Programmable interactions and emergent geometry in an array of atom clouds

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Interactions govern the flow of information and the formation of correlations between constituents of many-body quantum systems, dictating phases of matter found in nature and forms of entanglement generated in the laboratory. Typical interactions decay with distance and thus produce a network of connectivity governed by geometry—such as the crystalline structure of a material or the trapping sites of atoms in a quantum simulator1,2. However, many envisioned applications in quantum simulation and computation require more complex coupling graphs including non-local interactions, which feature in models of information scrambling in black holes3–6 and mappings of hard optimization problems onto frustrated classical magnets7–11. Here we describe the realization of programmable non-local interactions in an array of atomic ensembles within an optical cavity, in which photons carry information between atomic spins12–19. By programming the distance dependence of the interactions, we access effective geometries for which the dimensionality, topology and metric are entirely distinct from the physical geometry of the array. As examples, we engineer an antiferromagnetic triangular ladder, a Möbius strip with sign-changing interactions and a treelike geometry inspired by concepts of quantum gravity5,20–22. The tree graph constitutes a toy model of holographic duality21,22, in which the quantum system lies on the boundary of a higher-dimensional geometry that emerges from measured correlations23. Our work provides broader prospects for simulating frustrated magnets and topological phases24, investigating quantum optimization paradigms10,11,25,26 and engineering entangled resource states for sensing and computation27,28.

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Controllable quantum systems of cold atoms constitute powerful platforms for applications ranging from quantum simulation2 to entanglement-enhanced measurement12,14,15,29. Positioning atoms in optical tweezers and inducing local interactions by excitation to Rydberg states enables simulations of lattice spin models2 and bottom-up control of entanglement30. In a complementary top-down paradigm, collective entangled states have been generated in ensembles of up to $5 \times 10^4$ atoms with all-to-all interactions mediated by photons in optical cavities12–17 or by collisions in Bose–Einstein condensates29–34. For photon-mediated interactions, or analogous phonon-mediated interactions among trapped ions35,36, using multiple bosonic modes further enables tuning of the interaction range35–37.

Yet many objectives demand more versatile control of the graph of interactions3,6,8,24,38, including couplings that are neither local nor global. Engineering a wider range of non-local graphs opens up prospects for accessing spin-glass phases38,39 and new topologically ordered states40, exploring analogue approaches to combinatorial optimization10,11,25,26 and probing toy models of quantum gravity5,6,20–22. These goals have motivated proposals for programming the distance dependence of spin-exchange interactions in arrays of atoms or ions by tailoring the frequency spectrum of a drive field3,24,38, which couples the spins to a single mode of light or motion.

We realize programmable spin-exchange interactions in an array of atomic ensembles within an optical cavity. Our scheme (Fig. 1) produces a class of spin models described by an effective Hamiltonian

$$H_{\text{eff}} = - \sum_{\mu, \nu} J(r_{\mu\nu}) f_{\mu}^+ f_{\nu} + H_q$$

(1)

Here $f_{\mu}^+$ and $f_{\nu}^-$ denote the raising and lowering operators for the Zeeman spin of atoms $\mu$ and $\nu$, respectively; the distance between atoms is $r_{\mu\nu}$ and $H_q$ represents the quadratic Zeeman energy. The spin-exchange coupling $J$ arises from a process in which one atom flips its spin down while scattering a photon from a drive field into the cavity and a second atom rescatters this photon to flip its spin up3,13. We focus on a system of spin-1 atoms initialized in the $m = 0$ Zeeman state, in which the effect of this ‘flip-flop’ interaction is to produce correlated atom pairs in states $m = \pm 1$ (refs. 17,27,31,33,34,40).

Whereas the single-mode cavity ordinarily mediates interactions among all sites, we break this all-to-all connectivity by introducing a
magnetic field gradient along the cavity axis. The gradient introduces an energy difference $\Delta E = h\omega_B$ between the Zeeman splittings on adjacent sites, such that spin-exchange processes are off-resonant for spatially separated spins. To controllably reintroduce interactions between ensembles spaced by a distance of $r$ sites, we modulate the intensity of the drive field at frequency $\omega_B$, thus modulating the instantaneous spin-exchange coupling $J(t)$ as a function of time $t$. More generally, to obtain a specified set of couplings $J(r)$, we set a drive waveform

$$J(t) = \sum_r e^{-i\omega_B t}J(r)$$

according to the Fourier transform of the couplings. The drive waveform thus determines the dispersion relation $\chi(k) = -2\sqrt{\nu_k/\hbar\omega_B}$ for spin waves with momentum $k$, in dimensionless units of inverse sites, in a system of $N$ atoms per site.

We benchmark the interactions in a one-dimensional array of $M=18$ ensembles of $N=10^4$ rubidium-87 atoms, initializing all atoms in state $m=0$. This initial state with no average magnetization would classically have trivial dynamics. However, quantum fluctuations are amplified by the cavity-mediated interactions, producing correlated growth and macroscopic fluctuations in the populations of states $m = \pm 1$ (ref. 5). The spatial correlations of these amplified fluctuations reveal the connectivity of interactions.

To implement distance-dependent interactions, we optically drive the cavity for 100 to 200 $\mu$s, thereby transferring 30–50% of the atoms into the states $m = \pm 1$. Subsequently, we perform state-sensitive imaging to obtain the correlations

$$C_{ij}^{pm} = \frac{\text{Cov}(N_{i,m}, N_{j,n})}{\sqrt{\text{Var}(N_{i,m})\text{Var}(N_{j,n})}} \equiv \text{Corr}(N_{i,m}, N_{j,n}),$$

for each pair of sites $(i,j)$, in the populations $N_{i,m}, N_{j,n}$ of states $m = \pm 1$. Here, Cov and Var denote the covariance and the variance, respectively. Figure 1b shows the measured correlations for three different scenarios. For a monochromatic drive field, in a uniform magnetic field, we observe correlations of equal strength between all sites, indicating the expected all-to-all interactions. By contrast, adding a magnetic field gradient results in correlations being localized to individual sites. Finally, modulating the intensity of the drive light at frequency $\omega_B$ produces correlations between all pairs of sites separated by a distance $|i-j| = r$, as shown for $r = 10$.

The dependence of spatial correlations on modulation frequency is shown in Fig. 2a. There, we plot the average correlation intensity of the drive field at a frequency $\omega_B$, we generate correlated atom pairs in states $m = \pm 1$ at distance $r$. Measured correlations $C_{ij}^{pm}$ for integer values $r$ reveal correlations at distances $d = r \pm 1$ and $d = 0$. While the correlations at $d = 0$ indicate on-site pair creation that is resonant even for a single drive frequency, the correlations at $d = r$ confirm the presence of interactions at the distance set by the modulation frequency.
The interactions are spectrally well resolved as a function of drive frequency (Fig. 2a, inset), highlighting the precise control of the coupling distance.

To sensitively probe the growth and spreading of correlations, we examine the transverse magnetization, which provides an enhanced signal at early times. Specifically, we evaluate the normalized covariance \( C^{\text{xx}} = \text{Cov}(F^x_i, F^x_j)/N^2 \), where \( F^x_i = \sum_{\mu=1}^{\mu} f^x \mu \) denotes the collective magnetization on site \( i \) in a rotating frame set by the local magnetic field. Figure 2b shows \( C^{xx} \) as a function of time and distance \( d \), averaged over all sites \( i \), for a system programmed to interact at distance \( r = 3 \). Correlations first appear between nearest neighbours on the coupling graph and spread over time to further neighbours at multiples of the distance \( r \). We additionally compute the structure factor \( F^x_k = \sum_i e^{i k F^x_i}/\sqrt{M} \) plotting its root mean square value in Fig. 2c. We observe narrowing in momentum space as a function of time, complementary to the observed spreading of correlations in position space.

The growth of the structure factor is consistent with an analytical model in which spin waves of momentum \( k \) are amplified by a factor proportional to \( |\chi_k| \) per Bloch period of evolution. Equivalently, the growth in \( \bar{F}^x \) for each momentum mode \( k \) is proportional to the drive intensity \( J(t) \) at time \( t = k/\omega_B \). The amplification is notably strongest at minima of the dispersion relation \( \chi_k < 0 \). Pair creation thus drives the system towards states of minimal interaction energy, while increasing the quadratic Zeeman energy \( H_B \) to compensate.

Engineering the dispersion relation via the drive waveform enables the realization of periodic boundary conditions, despite the physical geometry of our array as an open chain. For a chain of \( M \) sites with periodic boundary conditions, the domain of the dispersion relation is a discrete set of points in momentum space, spaced by \( \Delta k = 2\pi/\mathcal{M} \). Correspondingly, we break the drive waveform into a train of short pulses with spacing \( \tau_p/\text{Min} \), where \( \tau_p = 2\pi/\omega_B \) is the Bloch oscillation period for spin excitations.

For an initial sinusoidal modulation designed to introduce interactions at distance \( r \), the pulsed variant has a frequency spectrum that includes peaks at both \( \omega_0 \) and \( (M-r)^{-1} \omega_0 \). The resulting correlations \( C^{xx} \), shown in Fig. 2d, are strongest at distances \( d = \pm r \) and \( d = \pm M - r \), indicating that the system now behaves as though the sites were situated on a ring.

We verify the periodic boundary conditions by directly reconstructing the effective geometry of the system from measured spin correlations \( C^{xx} \). Adopting an ansatz that correlations decay as a Gaussian function \( |C^{xx}| \propto \exp(-|p_p-p_j|/\mathcal{D}) \) of distance \( |p_p-p_j| \) in a \( D \)-dimensional space, we seek a mapping of the array sites to effective coordinates \( \rho \), that best fit the distances \( d_{ij} = \sqrt{\log |C^{xx}|} \) inferred from the correlations (Methods). We obtain the coordinates \( \rho \) from the distance matrix \( d_{ij} \) by metric multidimensional scaling. The result is shown in Fig. 3a for a system with nearest-neighbour interactions and periodic boundary conditions. We additionally calculate an inferred coupling matrix \( J_c = (C^{xx})^{-1} \). Colouring the edges between all pairs of sites according to \( J_c \) corroborates the ring-like coupling graph.

Tailoring the drive waveform enables versatile control over the geometry and topology of the coupling graph, as we illustrate by the same black-box reconstruction technique. We first observe that introducing interactions at a distance \( r > 1 \), with open boundary conditions, produces a set of \( r \) disjoint chains, as depicted in Fig. 3b for \( r = 3 \). Linking such chains with a second modulation frequency generates a two-dimensional graph, as shown by the triangular ladder in Fig. 3c, formed by interactions at distances \( r_1 = 1 \) and \( r_2 = 2 \). Furthermore, adding periodic boundary conditions enables the realization of non-trivial topologies. For example, Fig. 3d shows a square-lattice cylinder, whereas Fig. 3e shows a Möbius ladder. The characteristic twist of the Möbius strip is evident in the crossing of two bonds in the reconstructed geometry.
For a given coupling graph, the sign of the interaction $J(r)$ at each distance $r$ is set by the phase of the modulation at frequency $\omega$. We always choose the on-site interaction to be ferromagnetic, favouring a large spin polarization on each site, and choose a phase $\arg(J(r)) \in [0, \pi]$ to set either ferromagnetic or antiferromagnetic couplings at each non-zero distance $r$. Figure 3 includes examples with ferromagnetic (Fig. 3a, b), anti-ferromagnetic (Fig. 3c) and sign-changing (Fig. 3d, e) couplings.

The antiferromagnetic triangular ladder in Fig. 3c constitutes a fully frustrated $XY$ model. In the classical ground state, adjacent spins have a relative angle of approximately 120° (white arrows in Fig. 3c, top row). The measured spin correlations in Fig. 3c are consistent with the predicted ordering, with two degenerate minima in the structure factor (green squares) ordering of sites. The minimal correlation over all possible bipartitions (purple triangles) is peaked at $s = 1$, indicating a breakdown of locality. Error bars denote s.d. estimated by jackknife resampling.

Engineering non-local couplings enables radically different geometries to be explored, beyond those that can be visualized by an embedding of sites in a Euclidean space. Inspired by models of quantum gravity, for which the points on the real line are best viewed as leaves on an infinite regular tree graph. Such tree graphs feature in a version of the anti-de Sitter/conformal field theory correspondence ($p$-adic AdS/CFT), in models of information scrambling in black holes, and in tensor-network representations of strongly correlated states.

To access a treelike geometry, we engineer couplings

$$J(i - j) = \begin{cases} \frac{\varphi(i - j)}{\pi^4} & |i - j| = 2^n, \quad n \in \mathbb{Z} \\ 0 & \text{otherwise} \end{cases}$$

depicted in Extended Data Fig. 1, where the parameter $s$ enables tuning between Archimedean and non-Archimedean regimes. For $s < 0$ the system is approximately a one-dimensional chain (Fig. 4a (top)), whereas setting $s > 0$ theoretically produces the treelike geometry shown in Fig. 4a (bottom) (refs. 43, 44). Each leaf represents an array site, the position of which in the tree is determined by branching left or right at level $a$ if the $a$th bit of the site index $i$ is 0 or 1. Starting from the base of the tree, the first branching is governed by the least significant bit. Thus, the order of sites in the tree is rearranged from the physical order by the Monna map $\lambda(i)$, which reverses the order of bits in the site index $i$.

We confirm the transition from an Archimedean to a treelike geometry by measuring correlations $C^{\pm}$ for $s = 1$. We implement both models for $M = 16$ sites with periodic boundary conditions, using the drive waveforms in Fig. 4a (top), (bottom). For each value of $s$, we show $C^{\pm}$ as a function of physical positions $i, j$ (Fig. 4b) and as a function of positions $\lambda(i), \lambda(j)$ on the tree (Fig. 4c). Whereas for $s = -1$ the correlations decay smoothly as a function of physical distance, for $s = 1$ we observe
a non-monotonic dependence on physical distance due to the highly non-local structure of interactions. The Monna-mapped correlations for \( s = 1 \), however, are strongest near the diagonal—indicating a new sense of locality in the non-Archimedean geometry—and exhibit blocks consistent with the hierarchical structure of the tree.

To corroborate the realization of a non-Archimedean geometry, we plot the dependence of correlations on a treelike measure of distance in Fig. 4c. The natural metric for the treelike geometry is the 2-adic norm \( |a|_2 = 2^{-n} \), where \( n \) is the largest integer such that \( 2^n \) divides \( |a|_2 \). Intuitively, \( n \) represents the level of the tree—counting up from the base—at which the leaves representing sites 1 and connect. As a function of 2-adic distance, we observe a smooth decay of correlations.

A key feature of the tree graph is that only the vertices on the boundary represent physical sites, whereas the interior vertices constitute a holographic bulk geometry embodying the effective distance between sites. To investigate the validity of this holographic description, we perform a black-box reconstruction of the bulk geometry from spin correlations.

We begin by mapping the physical sites to effective coordinates in a Euclidean space, as before. Next, we draw bonds between pairs of maximally correlated sites and perform a coarse-graining procedure, treating each group of connected sites as a new larger site, and drawing new connections, until there is a path through the bulk between any two sites on the boundary.

The bulk reconstructions are shown in Fig. 4d for both the Archimedean (\( s = -1 \)) and non-Archimedean (\( s = 1 \)) cases. For \( s = -1 \), where interactions between physical neighbours dominate, the reconstruction produces only a one-dimensional loop. By contrast, for \( s = 1 \), a tree emerges from the reconstruction as a bulk geometry encapsulating the structure of spin correlations. This emergent geometry is analogous to the gravitational bulk in the \( p \)-adic AdS/CFT correspondence, in which the tree serves as a discretized version of hyperbolic space\(^{21,22}\).

The transition between two radically different geometries depending on the sign of the exponent \( s \) suggests that all sense of locality is lost as \( s \) approaches zero. To probe the breakdown of locality, we consider different possible bipartitions of the \( M = 16 \) sites into \( 8 \)-site subsystems \( I \) and \( J \) and examine correlations between the subsystems (Fig. 4e). Specifically, we plot a bipartite correlation \( C_{IJ} = \text{Corr}(F_I, F_J) \), where \( F_I = \sum_{i \in I} F_i \) denotes a coarse-grained spin, as a function of \( s \). For \( s < 0 \), the correlation \( C_{IJ} \) is smaller for a cut that is local according to the physical ordering of sites (blue circles) than for a cut that is local on the tree (green squares), whereas for \( s > 0 \) the situation is reversed, consistent with the change in effective geometry. Further plotting the minimum correlation \( C_{IJ} \) over all possible bipartitions (purple triangles) reveals a peak at \( s = 0 \), indicating the absence of any geometry providing a sense of locality.

The breakdown of locality at \( s = 0 \) paves the way for studies of fast scrambling\(^{1} \), the generation of system-wide entanglement at a conjectured maximal rate characteristic of black holes\(^{2} \). More broadly, our work provides a starting point for quantum simulations to investigate the conjecture that space–time geometry and gravity emerge from quantum mechanics at a fundamental level. The Monna-mapped correlations allow for the creation of quantum states that mimic many-body systems, offering a new perspective on the relationship between quantum mechanics and gravity.
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**Methods**

**Experimental sequence**
We begin by loading rubidium-87 atoms from a magneto-optical trap into an array of microtraps, in which we use optical pumping and adiabatic microwave sweeps to prepare the atoms in the \( F_l = 1, m_l = 0 \) state. We then transfer the atoms into a 1,560 nm optical lattice supported by the cavity, resulting in a set of \( M = 16 \) or 18 discrete ensembles. To generate programmable interactions between the ensembles, we apply a magnetic field gradient and drive the optical cavity along its axis with a modulated intensity. After the interaction time, we load the atoms back into the microtraps and use state-selective fluorescence imaging to measure the population in each Zeeman state. To measure the transverse magnetization, we apply a series of local spin rotations before the imaging sequence. Extended Data Figure 2a shows a schematic of the experimental sequence.

**Microtraps and lattice transfer**
Our experiments use a hybrid trapping scheme: whereas we perform cooling, internal state preparation and imaging in a microtrap array, we transfer the atoms to an intracavity optical lattice before inducing cavity-mediated interactions. The 1,560 nm intracavity lattice is in registry with the standing wave of 780 nm light used to drive interactions and thus maximizes the atom–light coupling. However, because the 1,560 nm light produces a strong and inhomogeneous a.c. Stark shift of the \( S^p_{1/2} \) state, we instead use the 808 nm microtrap array during the parts of the experimental sequence requiring near-resonant light, namely cooling, optical pumping and fluorescence imaging.

We initially turn on a two-dimensional array of \( M \times 2 \) optical microtraps at 808 nm during magneto-optical trap loading. The long axis of the array is aligned with the cavity axis, with 60 μm between traps. The two transverse arrays are designed to double the total trap volume and, correspondingly, the number of atoms loaded into the cavity for a fixed microtrap waist. Each microtrap has a waist of 6 μm and a depth of \( h \approx 4 \) MHz, where \( h \) is Planck’s constant. During the loading phase the transverse microtrap spacing is 100 μm. After loading the microtraps, the transverse spacing is reduced to 8 μm, so that both transverse arrays fit within the 25 μm waist of the intracavity lattice. We adiabatically transfer the atoms from the microtraps to the intracavity lattice, increasing the lattice power from an initial depth of \( h = 200 \) kHz to \( h = 3.5 \) MHz and then ramping off the microtraps. This preparation results in a one-dimensional array of \( M \) ensembles at a temperature of 100 μK, with each ensemble containing \( N = 10^4 \) atoms spread over 10 lattice sites.

For imaging, we transfer the atoms from the optical lattice back into the microtrap array by first switching on the \( M \times 2 \) microtraps before reducing the lattice depth to \( h = 200 \) kHz. Subsequently, we adiabatically move the microtraps away from the optical lattice by approximately 15 μm to avoid a.c. Stark shifts during imaging. The efficiency of this transfer from the lattice to microtraps is \( 80–90\% \). Additionally, the 808 nm trapping light causes 20% of the atoms to undergo state-changing scattering events before the end of the imaging sequence.

**Imaging and spin readout**
We detect the atoms in a sequence of four fluorescence images designed to independently measure the populations of all three Zeeman states within the \( F_l = 1 \) manifold and any residual atoms in \( F_l = 2 \). For each fluorescence image we apply a retro-reflected laser beam resonant with the microsecond transition of the D2 line for 100 μs and collect the resulting fluorescence signal on an electron multiplying charge-coupled device (EMCCD) camera. With the first imaging pulse, we measure the population in the \( F_l = 2 \) manifold, expelling these atoms from the microtraps by heating. For state-selective imaging of the \( F_l = 1 \) manifold, we sequentially apply three microwave sweeps that adiabatically transfer the atoms from each magnetic substate to \( F_l = 2 \) and perform fluorescence imaging after each sweep. A typical fluorescence signal of the atoms is shown in Extended Data Fig. 2b. For background subtraction we use a method from Xu et al.\(^{19}\) based on a principal component analysis of approximately 100 images without atoms. Technical noise in the measurement is approximately 1–2% of the total atom number.

To measure the transverse spin component \( F_{1z} \) we sequentially perform local spin rotations at each site \( i \) before the imaging sequence. For this purpose, we focus a circularly polarized laser onto each site by controlling the position of the beam with an acousto-optic deflector. By modulating the intensity of the laser at the local Larmor frequency, we induce a resonant Raman coupling between adjacent magnetic sublevels. We apply a 3 μs Raman pulse to produce a \( \pi/2 \) spin rotation. This locally maps \( F_{1z} \) onto the measurable population difference \( \Delta N_{\uparrow} - \Delta N_{\downarrow} \), illustrated in Extended Data Fig. 2c. Here, \( F_{1z} \) is defined in a rotating frame that depends on the local Larmor frequency at site \( i \). Shot-to-shot fluctuations in the Larmor frequency lead to a reduction in the measurable correlations between two sites, for which the reduction depends on the time between the corresponding Raman pulses. Thus, to suppress any bias in the measured correlations, we randomize the order of the local spin rotations in each experimental realization.

**Computation of correlations**
When visualizing the distance dependence of interactions, reconstructing effective geometries or probing bipartite correlations, we compute correlation functions \( C^{\text{mm}} \), \( C^{\text{mm}} \) and \( C_x \) from a minimum of 50 measurements (see Extended Data Fig. 3 and the Supplementary Information for the effects of finite statistics). Each correlation function is defined in the main text in terms of specified observables \( A \) and \( B \) as

\[
\text{Corr}(A, B) = \frac{\text{Cov}(A, B)}{\sqrt{\text{Var}(A)\text{Var}(B)}},
\]

where \( \text{Cov}(A, B) \equiv \langle AB \rangle - \langle A \rangle \langle B \rangle \) and \( \text{Var}(A) \equiv \text{Cov}(A, A) \). These correlations are normalized to the shot-to-shot variance, which provides the relevant spatial information while being agnostic to the total amount of pair creation. Effects of finite statistics on the measured correlations \( C^{\text{mm}} \) are examined in Extended Data Fig. 3.

To quantify pair creation dynamics, we measure in the \( \hat{x} \)-basis and normalize the covariance matrices to the population of atoms on each site rather than their variance,

\[
C_{\hat{x}i}^{\text{xx}} = \frac{\text{Cov}(F_{1i}^x, F_{m}^x)}{N^2} = \frac{(F_{1i}^x F_{m}^x)}{N}. 
\]

For this correlator, the measurement in the \( \hat{x} \)-basis provides a high sensitivity at early times and a large dynamic range for measurements over time. The normalization is chosen such that the extracted correlation is sensitive to the total amount of pair creation, allowing us to visualize the growth of correlations as a function of time.

**Interaction parameters**
To enable the programmable interactions, we apply a magnetic field gradient parameterized by the difference \( \omega_B \) in Zeeman splittings between adjacent array sites. This gradient is superposed on an overall bias field \( B_0 \) perpendicular to the cavity axis, which produces a Zeeman splitting of \( \omega_B/B_0 = 2 \pi \times 700 \text{ kHz G}^{-1} \) and a quadratic Zeeman shift of \( q/B_0^2 = 2 \pi \times 72 \text{ Hz G}^{-2} \). We work in a regime in which \( \omega_B/M > \omega_B > q \), that is, the variation in the magnetic field is small compared with the average field yet results in a Bloch oscillation frequency larger than the quadratic Zeeman shift. Specifically, we choose a magnetic field between 2 G and 4 G (as detailed in Extended Data Table I for each dataset) and a typical gradient \( \omega_B = 2 \pi \times 1.52(1) \text{ kHz per site} \), where paren-
theses denote one s.d. uncertainty on the last digit. For measurements of $C_{\text{rms}}$ in Figs. 1 and 2, we increase the ratio $\Omega_p/\Gamma$. This is accomplished either by increasing $\theta_p$ to $2\pi \times 12.47(2)$ kHz per site, or by reducing the effective quadratic Zeeman shift to $q = 2\pi \times 70$ Hz by applying an a.c. Stark shift to the $|L, 0\rangle$ state. To induce an a.c. Stark shift, we offset resonantly couple the states $|L, 0\rangle$ and $|2, 0\rangle$ via blue-detuned microwave radiation. The resulting energy shift is given by $\Delta_{\text{m}} = \Omega^2/4\delta_{\text{m}}$, where $\Omega_{\text{m}} = 2\pi \times 10$ kHz is the resonant Rabi frequency and $\delta_{\text{m}} = 2\pi \times 11.5$ kHz is the detuning.

We induce spin-exchange interactions among the atoms by applying a drive field detuned from cavity resonance. After accounting for a resonance shift due to the presence of atoms, we choose a typical drive-cavity detuning between $\delta_{\text{c}} = -2\pi \times 4$ MHz and $\delta_{\text{c}} = -2\pi \times 7$ MHz. The cavity mode itself has a large detuning of $\delta = -2\pi \times 11$ GHz from atomic resonance. The drive field is linearly polarized at an angle of $55^\circ$ with respect to the magnetic field, which is chosen to eliminate tensor light shifts. The instantaneous spin-exchange coupling is given by $J(t) = n_0 (\Omega^2/(2\delta))$, where $\Omega = 2\pi \times 13$ Hz is the vector a.c. Stark shift per circularly polarized photon in the cavity and $n_0(t)$ is the mean intracavity photon number due to control field driving the cavity. The average intracavity photon number over a full period of the drive waveform is typically $n_{\text{av}} = 2 \times 10^3$. This results in an average collective interaction strength of $2\sqrt{\Gamma} = 2\pi \times 0.6$ kHz between ensembles of $N = 10^4$ atoms.

To produce a set of couplings $J(r)$, we modulate the intensity of the drive field via an acousto-optic modulator as

$$J(t) = 2 \sum_{i,j} \{\cos(\omega_{\text{mod}} t + \phi_i) + 1\} / \sqrt{|\delta_{\text{mod}}|}$$

where we use the phases $\phi_i \in [0, \pi]$ to set the sign of the interactions. The coupling at $t = 0$ is given by $J(0) = 2 \sum_i \{J(0)\}$. To produce periodic boundary conditions in the system of $M$ sites, we additionally pulse the drive at a frequency of $M \omega_{\text{mod}}$. Each pulse has a duration of $0.3\tau_{\text{mod}} = 11$ ms. To keep the average interaction strength constant, we increase the drive strength during these pulses so that the number of drive photons in the cavity reaches a peak of about $n_{\text{av}} = 10^4$.

We choose the total duration of the interactions to be at least one Bloch period such that the dynamics are localized to single ensembles. In general, the Fourier limit for the resolution of the interactions (measured in sites) is given by the ratio $1/T$ of the Bloch period to the total interaction time. After a single Bloch period, interactions are thus localized to less than the spacing between sites, as shown in the inset to Fig. 2a for an interaction duration of $T = 2.5$ Bloch periods. A beneficial effect of Fourier broadening is to delocalize the interactions to more than the spatial extent of each ensemble. As the extent of each ensemble is approximately one-eighth of the distance between ensembles, we limit the duration of the drive pulse to $T < 8$ Bloch periods for all experiments.

Localization of interactions to specified distances also requires precise calibration of the magnetic field gradient such that the uncertainty in the target drive frequency $\omega_{\text{mod}}$ is less than the Fourier broadening of the drive field spectrum. In some measurements, the gradient can drift by up to 1.5%. This most strongly affects the resonance condition at longer distances, for example $r > 15$ in Fig. 2a. We model this effect in Supplementary Fig. 1.

Cavity parameters

The atoms are coupled to a near-concentric Fabry–Perot cavity with a length of 5 cm and an 18 μm waist at 780 nm. The cavity has vacuum Rabi frequency of $2\pi \times 21.6$ MHz and linewidth $\kappa = 2\pi \times 250(20)$ kHz, yielding a single-atom cooperativity $\eta = 4.5 \times 10^5$, where $\gamma = 2\pi \times 6.07$ MHz is the linewidth of the SP$_{1/2}$ state in rubidium. Our drive field is detuned by $\Delta = -2\pi \times 11$ GHz from the $|S|_{1/2}; F = 1\rangle \rightarrow |S|_{1/2}; F = 2\rangle$ transition, which produces a vector light shift per circularly polarized photon of $\Omega_p = -\frac{\gamma}{4\kappa} = 2\pi \times 26$ Hz on a maximally coupled atom at cavity centre. For an average atom this dispersive coupling is reduced to $\Delta = 2\pi \times 13$ Hz, primarily by thermal motion. This reduction is accounted for in our measurements of the collective cooperativity $\eta_c = 3 \times 10^4$. The Rayleigh range of the cavity is $z_R = 1.3$ mm, and each ensemble is within $0.4z_R = 520 \mu$m of the cavity centre. Displacement from the cavity centre contributes up to a 20% reduction in coupling for the most distant atoms.

The cavity cooperativity together with the detuning $\delta$, determines the relative strength of coherent interactions and two dissipation pathways: spontaneous emission into free space and collective emission into the cavity. Collective emission can be suppressed at fixed interaction strength by increasing the detuning $\delta$, from cavity resonance. This suppression is possible because the collective decay rate scales as $\Gamma_c \propto \delta_c^{-2}$, whereas the interaction strength for each mode scales as $\chi_c \propto \delta_c^{-1}$. However, decreasing collective decay comes at the cost of increasing the rate of free-space scattering $\Gamma_{\text{sc}}$, which is independent of $\delta_c$. Both interaction and decay via the cavity are enhanced with respect to free-space scattering by the collective cooperativity $\eta_c \gg 1$, which determines an optimal detuning for minimizing the combined rate of single-particle and collective decay. In our system, this optimal detuning is given by $\delta_{\text{opt}} / k = 1/\sqrt{\eta_c}$ (as derived in the Supplementary Information).

At the optimal detuning, the overall interaction-to-decay ratio depends on the collective cooperativity as $\eta_c^{1/2} (\Gamma_{\text{sc}} + \Gamma_c) \propto 1/\eta_c$. As a result, large collective cooperativity $\eta_c \gg 1$ enables entangled states to be generated. In the Supplementary Information, we analyse the role of the cooperativity in producing spin-nematic squeezed states, showing that the attainable metrological gain due to entanglement scales as $\eta_c$.

Our experiments operate at approximately twice the theoretical optimum detuning to reduce sensitivity to atom number fluctuations. These fluctuations change the frequency shift 4N MΩ of the cavity resonance and thus the detuning $\delta$, interaction strength $\chi_c$, and collective cooperativity $\eta_c$ within a single set of measurements, the total atom number varies by up to 5%, resulting in shot-to-shot fluctuations in the collective interaction strength of 20%. This is a small contribution to the variability of the pair creation process, which naturally produces fluctuations with standard deviations on the same order as the expected population in the states $m = \pm 1$ (ref. 3).

Working at a detuning that is larger than the optimum means that dissipation is primarily due to scattering to free space. For typical parameters, the scattering rate is 0.2 scattering events per atom per Bloch period, corresponding to a coherence time of 5 Bloch periods. The effect of this scattering is primarily evident in measurements of covariance and the structure factor. For example, at the latest times in Fig. 2b, c approximately 50% of the atoms have scattered at least once. This causes much of the reduction in the magnitude of the structure factor as compared to the idealized simulations using the truncated Wigner approximation seen in Extended Data Fig. 4. For the correlations that are shown in other figures, the normalization scheme chosen minimizes the impact of atom number and interaction strength fluctuations on the extracted spatial structure.

Interaction Hamiltonian

In equation (1), we describe the distance-dependent spin-exchange interactions by a static effective Hamiltonian $H_s$, with the spin on each site defined in a rotating frame set by the local magnetic field. Here we summarize the derivation of the effective Hamiltonian starting from the full time-dependent Hamiltonian $H_{\text{lab}}$ in the laboratory frame. The Hamiltonian $H_{\text{lab}}$ for the spin system, obtained by adiabatically eliminating the cavity mode $\omega_c$, is given by

$$H_{\text{lab}} = -J(t) \sum_{i,m} F_i F_m + \sum_i h_i F_i^+ + H_{\text{q}}$$

(8)
in terms of the collective spin $F_l = \sum_{\mu \neq \nu} f_{\mu \nu}$ on each site $l$, the local magnetic fields $h_l = \theta h$, and the quadratic Zeeman shift $H_q = q \sum_\mu \langle f_{\mu,\mu} \rangle^2$, in units where $\hbar = 1$. Moving to a rotating frame with $H_0^l = \sum_1^N h_l F_l^l$, the Hamiltonian becomes

$$H(t) = -\frac{1}{T} \sum_{l,m} e^{i(j - m)\omega q} F_{m}^l F_{m}^l + H_q$$

(9)

To obtain a form of the effective Hamiltonian that provides intuition about the spatial structure of the interactions, we consider the limit in which the collective interaction strength and quadratic Zeeman shift are weak compared with the gradient ($M$, $q \ll \omega q$). In this limit, the effective Hamiltonian is given to first order by the time average of equation (9). The interaction component of the resulting effective Hamiltonian $H_I + H_q$ is

$$H_I = -\frac{1}{T} \sum_{l,m} \langle f(1 - m) \rangle F_{m}^l F_{m}^l$$

(10)

where

$$J(r) = \frac{1}{T} \int_0^T dt e^{i\omega q r} J(t).$$

(11)

The dependence $J(r)$ of the couplings on distance is thus given by the Fourier transform of the drive waveform.

Even though the limits under which this effective Hamiltonian is derived, $M$, $q \ll \omega q$, are not strictly valid for the pulsed drive waveforms used to produce periodic boundary conditions, the intuition provided by this model is corroborated by the more generically valid model of the Floquet dynamics presented in the following section.

**Momentum-space dynamics**

To analytically compute the dynamics of the system, we write the Hamiltonian without approximation in terms of spin-wave operators

$$F_k^{\pm} = \frac{1}{\sqrt{2}} \sum_{l} e^{i(k \cdot r) \tau} k_l$$

as

$$H(t) = -M \langle f(t) \rangle F_{-\omega q}^l F_{\omega q}^l + H_q,$$

(12)

We can understand this Hamiltonian by recognizing that, in the laboratory frame, the magnetic field gradient causes spin waves to undergo Bloch oscillations at frequency $\omega q$. Only spin waves with momentum $k = 0$ in the laboratory frame couple to the cavity. In the rotating frame set by the gradient, the same physics can be visualized as spin waves remaining static over time, and the mode to which the cavity couples is given by $k = \omega t$. The quadratic Zeeman shift is left unchanged by the change of reference frames.

Because the system is finite and discrete, there are only $M$ orthogonal momentum modes. To obtain a discrete set of momentum-space couplings $J_k = -\lambda_k / 2N$, we drive interactions with a pulsed drive

$$J(t) = \sum_{k} x_k \delta(\omega q t - k)$$

that only takes on non-zero values $M$ times per Bloch period. We observe that the momentum modes decouple in the Hamiltonian,

$$H(t) = -\sum_k 2 (\delta_{k,0} - k) F_k^l F_k^l + H_q,$$

(13)

The evolution of any given momentum mode is discrete, with a short period of coupling to the optical cavity that induces spin–spin interactions, followed by a longer period of time when the state evolves only under the quadratic Zeeman shift. In the limit of a large collective interaction strength $| \lambda_k | > \omega q$, each momentum mode grows by a factor of

$$\Lambda = \left[ 2 \lambda_k \omega q \sin(\omega q t) \right]^{\frac{1}{2}},$$

after each Bloch period (Supplementary Information). This growth is reflected by the structure factor, with $F_k^l \propto \left| \chi_k \right|^2 \propto \left( N / k \right)^{\frac{1}{2}}$ after $T$ Bloch periods.

Even though our derivation of the growth of the structure factor assumes a pulsed drive field, which produces periodic boundary conditions, the same relation provides a good approximation in the case of a continuous drive field that produces open boundary conditions. In the latter case, we expect small deviations from the model because the cavity couples to a continuum of non-orthogonal momentum modes. We compare the continuous and pulsed cases in a numerical simulation presented in Extended Data Fig. 4.

A key feature of the evolution in momentum space is that the modes with minimum energy are maximally amplified in our system with $\chi_k < 0$. We can gain additional insight into this effect by considering the limit in which the dynamics are slow compared with the Bloch period and a time-averaged Hamiltonian is valid. In this case, the dynamics for each momentum mode are identical to the single-mode case that has been studied previously. The system is unstable to pair creation when the collective interaction strength $2 \delta = -4 M$ has a greater magnitude and opposite sign from the quadratic Zeeman shift $q$. This condition motivates our choice of ferromagnetic on-site interactions, such that $\chi_k < 0$, in our system with $q > 0$. The opposite signs of $\chi$ and $q$ allow the system to access low-energy states of the interaction Hamiltonian $H_l$ by transferring energy into $H_q$ via pair creation.

**Numerical modelling**

We numerically simulate the dynamics in our system using the semi-classical truncated Wigner approximation (TWA). To model quantum effects, which are essential for the pair-creation dynamics, the TWA simulation treats each ensemble of three-level atoms as three bosonic modes and samples the vacuum fluctuations of the initially unoccupied modes $m = \pm 1$. This corresponds to sampling the Wigner function of the initial state in phase space. Each sample is then propagated according to the classical equations of motion based on the Hamiltonian in equation (9). This enables us to incorporate experimental imperfections such as finite statistics (Extended Data Fig. 3) or finite system size (Extended Data Fig. 4) into the simulation, and to understand their effects on the measured data.

In the Supplementary Information we elaborate on the implementation of the TWA simulation and present additional simulations for comparison with our experimental data. In particular, we show effects of finite statistics for the reconstruction of the triangular ladder in Fig. 3c, and we examine the role of magnetic field fluctuations in the measurement of bipartite correlations in Fig. 4e.

**Euclidean reconstruction**

To reconstruct effective coordinates $\mathbf{r}$ and inferred couplings $J$ directly from measured correlations $C_{\mathbf{r}}$, we require an ansatz for the dependence of correlations on distance in the effective geometry. Here we apply our analytical model for the growth of the structure factor to derive the Gaussian ansatz for the decay of correlations. The dynamical evolution produces low energy states of the $XY$ Hamiltonian, which additionally allows us to relate the inverse correlation matrix and the inferred couplings.

To analytically motivate the Gaussian ansatz used for reconstructing effective geometries, we begin by relating the structure factor to the correlations we measure in the $x$ basis,

$$C_{\mathbf{r}}^{\text{xx}} \cdot \langle F_{\mathbf{r}}^{\text{xx}} F_{-\mathbf{r}}^{\text{xx}} \rangle = \frac{1}{M} \sum_{k_1, k_2} e^{i(\xi k_1 - \xi k_2)} \langle F_{\mathbf{r}}^{\text{xx}} F_{-\mathbf{r}}^{\text{xx}} \rangle = \frac{1}{M} \sum_{k} e^{i(k \mathbf{r})} \langle F_{\mathbf{r}}^{\text{xx}} F_{-\mathbf{r}}^{\text{xx}} \rangle.$$  

(14)

The final equality holds when the momentum modes are independent from one another, such that cross terms with $k_1 \neq k_2 \mathbf{r}$ to zero. This is true either when periodic boundary conditions are imposed or in the limit of an infinite system. Equation (14) enables the prediction of the form of spatial correlations from the dispersion relation $\chi_k$, which governs the growth of the structure factor.
As an illustrative example, we consider nearest-neighbour interactions created by the drive waveform \( \mathcal{J}(t) = (\cos \omega_d t + 1) \) corresponding to the dispersion relation

\[
\chi_r \propto (e^{ik/2} + e^{-ik/2})^2.
\]

(15)

As the correlations are the Fourier transform of the squared magnitude of the structure factor, we write an expansion of \( |\mathcal{F}_d|^2 \) in terms of powers of \( e^i \). Recalling \( |\mathcal{F}_d|^2 \propto |\chi_d|^2 \) after T Bloch periods of evolution, we compute

\[
|\mathcal{F}_d|^2 \propto (e^{ik/2} + e^{-ik/2})^2 = \sum_{d=2T}^{2T} 4T_{d+1} e^{-ikd}.
\]

(16)

The coefficients in this expansion are Fourier components corresponding to correlations at distance \( d \). Thus, we have \( C^{xx}(d) = (d + 2)^4 \). This binomial coefficient tends to a Gaussian function of distance \( d \) after several Bloch periods, analogously to a diffusion process.

More generally, a multifrequency drive leads to diffusion within the effective geometry set by the couplings. For a generic drive that produces a dispersion relation \( \chi_r \propto -\sum J(r)e^{i\omega r} \), correlations in position space are given by terms in the multinomial expansion of \( |\chi_d|^2 \). When \( J(r) > 0 \) this directly corresponds to a random walk within the effective geometry set by the couplings \( J(r) \). Motivated by the exact result for spreading in one dimension, we use a Gaussian ansatz for the correlation matrix to infer distances and hence the coordinates \( \mathbf{r} \) within the effective geometry.

We motivate the inferred coupling matrix \( J = (C^{xx})^{-1} \) by recalling that a population growth rate given by \( \chi_r \) generates a low energy state of the XY model, \( \mathcal{H}_r \). We approximate the final state as thermal, with large inverse temperature \( \beta \). We make use of the SO(2) symmetry to note that \( (F^T F) \) and \( (F^T F) \) are equivalent. Now, to compute \( (F^T F) \), we integrate over phase space, with a Boltzmann weighting \( \exp(-\beta\mathcal{H}) \). To constrain the overall spin length, we introduce a chemical potential \( \mu \), so that

\[
C^{XX} \approx \prod_i \int dF_i^T GF_i^T \exp(-\beta H_i - \mu F_i^2).
\]

(17)

The chemical potential can be incorporated into a modified coupling matrix \( J' = J - \mu \hat{\mathcal{G}}_J \). Evaluating the integral yields \( C^{xx} \propto J'^{-1} \). For the purposes of the reconstruction in Fig. 3, where we colour bonds between sites according to \( J' \), only the off-diagonal terms of \( J' \) are relevant.

The inverse correlation matrix, also known as the concentration or precision matrix, can also be interpreted as the partial correlation matrix, up to normalization. For a given set of variables \( x_i \), the partial correlation between \( x_i \) and \( x_j \) is the correlation after regressing out every \( x_{kl \neq ij} \). In a system with interactions at distance \( r \), sites spaced by \( r \) have a non-zero partial correlation, but sites at distances that are multiples of \( r \) have zero partial correlation, because the interactions between the sites at distance \( r \) mediate all the variance. Thus, the interpretation of the inverse correlation matrix as an inferred coupling matrix \( J = (C^{xx})^{-1} \) is well motivated even at early times, when correlations are still spreading across the system.

**Data availability**

All data displayed in Figs. 1–4 and Extended Data Figs. 1–4 are available from the corresponding author upon reasonable request.

**Code availability**

All code used for simulation and analysis is available from the corresponding author upon reasonable request.

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**Author contributions**

A.P., E.S.C., P.K. and M.S.-S. wrote the manuscript. All authors contributed to the discussion and interpretation of results.

**Competing interests**

The authors declare no competing interests.

**Additional information**

**Supplementary information**

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**Extended Data Fig. 1 | Coupling graphs.** Sketch of couplings $J(i-j)$ for the model in equation (4) with local interactions ($s = -1$, left) or treelike interactions ($s = 1$, right). The strengths of the interactions are indicated by the thickness and transparency of the red lines. For $s = 1$, reordering the sites according to the Monna map makes the couplings more local, corroborating the treelike geometry.
Extended Data Fig. 2 | Experimental sequence and imaging. a Schematic of experimental sequence for measurements of $F_i^x$. After driving the cavity to induce interactions, we apply spin rotations sequentially to the $M$ sites of the array and subsequently perform state-sensitive readout via fluorescence imaging. b Fluorescence images after spin rotation, showing the signal for the $F = 2$ manifold and the three magnetic substates for the case of interactions at distance $r = 3$ with periodic boundary conditions. c Transverse magnetization $F_i^x$ and structure factor $F_k^x$ extracted from the image in b.
Extended Data Fig. 3 | Effect of finite statistics. Left, correlation plot reproduced from Fig. 1, showing $C_{ij}^m$ obtained from 50 realizations of the experiment with interactions at distance $r = 10$. Right, simulation results obtained from a truncated Wigner approximation, where we either choose the same number of realizations as in the experiment or increase the number of realizations by a factor of 10 to reduce statistical uncertainty. The simulations indicate that residual correlations in the experimental data are mainly due to the finite sample size.
Extended Data Fig. 4 | Comparison between measured structure factor and simulation results. The left graph shows the measured structure factor after $T = 3$ Bloch periods of evolution, which is also shown in Fig. 2c. The two plots at right show results of a truncated Wigner simulation with and without periodic boundary conditions. For the simulated data we used 100 realizations of the TWA simulation, which is four times higher than the number of experimental realizations to reduce statistical fluctuations. For open boundary conditions, we find that the simulation has an offset with respect to the theoretical prediction (blue line). We attribute this offset to the finite system size, as the model is exact only for an infinite system or a system with periodic boundary conditions. Repeating the same simulation with a pulsed drive shown on the right shows that in this case the TWA simulation is consistent with the analytical model. The error bars indicate the standard error of the mean.
### Extended Data Table 1 | Experimental parameters

| Data set     | Magnetic field $B_0$ [G] | Gradient $\omega_B$ [kHz/site] | Quadratic Zeeman shift $q$ [Hz] | Interaction time $T\tau_B$ [ms] |
|--------------|-------------------------|--------------------------------|-------------------------------|-------------------------------|
| Fig. 1b all-to-all | 2.8                     | 0                              | $2\pi \times 580$             | 0.1                           |
| Fig. 1b localized    | 3.8                     | $2\pi \times 12.46$           | $2\pi \times 1100$           | 0.2                           |
| Fig. 1b distance     | 3.8                     | $2\pi \times 12.46$           | $2\pi \times 1100$           | 0.2                           |
| Fig. 2a             | 3.8                     | $2\pi \times 12.46$           | $2\pi \times 1100$           | 0.2                           |
| Fig. 2bc            | 2.0                     | $2\pi \times 1.53$            | $2\pi \times 290$            | up to 1.97                    |
| Fig. 2d             | 2.0                     | $2\pi \times 1.52$            | $2\pi \times 70$             | 3.95                          |
| Fig. 3              | 2.0                     | $2\pi \times 1.52$            | $2\pi \times 290$            | 1.32                          |
| Fig. 4              | 2.0                     | $2\pi \times 1.52$            | $2\pi \times 290$            | 1.32                          |

Magnetic offset field $B_0$, Bloch oscillation frequency $\omega_B = 2\pi/\tau_B$, effective quadratic Zeeman shift $q$, and interaction time $T\tau_B$ for each of the data sets presented in Figs. 1–4. The effective quadratic Zeeman shift is given by $q = B_0^2 \times 72\text{Hz}/G^2 + q_{\text{mw}}$, where $q_{\text{mw}}$ is the ac Stark shift of the state |1, 0⟩ induced via off-resonant microwave coupling to |2, 0⟩ (see Methods).