected as little as possible. The component $|000\rangle$ of the initial state corresponds to a state with zero, i.e. too few spin excitations, and thus to an error which is not corrected by the spin extraction procedure.

In Fig. 5d in the main text, the ideal and the measured populations in all four excitation number subspaces at the end of the extraction protocol are shown. From this information one can deduce that the protocol within experimental accuracy realizes the pumping of populations between the different excitation number subspaces as expected. To infer whether the coherences are preserved, we measured the three-qubit density matrices after the first excitation extraction attempt from the first site (step (3) in Fig. S5) and after the second extraction attempt from the second site (step (5)). The measured and the ideal density matrices are shown in Fig. S9b and c, where the relevant coherences within the $m_0 = 1$ subspace are highlighted in red color. From the comparison of measured and ideal density matrices it can be seen that the coherences in the $m_0$ excitation subspace are well-preserved.

The complete error correction protocol consists of the excitation extraction and injection procedures. For the same initial state $|\psi_0\rangle$, we implemented the injection procedure, which ideally only acts on the $|000\rangle$ state, and pumps the population from the $m = 0$ into the $m = 1$ subspace. Again, the coherences within the $m = 1$ subspace should be preserved under this procedure, yielding as a result of the injection protocol the ideal density matrix shown in the right part of Fig. S9d. Comparison with the measured density matrix shows that most of the population is pumped out of the $m = 0$ subspace, and that the initially present coherences within the $m = 1$ subspace are reasonably well conserved.

The ultimate goal of any error reduction protocol is certainly to increase the performance of a complex algorithm. As a step in this direction, we integrated the excitation removal protocol into the simulation sequence for dynamics according to composite dissipative maps with 3+1 ions. In Fig. S8 we compare the probabilities for all excitation numbers $m$ when (i) no error reduction technique is used (blue data points), (ii) with the post-selective QND measurement applied (red), and (iii) with the excitation removal procedure included in the simulation (green). It can be seen that the removal procedure has a higher overhead due to its considerable complexity. Nevertheless, as an indication of its usefulness, a slower decay of the probability of finding the population in the desired subspace for $m = 1$ can be observed, for the case in which the stabilization procedure is applied, compared to the case without any error correction. This indicates that the stabilization procedure indeed works qualitatively as intended, when it is incorporated into the actual simulation.

VI. ADDITIONAL EXPERIMENTS AND DATA ANALYSIS FOR COMPETING DISSIPATIVE AND COHERENT DYNAMICS

In Figs. 3 and 4 of the main text, the experimental results for systems of 3+1 and 4+1 ions are depicted. The attentive reader noticed that in Fig. 4a in the main text the data point for 5 maps is missing. This is due to the fact that in view of the length of the algorithm, the memory of the current experimental control system is not sufficient to generate the required pulse sequence for 4 elementary dissipative maps and one composite Hamiltonian dynamical map. It is however possible to generate the sequence for two elementary maps and one Hamiltonian dynamical map, and to repeat this sequence twice (data point for 6 maps).

Here, we extend the experimental analysis that was omitted from the main text due to space restrictions by showing different measures for the already presented data and also additional datasets: In Figs. S11 and S10 we add
FIG. S8 Experimental study of the error reduction protocol, incorporated into the actual quantum simulation of composite dynamical maps with 3+1 ions, with an initial and ideal excitation number of $m = 1$. The four plots show the fraction of the population in the subspaces with zero, one, two and three spin excitations, as a function of the number of elementary dissipative dynamical maps applied. For comparison, blue data points corresponds to the population under dissipative maps, without error detection or reduction technique applied. Red data points correspond to the case where after the final elementary dissipative map the QND post-selective measurement was applied. Green data points corresponds to the case where the extraction protocol has been applied at the end of the simulation. Bars are the theory, where depolarizing noise in the elementary dissipative maps is taken into account (cf. Sec. III.A).

the measures purity and off-diagonal order for the data already presented in Figs. 3 and 4 of the main text. The purity $\text{Tr} \rho^2$ is a measure for how close the measured state is to a pure state. Off-diagonal order measures the coherences between neighboring sites as the expectation value of the operator $\sum_j \sigma_{j}^{(j)} \sigma_{j+1}^{(j+1)}$ evaluated within the $m_0$-excitation subspace. This parameter emphasizes the effect of the competing Hamiltonian dynamics as it changes the sign from positive to negative after the application of a Hamiltonian map, as shown in figure S11.

Figure S12 shows an alternative dataset which demonstrates that the coherent competing Hamiltonian dynam-ical map in a system of three sites has - as physically expected - no effect if only a single excitation is present in the system. Comparison with the data of Fig. 4a in the main text, underlies the significance of the decrease in the overlap with the overlap with the target Dicke state $|D(2,3)\rangle$, and this effect is indeed caused by the competing Hamiltonian dynamics.

Figure S13 shows a dataset with competition and two spin excitations present in the system. Compared to the analysis in the main text, the competition strength is now set to $\phi = \pi/4$ instead of $\pi/2$. As expected, this leads to a reduced effect of the competing Hamiltonian maps on the dissipatively created order.

REFERENCES

FIG. S9 Reconstructed (left) and ideal (right) density matrices of the removal and injection process in a 3+1 ion experiment. Populations and coherences within the desired excitation number subspace with \( m = 1 \) are highlighted by red bars. 

a, The initial state is an equal-weight superposition of all eight computational basis states. 

b, The state after the system after a spin excitation removal attempt from the first site. 

c, The state of the system after the second swap operation to remove an excitation from the second site. After both steps, the coherences shown in red are well-preserved. 

d, The state of the system after a spin excitation injection attempt, starting again in the equal-weight superposition state shown in a. The population from the \(|DDD\rangle = |000\rangle\) state is depleted, and coherences within the \( m = 1 \) subspace are reasonably well preserved.
FIG. S10 Experimental results of dissipatively induced delocalization through composite dynamical maps with 3+1 ions. The results from an ideal model are shown in light-blue bars whereas those from a model including depolarization noise are indicated by dark-grey bars. Blue rectangles indicate the experimentally observed dynamics without any correction scheme whereas red rectangles include a post-selective error detection scheme (error bars, 1σ). Overlap fidelity, purity, population in the $m = 2$ subspace, and off-diagonal order in a 4-spin quantum simulation with 4+1 ions, studying purely dissipative dynamics that induces pumping towards the Dicke state $|D(2, 4)\rangle$ as shown in figure 3 in the main text.

FIG. S11 Experimental results for competing dissipative and coherent dynamics with 3+1 ions The results from an ideal model are shown in light-blue bars whereas those from a model including depolarization noise are indicated by dark-grey bars. Blue rectangles indicate the experimentally observed dynamics without any correction scheme whereas red rectangles include a post-selective error detection scheme (error bars, 1σ). Overlap fidelity, purity, population in the $m = 2$ subspace, and off-diagonal order in a 3-spin quantum simulation with 3+1 ions. The dynamics corresponds to dissipative maps and coherent Hamiltonian competition as shown in figure 4 in the main text.
FIG. S12 Experimental results for competing dissipative and coherent dynamics with 3+1 ions with a single excitation. The results from an ideal model are shown in light-blue bars whereas those from a model including depolarization noise are indicated by dark-grey bars. Blue rectangles indicate the experimentally observed dynamics without any correction scheme whereas red rectangles include a post-selective error detection scheme (error bars, 1σ). Data from simulated dynamics with Hamiltonian competition for 3 spins (3+1 ions), but only a single spin excitation present. As physically expected, the data confirms that in this case the Hamiltonian dynamical map has no effect.

FIG. S13 Experimental results for competing dissipative and coherent dynamics with 3+1 ions with weak interaction. The results from an ideal model are shown in light-blue bars whereas those from a model including depolarization noise are indicated by dark-grey bars. Blue rectangles indicate the experimentally observed dynamics without any correction scheme whereas red rectangles include a post-selective error detection scheme (error bars, 1σ). Data from simulated dynamics with weaker competing Hamiltonian dynamics (φ = π/4) in a 3-spin system with two spin excitations present in the system (3+1 ions).