Rabi oscillations and excitation trapping
in the coherent excitation of a mesoscopic frozen Rydberg gas

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We demonstrate the coherent excitation of a mesoscopic ensemble of about 100 ultracold atoms to Rydberg states by driving Rabi oscillations from the atomic ground state. We employ a dedicated beam shaping and optical pumping scheme to compensate for the small transition matrix element. We study the excitation in a weakly interacting regime and in the regime of strong interactions. When increasing the interaction strength by pair state resonances we observe an increased excitation rate through coupling to high angular momentum states. This effect is in contrast to the proposed and previously observed interaction-induced suppression of excitation, the so-called dipole blockade.

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Rydberg atoms, with their rich internal structure, have been in the focus of atomic physics for more than a century [1]. In the last decade Rydberg physics were extended to laser-cooled atomic gases which allowed the study of a frozen system with controllable, strong inter-actions and negligible thermal contributions. This has opened a wide field in both experiment and theory covering such diverse areas as resonant energy transfer [2,3], plasma formation [4,5], exotic molecules [6,7], and quantum random walks [8,9]. In addition, these frozen Rydberg systems have been proposed as a possible candidate for quantum information processing [10,11]. However, the coherent excitation of Rydberg atoms, an important prerequisite for quantum information protocols, has proven a challenging task. While Rabi oscillations between different Rydberg states have been demonstrated and thoroughly analyzed before [12], Rabi oscillations between the ground state and Rydberg states of atoms have not been observed directly so far, mainly owing to the small transition matrix element.

In this Letter we report on the experimental realization and observation of Rabi oscillations between the ground and Rydberg states of ultracold atoms. We demonstrate how strong interatomic interactions influence the coherent excitation of a mesoscopic cloud of atoms. We show that an interaction-induced coupling to a larger number of internal states can be used to trap the excitation. Employed in a controlled way, this effect offers future applications in the experimental realization of quantum random walks with exciton trapping [2].

Our experiments are performed with a magneto-optical trap (MOT) of about 10^7 ^87Rb atoms at densities of 10^{10} cm\(^{-3}\) and temperatures below 100 \(\mu\)K. Rydberg excitation is achieved with two counterpropagating laser beams at 780 nm and 480 nm (see Fig. 1). The laser at 780 nm is collimated to a waist of 1.1 nm ensuring a constant Rabi frequency of 2\(\pi\) \(\times\) 55 MHz over the excitation volume as determined from Autler-Townes splittings [13]. The laser at 480 nm is referenced to a temperature stabilized ZERODUR-resonator and its beam is shaped with a diffractive optical element that produces a flattop beam profile which is characterized with an adapted CCD camera with a spatial resolution of 5.6 \(\mu\)m. The measured flattop is depicted in Fig. 1. Further details will be described elsewhere [14]. This setup allows a tightly focussed beam with high intensity and at the same time overcomes the limits of a Gaussian beam shape. Gaussian beams have been used for stimulated adiabatic passage to high Rydberg states [13,15] but do not allow the observation of synchronous Rabi floppings in a mesoscopic sample due to the wide range of intensities across the beam [13]. In order to address only atoms in the flattop region we use spatially selective optical pumping (see Fig. 1): The magnetic field of the MOT is turned off 3 ms before excitation [27]. After turning off the trapping lasers, all atoms are pumped to the \(F=1\) hyperfine component of the 5S\(_{1/2}\) ground state within 300 \(\mu\)s.
A small tube of atoms perpendicular to the excitation beams is transferred to the $F=2$ hyperfine component with a 1 μs pulse of a tightly focused repumping beam (waist 10 μm, 10 nW). Only these atoms are resonant with the excitation lasers. They are optically pumped to the stretched $F=2, m_F=2$ state by a 1μs pulse of a $\sigma^+$-polarized pumper beam superimposed with the excitation lasers. From this state we can excite both stretched $nS$ and $nD$ states depending on the helicity of the excitation laser polarization; $n$ denotes the principal quantum number. By maintaining the two-photon resonance but detuning from the intermediate state $5P_{3/2}$ by typically $\delta / 2\pi = 140 \text{ MHz}$ this state experiences negligible population and atoms are directly transferred to the Rydberg state $^{13}. The overlap between the repumping laser and the 480 nm laser defines the excitation volume of about 10 μm × 10 μm × 100 μm, corresponding to ~100 atoms (see Fig. 2(b)). Rydberg atoms are detected by field-ionization and acceleration onto a microchannel plate detector. There is no signal for electric fields below the ionization threshold, therefore we can rule out the presence of spurious ions during excitation. After detection the MOT is turned back on and the whole cycle is repeated every 70 ms. Our setup allows to compensate electric fields $E_L$ in a direction perpendicular to the excitation lasers. Field components parallel to the field plates cannot be compensated and are determined from Stark shifts to be $E_0=160 \text{ mV/cm}$ $^{16}$. The field values given in the text represent the total field $E = (E_L^2 + E_0^2)^{1/2}$.

Fig. 2 shows the measured fraction of excited Rydberg atoms in the $31D_{5/2}$ state as a function of excitation time. Rabi oscillations are clearly visible. We observe the same temporal dependence for both $nS$ and $nD$ states with $n<40$ at both low and high atom densities. This indicates that for these small $n$ values the observed Rabi oscillations are not affected by Rydberg–Rydberg interactions and that all atoms perform identical Rabi oscillations simultaneously. The solid line in Fig. 2 is a theoretical prediction which takes the measured time dependence of the excitation pulses into account. It incorporates three sources of dephasing: the remaining admixture of the intermediate state, the residual intensity distribution of the flat-top beam profile, and a finite laser bandwidth of 2.4 MHz. With increasing importance these three sources lead to the observed damping of the measured Rabi oscillations. A detailed discussion together with quantitative comparisons between measured and calculated Rabi frequencies and other systematic verifications will be published elsewhere $^{14}$. For longer times ($\gtrsim 200 \text{ ns}$) the excitation probability reaches a steady state value of 1/2. This allows us to calibrate the detector: by overlapping the model with the measured data we obtain a scaling factor between single atom excitation probability and detector signal $^{28}$.

The observation of Rabi oscillations between ground and Rydberg states is a first step towards the implementation of quantum information protocols with cold Rydberg gases. The most promising protocols further rely on the so-called dipole blockade $^{11}$. The dipole-blockade describes a system in which the excitation of more than one atom is detuned from the single-atom excitation due to interaction-induced energy shifts, which suppresses multiple excitations. It allows to store a single excitation in a mesoscopic cloud of atoms and is accompanied by an increased collective Rabi frequency, evidence of which was observed only recently $^{17}$. The first experimental signatures of the blockade were observed as a suppression of excitation $^{18,19}$ due to the extra-ordinarily strong van der Waals interactions at large values $n$. Recently, a similar effect was observed at lower values of $n$ by using pair state resonances to switch from van der Waals to the stronger resonant dipole-dipole interaction with an external electric field $^{20}$.

For rubidium, $nD$ states offer similar resonances. Fig. 3 shows the electric field dependence of the involved pair states. At vanishing electric field two $47D_{5/2}$ atoms experience van der Waals interaction. At an electric field of $\sim 260 \text{ mV/cm}$ the $47D_{5/2}→47D_{5/2}$ pair state becomes energetically degenerate with the dipole-coupled $49P_{3/2}→45F_J (J=5/2, 7/2)$ pair state. This changes the interaction to long-range resonant dipole interaction. In contrast to the picture of the dipole blockade, however, the increased interaction strength does not translate into a reduced excitation probability at the atomic resonance: Fig. 4 shows the temporal evolution of the $47D_{5/2}$ population for two different densities. For the lower density of $4.7 \times 10^9 \text{ cm}^{-3}$ (gray points) we observe damped Rabi floppings similar to low-$n$ states. For the higher density ($10^{10} \text{ cm}^{-3}$, black points) we observe an increased excitation that clearly exceeds 0.5. This means that part of the atoms remain in Rydberg states without being stimulated back to the ground state. The effect is clearly density-
dependent and must therefore be interaction-induced.

As our detection cannot distinguish between 47D, 49P, and 45F, we assume an accumulation of atoms in the latter states. However, the increased excitation is in contrast to the previously observed excitation blockade at another pair state resonance [20]. We attribute the difference to the fact that the pair state resonance used in Ref. [20] comprises well-defined pair states (nP3/2-nF3/2 and nS1/2-(n+1)S1/2 with |m|=1/2) while in our case the involved 45F,J state is highly degenerate. As each sub-state has a different coupling strength this leads to a dephasing and a coupling back to the original 47D state is suppressed. The residual electric field in our setup enhances this dephasing as it mixes different m,J-levels and thus leads to a dipole coupling with all possible values of m,J, comprising 14 sub-states of the two values of J=5/2 and 7/2 for the 45F,J state and 6 sub-states for the 47D3/2 state.

We can heuristically model the dephasing between different pair states with an analytical model illustrated in Fig. 3. It incorporates the coherent excitation of a pair |gD⟩ of a ground state atom |g⟩ and a 47D atom |D⟩ to a pair |DD⟩ of two 47D atoms as well as the coupling to the |PF⟩ state with a field-dependent energy shift relative to |DD⟩. The dephasing is introduced phenomenologically by a decay rate Γ out of the three level system. This system is an analog of the excitation of an optical three level system and can be fully described by the according optical Bloch equations [21]. In the case of coherent couplings (Γ=0), this system describes the dipole blockade as an analog to the Autler-Townes splitting in the optical system. By including incoherent coupling (Γ>0), we reproduce the increased excitation measured in Fig. 4. The solid line corresponds to model calculations with parameters for the excitation that are scaled from measured low-n Rabi oscillations. We have set Δ to the theoretical value Δ(E=160 mV) = −2π × 150 MHz (see Fig. 3). This leaves two free parameters: the coupling strength Ω_F = 2π × 40 MHz and the dephasing rate Γ = 2π × 30 MHz where chosen to achieve a good overlap with the experimental data. Both values are consistent with the theoretical value Ω_F^max = 2π × 64 MHz for the strongest coupling between the stretched states (m,J = J = ℓ+1/2) at the most probable interatomic distance of 2.6 μm. The dashed line corresponds to a low density model where Ω_F and Γ are scaled down proportional to the density (as expected for an 1/R^3-dependent interaction).

With the same values for Ω_F and Γ our model also explains spectral measurements shown in Fig. 5. By changing the electric field we can tune the energy difference between the |DD⟩ and |PF⟩ pair states. Fig. 5(a) shows a measured spectrum of the 47D5/2 line at E=160 mV/cm (E⊥=0) together with model calculations. All model parameters are the same as those for the excitation time scan in Fig. 4. Fig. 5(b) shows a spectrum at E=260 mV/cm corresponding to Δ=0. The dashed line shows the model result in the low-density limit (Ω_F = 0), the solid line includes a coupling with Ω_F = 2π × 5 MHz and Γ = 2π × 30 MHz, both chosen to yield the best
overlap with the experiment. We attribute the reduced coupling strength \( \Omega_F \) to the increased electric field which splits the involved states, effectively reducing \( \Omega_F \) from 40 MHz at 160 mV to 5 MHz at 260 mV. There are three distinct features of the measured spectra at higher densities: line broadening, an enhanced excitation probability, and an electric field dependent shift. The model qualitatively describes all of these features.

We want to emphasize that the excitation trapping described here is not due to state redistribution by collisions with ions \[22\] since we have verified that there are no ions produced during the short excitation time. Furthermore the excitation volume is too small to support avalanche ionization. The excitation enhancement is also thermore the excitation volume is too small to support avalanche ionization. We thank T. Pohl for fruitful discussions and C. Pruss and W. Osten from the Institute for Technical Optics in Stuttgart for the production of the beam shaping element. The project is supported by the Landesstiftung Baden-Württemberg within the "Quantum Information Processing" program, and by a grant from the Ministry of Science, Research and Arts of Baden-Württemberg (Az: 24-7532.23-11-11/1).

After submission of this article we became aware of a complementary work which demonstrates Rabi oscillation with an interaction-dependent damping in microscopic samples of 1 to 10 atoms \[26\].

This can result in pair states which are not dipole-coupled to other pair states thus stopping spatial diffusion. This effect can be exploited as an exciton trap for continuous time quantum random walk experiments \[9\]. It also constitutes another constraint to states that are suited for dipole blockade experiments besides the already identified zeroes in Rydberg–Rydberg interactions for certain internal states \[24\] and atom pair alignments \[25\].

We want to emphasize that the excitation trapping described here is not due to state redistribution by collisions with ions \[22\] since we have verified that there are no ions produced during the short excitation time. Furthermore the excitation volume is too small to support avalanche ionization. The excitation enhancement is also different from the antiblockade for the case where the Autler-Townes splitting of the lower transition matches the Rydberg–Rydberg interaction energy \[23\]. The mechanism presented here can be seen as an energy diffusion inside the atom rather than a spatial diffusion which relies on internal state swapping between atoms and is the source of a different dephasing mechanism for energy transfer processes \[2, 3\]. The internal energy dissipation can even counteract spatial diffusion by transferring the excitation into states with different angular momenta.

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**References**

[1] T. F. Gallagher, *Rydberg Atoms* (Cambridge University Press, Cambridge, 1994).
[2] W. R. Anderson et al., Phys. Rev. Lett. 80, 249 (1998).
[3] I. Mourachko et al., Phys. Rev. Lett. 80, 253 (1998).
[4] M. P. Robinson et al., Phys. Rev. Lett. 85, 4466 (2000).
[5] T. Amthor et al., Phys. Rev. Lett. 98, 023004 (2007).
[6] C. Boissseau et al., Phys. Rev. Lett. 88, 133004 (2002).
[7] C. H. Greene et al., Phys. Rev. Lett. 85, 2458 (2000).
[8] R. Côté et al., New J. Phys. 8, 156 (2006).
[9] O. Mühlken et al., Phys. Rev. Lett. 99, 090601 (2007).
[10] D. Jaksch et al., Phys. Rev. Lett. 85, 2208 (2000).
[11] M. D. Lukin et al., Phys. Rev. Lett. 87, 037901 (2001).
[12] T. R. Gentile et al., Phys. Rev. A 40, 5103 (1989).
[13] J. Deiglmayr et al., Opt. Comm. 264, 293 (2006).
[14] M. Reetz-Lamour et al., arXiv:0801.3999 (2008).
[15] T. Cubel et al., Phys. Rev. A 72, 023405 (2005).
[16] K. Singer et al., J. Phys. B 38, S321 (2005).
[17] R. Heidemann et al., Phys. Rev. Lett. 99, 163601 (2007).
[18] D. Tong et al., Phys. Rev. Lett. 93, 063001 (2004).
[19] K. Singer et al., Phys. Rev. Lett. 93, 163001 (2004).
[20] T. Vogt et al., Phys. Rev. Lett. 97, 083003 (2006).
[21] R. M. Whitley and C. R. Stroud Jr., Phys. Rev. A 14, 1498 (1976).
[22] S. K. Dutta et al., Phys. Rev. Lett. 86, 3993 (2001).
[23] C. Ates et al., Phys. Rev. Lett. 98, 023002 (2007).
[24] T. G. Walker and M. Saffman, J. Phys. B 38, S309 (2005).
[25] A. Reinhard et al., Phys. Rev. A 75, 032712 (2007).
[26] T. A. Johnson et al., arXiv:0711.0401 (2007).
[27] We use the thermal expansion of the cloud to change the ground state density by increasing this time up to 15 ms.
[28] This factor is inverse proportional to the number of atoms in the excitation volume and therefore also to the density. By determining this factor we can thus calibrate relative densities. We do this for low-n states. Absolute densities are determined by absorption imaging.