We investigate the possibility of a bistable phase in an open many-body system. To this end, we discuss the microscopic dynamics of a continuously off-resonantly driven Rydberg lattice gas in the regime of strong decoherence. Our experimental results reveal a prolongation of the temporal correlations exceeding the lifetime of a single Rydberg excitation and show strong evidence for the formation of finite-sized Rydberg excitation clusters in the steady state. We simulate the dynamics of the system using a simplified and a full many-body rate-equation model. The results are compatible with the formation of metastable states associated with a bimodal counting distribution as well as dynamic hysteresis. However, a scaling analysis reveals that the correlation times remain finite for all relevant system parameters, which suggests the formation of many small Rydberg clusters and finite correlation lengths of Rydberg excitations. These results constitute strong evidence against the presence of a global bistable phase previously suggested to exist in this system.

I. INTRODUCTION

Recent progress in experimental control offers unique possibilities to study the interplay between pure quantum mechanical systems and their coupling to reservoirs. In particular, the competition between external drive and dissipation can result in interesting phases in the steady state [1–4]. One fascinating avenue is to study the properties of these phases and the possibility of phase transitions in the context of an open quantum system [1,5–8]. In addition, including strong and nonlocal interactions may allow us to prepare and study strongly correlated many-body systems [9–11].

Rydberg atoms are particularly interesting as they provide strong and quasi-long-range interactions in an inherently dissipative environment. By means of an optical drive, ground-state atoms can be excited to states with high-principal quantum numbers. These Rydberg states have a large polarizability, resulting in a strongly interacting quantum gas. Therefore, these systems are an ideal platform to study nonequilibrium many-body physics with strong dissipation [2,12–17].

Depending on the driving scheme to the excited Rydberg state and the geometry of the atomic ensemble, Rydberg gases can exhibit very different phases. In the case of the so-called blockade regime, i.e., for resonant driving, the strong interaction shift of Rydberg states induced by an already-present Rydberg excitation suppresses further excitations in a mesoscopic ensemble of surrounding atoms. This leads to the concept of a Rydberg superatom [18–22] and was initially proposed to realize quantum gates [23]. Here, interesting phases with antiferromagnetic, crystalline long-range order have been predicted [2], and phases with finite-length spatial correlations have been observed [24]. In the case of the so-called antiblockade regime, i.e., for off-resonant driving, the interaction shift may be compensated by detuning the frequency of the excitation laser from resonance. Now, instead of suppressing excitations within a certain volume, further Rydberg excitations are facilitated at a specific distance from an already-excited atom. This leads to the formation of Rydberg excitation clusters [15,25,26].

One interesting aspect observed in recent antiblockade experiments using a full counting analysis is a bimodal excitation number distribution [27,28] and a hysteresis in the excitation dynamics [13,17]. It has been under debate whether this indicates a transition to a bistable steady state of the open many-body system [8,9,13,16,29,30]. We will show that both hysteresis and bimodal counting statistics are features of metastable states.

A metastable state manifests itself in a separation of time scales, where the relaxation time $T$ is much larger than all other time scales in the system [6,31]. This may lead to a hysteresis behavior upon temporal changes of system parameters on a time scale $\tau$ (sweep time) that is small compared to $T$. When the sweep time becomes much larger than $T$, the hysteresis disappears. Associated with a finite value of $T$ is an upper limit for all spatial correlation lengths, $\xi \sim T$. We argue that metastable systems with
correlation length similar to the system size allow for a bimodal distribution. If $T$ increases with the system size $L$ and eventually diverges in the thermodynamic limit, the metastable states become degenerate, truly stationary states of the system. In this case, the steady state is called bistable (multistable). As will be shown later, bistability in an infinite, translationally invariant system implies long-range spatial order. Thus, different from metastability, the emergence of bistability can be associated with a phase transition to an ordered state.

In this paper, we experimentally investigate and theoretically model the excitation dynamics of a Rydberg lattice gas with off-resonant continuous driving and strong dissipation [see Fig. 1(a)]. We continuously monitor the laser-induced excitation and subsequent ionization of Rydberg atoms and analyze its temporal correlations [Fig. 1(b)] in the antiblockade regime, scanning the frequency and intensity of the excitation laser. Although there is a constant loss of atoms, our measurement technique allows us to access steady-state properties. The results indicate the formation of small clusters with a long but finite lifetime, which originate from a correlated excitation cascade. We compare the experimental results to numerical simulations based on a simplified single-cluster model [Fig. 1(c)] and a more advanced rate-equation model of extended systems.

To check for the existence of bistability, we perform a finite-size extrapolation in the numerical simulations. Our results indicate that the correlation time and length remain finite in the experimentally relevant regime amenable to a rate-equation description. The bimodal character of the full counting distribution [28] disappears when the observed system size exceeds the correlation length, while a dynamic hysteresis can still be observed, provided the sweep time is sufficiently small [32]. This suggests that the Rydberg aggregate in the antiblockade regime consists of many small independent clusters and is incompatible with a global bistable regime.

II. BISTABLE AND METASTABLE STEADY STATES

Before discussing Rydberg lattice gases, let us start with a general introduction to bistability and metastability in open quantum systems. Quantum optical systems coupled to external reservoirs and subject to external drives can often be described in terms of a Lindblad master equation for the many-body density matrix $\rho$ [33–35],

$$\frac{d}{dt}\rho = \mathcal{L}\rho. \quad (1)$$

The Lindbladian superoperator $\mathcal{L}$ is the generator of the dynamics, determined by the Hamiltonian $H$ describing the unitary evolution and the jump operators $L_\mu$ describing the Markovian reservoir couplings,

$$\mathcal{L}(\rho) = -\frac{i}{\hbar}[H,\rho] + \sum_\mu \Gamma_\mu \left( 2(L_\mu^\dagger L_\mu - L_\mu L_\mu^\dagger) - \rho L_\mu L_\mu^\dagger + L_\mu^\dagger L_\mu \rho \right). \quad (2)$$

Note that $\mathcal{L}$ has complex eigenvalues $\lambda_k$ and corresponding right eigenvectors $\rho_k$, i.e., $\mathcal{L}\rho_k = -\lambda_k \rho_k$. The non-negative real parts of the eigenvalues, $\text{Re}[\lambda_k] \geq 0$, determine the characteristic relaxation rates towards a stationary state $\rho_0$ or a manifold of stationary states $\{\rho_0^{(n)}\}$ with $\lambda_0 = 0$. It should be noted that all “excited” eigenvectors $\rho_k$ with $\lambda_k \neq 0$ are not density matrices since $\text{Tr}\{\rho_k\} = 0$. If the
steady state is unique, it is called stable; if two orthogonal stationary solutions, \( \rho_0^{(1)} \) and \( \rho_0^{(2)} \), exist, the steady state is called bistable. In the first case, any initial state will eventually evolve into the unique steady state; in the second case, the asymptotic state for large times is a mixture of the two solutions with coefficients determined by the initial state \( \rho(t = 0) \):

\[
\rho_{ss} = p \rho_0^{(1)} + (1 - p) \rho_0^{(2)}, \quad p = \text{Tr}\{\tilde{\rho}_0^{(1)} \rho(t = 0)\}. \tag{3}
\]

Here, \( \rho_0^{(n)} \) is the left eigenvector of \( \mathcal{L} \) corresponding to \( \rho_0^{(n)} \) and \( \text{Tr}\{\tilde{\rho}_0^{(n)} \rho_0^{(m)}\} = \delta_{nm} \).

In the cases we are interested in here, both the Hamiltonian \( H \) and the jump operators \( L_\mu \) are local; i.e., they can be written as a sum of terms that act only locally. In such a case, the steady state is generically unique if the system is finite, unless the reservoir interactions are fine-tuned. General conditions for the uniqueness of steady states can be found, e.g., in Refs. [36–38].

Since experiments can be performed only in finite time, it is interesting to ask if states exist that, while not being true stationary states, appear to be stable over long but finite time scales. These states are called metastable, and they can be written as a sum of terms that act only locally. Without loss of generality, we assume \( L_\mu = 0 \) and the systems are finite-dimensional. General conditions for the uniqueness of steady states can be found, e.g., in Refs. [36–38].

![Figure 2](image)

**FIG. 2.** Generic spectrum of relaxation (damping) rates as a function of system size \( L \) for (a) a metastable system and (b) a truly bistable system.

Configurations. However, each of these two configurations corresponds to a metastable state, which, as shown in Ref. [6], is an admixture of the true steady state \( \rho_0 \) and the first excited eigenvector \( \rho_1 \). The metastable states eventually relax to the true steady state on a time scale

\[
T \sim L^{a=0}.
\]

Approaching the thermodynamic limit, there is a critical slow down, and the system becomes truly bistable.

The transition from a unique to a bistable steady state is a true phase transition, which can be characterized by an order parameter. Bistability and metastability cannot be distinguished, however, in finite systems, and one has to investigate the scaling behavior of the characteristic time scales for \( L \to \infty \). We now show that bistability is always associated with long-range order. To this end, we consider a translationally invariant system and assume that two orthogonal steady states, \( \rho_0^{(1)} \) and \( \rho_0^{(2)} \), exist. Then, the asymptotic state at large times is given by Eq. (3). In order to detect bistability, there must be a local observable \( \hat{x}_j \) that distinguishes the two states. Here, the index \( j \) labels positions (or some compact part of space) that are acted on by \( \hat{x}_j \). Without loss of generality, we assume

\[
\langle \hat{x}_j \rangle_1 = \text{Tr}\{\rho_0^{(1)} \hat{x}_j\} = +1\quad \text{and} \quad \langle \hat{x}_j \rangle_2 = \text{Tr}\{\rho_0^{(2)} \hat{x}_j\} = -1.
\]

Making use of translational invariance, we find, in the asymptotic state \( \rho_{ss} \), Eq. (3):

\[
\langle \hat{x}_j \rangle_1 = 2p - 1.
\]

Let us now consider \( \hat{Y} = \sum_{j=1}^L (\hat{x}_j - \langle \hat{x}_j \rangle_1) \sum_{k=1}^L (\hat{x}_k - \langle \hat{x}_k \rangle_2) \). Obviously, \( \langle \hat{Y} \rangle_1 \geq 0 \) in any state. In both steady states, \( \rho_0^{(\mu)} \), one finds

\[
\sum_{j,k} \langle \hat{x}_j \hat{x}_k \rangle_\mu \geq \sum_{j,k} \langle \hat{x}_j \rangle_\mu \langle \hat{x}_k \rangle_\mu = L^2.
\]

For the general asymptotic state \( \rho_{ss} \), this yields

\[
\sum_{j,k} \langle \hat{x}_j \hat{x}_k \rangle_{ss} = p \sum_{j,k} \langle \hat{x}_j \hat{x}_k \rangle_1 + (1 - p) \sum_{j,k} \langle \hat{x}_j \hat{x}_k \rangle_2 \geq L^2.
\]

Now, considering correlations of the local observable \( \langle \hat{x}_j \hat{x}_k \rangle = \langle \hat{x}_j \rangle_1 \langle \hat{x}_k \rangle_2 \), which, because of translational invariance, only depend on \( d = j - k \), one finds

\[
\sum_d \langle \hat{x}_0 \hat{x}_d \rangle_{ss} \geq L(1 - (2p - 1)^2).
\]

Unless \( p = 0 \) or 1, this inequality can only be fulfilled in the thermodynamic limit \( L \to \infty \) if

\[
\langle \hat{x}_0 \hat{x}_d \rangle_{ss} \xrightarrow{d \to \infty} c \neq 0.
\]

Thus, bistability is only possible if there is long-range order. This result is very intuitive. If only two (or some
finite number of) stationary states exist, which differ in the expectation value of some local observable, there must be long-range correlations in the system since one end of the system has to “know” in which state the other end is. For any system that is bistable in the thermodynamic limit, we thus expect a scaling of the correlation length

$$\xi \sim L^{1/2}.$$  

Since the speed at which correlations can spread in the system is finite, the transition into a bistable state is associated with a critical slow down, i.e., with characteristic time scales that diverge with the system size. If the correlation length $\xi$ is finite, the number of stationary configurations is $2^N$, with $N \sim \xi/L$. For $L \gg \xi$, the central limit theorem holds, and global probability distributions become single-peaked Gaussians. In the following, we argue that under realistic experimental conditions, correlation length and characteristic time scales in off-resonantly driven Rydberg lattice gases remain finite, and there is no bistability in this system.

### III. EXPERIMENT

Experimentally, we continuously probe the Rydberg excitation dynamics in the antiblockade regime. To this end, we prepare $N_{\text{at}} = 20,000$ $^8$Rb atoms in a 3D optical lattice in the Mott insulating phase at unit filling. The lattice constants in the $x$ and $y$ directions are $a_{x,y} = 374$ nm, and in the $z$ direction, $a_z = 529$ nm. We couple ground-state atoms in the $5S_{1/2}$ state continuously via a one-photon transition at a wavelength of $\lambda = 297$ nm to the excited Rydberg state $25P_{1/2}$ with Rabi frequency $\Omega$ and blue detuning $\Delta > 0$. See Fig. 1(a) for a sketch of the experiment. The spontaneous decay rate from the Rydberg state is given by $\Gamma_{\text{sp}} = \tau_{\text{sp}}^{-1} = 50$ kHz, and we estimate a bare decoherence rate of $\gamma_0 \approx 300$ kHz, which primarily originates from the laser line width.

The dipole trap lasers provide a weak photoionization rate $\Gamma_{\text{ion}} \approx 2$ kHz, which we use to observe the excitation dynamics in the system. The ions are guided towards a detector by a small electric field of 90 mV cm$^{-1}$. The detector efficiency is 40%. The retrieved ion signal with rate $R_{\text{ion}}(t) = \Gamma_{\text{ion}} N_{\text{Ry}}(t)$, corrected by the detector efficiency, serves as a weak probe for the full excitation number $N_{\text{Ry}}(t)$. We experimentally determine the Rabi frequency by analyzing the temporal statistics of the first detected ion. A more detailed discussion of the experimental setup and preparation can be found in Ref. [39] and in Appendix A. In the inset of Fig. 1(b), the measured ion rate $R_{\text{ion}}(t)$ for a Rabi frequency of $\Omega = 2\pi \times 77$ kHz and a detuning of $\Delta = 2\pi \times 13$ MHz is shown. The recorded signal extends over several tens of ms, which is 3 orders of magnitude larger than the effective natural lifetime $\tau_{\text{sp}} = 20$ $\mu$s (reduced by blackbody transitions into neighboring states) of the $25P_{1/2}$ state. Therefore, in contrast to previous studies [27,28], our measurements allow us to study the steady-state of the system [40].

The experimentally accessible quantities, which we use to characterize the formation and the dynamics of Rydberg clusters, are the average ion rate and the time-dependent two-particle correlation function

$$\langle R_{\text{ion}}(t) \rangle \text{ and } g^{(2)}(\tau) = \frac{\langle R_{\text{ion}}(t+\tau)R_{\text{ion}}(t) \rangle}{\langle R_{\text{ion}}(t+\tau) \rangle \langle R_{\text{ion}}(t) \rangle},$$

respectively. In a single experimental realization [see inset of Fig. 1(b)], strongly bunched excitations are already visible, indicating strong excitation number fluctuations. To quantify them, we extract $g^{(2)}(\tau)$ of the measured ion signals [see Fig. 1(b)]. For typical parameters, we find a value of $\langle g^{(2)}(0) \rangle = 1$. Moreover, we find an exponential decay of the correlation signal with time scale $\tau(\tau)$ much larger than the lifetime of the $25P_{1/2}$ Rydberg state. Crucially, the strong excitation bunching reflects two different time scales arising in the antiblockade regime: While a first seed excitation is strongly suppressed [indications thereof can be seen in the inset of Fig. 1(b) as large intervals of zero signal in a single experimental run], the excitation rate for a second atom can be enhanced in the presence of the first [25,26,41]. In addition, further facilitated excitations lead to the correlated growth of Rydberg excitations which form a cluster. This leads to a large bunching amplitude $g^{(2)}(0)$ and a long correlation time $\tau(\tau)$, which measures the lifetime of a single cluster $\tau_{\text{cl}}$, i.e., $\tau_{\text{cl}} = \tau(\tau)$. The ion rate $R_{\text{ion}}$ and the cluster lifetime $\tau_{\text{cl}}$ form the basis of our further analysis.

### IV. SINGLE-CLUSTER DYNAMICS

To gain insight into the cluster dynamics, we introduce a simplified cluster model and compare it to our experimental results. The model is capable of describing the growth of the excitation number in a single cluster, and it qualitatively captures all relevant features observed in the experiment. It characterizes the excitation cascade of independent, noninteracting excitation clusters, each of them being described by an effective-rate-equation model.

In the case of strong decoherence, it has been shown that rate-equation models are a good approximation [42–45] and were already successfully used to describe current experiments [15,20,27]. The effective cluster model is illustrated in Fig. 1(c). It only considers the total number of excitations $m$ and does not give insight into the internal microscopic structure of a cluster. The time evolution of the probability $p_m$ to be in a cluster of size $m$ is governed by the equations

$$\partial_t p_0 = -N_{\text{at}} \Gamma_{\text{seed}} p_0 + \Gamma_{\text{sp}} p_1,$$
\[ \begin{align*}
\partial_t p_1 &= +N_{at}\Gamma_{\text{seed}}p_0 - (\Gamma_{\text{sp}} + \Gamma_\uparrow)p_1 \\
&\quad + (\Gamma_\uparrow + 2\Gamma_{\text{sp}})p_2, \\
\partial_t p_m &= -(2\Gamma_\uparrow + m\Gamma_{\text{sp}})p_m + \Gamma_\uparrow p_{m-1} \\
&\quad + (\Gamma_\uparrow + (m + 1)\Gamma_{\text{sp}})p_{m+1} \quad (m \geq 2).
\end{align*} \]  

Within the single-cluster model, we can identify three main contributions to the dynamics:
(i) a slow seed with rate \( \Gamma_{\text{seed}} \) for generating new clusters,
(ii) a fast rate \( \Gamma_\uparrow \) for increasing and decreasing the size of a cluster \( m \), and
(iii) the spontaneous decay rate \( \Gamma_{\text{sp}} = \tau_{\text{sp}}^{-1} \) limiting the size of a cluster.

The seed excitation with rate
\[ \Gamma_{\text{seed}} = \frac{2\Omega^2\gamma_0}{\gamma_0^2 + \Delta^2} \]  

is strongly suppressed because of the large detuning \( \Delta \) in the antiblockade regime. For a system with \( N_{at} \) atoms, we expect independent seed events to occur on a time scale \( (N_{at}\Gamma_{\text{seed}})^{-1} \). However, the presence of already-excited Rydberg atoms strongly alters the summed excitation rate of all neighboring atoms,
\[ \Gamma_{\uparrow} = \frac{\gamma_{\text{eff}}^22\Omega^2\gamma_0}{\gamma_0^2 + (\Delta - \Delta_{\text{int}})^2}, \]  

where \( \Delta_{\text{int}} \) is the interaction shift due to surrounding excited atoms. The competition between the detuning \( \Delta \) and interaction shift \( \Delta_{\text{int}} \) may lead to facilitated excitations [26]. The ratio between seed rate \( \Gamma_{\text{seed}} \) and the rate for cluster growth \( \Gamma_{\uparrow} \) already allows us to identify different regimes of correlated excitation dynamics. For \( \Gamma_{\uparrow} \gg \Gamma_{\text{seed}} \), we expect a strong excitation cascade, creating a cluster. In this regime, each excitation triggers further excitations, and we expect a substantial bunching amplitude \( g^{(2)}(0) \). For \( \Gamma_{\uparrow} \ll \Gamma_{\text{seed}} \), we expect primarily uncorrelated excitations, leading to a bunching amplitude \( g^{(2)}(0) \approx 1 \) and clusters of size one. Finally, we include spontaneous decay with a rate \( m\Gamma_{\text{sp}} \) increasing linearly with the cluster size \( m \). Now, starting from an initial seed excitation, the cluster grows and shrinks with rate \( \Gamma_{\uparrow} \), similar to a random walk along the cluster size \( m \geq 0 \). Although the single-atom spontaneous decay rate \( \Gamma_{\text{sp}} \) can be small, this is the limiting factor for large Rydberg excitation clusters. Microscopically, a decay event creates a defect in a consecutive chain of excited atoms and splits the cluster into smaller parts. The strong interactions prevent these parts from merging again. Besides the geometry, we account for the splittings introducing an effective coordination number \( \bar{z}_{\text{eff}} \). In Appendix B, we discuss the microscopic model in more detail.

The effective model allows us to quantitatively compute the typical size \( \bar{m} \) of a cluster. Moreover, it gives us an intuitive understanding of \( g^{(2)}(0) \) and permits us to estimate the number of independent clusters \( N_{\text{cl}} \). Note that our further experimental analysis does not rely on the explicit evaluation of Eq. (13) and is thus independent of the exact interaction potential between two Rydberg atoms.

### A. Cluster size

To begin with, we discuss the typical size of a cluster \( \bar{m} \). It is given by the ratio of the overall lifetime of a single cluster \( \tau_{\text{cl}} \) and the spontaneous lifetime of a single Rydberg excitation \( \tau_{\text{sp}} \),
\[ \tau_{\text{cl}} = \bar{m}\tau_{\text{sp}}. \]  

The above expression can be understood in the following way: When a cluster contains \( \bar{m} \) excitations, it requires \( \bar{m} \) consecutive decay events of time scale \( \tau_{\text{sp}} \) before the whole cluster vanishes. The facilitation mechanism leads to a correlated excitation growth. This is in contrast to a noninteracting system, where all excitations are independent and decay on the same single-atom time scale \( \tau_{\text{sp}} \). We verified this intuitive picture by comparing full simulations to the simple cluster model and an analytic approach to the corresponding cluster size (see Appendix B). We use Eq. (14) to extract the cluster size from the experimental data. Within the single-cluster model, the cluster lifetime \( \tau_{\text{cl}} \) and thus the typical cluster size \( \bar{m} \) only depend on the ratio \( \Gamma_{\uparrow}/\Gamma_{\text{sp}} \).

In Fig. 3, the measured cluster size \( \bar{m} \) (corresponding to a certain cluster lifetime \( \tau_{\text{cl}} \)) is shown for the full spectrum in the antiblockade regime \( \Delta > 0 \) and different driving strength \( \Omega \). The data reveal a strong increase in the cluster size by increasing the driving strength \( \Omega \) as one would naively expect. In the limit of large detuning \( \Delta \gg \Delta_{\text{int}} \), all excitation rates are strongly suppressed, and we obtain a cluster size of one, corresponding to uncorrelated Rydberg excitations. In this case, the lifetime \( \tau_{\text{sp}} \) is the only relevant time scale. Close to the resonance, cascaded excitations lead to the formation of finite clusters of size \( \bar{m} \lesssim 10 \). Note that for strong driving and/or small detuning, the lifetime of the entire sample approaches the cluster lifetime, and no reliable values for \( \tau_{\text{cl}} \) can be extracted. The preferred generation of clusters with \( \bar{m} > 1 \) in this regime is associated with an excitation bunching. In the inset of Fig. 3, we plot \( g^{(2)}(0) - 1 \). Experimentally, we find substantial bunching amplitudes up to \( g^{(2)}(0) \sim 13 \). The measurements show a peak in \( g^{(2)}(0) \), which coincides with a cluster size of \( \bar{m} = 2 \). The peak shifts towards larger detuning with increasing Rabi frequency \( \Omega \). The presence of a peak and its shift can be understood in the following way: For larger detuning, where the typical cluster size is smaller than \( \bar{m} = 2 \), the influence of an uncorrelated
background noise signal becomes important and diminishes the amplitude. For smaller detunings, the cluster size, as well as the rate of creating new independent clusters, increases drastically. Both reduce the value of $g^{(2)}(0)$ because of an increase of uncorrelated ionization events.

**B. Number of clusters**

Next, we experimentally determine the seed rate $\Gamma_{\text{seed}}$ by an analysis of the first ionization event in each single measurement (details can be found in Appendix A). Comparing the time scale with which new clusters are produced $(N_{\text{at}}\Gamma_{\text{seed}})^{-1}$ to the lifetime of a single cluster $\tau_{\text{cl}}$, we can approximately deduce the number of clusters $N_{\text{cl}}$,

$$N_{\text{cl}} \approx N_{\text{at}}\Gamma_{\text{seed}}\tau_{\text{cl}}. \quad (15)$$

The results are shown in Fig. 4. Similar to the cluster size $\bar{m}$, the number of clusters $N_{\text{cl}}$ increases with increasing driving strength $\Omega$ and decreasing detuning $\Delta$. This shows that for stronger driving and/or smaller detuning, the Rydberg aggregate evolves into a steady state, which is characterized by the presence of a large fluctuating number of independent clusters with rather small size. This already suggests that in the experimentally observed parameter regime, there is no global bistable phase with a large correlation length. For such a phase, we expect the system to be in a single-cluster state with extent over the full system.

**C. Validity of single-cluster model**

The extraction of the cluster size and number from the experimental data performed in the previous section partially relies on the validity of the cluster model. In the following, we make a consistency check to test the cluster model. To this end, we compare the fraction of Rydberg excitations retrieved from the cluster model, $\rho_{\text{cl}} = N_{\text{cl}}\bar{m}/N_{\text{at}} = \tau_{\text{cl}}^2\Gamma_{\text{sp}}\Gamma_{\text{seed}}$, to the excitation fraction $\rho_{e}$ directly extracted from the measured ionization rate $R_{\text{ion}}(t)$: $\rho_{e} = R_{\text{ion}}^{\text{max}}/(N_{\text{at}}\Gamma_{\text{ion}})$. While the first expression incorporates Eqs. (12) and (13) derived from the cluster model, the second solely relies on experimental data. Both quantities should be identical, independent of the Rabi frequency of the driving field and the detuning. The comparison is shown in Fig. 5. The dashed line indicates the ideal agreement $\rho_{\text{cl}} = \rho_{e}$. We see that all experimental curves obtained for different driving strengths fall on top of each other. For small excitation fractions, we furthermore find good agreement between the results from the effective cluster model and the experimental ionization signal. In the strong driving regime, the experimentally retrieved ion signal results in a smaller excitation fraction than predicted by the cluster model. There may be two reasons for this result: First, we monitor the ionization signal in a single experimental run with an exponential atom loss and take the maximal ionization signal to reflect the steady-state excitation fraction without atom loss. Especially in the strong driving regime, atom loss may lead to smaller maximum ionization rates, reducing the extracted excitation fraction $\rho_{e} < \rho_{\text{cl}}$. Second, our simple single-cluster model relies on the assumption that clusters are
independent. However, in Appendix B, we show, by using a full rate-equation simulation, that cluster collisions become important for strong driving. Cluster collisions limit, in particular, the independent growth of clusters. This would also lead to a reduced excitation fraction \( \rho_c < \rho_{cl} \). In summary, we conclude that the cluster model is an appropriate description for the cluster dynamics, especially for small excitation fractions.

V. BEYOND THE SINGLE-CLUSTER MODEL

The experimental protocol and the single-cluster model introduced above only allow us to measure and describe volume-integrated quantities. To also study finite-size effects, in this section, we perform numerical simulations using a many-body rate-equation approach, which goes beyond the simple cluster model and can be extended to large system sizes. In particular, we numerically perform a finite-size extrapolation of the correlation time. In contrast to the cluster model, which describes “pointlike” multilevel systems, here we take into account the spatial distribution of the atoms in the experiment and calculate the level shift for each atom individually. In this way, we have access to the full spatial distribution of excitations and are able to track the growth and decay of every single cluster. The microscopic description and numerical details of the system are discussed in Sec. VI, while we here present the main results.

A. Finite-size scaling of the correlation time

We use the rate-equation model to study the finite-size scaling of the cluster lifetime. The results are shown in Fig. 6. We increase the linear system size \( L \) of a 3D Rydberg lattice gas and extract the cluster lifetime from the \( g_2^{(2)}(\tau) \) correlation function for three different detunings and fixed Rabi frequency \( \Omega = 2\pi \times 160 \) kHz, using the same evaluation as for the experimental data. The data points strongly suggest a saturation in the limit \( L \to \infty \). The dashed lines in Fig. 6 correspond to an exponential saturation fit with a characteristic length scale matching the cluster size \( \bar{m} \). This is a first evidence that clusters, in general, do not extend over the whole system.

B. Full counting distribution and hysteresis

In a system where all time scales are finite, all correlation lengths must be finite unless the interactions are infinitely long ranged. As a consequence, the bimodality of the full counting distribution resulting from the presence of two distinguishable metastable states should only persist if the system size \( L \) is smaller than the correlation length \( \xi \). When \( L \) exceeds \( \xi \), averages are taken over independent regions, and the bimodal distribution starts to wash out. In the limit \( L \gg \xi \), the system behaves as \( N = (L/\xi)^3 \) independent subsystems, and the central limit theorem applies, eventually leading to an overall single-peaked Gaussian distribution. Figure 7(a) shows the distribution function of the excitation number for different values of \( L \) after 1-ms excitation time. In a small system (\( L = 3 \)), seed events are rare, and the cluster size is limited by \( L \). Therefore, the excitation number distribution is peaked at zero excitations. When the system size is comparable to the cluster extent (\( L = 6 \)), a bimodal excitation number distribution occurs. Here, a dynamical switching between low and high total excitation number is observable. However, in the limit of large systems (\( L = 10 \)), the bimodal structure disappears. Instead, a Gaussian distribution emerges with a peak at a large excitation number. Therefore, the bimodality seen in Ref. [28] may be a finite-size effect. Our experimental results and theoretical investigations suggest that the correlation lengths remain short, on the order of the cluster extent, and there is no phase transition to a global bistable regime.

Now, let us discuss dynamic hysteresis features present in metastable systems [32]. While finite-size effects vanish
FIG. 7. (a) Excitation number probability $p_{N_{Ry}}$ for $\Omega = 2\pi \times 185$ kHz and $\Delta = 2\pi \times 15$ MHz for three different linear system sizes $L$ after 1-ms propagation. For intermediate system size ($L = 6$), the distribution shows a bimodal behavior. (b) Hysteresis scan in the detuning parameter $\Delta$ for $\Omega = 2\pi \times 160$ kHz with linear system size $L = 10$. The sweep times in the forward and backward directions are $\tau = 0.88$ ms and $\tau = 88$ ms. In the case of a large sweep time $\tau = 88$ ms, the dynamic hysteresis disappears. The mean Rydberg excitation number in the backward and forward scans are averaged over 500 trajectories. The standard deviation is indicated by the shaded area.

whenever the system size exceeds the correlation length $L \gg \xi$ [see Fig. 7(a)], a hysteresis behavior can still be seen, depending on the ratio between relaxation time $T$ and sweep time $\tau$. Here, the sweep time $\tau$ is the duration of a continuous parameter change in the detuning $\Delta$ from an initial to a final value. In our system, the relaxation towards the stationary state is determined by the growth of finite clusters. Therefore, we expect and numerically verified that $T$ is on the order of the cluster lifetime $t_{cl}$. For example, we show in Fig. 7(b), for two different sweep times $\tau$, the dynamic hysteresis. In the case of a small sweep time $\tau = 0.88$ ms, a large dynamic hysteresis area can be identified, while for $\tau = 88$ ms, the hysteresis area vanishes. The hysteresis behavior persists on a rather large time scale compared to the relaxation time, which is consistent with a simple model that describes the sweep as a sequence of successive small parameter quenches. Since the steady-state excitation number $\bar{N}_{Ry}(\Delta)$ is different for different values of $\Delta$, the coarse-grained relaxation of the excitation number is no longer exponential but attains algebraic corrections. In the limit of many small-parameter quenches with approximately constant relaxation time $T$ starting from an initial detuning $\Delta_0$ to $\Delta_1$ in time $\tau$, the Rydberg excitation number is given by

$$N_{Ry}(\tau) = N_{Ry}(0)e^{-\tau/T} + \tau/T \int_0^\tau dx \bar{N}_{Ry}(x)e^{-x/Ts},$$  \hspace{1cm} (16)

with $x = (\Delta_1 - \Delta)/(\Delta_1 - \Delta_0)$. For $\bar{N}_{Ry} = \text{const}$, we obtain the known exponential relaxation with time scale $T$. However, as seen in Fig. 7(b), the mean Rydberg excitation number is typically not constant. Linearization of $\bar{N}_{Ry}$ directly results in algebraic correction scaling with $(\tau/T)^{-1}$. This may have a tremendous effect on the hysteresis relaxation. To give an example, in our system, the relaxation time is on the order of $T \sim 100$ ms, comparable to the cluster lifetime. However, the hysteresis area vanishes only on a time scale of about 100 ms, which is 3 orders of magnitude larger. The algebraic relaxation of the hysteresis area is consistent with other simulations [16,32] and agrees well with the findings in our system. For more details on the hysteresis relaxation, see Appendix E.

VI. MICROSCOPIC DESCRIPTION AND SIMULATION

Before we conclude, we give some more details about the microscopic description of our experiment. For the relevant length scales, we use an approximate interaction potential $V(r) \approx C_9/r^9$ with $C_9 = 2\pi \times 2.1$ kHz $\mu$m$^9$, which is motivated by exact diagonalization of a subspace of the two-body-interaction Hamiltonian, including dipole-dipole, dipole-quadrupole, and quadrupole-quadrupole interactions [46]. For details regarding the calculation, see Appendix C. We do not take into account possible blackbody induced transitions into neighboring Rydberg states, which have recently been suggested to explain anomalous line broadening in dense samples [47] and which would introduce different types of interactions. While this effect is probably present on the time scales on which our experiments are performed, we believe that because we continuously pump atoms into the 25P state, the above given potential still dominates the cluster growth dynamics. We also checked that the qualitative behavior of the system discussed here stays the same for a different type of interaction (e.g., for van der Waals–type potentials).

Besides the decoherence stemming mainly from laser noise $\gamma_0$, we also include an inhomogeneous broadening $\gamma_{\text{inh}}$ motivated by the coupling to motional states in a lattice of finite width [48,49]. Specifically, we use an additional approximate decoherence rate

$$\gamma_{\text{inh}} = \sum_\langle j \rangle |\partial_r V(r_j)|\sigma/\sqrt{\pi},$$  \hspace{1cm} (17)

where $\sigma = 60$ nm is the width of the localized wave packet in the optical lattice and the sum runs over all excited atoms $\langle j \rangle$ with distance $r_j$. Therefore, the total decoherence rate is $\gamma = \gamma_0 + \gamma_{\text{inh}}$. For our interaction potential, this is a good approximation for $\Delta > 2\pi \times 18$ MHz. (For a detailed discussion, see Appendix D.) Additionally, we take into account the ionization process by a rate $\Gamma_{\text{ion}}$, which corresponds to a loss of atoms. Each ionization event alters the geometry of the sample of Rydberg atoms. To properly
account for the detection background, we superimpose the extracted Rydberg excitation dynamics with an uncorrelated noise signal with rate $\Gamma_n$. It originates from atoms that are trapped on the outer region of the optical lattice. Since the atomic density drops exponentially, these atoms do not contribute to the cascaded excitation dynamics. Although the background noise rate $\Gamma_n$ may depend on the detuning $\Delta$ and driving strength $\Omega$, it is sufficient to approximate it with a constant value $\Gamma_n = 1$ kHz here. Note that all parameters used in our simulations are estimated from or determined by our experimental measurements. For a typical simulation using $N_\text{at} = 1000$ atoms, we already see strongly reduced finite-size effects.

The numerical results obtained for the temporal correlation function $g^{(2)}(\tau)$ are shown in Fig. 8. Although the simulations do not reproduce the experimental results on a quantitative level, we find good qualitative agreement. First, the typical cluster sizes are consistent with the values obtained in the experiment and have approximately the same behavior when varying $\Delta$ and $\Omega$. Second, the structure of the numerically evaluated bunching amplitude $g^{(2)}(0)$ is similar to the one extracted from experiment. In particular, the $g^{(2)}(0)$ peak coincides with a cluster size of $\tilde{m} = 2$ as seen experimentally. However, the actual size of $g^{(2)}(0)$ depends on the system size and therefore strongly deviates from the experiment. We believe this originates from, e.g., the motion of the excited atoms due to the interaction or the decay to other Rydberg states due to blackbody radiation. While we are aware that these effects could play a role, it is numerically very challenging to incorporate all of them in a full many-body simulation. From the results obtained, we infer that the rate-equation model used is nevertheless a sufficiently good approximation of the Rydberg excitation dynamics seen in the experiment.

After checking the validity of the numerical simulation, we can employ it to theoretically study the spatial cluster dynamics. In the case of a van der Waals interaction $V(r) = C_6/r^6$, we observe directional excitation dynamics as indicated in Fig. 1(a) along a certain lattice direction. The directionality stems from the competition between the quasi-long-range interaction and the finite line width $\gamma$. To understand this, let us discuss the case of two neighboring Rydberg excitations on a lattice where a single excitation facilitates its nearest neighbors, i.e., $\Delta = V(a)$. Now, for a third particle along the direction of the two excitations, the residual interaction shift in the antiblockade configuration is $V(a)/64$, while in the orthogonal direction it is $V(a)/8$. Comparing to the line width $\gamma$, this may suppress the growth of the cluster in orthogonal directions while promoting the growth along the lattice direction. Crucially, the directionality depends on the range of the interaction. In the case of short-range interaction, e.g., the interaction potential $V(r) = C_9/r^9$ relevant for our experiment, the impact on the next-nearest neighbors is typically negligible. Therefore, no directional Rydberg excitation dynamics is seen in the full numerical simulations using this potential. However, we believe that directional growth should be experimentally observable in different systems.

VII. CONCLUSION AND OUTLOOK

To conclude, we discussed the excitation dynamics of a large dissipative Rydberg lattice gas probing the full antiblockade regime. We showed that the increase in the relaxation time corresponds to the formation of small clusters within the Rydberg aggregate. Using a simplified cluster model, we could estimate the size of individual clusters, $\tilde{m} \lesssim 10$, as well as the number of independent clusters, $N_\text{cl} \lesssim 500$. Furthermore, we experimentally tested the validity of the effective cluster model. Using a many-body rate-equation model, we find qualitative agreement between our theoretical model and the experimental results. We performed an extrapolation of the cluster size with the linear system size, indicating that the clusters remain small. The absence of long-range correlations indicates the absence of a true phase transition to a bistable regime. Nevertheless, a bimodal excitation number distribution can be observed for small system sizes, as well as a dynamic hysteresis for small sweep times.

Our results show that the experimental and theoretical identification of phase transitions in open quantum systems require a careful analysis of the system-size scaling and of the resulting correlation functions. To fully understand the thermodynamic limit of such systems, numerical simulations should be benchmarked with experimental data. Given the huge numerical effort to calculate and the experimental challenge to measure higher-order correlation
functions with good spatial and temporal resolution, this is, in general, a nontrivial task. We believe that temporal correlations are very helpful in this respect, as they directly reflect the intrinsic dynamics and fluctuations of the steady state. Spatial correlations are instead typically limited to single destructive snapshots of the system and are insensitive to the dynamics.

In our system, a transition to a truly bistable stationary phase is expected to be of first order [9]. An experimental realization would allow us to study these types of phase transitions in stationary states of open systems. We have shown that antiblocked Rydberg gases with van der Waals–type interactions do not lead to diverging correlation times and long-range order in the experimentally relevant regime of large decoherence. Thus, there is no phase transition to a bistable phase. Whether or not long-range order can occur in a coherent regime, taking into account atomic motion, remains an open question. Numerical simulations in this regime can be performed only for very small system sizes, as the coherent many-body dynamics is no longer accessible by classical Monte Carlo simulations. Here, however, experiments are needed that require a substantially reduced decoherence level.

ACKNOWLEDGMENTS

We acknowledge financial support by the DFG within the SFB/TR185. We thank Carsten Lippe for discussing and revising the manuscript. We thank Dominik Linzner and David Petrosyan for stimulating discussions. F. L. and O. T. are recipients of a fellowship through the Excellence Initiative MAINZ (DFG/GSC 266). This work was supported by the Allianz für Hochleistungsrechnen Rheinland-Pfalz (AHRP).

M. F. and H. O. conceived the study of off-resonant Rydberg excitation dynamics. O. T. and T. N. performed the experiment and analyzed the data. H. O. supervised the experiment. F. L. and M. F. developed the theoretical model. F. L. performed the numerical simulations of the many-body rate-equation model. F. L. and O. T. performed numerical simulations of the single-cluster model. M. F. supervised the numerical simulations. F. L. and O. T. prepared the early version of the manuscript. All authors contributed to the data interpretation and manuscript.

APPENDIX A: EXPERIMENTAL METHODS

An experimental cycle consists of loading a 3D magneto-optical trap of $^{87}$Rb for 2.5 s from a 2D magneto-optical trap. The precooled atoms are loaded into a crossed optical dipole trap, generated from a Nd:YAG fiber amplifier at 1064 nm, and they undergo forced evaporation within 4 s, resulting in a Bose-Einstein condensate of 20,000 atoms in an isotropic trap with a trap frequency of $2\pi \times 64$ Hz. After evaporation, we perform a 2-ms-long linear scan with a scanning electron microscope across the cloud, yielding an ion signal proportional to the atom number while only marginally influencing the atomic sample. This allows us to check for atom-number fluctuations during preparation and to correct for long-term instabilities. Because of a finite magnetic-field gradient present during evaporation, the sample is spin polarized in the $F = 1, m_F = +1$ state, from which we transfer it to the $F = 2, m_F = +2$ state by a microwave Landau-Zener sweep. We load the prepared condensate into a 3D optical lattice, created from a Ti:sapphire laser at 748 nm, with lattice constants $a_x, y = 374$ nm and $a_z = 529$ nm in an exponential ramp with time constant $\tau = 20$ ms to a final lattice depth of $S = 20E_{\text{rec}}$. This ensures that we are in the Mott insulating phase regime and have a maximum of one atom per lattice site [50].

To measure the cluster dynamics of Rydberg aggregates, we couple the atomic cloud with a one-photon transition at 297 nm to the Rydberg state $25P_{1/2}$ with fixed detuning $\Delta$ and intensity $I$ for 100 ms. The laser light is produced from a frequency-doubled dye laser at 594 nm. During the light-matter interaction, Rydberg excitations generated in the cloud are photoionized by the trapping light and blackbody radiation with a rate of $\Gamma_{\text{ion}} \approx 2$ kHz and guided by a small electric field of 90 mV cm$^{-1}$ to a discrete dynode detector. The underlying lattice structure hereby ensures that effects from associative ionization [51] or molecular Rydberg states [50] are negligible. Nevertheless, we only probe with blue detuning where molecular states are only addressable through a spin flip in the second atom [52], which is strongly suppressed compared to a normal molecular excitation. We analyze the detected ion signal by calculating the $g^{(2)}$-correlation function binned with $1 \mu$s. We correct for the overall decay of the sample by normalizing with the averaged ion rate of multiple runs. Because of detector ringing and ion repulsion during the time of flight to the detector, we cannot analyze the first two microseconds of the correlation function.

![Graph](image)

**FIG. 9.** We retrieve the seed rate $\Gamma_{\text{seed}}$ by fitting an exponential decay $\exp(-\Gamma_{\text{seed}} t)$ to the probability of detecting a first ion after the time of flight $t$ to our ion detector (inset). From a $\sim \Delta^2$ fit to the extracted seed rates, we can calculate the Rabi frequency $\Omega$, taking into account the finite detection and ionization efficiency, as well as the bare decoherence rate.
The seed rate $\Gamma_{\text{seed}}$ and Rabi frequency $\Omega$ for each experimental parameter are determined by an analysis of the arrival time of the first detected ion (see Fig. 9). Note that the first ion arrival time is independent of the cluster-dynamics analysis. By doing a statistical analysis for each laser power and detuning over multiple experimental runs, we can extract the multiparticle seed rate, taking into account our finite detection and ionization efficiency. Additionally, we extract the underlying Rabi frequency by fitting the seed rates for different detuning with $\Gamma_{\text{seed}} = [(2\Omega^2/\gamma)/\Delta]^2$ (for $\Delta \gg \gamma$). We checked that the extracted Rabi frequencies approximately follow the expected $\sqrt{I}$ scaling.

**APPENDIX B: CLUSTER MODEL**

Here, we study a simple 1D lattice of Rydberg atoms in the antiblockade configuration. Specifically, we set the lattice constant $a$ equal to the facilitation radius $r_{\text{fac}} = (C_0/\Delta)^{1/6}$ assuming that Rydberg atoms interact via a typical van der Waals potential $V(r) = C_6/r^6$. We compare a full numerical simulation based on a many-body rate-equation model [42–45] to our simplified cluster model introduced in Sec. IV.

As pointed out before, the first seed excitation is produced with a rate $N_a\Gamma_{\text{seed}}$. Now, this seed excitation triggers further excitations with a rate proportional to $\Gamma_{\text{fac}} \gg \Gamma_{\text{seed}}$, where

$$\Gamma_{\text{fac}} = \frac{2\Omega^2}{\gamma_0}. \quad (B1)$$

In the case of a 1D system, the first excitation enhances the excitation rate of its two neighbors [see Fig. 10(a)]. We introduce a geometric coordination number $z$ counting the number of atoms with enhanced excitation rate, Eq. (B1). In the 1D case, we identify a geometric coordination number $z = 2$ for the transition between cluster size $m = 1$ to $m = 2$ [see Fig. 10(b)]. First, let us neglect the effect of long-range interactions beyond the lattice constant $a$. Then, a cluster grows and shrinks with rate $z\Gamma_{\text{fac}}$. This is in analogy to a random walk along the cluster size axis $m$ with spreading $\propto \sqrt{\Gamma_{\text{fac}} t}$. However, because of spontaneous decay with rate $\Gamma_{\text{sp}}$, the size of a cluster $m$ is limited. The competition between drive and decay leads to a finite size. A decay event may split the cluster into two parts [see Fig. 10(b)], leading to a doubling of the coordination number $z$. However, this requires that the individual parts of the cluster are spatially well separated and do not influence each other. For instance, the last configuration in Fig. 10(a) shows that the two parts of the cluster cannot merge into a single one. Without the effect of two atoms blocking each other, the coordination number $z = 2n + 1$ increases along the axis of splittings $n$. In Fig. 10(b), the transition rates between all possible configurations labeled by cluster size $m$ and number of splittings $n$ are shown.

In the following, we discuss assumptions, which allow us to estimate the size and the lifetime of a cluster. Since we are mostly interested in the cluster size $m$, we first neglect all splitting processes, fixing the dynamics to the case $n = 0$. This is correct in the regime of weak driving $\Gamma_{\text{fac}} \ll \Gamma_{\text{sp}}$ with coordination number $z = 2$ due to the geometry. For strong driving $\Gamma_{\text{fac}} \geq \Gamma_{\text{sp}}$, we have to account for an increase in the coordination number $z$. To do so, we determine an effective coordination number $z_{\text{eff}}$ self-consistently. This results in the simplified cluster model discussed in Sec. IV with $\Gamma_{\text{fac}} = z_{\text{eff}} \Gamma_{\text{fac}}$ [see Fig. 1(c)].

Now, using a detailed balance ansatz and projecting to the case of having a cluster ($m \geq 1$), we calculate the cluster size distribution $p_m$. This yields the iterative formula

$$p_m = \frac{z_{\text{eff}} \Gamma_{\text{fac}}}{z_{\text{eff}} \Gamma_{\text{fac}} + m \Gamma_{\text{sp}}} p_{m-1} \quad (B2)$$

with normalization condition $\sum_{m=1}^\infty p_m = 1$. Using the full-width half-maximum value $\bar{m}$ of the cluster-size distribution $p_m$, we can estimate the cluster lifetime $\tau_{\text{cl}}$ to be equal to Eq. (14):

![Figure 10](image1.png)

**FIG. 10.** (a) Different configurations during the time evolution of a single cluster starting from an initial seed excitation. In the last configuration, the two excited atoms prevent the excitation of the atom in the middle. A cluster, which is split apart, cannot merge. (b) The configuration space of the cluster dynamics is spanned by the cluster size $m$ and the number of splittings $n$. 

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\[ \tau_{\text{cl}} = \frac{\bar{m}}{\Gamma_{\text{sp}}} \]  

Note that the cluster lifetime agrees with the lifetime of a single Rydberg excitation in the limit \( \Gamma_{\text{fac}} \ll \Gamma_{\text{sp}} \), i.e., \( \bar{m} = 1 \).

Now, we compare the dynamical properties of a large lattice system using a many-body rate-equation method to the simplified cluster model. As discussed in Sec. III, to obtain the lifetime of a single cluster, we calculate the second-order temporal correlation function \( g^{(2)}(\tau) \). We extract the relaxation time, which we identify with the cluster lifetime, and the bunching parameter \( g^{(2)}(0) \). The results for different detunings \( \Delta \) and fixed facilitation radius \( r_{\text{fac}} = a \) are shown in Fig. 11. As we would expect, with increasing driving strength \( \Omega/\Gamma_{\text{sp}} \), the lifetime of the cluster increases. This allows us to determine the effective coordination number \( z_{\text{eff}} \) used in the simplified cluster model. Besides the geometric coordination number \( z = 2 \), we account for the number of splittings, which occur with increasing cluster lifetime, by setting

\[ z_{\text{eff}} = z + (\tau_{\text{cl}} - \tau_{\text{sp}})/\tau_{\text{sp}}, \]

where \( \tau_{\text{sp}} = \Gamma_{\text{sp}}^{-1} \) is the lifetime of a single Rydberg excitation. Comparing the effective cluster model (dashed line) with the full rate-equation simulations (bullet points), we find excellent agreement for small driving strength \( \Omega/\Gamma_{\text{sp}} \). However, the full simulations show that the cluster lifetime decreases again with \( \Omega/\Gamma_{\text{sp}} \) depending on the detuning \( \Delta \). We can understand this as an effect of interacting clusters in the high-excitation density regime, which is beyond the simple cluster model. Increasing the detuning \( \Delta \), we decrease the seed rate \( \Gamma_{\text{seed}} \) with which new clusters are produced. Therefore, the Rydberg excitation density \( \rho_{\text{e}} \) decreases [see inset in Fig. 11(a)]. In our numerical simulation, we identify the regime where cluster collisions are important near an excitation density \( \rho_{\text{e}} = 0.2 \). These collisions strongly reduce the independent growth of a cluster.

Moreover, using the effective cluster model and including the first seed excitation in the cluster-size distribution, we can estimate the bunching parameter \( L \times (g^{(2)}(0) - 1) = \langle m(m-1) \rangle / \langle m \rangle^2 \). Comparing the full simulations with the simplified cluster model, we find good agreement. With increasing driving strength \( \Omega/\Gamma_{\text{sp}} \), the bunching parameter, which measures deviations from Poisson counting statistics, decreases. Therefore, with growing \( \Omega/\Gamma_{\text{sp}} \), the number of independent clusters \( N_{\text{cl}} \) and the lifetime \( \tau_{\text{cl}} \) increase. Both contribute to a reduction of the bunching parameter.

**APPENDIX C: RYDBERG-RYDBERG INTERACTION POTENTIAL**

To calculate the interaction potential between two Rydberg atoms, we perform an exact diagonalization of the interaction Hamiltonian as shown in Ref. [46]. We use a

![FIG. 11. Numerical simulation of the dynamics using a many-body rate-equation method in a large 1D lattice system with size \( L \gg 100 \) for different detunings \( \Delta \) over the line width \( \omega = \gamma_0 \sqrt{\Omega^2/\gamma_0^2 \Gamma_{\text{sp}} + 1} \) and fixed facilitation radius \( r_{\text{fac}} = a \). The decoherence rate is \( \gamma_0/\Omega = 2.05 \) and decay rate is \( \Gamma_{\text{sp}}/\Omega = 0.1 \). (a) Cluster lifetime \( \tau_{\text{cl}} \) and (b) bunching parameter \( g^{(2)}(0) - 1 \). The dashed lines are the results from the cluster model using \( z_{\text{eff}} = 2 + (\Omega/\Gamma_{\text{sp}})^2/8 \) as the effective coordination number.](#)

![FIG. 12. Interaction potentials of the pair state \( |25\text{P}_{1/2}, 25\text{P}_{1/2}\rangle \) obtained from a diagonalization of the interaction Hamiltonian including up to quadrupole-quadrupole interactions. The different potentials correspond to different superpositions of the \( m_j \) states. The green dashed line is a fit to the only repulsive potential with \( V(r) = C_q/r^9 \) and \( C_q = 2\pi \times 2.1 \text{ kHz} \cdot \mu \text{m}^9 \).](#)
basis of pair states $|n_1l_1j_1m_{j_1}n_2l_2j_2m_{j_2}\rangle$ in the energetic vicinity of the state of interest and include all interaction terms up to quadrupole-quadrupole interactions as well as a small finite magnetic field of 1 Gs. For the calculations shown here, the basis consists of 1604 pair states with a maximum angular-momentum quantum number of $l = 3$ in an energetic vicinity of 50 GHz to the pair state $|25P_{1/2}, 25P_{1/2}\rangle$. The results of the diagonalization are shown in Fig. 12. For the $|25P_{1/2}, 25P_{1/2}\rangle$ state, there are four different interaction channels depending on the $m_j$ state of the pair state. The symmetric superposition $(1/\sqrt{2})(|+1/2,-1/2\rangle + |-1/2, +1/2\rangle)$ shows the strongest attraction, the two fully stretched states are nearly degenerate but still attractive, while the antisymmetric superposition gets blueshifted by a dipole-quadrupole interaction with the $|22F_{5/2}24D_{3/2}\rangle m_j$ manifold. We fit the relevant repulsive interaction potential with a $C_0/r^9$-type potential and use it in the simulations presented here.

**APPENDIX D: INHOMOGENEOUS BROADENING**

Here, we discuss an inhomogeneous broadening mechanism resulting from the finite width $\sigma$ of a localized wave packet and the steep slope of an interaction potential $V(r) = C_0/r^6$. The excitation rate $\Gamma_{ex}$ for a ground-state atom in the presence of an atom already excited to the Rydberg state is strongly suppressed and broadened. As already discussed in Refs. [48,49], this stems from the coupling to motional states.

First, using the rate-equation approximation [42], the bare excitation rate of an atom at distance $r_0$ to an excited Rydberg state is

$$\Gamma_{ex}^0(r_0) = \frac{2\Omega^2\gamma_0}{r_0^2 + (\Delta - V(r_0))^2}. \quad (D1)$$

However, this is only true in the limit of vanishing width $\sigma \to 0$. Now, consider an atom localized in an optical lattice. Using the harmonic oscillator approximation, we can approximate the extent of the localized Wannier wave function

$$\Psi(r) = \left(\frac{1}{\pi\sigma}\right)^{1/4} \exp\left(-\frac{r^2}{2\sigma^2}\right), \quad (D2)$$

with the harmonic oscillator width $\sigma$. In a semiclassical approximation, we interpret $|\Psi(r)|^2$ as the probability distribution of finding an atom at position $r$. Then, the total excitation rate $\Gamma_{ex}$ for an atom localized with width $\sigma$ is

$$\Gamma_{ex} = \int_{-\infty}^{\infty} dr |\Psi(r)|^2 \Gamma_{ex}^0(r - r_0), \quad (D3)$$

where $r_0$ is the average distance to the already-excited Rydberg atom. Note, in general, that here we have to evaluate a 3D integral over the positions of all excited atoms. In this case, the effective detuning $\Delta - \sum_j C_\alpha/|r_j|^6$ describes the interaction-shift landscape generated from all excited Rydberg atoms.

In Fig. 13, we plot the excitation rate $\Gamma_{ex}$ for three different experimental relevant distances $r_0$. For two neighboring atoms in a lattice ($r_0 = a_x$), the excitation rate is strongly suppressed and broadened compared to the uncorrelated excitation rate $\Gamma_{ex}$ already occurring at $r_0 = 2a_x$. However, for excitation rates $\Gamma_{ex}/\Gamma_{sp} \gg 1$ larger than the lifetime of a single Rydberg excitation, we can still expect cascaded excitations. Here, we identify a large region up to $\Delta/2\pi = 40$ MHz, where an excitation cascade is possible.

The exact numerical evaluation of the excitation rate (D3) in a large-scale 3D system using a stochastic many-body rate-equation model is numerically challenging. However, to include the strong impact of the inhomogeneous broadening, we use an approximate decoherence rate

$$\gamma = \gamma_0 + \frac{1}{2}\sqrt{2}\gamma_0 \frac{\sigma}{\sqrt{\pi}}, \quad (D4)$$

which can be easily extended to many atoms (see Sec. V). The approximate decoherence rate is motivated by the reduced excitation rate $\Gamma_{ex} = 2\Omega^2/\gamma$ at $\Delta = V(r_0)$ [48] embedded in a Lorentzian excitation profile $\Gamma_{ex}^0 = 2\Omega^2/\gamma^2 + (\Delta - V(r))^2$. This decoherence rate includes the energy width of the linearized interaction potential over the width $\sigma$.

In Fig. 14, we compare the full excitation rate (D3) with the excitation rate $\Gamma_{ex}^0$ using the decoherence rate (D4) for $r_0 = a_x$. Moreover, we plot the bare excitation rate $\Gamma_{ex}^0$...
without additional broadening. The approximate rate $\Gamma_{\text{ex}}^\dagger$ nicely describes the suppression of excitation and the broadening compared to the bare rate $\Gamma_{\text{ex}}^0$. In particular, the regime $\Delta/2\pi \gtrsim 18$ MHz agrees well with the full model (D3). However, for smaller detunings $\Delta/2\pi \lesssim 18$ MHz, the simple approximate model is not valid anymore. Therefore, we believe that the approximate decoherence rate is an efficient and good approximation at least for detunings $\Delta/2\pi \gtrsim 18$ MHz.

APPENDIX E: HYSTERESIS RELAXATION

Here, we discuss a simple model for the relaxation of the hysteresis area. In the case of a single quench from a parameter $\Delta_0$ to $\Delta_1$, we typically have an exponential relaxation of some observable $n$ with relaxation time scale $T_1$:

$$n(t) = \bar{n}_1 + (n_0 - \bar{n}_1)e^{-t/T_1}. \quad (E1)$$

Here, $\bar{n}_1$ is the equilibrium value of the observed parameter and $n_0$ its initial value, which is not necessarily at equilibrium.

In the case of a hysteresis, we perform $N$ small quenches $\Delta_j$ to $\Delta_{j+1}$ on a time scale $\tau$. The total sweep time is $\tau_{\text{sweep}} = N\tau$. We denote $\bar{n}_j$ the equilibrium value at $\Delta_j$ and $n_j = n(\bar{n}_j)$ the nonequilibrium value after time $j\tau$. The relaxation time $T_j = T(\Delta_j)$ might depend on the tuning parameter $\Delta_j$. Then, after $N$ consecutive quenches of the form Eq. (E1), the nonequilibrium value of the observable $n$ is given by

$$n_N = n_0 \exp \left( - \sum_{j=1}^{N} \frac{\tau}{T_j} \right) + \sum_{j=1}^{N} \bar{n}_j (1 - e^{-\tau/T_j}) \exp \left( - \sum_{k=j}^{N-1} \frac{\tau}{T_k} \right). \quad (E2)$$

Now, let us perform a continuum limit, where we keep the sweep rate $\dot{\Delta} = (\Delta_N - \Delta_0)/\tau_{\text{sweep}}$ constant and increase the number of steps $N \to \infty$. We assume equidistant steps in the quench parameter $\delta = \Delta_{j+1} - \Delta_j$. Since the sweep time $\tau_{\text{sweep}}$ and the parameter regime $\Delta_N - \Delta_0$ are constant, we have $\delta, \tau \to 0$. This allows us to approximate the exponential

$$1 - e^{-\tau/T_j} = \tau/T_j \quad (E3)$$

and replace the sum with an integral. We obtain the continuum counterpart of Eq. (E2):

$$n(\tau_{\text{sweep}}) = n_0 \exp \left( - \frac{1}{\Delta} \int_{\Delta_0}^{\Delta_N} \frac{\text{d}\Delta}{T(\Delta)} \right) + \frac{1}{\Delta} \int_{\Delta_0}^{\Delta_N} \text{d}\Delta \left[ \frac{\bar{n}(\Delta)}{T(\Delta)} \exp \left( - \frac{1}{\Delta} \int_{\Delta}^{\Delta_N} \frac{\text{d}\Delta}{T(\Delta)} \right) \right]. \quad (E4)$$

For a constant relaxation time $T = T(\Delta)$, we obtain Eq. (16).

In Fig. 15, we show the relaxation of the Rydberg excitation number with increasing sweep time $\tau_{\text{sweep}}$. The result shows the expected power-law relaxation $\tau_{\text{sweep}}^{-1}$.


[40] The assumption of a steady state breaks down whenever the time scale for the loss of atoms becomes of similar size as the correlation time $\tau^{(2)}$. In our analysis, we discard such data points.


