Spin Squeezing by Rydberg Dressing in an Array of Atomic Ensembles:
Supplemental Material

In this supplement, we provide additional information about our experimental methods and theoretical models. In Sec. I, we describe the experimental setup, calibrations, and data analysis methods. In Sec. II, we derive theoretical models for the interactions induced by Rydberg dressing and the resulting spin squeezing, and we apply these models to examine limits to squeezing imposed by the finite interaction range. We also present supporting information regarding the contaminant atoms that require us to perform the Rydberg dressing stroboscopically.

I. EXPERIMENTAL DETAILS

A. Projection noise calibration

An accurate calibration of the atom number \( N \) is crucial to quantifying spin squeezing. We calibrate the number of atoms by measuring the quantum projection noise of coherent spin states with variable initial tilt \( \theta \). These data generically allow for distinguishing quantum projection noise from any technical noise sources via the dependence on \( \theta \). Fitting the projection noise with the known variance of the binomial distribution then provides a precise calibration of the atom number. Specifically, for a coherent state prepared at a polar angle \( \theta \) on the Bloch sphere, the probability that an atom is measured in state \( | ↑ \rangle \) is \( p = \cos^2(\theta/2) \). Accordingly, the fraction \( f_\uparrow \) of atoms measured in \( | ↑ \rangle \) follows a binomial distribution with mean \( p \) and variance \( \sigma^2_{f_\uparrow} = p(1-p)/N \). The variance of the fractional difference \( f_\uparrow - f_\downarrow \) in populations of the two spin states, where \( f_\downarrow = 1 - f_\uparrow \), is then \( V[f_\uparrow - f_\downarrow] = 4p(1-p)/N \).

We prepare different initial states \( | \theta \rangle \) by applying a resonant microwave drive pulse of varying length. We perform 100 measurements of \( f_\uparrow \) at each initial state to determine the expectation value \( E[f_\uparrow] \) and variance \( V[f_\uparrow - f_\downarrow] \). The results of a typical projection noise calibration are shown in Fig. S1. We post-process our measurements of \( f_\uparrow \) by performing a linear spatial regression across microtraps. This removes correlated noise that arises as a result of common-mode experimental errors, such as microwave power fluctuations. The average amounts of technical noise removed by the spatial regression across microtraps, normalized to the maximum variance, are on the 10% level. We fit the remaining noise with the functional form

\[
V[f_\uparrow - f_\downarrow] = \frac{4p(1-p)}{N} + ap^2 + \left( \frac{b}{N} \right)^2 \tag{S1}
\]

with \( N \), \( a \), and \( b \) as free parameters. The term \( ap^2 \) accounts for atom-related technical noise during imaging: a small fraction of the atoms decay into \( |F = 3\rangle \) as they are imaged in \( |F = 4\rangle \), and \( a \) quantifies any fluctuations in this fraction. The constant term \( b \) accounts for detection noise from our EMCCD camera. For the data plotted in Fig. S1(a), we obtain mean values of \( N = 220(6) \) atoms, \( a = 0(2) \times 10^{-5} \), and \( b = 2.3(1) \). The fit value for \( b \) represents an atom number resolution equivalent to 3% of the projection noise variance, while the remaining technical noise is consistent with zero.

![Projection noise calibration](image)

FIG. S1. **Projection noise calibration.** (a) Variance of \( f_\uparrow - f_\downarrow \) as a function of \( E[f_\uparrow] \) for microtrap 5 (purple dots), and fit to functional form of Eq. S1 (purple line). (b) Summary of the fitted atom number \( N \) for all nine microtraps. (c) Summary of the integrated fluorescence signal, measured in EMCCD camera counts, per atom.
We calibrate the contrast $C$ used to calculate the Wineland squeezing parameter by Ramsey interferometry, as shown in Fig. S2(a). This calibration is performed both with and without applying the dressing light [Fig. S2(b)]. In both cases, we fix the Ramsey time to 4.8 ms and include spin-echo pulses, thereby calibrating the contrast under the same conditions as in the squeezing measurement. We attribute the non-unity baseline contrast to inhomogeneous trap light shifts that are imperfectly canceled by spin echo due to atomic motion. This calibration is carried out both before and after the squeezing measurement, with results for the contrast consistent within error.

In calibrating the contrast with dressing pulses, we make a small correction to the fringe amplitude obtained from the fit to account for atom loss from the microtraps. We determine the average loss directly from the atomic signal in the data used for squeezing by comparing measurements with and without dressing light taken in alternate shots of the experiment. The resulting ratio $N/N_0$ of atom number in each microtrap with and without dressing determines the loss $L = 1 - N/N_0$ plotted in Fig. S2(c). We then multiply the amplitude of the Ramsey fringe measured after dressing by $N/N_0$ to obtain the contrast $C$ including loss, which we apply in calculating the squeezing parameter. The data shown in Fig. S2(b) yield a loss-adjusted contrast $C = 0.95(2)$ with the dressing light and a baseline contrast $C_0 = 0.96(2)$ in the absence of dressing light.

C. Atomic state preparation and detection

After initial cooling in a magneto-optical trap and optical molasses, cesium atoms are loaded into a 1064 nm optical dipole trap with a $\sim 50$ $\mu$m waist and trap depth $h = 5(1)$ MHz. The atoms are then transported to a science chamber by scanning the focus of an electrically tunable lens (Optotune). A bright molasses stage loads the atoms into a one-dimensional array of 1064 nm microtraps, which is generated by a crossed acousto-optic deflector (AOD) system. This array consists of a set of nine microtraps spaced 25 $\mu$m apart, each with a waist of 6 $\mu$m. The dipole trap is briefly turned off to allow any atoms not confined in microtraps to escape, and is then turned back on to provide additional confinement along the microtraps’ axial dimension. After state preparation, the measured temperature of the atoms in each trap is $T = 22(1) \mu$K. The resulting rms cloud sizes are $\sigma_{(x,y,z)} = [1.7(2), 1.7(2), 19(2)] \mu$m, where $\hat{z}$ denotes the vertical direction in the lab and $\hat{x}$ denotes the axis of the array. With an average of $N = 200$ atoms per trap, the peak atomic density is $\rho_0 = 2.3(3) \times 10^{11}$ $\text{cm}^{-3}$.

The science chamber contains an array of eight stainless steel electrodes with independently tunable voltages. We calibrate these voltages to minimize the electric field along three orthogonal axes by measuring the quadratic Stark shift of the $|60P_{3/2}\rangle$ Rydberg state. These calibrated voltages are fixed for the duration of the experimental sequence.

We perform state-sensitive fluorescence imaging on the $D_2$ line using 852 nm light. Prior to imaging, we change the spacing of the microtraps from 25 $\mu$m to 50 $\mu$m apart by ramping AOD frequencies, phases, and amplitudes with
a jerk-minimizing polynomial ramp over 5–6 ms, such that fractional fluorescence from one microtrap detected in an adjacent trap is at most 1%. During imaging, we first use light tuned to the $|F = 4\rangle \rightarrow |F' = 5\rangle$ transition to image only the $|F = 4\rangle$ atoms. After this, we reapply the same pulse to resonantly expel any remaining $|F = 4\rangle$ atoms. We then apply a microwave $\pi$ pulse to transfer atoms from $|F = 3, m_F = 0\rangle \rightarrow |F = 4, m_F = 0\rangle$ and perform resonant fluorescence imaging once again. Typically, around 5% of all atoms are in $|F = 3, m_F \neq 0\rangle$ states due to imperfect optical pumping; these atoms do not contribute to the experiment, as they are not affected by microwave pulses or Rydberg dressing light and do not contribute to detected fluorescence during the imaging sequence. We measure a 3% decrease in our imaging efficiency for $|F = 3\rangle$ atoms relative to $|F = 4\rangle$. We attribute this to an imperfect $\pi$ pulse during $|F = 3\rangle$ readout and calibrate our fluorescence signals to account for this effect.

### D. Microwave parameters, sequences, and calibration

We measure an average microwave Rabi frequency $\Omega_{\text{MW}} = 2\pi \times 18.9(2)$ kHz, with percent-level inhomogeneity in $\Omega_{\text{MW}}$ across microtraps. One small but important source of calibration error and drift is the differential ac Stark shift from the trapping light, which shifts the clock state resonance by approximately 500 Hz. We initially calibrate our microtrap light intensities such that the ac Stark shift of the clock transition is the same to within 5 Hz across all microtraps. During a typical measurement, the center of the dipole trap (which provides confinement in the axial dimension of each microtrap) can drift by $\pm 3$ μm, which results in a measured microwave resonance drift of approximately $\pm 25$ Hz, a non-negligible shift compared to the $\sim 200$ Hz frequency scale set by typical Ramsey sequence times.

We implement spin echoes primarily to cancel the average ac Stark shift from the Rydberg dressing light, as discussed in Sec. 1E, but secondarily to (a) cancel any remaining systematic drifts that result in microwave calibration error, and (b) mitigate sensitivity to the percent-level gradient in microwave Rabi frequency. Borrowing spin-refocusing techniques from NMR, we use the sequence MLEV-4 [1], in which every “unit” of 4 spin echoes takes the form of rotations around (X, X, -X, -X), and where each pulse is a single $\pi$ pulse rather than a composite pulse. This sequence provides the best Ramsey contrast for our typical dressing Ramsey sequence times of $\sim 4.8$ ms. The microwave pulse train consists of 3 MLEV-4 “units” for a total of 12 $\pi$-pulses, one every $\sim 400$ μs, and is used in every sequence in this paper that requires spin echoes.

### E. Rydberg dressing laser system, parameters, and pulse shaping

To generate the 319 nm Rydberg dressing light, we start with 1275 nm light from an external-cavity diode laser (LEOS Solutions) that is used to seed a pre-amplifier (Thorlabs BOA1130P) followed by a Raman fiber amplifier (RFA, MPB Communications). The seed light is frequency-stabilized by a Pound-Drever-Hall lock to a ULE reference cavity (Stable Laser Systems). Light from the RFA is resonantly doubled in two stages (LEOS Solutions), each consisting of an LBO crystal in a bow-tie optical cavity. We use an electro-optic polarization modulator (QUBIG GmbH) and an acousto-optic modulator (AOM) for generating and shaping the Rydberg dressing pulses. The dressing light is then focused down to a waist of 55 μm and intersects the 1D array of microtraps at a 30° angle of incidence [Fig. 1(a)], such that the beam addresses most of the microtraps but with a spatially varying intensity that allows us to explore different Rabi frequencies within a single measurement.

We dress with the $60P_{3/2}$ Rydberg state, which has an attractive van der Waals interaction characterized by the coefficient $|C_6| = 2\pi \times 359$ GHz μm$^6$. We use $\sigma^+$-polarized light to couple from the ground state $|6S_{1/2}, F = 4, m_F = 0\rangle$ to the $|60P_{3/2}, J = 3/2, m_J = 3/2\rangle$ and $|60P_{3/2}, J = 3/2, m_J = 1/2\rangle$ states. At our typical power of 300 mW and beam waist of 55 μm, we measure Rabi frequencies $\Omega/(2\pi) = 1.3$–1.5 MHz. As the sign of $C_6$ is negative, we work at a positive (blue) detuning $\Delta_s \approx 2\pi \times 8$ MHz from the $|6S_{1/2}, F = 4, m_F = 0\rangle \rightarrow |60P_{3/2}\rangle$ resonance to avoid crossing the pair-state energy of two Rydberg atoms at any length scale. In prior work we operated at $n = 43$ due to an enhanced $C_6$ coefficient from a nearby Förster resonance [3], but we have since found that this near-Förster-resonant dressing causes increased loss compared to $n = 60$ under similar experimental conditions.

We shape the Rydberg dressing pulses by applying a 2.5 MHz low-pass filter (Mini-Circuits BLP-1.9+) to a train of square pulses produced by an arbitrary waveform generator, which is then used to modulate the RF power driving the AOM. These shaped pulses allow us to avoid $\sim 2$ MHz sidebands produced by square pulses of approximately 500 ns in length; the red-detuned 3rd and 4th-orders of these unwanted sidebands address the atoms at a frequency at or very near resonance compared to our chosen +8 MHz detuning, leading to measurable excess loss. We characterize the pulse shape by the intensity profile $h(t)$ of our light as measured on a photodiode, normalized to a peak value.
When $\hbar_{\text{max}} = 1$, the corresponding Rabi frequency is given in terms of the peak Rabi frequency $\Omega_p$ as

$$\Omega(t) \equiv \Omega_p \sqrt{\hbar(t)}.$$  \hfill (S2)

For terms of order $k > 0$ in $\Omega^2$, such as the $k = 1$ and $k = 2$ terms in the expansion for $U$ found in Eq. S11, we define a corrected effective interaction time

$$T_k \equiv \int_{-\infty}^{\infty} \hbar^k(t) dt$$ \hfill (S3)

for a single shaped pulse. We additionally define $T_0 = T_p$, representing the interaction time for an idealized square pulse; for pulses typically used in our twisting and squeezing sequences, we find $T_1/T_0 \approx 0.95$ and $T_2/T_0 \approx 0.83$. We account for these corrections in all fits that involve $\Omega$.

## F. Twisting and dressed pulse length calibrations

Once we determine a dressing pulse delay $\tau_d = 100$ $\mu$s and fix our spin echo pulse separation at 400 $\mu$s [Fig. S2(a)], we still have the freedom to choose a pulse length $\tau_p$ and total interaction time $\tau_{\text{int}}$. Fixing the number of pulses $M = 48$ and measuring the one axis twisting-like phase precession $\phi$ of states initialized at $\theta = 3\pi/4$ and $\pi/2$ on the Bloch sphere, we calculate the twisting strength $Q$ via $\phi = -Q \cos \theta$, as in Ref. [3]. In Fig. S3(a), $Q$ is plotted as a function of the total dressed interaction time $\tau_{\text{int}} = M\tau_p = 48\tau_p$. A linear fit gives a mean-field interaction strength of $\chi = 2\pi \times 6.4(8)$ kHz for these data. We generally choose $\tau_p$ such that one or more microtraps has a twisting strength $Q \geq 1$ rad, but also such that our contrast does not significantly decay.

After scanning $\tau_p$ at a fixed pulse number and thus a varying total interaction time, we may also ask whether the contrast decay we observe for larger $\tau_p$ vanishes at fixed interaction time. In Fig. S3(b) we plot $Q$ versus contrast (normalized to the contrast with no dressing light) for three different pulse lengths $\tau_p = (630$ $\text{ns}, 940$ $\text{ns}, 1.9$ $\mu$s) and corresponding pulse numbers $M = (48, 32, 16)$, chosen such that the interaction time $\tau_{\text{int}} = M\tau_p$ remains constant. The pulse delay remained fixed at $\tau_d = 100$ $\mu$s, and as a result the number of total spin echoes (12, 8, 4) and Ramsey times (4.8, 3.2, 1.6) $\text{ms}$ were allowed to vary between datasets. Here we observe that longer dressing pulses lead...
**FIG. S4. Dicke ladders for a fully-blockaded system.** The dressed ac Stark shift and its suppression due to Rydberg blockade may be modeled as two coupled Dicke ladders in the Hilbert space of states \(|N_{\uparrow}, N_{\downarrow}, N_r\rangle\). The left ladder represents states with no Rydberg excitations \((N_r = 0)\), while the right ladder considers those with \(N_r = 1\). States with equal population in \(N_{\downarrow}\) are coupled by the collective Rabi frequency \(\sqrt{N_\uparrow}\Omega\), which represents the coherent drive of a single excitation in \(|r\rangle\) shared between the \(N_\uparrow\) atoms in \(|\uparrow\rangle\).

to faster contrast decay at comparable values of \(Q\); fitting the normalized contrast to a Gaussian decay of the form 
\[ \exp\left(-\frac{Q^2}{2\sigma_Q^2}\right) \]

yields \(\sigma_Q = [4.1(8), 2.1(4), 1.2(1)]\) rad, respectively. For the shortest pulses, the value \(\sigma_Q = 4.1(8)\) rad is consistent with the prediction of the one-axis twisting Hamiltonian with \(N_c = 16(6)\) atoms within the interaction range (Sec. II C 1).

To accompany every noise measurement shown in this Letter, we took independent measurements of the quantities \(Q\), dressing Rabi frequency \(\Omega_p\), and atom number \(N\) for each microtrap. Plotting \(Q\) as a function of \(N \Omega_p^4 \tau_{\text{int}}/(16\Delta^3)\) [Fig. S3(c)] allows us to observe an approximately linear relationship, with the slope describing the fraction of interacting neighbors within each trap, \(N_c/N\). While we cannot continuously measure \(Q\) and take other data simultaneously, we calibrate \(Q\) both before and after every noise measurement.

## II. THEORY

### A. Collective interactions via Rydberg dressing

Here we derive an analytical expression for the ac Stark shift \(U\) induced by the Rydberg dressing light and the resulting collective interaction strength \(\chi\). Consider a cloud of \(N\) Rydberg-dressed atoms in the fully blockaded regime, where \(|C_0\rangle r_i^6 \gg \Delta, \sqrt{N}\Omega\) for all pairwise distances \(r_i\) in the cloud. The system will be approximately described by a permutation-symmetric Hamiltonian of the general form
\[
H_{\text{eff}} = \sum_{p=0}^{\infty} c_p S_z^p
\]

where \(S_z = (N_{\uparrow} - N_{\downarrow})/2\). From this Hamiltonian we can calculate the light shift
\[
U \equiv \frac{\partial H_{\text{eff}}}{\partial S_z},
\]

which represents the ac Stark shift due to the dressing light, and the interaction strength
\[
\chi \equiv -Nc_2 = -\frac{N}{2} \left. \frac{\partial^2 H_{\text{eff}}}{\partial S_z^2} \right|_{S_z=0},
\]

which parametrizes one-axis twisting. To arrive at \(H_{\text{eff}}\), we diagonalize the full Hamiltonian in the Hilbert space of states \(|N_{\uparrow}, N_{\downarrow}, N_r\rangle\), where we fix the total number of atoms \(N = N_{\uparrow} + N_{\downarrow} + N_r\) and allow only \(N_r = 0\) or \(N_r = 1\) excitations to \(|r\rangle\). Schematically, we have two coupled Dicke ladders, as shown in Fig. S4. This system is described by a block-diagonal Hamiltonian with blocks
\[
H(N_{\uparrow}) = \begin{pmatrix}
0 & \sqrt{N_{\uparrow}\Omega/2} \\
\sqrt{N_{\uparrow}\Omega/2} & -\Delta
\end{pmatrix}
\]
whose eigenvalues are

$$E_{\pm} = -\frac{\Delta}{2} \left[ 1 \pm \sqrt{1 + N_\uparrow (\Omega/\Delta)^2} \right]. \tag{S8}$$

Using the constraint $S_z = N_\uparrow - N/2$, we calculate the light shift

$$U = \frac{\partial E_+}{\partial N_\uparrow} = \frac{\Omega^2}{4\Delta} \frac{1}{\sqrt{1 + N_\uparrow (\Omega/\Delta)^2}} \tag{S9}$$

and the interaction strength

$$\chi = -\frac{N}{2} \frac{\partial^2 E_+}{\partial N^2} \bigg|_{N_\uparrow=N/2} = N \frac{\Omega^4}{16\Delta^3} \frac{1}{\left[ 1 + (N/2) (\Omega/\Delta)^2 \right]^{3/2}}. \tag{S10}$$

These results are consistent with the perturbative treatment of Refs. [4, 5] in the weak dressing limit $N (\Omega/2\Delta)^2 \ll 1$. Here we can approximate the light shift as

$$U \approx \frac{\Omega^2}{4\Delta} \left[ 1 - \frac{N_\uparrow}{2} \left( \frac{\Omega}{\Delta} \right)^2 \right], \tag{S11}$$

leading to a collective interaction strength

$$\chi \approx N_\uparrow \frac{\Omega^4}{16\Delta^3}. \tag{S12}$$

The derivation of the dressing light shift and collective interaction strength is generalizable to a system of number density $\rho$ extending over more than one blockade radius $r_c = (C_6/2\sqrt{\Omega^2 + \Delta^2})^{1/6}$ by replacing $N_\uparrow \rightarrow N_\uparrow^{(c)}$, where $N_\uparrow^{(c)}$ is the number of (interacting) atoms within a single blockade radius:

$$N_c = \rho \cdot \frac{4\pi}{3} r_c^3 = \rho \cdot \frac{4\pi}{3} \left( \frac{C_6}{2\sqrt{\Omega^2 + \Delta^2}} \right)^{1/2}. \tag{S13}$$

For atoms prepared in $|\theta\rangle = \cos(\theta/2) |\uparrow\rangle + \sin(\theta/2) |\downarrow\rangle$ we have $N_c^{\uparrow} = N_c \cos^2(\theta/2)$. In fitting the dependence of the light shift on detuning to extract $N_c$ in Fig. 3 of the main text, we must account for the fact that $N_c$ itself depends on detuning. We define the constants $W \equiv \sqrt{C_6 \cdot 4\pi \rho/3}$ and $W_\uparrow = W \cos^2(\theta/2)$ such that

$$N_c^{\uparrow} = \frac{W_\uparrow}{\sqrt{2(\Omega^2 + \Delta^2)}^{1/4}}. \tag{S14}$$

This allows us to rewrite Eq. S9 as

$$U = \frac{\Omega^2}{4\Delta} \left[ 1 + \frac{\Omega^2}{\Delta^2} \frac{W_\uparrow}{\sqrt{2}(\Omega^2 + \Delta^2)^{1/4}} \right]^{-1/2}, \tag{S15}$$

which we use to fit the data shown in Fig. 3(b) with $\Omega$ and $W_\uparrow$ as free parameters. We account for the pulse shape, as discussed in Sec. I E, when stating values for $\Omega_p$, $N_\uparrow^{\uparrow}$, and $N_c$ in Fig. 3(c-d).

### B. Rydberg-dressed ground state interaction potential and interaction strength distribution

To compare the measured interaction parameters ($\chi$, $N_c$) with theoretical predictions, we calculate the dressed ground-state interaction potential using Rydberg pair potentials [6], Rabi frequency $\Omega = 2\pi \times 1.305$ MHz, and detuning $\Delta = 2\pi \times 8$ MHz, as described in Ref. [3]. The interaction potential is shown in Fig. S5(a-b), where $\theta = 0$ is referenced to the propagation direction of the Rydberg dressing beam, which is orthogonal to the axial (z) direction of the traps as defined in Sec. 1C. From this interaction potential, we generate a matrix of pairwise interactions $J_{ij}$ by simulating the spatial distribution of the atoms in a single microtrap, sampling atom positions from Gaussian
distributions with experimentally calibrated standard deviations $\sigma^{(x,y,z)} = [1.7(2), 1.7(2), 19(2)]$ µm (Sec. I C). An atom $j$ then experiences a collective interaction strength

$$\chi_j = -\frac{1}{2} \sum_i J_{ij}. \quad (S16)$$

Figure S5(c) shows a histogram of the resulting values $\chi$, normalized to the depth of the interaction potential $|J_0| = 2\pi \times 2.1$ kHz, for a cloud of $N = 200$ atoms. The average value $\langle \chi \rangle / |J_0| = 19(1)$ is approximately consistent with the values $N_c$ determined from the suppression of the ac Stark shift in Fig. 3 of the main text.

C. Squeezing dynamics

In Fig. 4 of the main text we compare the measured squeezing with a model of one-axis twisting with total spin $S = N_c/2$. Here we give expressions for this model, including the effect of finite baseline contrast $C_0$. We also compare the model of one-axis twisting with $N_c$ neighbors to an exact calculation of the Ising dynamics with pairwise interactions given by the Rydberg-dressed potential, and to an opposite limit of ideal one-axis twisting with global interactions among all $N$ atoms.

1. Spin squeezing by one-axis twisting

Analytic expressions for the squeezing arising from the one-axis twisting Hamiltonian were derived by Kitagawa and Ueda in Ref. [7]. Briefly, the approach is to calculate the time evolution of the variance $\langle S_y^2 \rangle$ and covariance $\langle S_y S_z + S_z S_y \rangle$ under the Hamiltonian $H_{\text{twist}} = -\chi S_z^2 / N$ for a system initialized in a coherent spin state along $\hat{x}$. From these quantities, as well as $\langle S_z^2 \rangle = S/2$ which remains invariant, one obtains the spin variance

$$\langle S_y^2 \rangle = \langle S_z^2 \rangle \cos^2 \alpha + \langle S_y^2 \rangle \sin^2 \alpha + \langle S_y S_z + S_z S_y \rangle \sin \alpha \cos \alpha \quad (S17)$$

as a function of rotation angle $\alpha$. By symmetry $\langle S_y \rangle = \langle S_z \rangle = 0$, so extremizing Eq. S17 with respect to $\alpha$ yields the minimum and maximum variances

$$\langle S_y^2 \rangle_{\text{min/\max}} = \frac{V_+ \pm \sqrt{W^2 + V^2}}{2}, \quad (S18)$$

where

$$W = \langle S_y S_z + S_z S_y \rangle, \quad (S19a)$$

$$V_\pm = \langle S_y^2 \pm S_z^2 \rangle. \quad (S19b)$$

The orientation $\alpha_{\text{opt}}$ of the squeezed quadrature is obtained from the minimization of $\langle S_y^2 \rangle$ as

$$\alpha_{\text{opt}} = -\frac{1}{2} \arctan \left( \frac{W}{\langle S_y^2 \rangle - \langle S_z^2 \rangle} \right). \quad (S20)$$
The variances in Eq. S18, together with the length $|\langle S \rangle| = \langle S_x \rangle$ of the Bloch vector, determine the squeezing parameters

$$\xi_{\text{min/max}}^2 = \frac{N(S_{\text{min/max}}^2)}{|\langle S_x \rangle|^2}$$  \hspace{1cm} (S21)

for the squeezed and antisqueezed quadratures. For the data presented in Fig. 4 we fit the measured $\xi^2$ versus $C$ with the sinusoidal function

$$\xi^2(C) = (\xi_{\text{max}}^2 - \xi_{\text{min}}^2) \sin^2 (C - C_{\text{opt}}) + \xi_{\text{min}}^2$$  \hspace{1cm} (S22)

where $C_{\text{opt}}$, $\xi_{\text{min}}^2$, and $\xi_{\text{max}}^2$ are free parameters.

In modeling the squeezing in our experiment we account for imperfect initial contrast $C_0$. We assume that the imperfect contrast is due to uncorrelated phase shifts $\phi_j$, with $\langle \phi \rangle = 0$, applied to the atoms indexed $j$. Correspondingly, the spin raising operator for the $j^{th}$ atom in the Heisenberg picture is modified as

$$s_j^+ \rightarrow s_j^+ e^{i\phi_j},$$  \hspace{1cm} (S23)

leading to a reduced length of the collective Bloch vector

$$\langle S_x \rangle \rightarrow C_0 \langle S_x \rangle,$$  \hspace{1cm} (S24)

where $C_0 = \langle \cos \phi \rangle$. In addition to shortening the Bloch vector, the imperfect contrast reduces the correlations $W = \langle S_y S_z + S_z S_y \rangle$ responsible for squeezing and modifies the variance $\langle S_y^2 \rangle$ as

$$W \rightarrow C_0 W$$  \hspace{1cm} (S25a)

$$\langle S_y^2 \rangle \rightarrow C_0^2 \langle S_y^2 \rangle + (1 - C_0^2) \frac{S}{2}$$  \hspace{1cm} (S25b)

The resulting analytic expressions for one-axis twisting dynamics, building on Ref. [7], are given in terms of the twisting strength $Q = \int \chi(t) \, dt$:

$$\langle S_x \rangle = C_0 S \cos^{2S-1} \left( \frac{Q}{2S} \right),$$  \hspace{1cm} (S26a)

$$\langle S_y^2 \rangle = \frac{S}{2} + C_0^2 S (S - 1/2) \left[ 1 - \cos^{2S-2} \left( \frac{Q}{2S} \right) \right],$$  \hspace{1cm} (S26b)

$$\langle S_y S_z + S_z S_y \rangle = C_0 S (2S - 1) \sin \left( \frac{Q}{2S} \right) \cos^{2S-2} \left( \frac{Q}{2S} \right).$$  \hspace{1cm} (S26c)

We apply Eqs. S26(a-c) to model the squeezing and antisqueezing in the main text.

In addition, we compare Eq. S26a to the measured dependence of contrast on twisting strength in Sec. IF of the supplement. In the experimentally relevant regime $Q \ll S$, the length of the collective spin vector decays as a Gaussian function of the twisting strength: $\langle S_x \rangle = S e^{-Q^2/(2\sigma_Q^2)}$, where $\sigma_Q = 2S/\sqrt{2S-1}$. Assuming an effective spin length $S = N_c/2$ set by the number of neighbors in the interaction ellipsoid, a Gaussian fit to the contrast decay reveals $N_c$ via

$$N_c = \sigma_Q^2 \left( 1 + \frac{1}{\sqrt{1 - 4/\sigma_Q^2}} \right).$$  \hspace{1cm} (S27)

2. Effects of finite interaction range

While all-to-all coupled systems preserve a permutation symmetry that conserves the total system spin $S$, no such symmetry exists for systems with finite-range interactions, and we thus expect the squeezing $\xi_{\text{min}}^2$ ultimately to be limited by a decrease in signal $|\langle S \rangle|$ with increasing interaction time. This limitation is particularly fundamental for the case of Ising interactions: due to the absence of non-commuting terms in the Hamiltonian, correlations form only between pairs of atoms that interact directly and thus, in the idealized limit where the atomic positions are pinned, the correlations cannot spread beyond the interaction range. An additional limitation arising from the locality of interactions is that the interaction strength is sensitive to the local density.
FIG. S6. Effects of finite interaction range and inhomogeneous density. Simulated squeezing dynamics for an inhomogeneous cloud matching our experimental trap geometry and Rydberg interaction parameters from Fig. S5. Solid blue line denotes average value over 1000 instances, while blue shading denotes ± 1σ values. The olive curves and shading show analytical predictions for the range of expected squeezing for all-to-all coupled systems with \( N_c = 13 \) atoms (light olive dotted, our empirical value) to \( N_c = 19 \) atoms (dark olive dash-dotted, based on the estimate \( N_c = \langle \chi \rangle / |J_0| \) from the simulated distribution of interaction strengths in Fig. S5). We additionally show the analytical all-to-all prediction for \( N = 63 \) atoms (red dash-dot), which would be achieved by extending the system to be larger than the interaction range in all three dimensions at fixed density \( \rho = 2 \times 10^{11} \text{ cm}^{-3} \), as well as simulated results for \( N = 200 \) (yellow dashed) and \( N = 1000 \) (orange dashed) on a 3D lattice with the same density. The purple dash-dotted curve shows the analytical all-to-all prediction for our full system size of \( N = 200 \) atoms.

To analyze the limits to squeezing with the local Ising interactions produced by Rydberg dressing, we perform a full simulation of the squeezing dynamics in a Gaussian cloud of atoms. Because the Ising interactions contain no non-commuting terms, one can efficiently compute the time evolution for any specified set of pairwise interactions \( J_{ij} \) [8]. Following the example of Ref. [8], we compute correlation functions of the form \( \langle s_i^x s_j^z \rangle \) between pairs of individual spins \((i, j)\). From these pairwise correlations we calculate moments of collective observables, and thus the squeezing parameter, following Eqs. S17-S21.

The results of the simulation are shown in Fig. S6. Each instance of the simulation samples \( N = 200 \) atoms from a three-dimensional Gaussian distribution with standard deviations \( \sigma(x, y, z) = (1.7, 1.7, 19) \text{ μm} \) as measured in our experiment, corresponding to a peak density \( \rho_0 = 2.3 \times 10^{11} \text{ cm}^{-3} \). For simplicity, the atoms are assumed to be at rest during the simulation. From the atomic coordinates and the calculated Rydberg-dressed potential shown in Sec. II B, we generate a pairwise interaction matrix and calculate the corresponding local mean-field interaction strengths \( \chi \) (Fig. S5). We then calculate the average twisting strength \( Q = \langle \chi \rangle \tau_{int} \) as a function of the interaction time \( \tau_{int} \). In Fig. S6, we plot the squeezing parameter \( \xi_{min}^2 \) averaged over 1000 instances as a function of \( Q \) in solid blue, with shading at ±1σ.

To examine the impact of the finite interaction range and the spatial inhomogeneity on the squeezing, we compare the full simulation of the short-range interactions with models of ideal one-axis twisting. The olive curves and shading in Fig. S6 account for the finite interaction range by showing analytical predictions for all-to-all coupled one-axis twisting with total spin \( S = N_c / 2 \), where we set either (1) \( N_c = \langle \chi \rangle / J_0 = 19 \) atoms (dark olive dash-dot), the average number of neighbors predicted for our atomic cloud based on the calculated interaction potential; or (2) \( N_c = 13 \) atoms (light olive dotted), the average extracted value across microtraps from the fits in Fig. 3. The purple dash-dotted curve compares these to an all-to-all coupled system with \( N = 200 \) atoms, the number typically found within a microtrap. All simulations obey the limiting early-time behavior \( \xi_{min}^2 \approx 1 - Q \). At the twisting strengths \( Q < 0.7 \text{ rad} \) accessed in Fig. 4 of the main text, the three models examined here are indistinguishable within experimental error, showing that the short-range character of the interactions is not the limiting factor in the present experiment.

For stronger twisting, both the finite number of neighbors and the inhomogeneous twisting strength pose limitations to the squeezing. At fixed density and dressing parameters, the number of neighbors could be increased by modifying the microtrap geometry. In our present experiment with rms cloud sizes \( \sigma(x, y, z) = (1.7, 1.7, 19) \text{ μm} \) (Sec. I C) and interaction radii \( \rho_c(x,y,z) = (3, 5, 5) \text{ μm} \) (Sec. II B), the cloud is larger than the interaction radius in only one dimension. Increasing the radial cloud sizes at fixed density \( \rho = 2 \times 10^{11} \text{ cm}^{-3} \) would produce \( N_c = 63 \) neighbors within the interaction ellipsoid. In Fig. S6, we plot the analytical prediction (red dash-dotted curve) for an all-to-all coupled system with \( N = 63 \) which achieves a minimum squeezing parameter \( \xi_{min}^2 = 0.09 \) (equivalently 10.5 dB).
To confirm that the one-axis twisting model provides a good approximation of the attainable squeezing in a realistic system with uniform density, we additionally simulate the full Ising dynamics on a 3-dimensional lattice of spacing $\rho^{-1/3} = 1.7 \, \mu m$ with unity filling within a spherical region containing $N = 200$ (yellow dashed curve) or $N = 1000$ (orange dashed curve) atoms. These simulations are consistent with the squeezing approaching 10 dB in an ordered array of dimensions larger than the interaction range.

3. Comparison with alternative platforms

The predicted squeezing attainable by Rydberg dressing in a large system at uniform density, at the level of 10 dB, is less than the record spin squeezing attained in optical cavities [9, 10] but comparable to the largest enhancements in spectroscopic sensitivity observed to date. For reference, a clock stability 10.5 dB beyond the standard quantum limit has been demonstrated by Ramsey spectroscopy using input states squeezed by 20 dB via cavity-based quantum non-demolition measurement [9]. A larger spectroscopic enhancement of 11.8 dB has been accessed by an echo protocol employing cavity-mediated interactions both to generate a non-Gaussian entangled state and facilitate its detection [11]. This protocol, which achieves a Heisenberg scaling in metrological gain through one-axis twisting, could also be applied in future experiments with Rydberg-dressed atoms to access a larger benefit than from squeezing alone [12, 13].

Compared with alternative methods of generating entanglement, Rydberg dressing is particularly well suited to applications that benefit from independent control of squeezing in multiple ensembles. Prior work in this context has leveraged collisional interactions in BECs to access 3.4 dB of metrological squeezing across an array of up to 30 ensembles and a 24% enhancement in magnetic field gradiometry [14]. Rydberg dressing offers promise for achieving larger metrological gain, with the added benefit of optical control for switching off interactions during the operation of a sensor or clock.

D. Contaminant Rydberg states

In Fig. 2 of the main text, we show that introducing a delay between Rydberg dressing pulses significantly reduces super-Poissonian loss that we attribute to the presence of atoms in contaminant Rydberg states. Here we identify the dominant contaminant states based on their branching ratios, lifetimes, and range of influence within the system of Rydberg-dressed atoms. We also explore how the creation of contaminant atoms and their influence on the larger system depend on system size and dimensionality.

1. Decay channels and lifetimes

We calculate decay rates from $|r\rangle = |60P_{3/2}, m_J = 3/2\rangle$ to nearby S and D states, as well as $C_3$ coefficients and lifetimes for these contaminant states, using the Alkali Rydberg Calculator (ARC) [6]. We focus our attention on states with $C_3$ coefficients sufficiently large to shift atoms in our ensembles into resonance with the dressing beam. We estimate a threshold $C_3$ coefficient from our dressing detuning ($\Delta_\ast = 2\pi \times 8 \, MHz$) and typical interatomic distance of $r_i = 1.8 \, \mu m$ as

$$C_3 \geq \Delta_\ast r_i^3 = 2\pi \times 47 \, MHz \, \mu m^3.$$  \hfill (S28)

Figure S7(a) shows the branching ratio into a given contaminant state plotted against the characteristic distance, $d = (C_3/\Delta_\ast)^{1/3}$, at which the interaction shift equals the detuning of the dressing light. The branching ratio is calculated by taking the product of the decay rate into a given state from $|r\rangle$ and the lifetime of $|r\rangle$. For the states shown here, the lifetimes of the S and D states are on the order of 75 $\mu s$ and 100 $\mu s$, respectively. The lifetime of $|r\rangle$ is 150 $\mu s$.

When an atom transitions into a contaminant Rydberg state, a natural question to ask is whether the lifetime of the resulting excitation is set by the radiative lifetime of the state, or is instead set by the antitrapping experienced by the Rydberg atom in the presence of 1064 nm light. To estimate the antitrapping lifetime, we neglect the small thermal velocity for atoms at 22 $\mu K$ and assume atoms start at rest with an initial spatial distribution corresponding to our measured cloud $\sigma_{\text{rms}}$ values from Sec. I.C. We calculate the force an atom would feel were it to decay to $|60S_{1/2}\rangle$ at $t = 0$, an illustrative example contaminant state which has both a high branching ratio from $|60P_{3/2}, m_J = 3/2\rangle$ and a large $C_3$ coefficient. We then calculate the time required for the atom to be forced outside of the $2\sigma_{\text{rms}}$ cloud radii in any direction. The results of 2000 independent simulation instances are shown in Fig. S7(b). We find that the mean...
thermal timescale for trap expulsion is 115 µs, which is comparable to the radiative lifetimes of the contaminant states. We conclude that both effects must be contributing to the effective lifetimes of any contaminant state excitations in the cloud.

2. Effects of system size and dimensionality

The stroboscopic dressing technique introduced in this work is of particular importance for systems with large atom number $N$. To understand why, we consider how system size affects the typical interaction time needed to create a contaminant atom in a simple model that assumes adiabatic dressing pulses but allows for an enhanced excitation rate due to laser phase noise. The rate $\gamma_{\text{exc}}$ of incoherent excitation from the ground state $|\uparrow\rangle$ to the Rydberg state $|r\rangle$ is given by

$$\gamma_{\text{exc}} = \frac{(\gamma_L + \Gamma_r)\Omega^2}{(\gamma_L + \Gamma_r)^2 + 4\Delta^2},$$

where $\Gamma_r$ is the linewidth of the Rydberg state $|r\rangle$ with which we are dressing and the $\gamma_L$ is the linewidth of the dressing laser, presumed to be Lorentzian. Atoms also undergo two-photon scattering directly from $|\uparrow\rangle$ into contaminant Rydberg states at a rate $\left(\Omega/2\Delta\right)^2\gamma_{\text{BB}}$, where $\gamma_{\text{BB}}$ is the blackbody decay rate from $|r\rangle$. However, this rate is insufficient to explain the loss in our experiments, suggesting that the dominant pathway is excitation to $|r\rangle$ at a rate $\gamma_{\text{exc}}$ that is governed by laser noise, followed by decay to contaminant states at rate $\gamma_{\text{BB}}$. For sufficiently short interaction times $\tau_{\text{int}}$ we may take the total number of atoms $N^\uparrow$ in the state $|\uparrow\rangle$ to be constant, and the expected number of excitations to $|r\rangle$ is then $N_r = \gamma_{\text{exc}}\tau_{\text{int}}N^\uparrow$. For a decay rate $\gamma_{\text{BB}}$ from $|r\rangle$ to a contaminant Rydberg state $|c\rangle$, the expected time $T_c$ to produce a single atom in $|c\rangle$ is then

$$T_c = \frac{2}{\gamma_{\text{exc}}\gamma_{\text{BB}}N^\uparrow}.$$

For the 60P$_{3/2}$ Rydberg state, the room-temperature blackbody transition rate is $\gamma_{\text{BB}} \approx 2\pi \times 800$ Hz and the linewidth is $\Gamma_r \approx 2\pi \times 1$ kHz. For a representative laser linewidth $\gamma_L = 2\pi \times 10$ kHz, Rabi frequency $\Omega = 2\pi \times 1.2$ MHz, detuning $\Delta = 2\pi \times 8$ MHz, and system with $N^\uparrow = 100$ atoms in $|\uparrow\rangle$, the continuous interaction time expected to produce a single contaminant atom is $T_c \approx 100$ µs. Note that $T_c$ decreases with larger system sizes as $T_c \propto 1/\sqrt{N^\uparrow}$, a scaling that highlights the particular importance of stroboscopic dressing techniques in larger systems.

We now consider how the influence of a contaminant atom on neighboring ground-state atoms depends on system size and dimensionality. As shown in Fig. S7(a), the characteristic distance $d = (C_3/\Delta)^{1/3}$ over which a contaminant...
atom strongly affects ground-state atoms is \( d \approx 10 \mu m \). We define \( N_d \) to be the number of atoms within a distance \( d \) of a contaminant atom. For simplicity, we consider the worst-case condition that all \( N_d \) atoms are lost from the system, corresponding to a super-Poissonian loss process in which atoms are lost in groups of size \( g = 1 + N_d \). For a small fractional atom loss \( \ell \), this process adds an amount \( g\ell \) of noise to the normalized variance \( \sigma^2 \), as observed in Fig. 2(d).

If a system of \( N \) atoms is small compared to \( d \) in all dimensions, a single contaminant atom interacts with all ground-state atoms and we have \( N_d = N^\uparrow = N/2 \). This is approximately the case in our present work, where \( d \) for relevant contaminant states is larger than the radius \( \sigma_x, \sigma_y \) of each cloud and comparable to the length \( \sigma_z \). For systems of density \( \rho \) that are larger than \( d \) in \( D \) dimensions, however, the number of ground-state atoms affected by a contaminant atom becomes

\[
N_d \propto \rho \times \left( \frac{C_3}{\Delta} \right)^{D/3}
\]

This adverse scaling with density \( \rho \) and dimensionality \( D \) corroborates the myriad of experimental results that have observed avalanche decay when implementing Rydberg-dressed interactions in large systems [15–18].