An experimental approach for investigating many-body phenomena in Rydberg-interacting quantum systems

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Received July 3, 2013; accepted September 30, 2013

Recent developments in the study of ultracold Rydberg gases demand an advanced level of experimental sophistication, in which high atomic and optical densities must be combined with excellent control of external fields and sensitive Rydberg atom detection. We describe a tailored experimental system used to produce and study Rydberg-interacting atoms excited from dense ultracold atomic gases. The experiment has been optimized for fast duty cycles using a high flux cold atom source and a three beam optical dipole trap. The latter enables tuning of the atomic density and temperature over several orders of magnitude, all the way to the Bose–Einstein condensation transition. An electrode structure surrounding the atoms allows for precise control over electric fields and single-particle sensitive field ionization detection of Rydberg atoms. We review two experiments which highlight the influence of strong Rydberg–Rydberg interactions on different many-body systems. First, the Rydberg blockade effect is used to pre-structure an atomic gas prior to its spontaneous evolution into an ultracold plasma. Second, hybrid states of photons and atoms called dark-state polaritons are studied. By looking at the statistical distribution of Rydberg excited atoms we reveal correlations between dark-state polaritons. These experiments will ultimately provide a deeper understanding of many-body phenomena in strongly-interacting regimes, including the study of strongly-coupled plasmas and interfaces between atoms and light at the quantum level.

Keywords ultracold Rydberg gases, ultracold plasmas, Bose–Einstein condensation, atom–light interactions, many-body interactions

PACS numbers 32.80.Ee, 32.80.Qk, 34.80.Dp, 37.10.De, 67.85.-d, 67.85.Hj

1 Introduction

The experimental and theoretical investigation of ultracold atomic gases involving highly excited Rydberg states has recently attracted a great deal of interest [1–3]. One reason is that the strong interactions between Rydberg atoms can dramatically influence the laser ex-
citation process (Rydberg blockade effect), giving rise to intriguing new many-body phases and strong spatial correlations between the atoms [2, 4–9]. For example, chirping the excitation laser field may enable the formation of new crystalline states of Rydberg atoms [10–12]. Or, by weakly admixing Rydberg character to ground states with off-resonant laser light it might be possible to introduce new types of interactions between dressed atoms [13–16]. The Rydberg blockade effect can likewise influence the formation of ultracold plasmas, where spatial correlations between Rydberg atoms prior to their ionization may enable the production of ultracold plasmas in the strongly-coupled regime [17, 18]. Interestingly, the strong Rydberg–Rydberg interactions can also have a back-action on the laser fields coupling to the Rydberg state. Very recently, electromagnetically induced transparency using Rydberg states was shown to create effective interactions between photons [19–22], paving the way to the generation of high-repetition-rate sources of nonclassical light as a resource for quantum information processing or for nonlinear optics involving ultracold Rydberg gases [3, 23].

To fully explore these new prospects, modern Rydberg atom experiments require a high level of experimental sophistication in both preparation and detection of the excited atoms. High atomic densities and flexible trapping geometries are a key ingredient, providing large optical densities to enhance nonlinear optical effects, and are used to introduce and control collective many-body effects. Additionally, the development of sensitive detection techniques is required to study the physics of strongly-correlated Rydberg interacting systems, providing direct access to their microscopic details.

In this paper we describe in detail our experimental system tailored for the investigation of strongly-interacting Rydberg gases and the detection of atomic correlations. In addition to good electric field control, our experiment offers fast experimental cycle times, good magnetic field control, and excellent optical access. This is possible through the use of a high flux two-dimensional magneto-optical trap (2D-MOT) cold atom source and a three-beam optical dipole trap. The latter allows for varying the atomic density over three-orders of magnitude, including all-optical preparation of $^{87}$Rb Bose–Einstein condensates, and for generating quasi-one-dimensional and three dimensional Rydberg systems. A special electrode structure is used to control electric fields and to field ionize Rydberg atoms for detection, without limiting the optical access. The paper is structured as follows: Section 2 contains a detailed description of our experimental apparatus, focusing on the special features used to study Rydberg-interacting quantum systems. In Section 3 we review two experiments performed in the strongly-interacting regime using this apparatus: the spontaneous formation of an ultracold plasma from a Rydberg blockaded gas [17], and the emergence of correlations between dark-state polaritons in Rydberg state electromagnetically induced transparency [22].

2 Experiment

We start by reviewing the technical details of our experimental apparatus. Laser cooling and the used trapping techniques which are common to many ultracold atom experiments are described in detail in reviews by Ketterle et al. [24, 25]. Here we focus specifically on the key features of our experimental system (Fig. 1) which make it ideally suited for the investigation of Rydberg-interacting quantum systems. The primary goal is to maximize optical access to the atoms, while providing excellent electric field control. The latter is a main requirement of any Rydberg apparatus, because Rydberg atoms are extremely sensitive to electric fields [26, 27].

Rydberg atoms experience large DC-Stark shifts $\Delta E = -\alpha E^2/2$, since their atomic polarizability $\alpha$ scales with the seventh power of the principal quantum number $n$. For example, the $|55S_{1/2}\rangle$ state has a polarizability of $\alpha \approx 100 \text{MHz}/(\text{V/cm})^2$ [28], while the lifetime limited
transition linewidth is only $\approx 30$ kHz [29]. Thus, a small electric field of $E = 1$ V/cm can shift the Rydberg state by several hundred linewidths. On the other hand, moderate electric fields can be used to manipulate Rydberg atom properties, including the use of Förster tuning of interactions [30–34] or spatially resolved excitation using inhomogeneous electric fields [35]. Another important consideration is stray fields, caused for example by the presence of a few spurious ions [2]. To minimize this possibility we isolate the cold atom source from the main experimental chamber to eliminate ions emitted from the alkali metal dispensers. For this we use a 2D-MOT, which loads a 3D-MOT in the main science chamber [36–39]. Since the cooling mechanism and hence the transmission through the differential pumping tube are exclusive to the designated neutral atomic species, the source is ion free, while providing high atom fluxes which are comparable to Zeeman slowers. Further advantages of the 2D-MOT are that it does not reduce optical access to the science chamber and that it provides a slow atomic beam with small divergence and high beam luminosity which sets the basis for experiments with high repetition rates.

Once the 3D-MOT is loaded from the cold atom beam, the pre-cooled atoms are directly transferred into an optical dipole trap (ODT). We use all-optical evaporative cooling, in order to precisely control the density and temperature of the atoms and to produce a Bose–Einstein condensate (BEC) if required. Compared to magneto-static traps [26, 40], optical dipole traps are unrestricted to the used spin states and allow for almost instant trap switch-off. Furthermore, they minimally affect the optical access to the atoms and their operation is technically less demanding than hybrid optical and magnetic traps [41, 42]. Finally the realization of fast experimental duty cycles using ODTs is comparable to chip based traps [43, 44].

The good optical access in our experiment is exploited for optical absorption imaging using a charged coupled device (CCD) camera. By imaging the absorption of a weak probe beam transmitted through the atomic cloud in the presence of a second strong coupling laser to the Rydberg states it is possible to observe electromagnetically induced transparency (EIT) in a spatially resolved manner. This serves to map the properties of the Rydberg states onto the probe light field, providing a local probe of the system and the possibility to analyze hundreds of EIT-systems in parallel (mapped onto hundreds of pixels), each with different parameters, within a single experimental run. By combining EIT-based imaging with field ionization detection, one can access both coherences and populations of the interacting many-body system simultaneously [45].

### 2.1 High flux cold atom source

Our 2D-MOT is housed in a $140 \times 35 \times 35$ mm$^3$ glass cell and is pressure separated from the science chamber by a differential pumping tube. To maintain the good optical access to the science chamber, we attach the 2D-MOT under an angle of 20° with respect to the horizontal plane of the experiment. To minimize its space consumption we designed a compact (200 $\times$ 200 $\times$ 350 mm$^3$) fiber based cage system which provides the 780 nm cooling light and the magnetic quadrupole fields required for two-dimensional cooling and trapping [see Fig. 2(a)]. The cage system can be divided in three planes that intersect in the symmetry axis of the glass cell. Two perpendicular planes hold one inch polarization optics for...
three equally spaced 2D cooling regions, that are generated by retro-reflected Gaussian laser beams with 20 mm diameter (1/e^2) each. Two units with six permanent neodymium bar magnets [46] are oriented in the third bisecting plane and realize two-dimensional magnetic trapping fields with a field gradient of 16 G/cm. A Gaussian pushing beam with 7 mm diameter (1/e^2) propagates along the symmetry axis of the glass cell and guides the pre-cooled atoms through the 800 μm diameter exit hole of the differential pumping stage into the science chamber. To minimize vacuum contamination in the glass cell, we use 98% enriched ⁸⁷Rb dispensers as our atom source.

For routine operation of the 2D-MOT we use the following parameters which were found to maximize the 3D-MOT loading rate. The loading rate of the 3D-MOT was measured using absorption images taken at different times during 2D-MOT operation. We typically use 80 mW total cooling laser power and 3 mW total repumping laser power. The circularly polarized cooling laser is 9 MHz red detuned from the (5S₁/₂, F = 2) → (5P₃/₂, F = 3) cooling transition and the co-propagating repumping laser is resonant with the (5S₁/₂, F = 1) → (5P₃/₂, F = 2) transition. The atom flux is further increased by separately adjusting a pushing beam to direct atoms through the differential pumping tube. The power of the pushing beam is 40 mW and the red detuning with respect to the cooling transition is 43 MHz. This allows us to address atoms with velocity classes in the range of [0, −45 m/s] that would otherwise not contribute to the atom flux. The use of the pushing beam is found to improve the flux by a factor of 12 ± 2 as illustrated in Fig. 2(b). Varying the dispenser current offers a simple way to control the rubidium vapor pressure, which strongly influences the atomic flux and the MOT loading rate [see Fig. 2(b)]. A maximum 3D-MOT loading rate of (3.9 ± 0.3) × 10⁹ s⁻¹ is obtained for a dispenser current of 3.2 A [47]. For higher currents background gas collisions become important such that the flux decreases again [36–39]. Under optimum conditions we trap 3.5 × 10⁸ atoms in 200 ms. Further details about the design and construction of the 2D-MOT can be found in Ref. [48].

### 2.2 Optimized three beam optical dipole trap

The optical dipole trap (ODT) is designed as a horizontally oriented three beam trap and spatially overlapped with the 3D-MOT using a geometry which provides a large trapping volume and tight confinement. To fulfill these seemingly incompatible requirements, we decouple the loading from the evaporation process using a separate reservoir and dimple trap [49–51]. The large reservoir trap consists of two weakly-focused beams crossing at a small angle, while the dimple trap is tightly focused to enhance the thermalization rate. The ODT is generated from a 50 W single frequency fiber amplifier laser with a wavelength of at 1064 nm (approximately 22 W at the position of the atoms).

The cigar shaped reservoir trap is realized by imaging the Bragg diffraction pattern of an acousto-optic modulator (AOM) onto the atoms [see Fig. 3(a)]. In this way we produce two horizontally polarized Gaussian laser beams of 120 μm waist crossing at an angle of 5.4°, and frequency shifted by the acousto-optical modulator (AOM) frequency νAOM (typically 80 MHz) which prevents laser beam interference. Imaging the AOM not only simplifies the crossing alignment but also allows for dynamical power balancing of the two intersecting reservoir arms through the radio frequency power. For a typical laser power of 10.7 W in each reservoir beam the trap depth is \( U_{0, \text{res}} \approx k_B \times 145 \mu \text{K} \). Most importantly the crossed geometry leads to a significant enhancement of confinement along the symmetry axis of the reservoir. A dramatic consequence of this is illustrated in Figs. 3(b) and (c), where we compare the performance of a single beam reservoir to a crossed beam reservoir, both with the same trap depth and the same radial confinement. As compared to the single beam trap, we observe an

![Fig. 3 Atom trapping in a crossed 1064 nm reservoir trap. (a) Schematic of the reservoir trap. An adjustable (double arrow) imaging system with a ≈ 0.3 magnification is used to image the diffraction pattern of an acousto–optic modulator (AOM) onto the atom cloud. (b) and (c) show absorption images of a single beam (AOM off) and crossed (AOM on) reservoir trap, respectively after 2 ms of free expansion. The lower panels show the according integrated density distributions. Using the same all over laser beam power (21.5 W) in both scenarios, the crossed beam trap confines almost one order of magnitude more atoms.](image-url)
increase in atom number by approximately one order of magnitude: from $4 \times 10^5$ to $3 \times 10^6$. This approach seems to be comparable to or even more efficient than the loading improvement using hybrid magnetic and optical traps [52].

The dimple beam is a third independently controllable, vertically polarized laser beam focused to a waist of 23 µm, crossing the reservoir axis under an angle of 45°, thereby creating an anisotropic ellipsoidal trapping potential. With a typical dimple laser power of 2 W, we increase the depth of the potential by $U_{0,\text{dimple}} \approx k_B \times 350 \mu$K such that a major fraction of the atoms initially collected in the reservoir are transferred into the dimple trap. The reservoir beams also serve to increase the confinement along the weak axis of the dimple trap.

2.3 Preparation of dense atomic samples

Experimental loading cycle — Our experimental loading cycle is optimized to maximize the atom number in the reservoir trap and usually begins with a 550 ms loading phase at a dispenser current of 2.3 A. We typically load $3 \times 10^8$ atoms in the 3D-MOT while the three beam ODT is already switched on. The optimized parameters are: 125 mW total cooling power distributed over six Gaussian MOT beams with 20 mm diameter ($1/e^2$), a red detuning from the cooling transition of 19 MHz, and a magnetic field gradient of 10.5 G/cm. The 2D-MOT is then switched off and the 3D-MOT is slightly compressed within 200 ms by increasing the magnetic field gradient to 14 G/cm and by ramping the cooler red detuning to 25 MHz. To further boost the density of the atomic cloud and hence optimize the loading of the ODT [53], we use a 150 ms temporal dark MOT phase [54]. During this phase, the red detuning of the cooling light is ramped to 48 MHz and the total cooling power is reduced to 11 mW. At the same time the power of the repumping laser is reduced from 140 µW to 1 µW, resulting in a sizable population in the lower hyperfine ground state $|5S_{1/2}, F = 1\rangle$. To minimize atom loss due to hyperfine changing collisions in the ODT, we pump all atoms into the $|5S_{1/2}, F = 1\rangle$ manifold by turning off the repumping laser 2 ms before the end of the dark MOT phase. At this point, about $5 \times 10^8$ atoms have been transferred into the combined trap. After thermalization of the atoms around $T \approx 50 \mu$K and given our trap parameters, we estimate a collision rate $\Gamma_{\text{coll}} \approx 18$ kHz, which constitutes a favorable starting point for the evaporative cooling phase.

Characterization of the evaporation ramp — After 40 ms of free evaporation at constant trap depth we start a forced evaporation phase by adiabatically lowering the depth of the reservoir and the dimple trap. We found that active intensity stabilization of the dipole trap laser system during the evaporation ramp is not necessary. As the re-thermalization rate decreases during evaporation due to the reduced confinement, it is favorable to lower the trap depth at decreasing rate $dU_{0}/dt$ to keep the evaporation efficiency approximately constant [55]. We achieve this with a sequence of three exponential ramps ($A \rightarrow C$, $C \rightarrow E$, $E \rightarrow G$) with durations of 0.4 s, 4 s, and 1 s respectively [see Fig. 4(a)]. The time constants for each ramp were experimentally optimized to obtain the highest gain in phase space density $\rho_0$. Optimum results are found for a dimple power reduction by two orders of magnitude, while the reservoir power is only reduced by a factor of $\approx 25$. The presence of the reservoir beams maintain strong confinement along the weak axis of the dimple throughout the evaporation process.

To characterize the evaporation ramp we first investigate the temperature decrease as a function of the number of atoms remaining in the dimple trap. We find that the temperature decreases by a factor of $\approx 100$ while the atom loss only amounts to a factor of $\approx 10$. The measured temperature rate of change is $\dot{T}/T = (2.3 \pm 0.3) \dot{N}/N$. Assuming harmonic confinement we calculate the atomic density during the evaporation sequence,

$$\rho_0 = N T^3 (m \lambda B/h)^3$$
where $\varpi$ is the geometrical mean of the calculated trapping frequencies and $\lambda_{AB}$ is the thermal de Broglie wavelength. With the temperature $T$ and particle number $N$ it is possible to deduce the phase space density

$$\varrho_0 = \rho_0 \lambda_{AB}^3 \propto NT^{-3}$$  \hspace{1cm} (2)

The black data points (A-G) in Fig. 4 (b) show $\varrho_0$ as a function of $N$. As compared to the exclusive use of the reservoir [blue square (R)], we win approximately two orders of magnitude in phase space density when confining the atoms in the three-beam trap. For later evaporation times (B-G), we observe an increase in phase space density with decreasing atom number, with a constant cooling efficiency

$$\gamma_{evap} = \frac{\ln[\rho_0/\rho_0']}{\ln[N/N']} = 3.3 \pm 0.3$$  \hspace{1cm} (3)

as shown by the solid line in Fig. 4 (b). This compares favorably with other recent experiments: $\gamma_{evap} = 3.5$ [56], $\gamma_{evap} = 2.8 \pm 0.5$ [50]. Furthermore, we estimate the truncation parameter during evaporation $U_0/(k_B T) = 5.8$, which is typical for all-optical evaporation schemes. The critical phase space density is reached with $N \approx 1 \times 10^5$ atoms in the dipole trap. Figure 4 (c) shows the evolution of the atomic density (in situ) during the evaporation ramp. The three-beam trap allows us to maintain a high density throughout the evaporation, since the confinement along the symmetry axis of the trap is almost kept constant via a moderate reduction of the reservoir power. After evaporation the trap can be switched off and by varying the expansion time the atomic density is then varied over several orders of magnitude.

**BEC-transition** — To produce a Bose–Einstein condensate, the dimple trap depth is decreased further. Measurements for different final beam powers of the dimple trap are shown in Fig. 5. At a trap depth of approximately 700 nK, the gas crosses the transition to a BEC. The expected critical temperature with $N \approx 1 \times 10^5$ is $T_c = 260$ nK. Reducing the trap depth further leads to an almost pure condensate with $\approx 2 \times 10^4$ atoms. By Stern–Gerlach separation we measure the relative population of the three magnetic sub-levels to be 0.5, 0.3, and 0.2 in the $|m_F = -1\rangle$, $|m_F = 0\rangle$, and $|m_F = +1\rangle$ magnetic states of the $|5S_{1/2}, F = 1\rangle$ manifold, respectively. Using slower evaporation ramps, the best number achieved was $\approx 6 \times 10^4$ atoms in a 70% pure condensate. The shortest total cycle time which still produced a BEC was around 4.5 s.

2.4 **Rydberg atom detection**

**Rydberg detector** — To manipulate and detect the Rydberg atoms we use a specially designed stainless steel electrode assembly [shown in Figs. 6 (a) and (b)]. It is designed in such a way that it does not obscure the optical access to the atom cloud. I.e., the electrode structure allows the use of beams with maximum beam diameters of 28 mm and provides optical access through four DN63CF viewports with full opening angles of $\approx 26^\circ$ and via three DN40CF viewports with full opening angles of $\approx 13^\circ$, such that optical probing of Rydberg states and optical lattices can easily be implemented. The electrode structure can be used to null electric fields, to apply offset fields and field gradients, or to ionize Rydberg atoms (with fields up to $\gtrsim 1.2$ kV/cm) and guide the resulting ions to a charged particle detector.

The electrode structure is attached to the science chamber with a DN160CF to DN63CF reducing flange with 18 welded SHV connections (8) as shown in Fig. 6 (b). An electrically grounded supporting structure (1) holds in total ten individually addressable electrodes in place, which are used to control the electric field landscape at the position of the atoms. The electrodes are divided into two deflection half rings (2) and eight field plates as illustrated in Fig. 6 (a). The field plates are situated at the extremes of a virtual cylindrical volume of height $44$ mm and diameter $34$ mm and are arranged such that four field plates are situated “above” (3) and four field plates are situated “beneath” (3) the atoms. Each field plate resembles a quarter cylinder segment with an inner radius of $15$ mm, a height of $8$ mm and a wall thickness of $2$ mm.
Fig. 6  Electrode assembly for electric field control and for field ionization of Rydberg atoms. (a) Schematic drawing of the electrode assembly for electric field control and for field ionization of Rydberg atoms. A grounded supporting structure ① holds two deflection rings ② as well as four upper ③ and four lower ④ field plates in place, that are used to control the electric field landscape at the position of the atoms. Charged particle detection is possible with either a micro-channel plate detector (MCP) ⑤ or a channel electron multiplier detector (CEM) ⑥. (b) Photograph of the stainless steel structure which also shows the Faraday cages ⑦ that surround the detectors and the vacuum flange ⑧ with 18 high voltage connections. The position of the atom cloud and the orientation of the dipole trap are sketched on the photograph.

The shape and separation of the field plates and deflection rings were optimized with extensive SIMION simulations to produce the lowest electric field inhomogeneities at the position of the atoms. An example of such a simulation based optimization is shown in Fig. 7. Here we approximated the final shape of the eight field plates by two ring-shaped electrodes as shown in Fig. 7(a).

In the SIMION simulations we varied the geometry (with $A \equiv$ separation, $B \equiv$ thickness, $C \equiv$ inner diameter, and $D \equiv$ outer diameter) of the “test field plates” and computed the normalized electric field vector components. Figure 7(b) shows the normalized modulus of the $z$-component of the electric field as a function of the radial distance from the center of the chamber for five different geometries. Similar results were obtained for the radial field components. In general, we observe that a prolate geometry results in smaller electric field inhomogeneities along the $z$-axis than an oblate geometry. Accounting for the grounded vacuum chamber, the exact course of the electrical wiring and the true shape of the field plates of the final geometry of the electrode structure depicted in Fig. 6, the simulated field deviations only amount to $< 0.5\%$ over a volume of $1\text{mm}^3$. Furthermore, our simulations predict an electric stray field compensation down to $5\text{mV/cm}$ in a volume of $1\times 1\times 1\text{mm}^3$.

We use bipolar high voltage switches with an effective rise time of $\approx 300\text{ns}$ to dynamically control the electric fields and to perform field ionization of Rydberg atoms. We realize this by applying large state-dependent ionization fields. Half-ring-shaped deflection electrodes (②) are used to guide the produced ions on a curved trajectory to either a micro-channel plate detector (MCP; ⑤) or a channel electron multiplier detector (CEM; ⑥) as shown in Fig. 6(a). The CEM and the MCP both have a circular detection area of $165\text{mm}^2$. Contrary to the design described in Refs. [26, 27], the two detectors are mirror-symmetrically installed above the upper field plates to maintain optical access in the horizontal plane of the experiment. To minimize the stray fields produced by the $\approx 2\text{kV}$ bias voltages, both detectors are mounted.

![Fig. 7](image-url) Optimization of the field plate geometry. (a) Basic field plate geometry with $A \equiv$ separation, $B \equiv$ thickness, $C \equiv$ inner diameter, and $D \equiv$ outer diameter of the field plates. (b) The normalized modulus of the $z$-component of the electric field is plotted as a function of the radial distance from the center of the chamber ($x = y = z = 0$). The optimization is shown for five different parameter sets, which are specified in the inset of (b).
inside Faraday cages [see (7) in photograph in Fig. 6(b)]. As the detector has no direct line of sight to the ion’s “place of birth” we obtain absolutely background-free detection signals with single-ion resolution. From the measured ion-pulse-height-distribution we infer a counting efficiency of 0.9, limited by the electrical noise. Combined with the MCP detection efficiency (≈ 0.6), the angular responsivity (≈ 0.9) and the guiding efficiency (≈ 0.9) this gives an estimate for the overall Rydberg atom detection efficiency of ηMCP ≈ 0.4, when using the MCP detector. Using similar arguments and accounting for the CEM detection efficiency of ≈ 0.4, we estimate a overall detection efficiency of ηCEM ≈ 0.3.

Optical probing of Rydberg states — Another possibility to probe Rydberg state properties is the use of Rydberg electromagnetically induced transparency (EIT) as a means for spatially resolved optical detection with a CCD camera [22, 57, 58]. EIT is a quantum interference effect in which an optical control field renders an otherwise absorbing gas transparent [59]. The basic principle involves a three-level system composed of two long-lived states |1⟩ and |3⟩ that are coupled via a short-lived state |2⟩ which spontaneously decays with rate Γ. In Rydberg state EIT the system is a ladder system with |3⟩ being a Rydberg state. A strong laser coupling Ωc results in an Autler–Townes doublet of dressed states, such that the transition amplitudes for the |1⟩ → |2⟩ resonance interfere destructively. This gives rise to a narrow transparency resonance which is mapped onto the light field and which can therefore be detected on a CCD camera.

In the experiments detailed here, we chose the three-level ladder system depicted in Fig. 8(a), with: |1⟩ ≡ |5S1/2, F = 2⟩, |2⟩ ≡ |5P3/2, F = 3⟩, and |3⟩ ≡ |40S1/2⟩. For simplicity we moreover restrict ourselves to a regime of experimental parameters in which Rydberg–Rydberg interaction effects on the EIT can be neglected [60–62]. We prepare a cigar shaped Gaussian cloud (σradial = 75±3 μm, σaxial = 280±10 μm) with N ≈ 2.4×10^5 atoms and a peak atomic density ρ = 2.2×10^6 cm⁻³. The atoms are subsequently excited to |40S1/2⟩-Rydberg state, which does not exhibit significant Rydberg–Rydberg interaction at these densities. The Rydberg state is addressed via resonant two photon excitation, realized by a collimated (waist = 15 mm) 780 nm probe beam and a counter propagating focused (waist = 25 μm) 480 nm coupling beam. The σ⁺ polarized probe laser has a Rabi frequency Ωp/(2π) = (2.2 ± 0.3) MHz, while the σ⁻ polarized coupling laser has a peak Rabi frequency Ωc/(2π) = (4.8 ± 0.2) MHz. Both lasers are switched on for an excitation time of texc = 100 μs. During this time we collect the probe light, which uniformly illuminates the cloud, on a CCD camera for optical detection [see Fig. 1]. The imaging system has an optical resolution of ≈ 12 μm (Raleigh criterion). The fact that EIT allows us to map the properties of the Rydberg state onto the light field is directly seen on the recorded absorption images: on the two photon resonance we observe a local transparency due to the presence of the coupling beam [see Fig. 8(b)]. From a detailed image analysis we infer that the latter causes a ≈ 75% reduction of the two level absorption in the excitation volume.

![Image](image)

**Fig. 8** Optical probing of Rydberg states using electromagnetically induced transparency. (a) Energy level diagram with Ωc (Ωp) the Rabi frequency and Δp, Δc the laser detuning of the 780 nm probe (480 nm coupling) laser, respectively. Γ is the spontaneous decay rate of the intermediate |5P⟩ state. (b) Absorption image of the atomic cloud taken after a free expansion of 600 μs, using the laser parameters Δp = Δc = 0, Ωp/(2π) = (2.2 ± 0.3) MHz, and Ωc/(2π) = (4.8 ± 0.2) MHz. The induced transparency inside the excitation volume is clearly visible in the center of the absorption image.

3 Emergence of correlations in strongly-interacting ultracold Rydberg gases

For neutral atoms, Rydberg–Rydberg interactions possess the strongest inter-particle interaction, exceeding typical interactions in cold gases by several orders of magnitude [1]. In these systems the laser excitation process itself has the potential to create strongly correlated many-body states. The hallmark is the Rydberg blockade [2], in which the interaction between Rydberg atoms inhibits the simultaneous laser excitation of multiple atoms within the blockade radius RBL. This radius is defined as the distance between two atoms at which the Rydberg interaction energy equals the excitation bandwidth Ω:

\[
R_{\text{BL}} = \sqrt{\frac{|C_6|}{\hbar \Omega}}
\]  

(4)

The strong Rydberg–Rydberg interactions typically results in blockade radii on the scale of several micrometers. For instance the |55S1/2⟩ state has C6 = 2πℏ × 50 GHz·μm⁵ and for Ω ≈ 2π × 1 MHz this results in a blockade radius of RBL ≈ 5 μm. The blockade effect determines the excitation dynamics of an ultracold Rydberg...
gas. Since the excitation is coherently shared amongst all atoms inside each blockade sphere, collective excitation effects can arise [63–67]. Bulk properties of Rydberg gases are likewise influenced by the blockade phenomenon as evidenced by a strong suppression of excitation [30, 68–70] or sub-Poissonian counting statistics [22, 71, 72], as will be discussed in Section 3.2.

Figure 9 shows a typical measurement of the suppressed excitation of |55S⟩ Rydberg atoms as a function of atomic density. The latter was varied by controlling the population in the |5S_{1/2}, F = 2⟩ state by optical pumping. The two measurements in Fig. 9 were performed after different expansion times of the gas and the difference in the saturation Rydberg numbers reflects the relative sizes of the excitation volumes. The squares were obtained for an ellipsoidal excitation volume with Gaussian widths $\sigma_x \approx 40 \mu m$, $\sigma_y \approx 27 \mu m$, $\sigma_z \approx 27 \mu m$ after 200μs expansion time, whereas the circles were obtained for a volume defined by $\sigma_x \approx 40 \mu m$, $\sigma_y \approx 16 \mu m$, $\sigma_z \approx 16 \mu m$ and 50μs expansion time. The Rydberg atom number is corrected assuming a detection efficiency of $\eta = 0.4$ and is in good agreement with the stationary solution of the rate equation model described in Ref. [17], using $R_B = 5 \mu m$ (solid lines). The dashed line indicates the anticipated Rydberg atom number neglecting interactions.

![Fig. 9](image)

**Fig. 9** Measurement of interaction-induced Rydberg excitation suppression. The plot shows the |55S⟩ Rydberg atom number (corrected for the 40% detection efficiency) as a function of the atomic density in the excitation volume. It compares two sets of experiments done with different excitation volumes (see text). The solid lines are fits to the data using the stationary solution of a simple rate equation model. The dashed line indicates the Rydberg atom number for the neglect of interactions.

Having discussed the preparation and detection of strongly-interacting Rydberg gases in detail, we now review two experiments in which many-body effects emerge in the strongly-interacting regime. Section 3.1 discusses the application of correlations between Rydberg atoms as a new route to strongly coupled ultracold plasmas. Section 3.2 then describes the observation of spatial and temporal correlations between Rydberg-interacting dark-state polaritons revealed by their sub-Poissonian counting statistics.

### 3.1 Towards strongly coupled ultracold plasmas via Rydberg blockade

In dense samples, ionizing collisions between Rydberg atoms can produce an ion cloud which traps electrons produced during later collisions. Subsequently the trapped electrons rapidly collisionally ionize the remaining Rydberg atoms, resulting in an ensemble of unbound charged particles, referred to as an ultracold plasma [73, 74]. This spontaneous evolution of a Rydberg-interacting gas to an ultracold plasma has been observed in several experiments [17, 75–77], providing a novel approach to study ultracold plasmas. Ultracold plasmas constitute an ideal laboratory system to model exotic phases of matter such as laser-induced plasmas [78], dense astrophysical plasmas [79] or even quark gluon plasmas [80]. Research in this field is fueled by the potential to access the so-called strongly coupled regime, in which the Coulomb interaction energy exceeds the kinetic energy of the ions, giving rise to many-body effects and strong spatial correlations between particles [81, 82]. This regime is quantified by the Coulombic coupling parameter

$$G = \frac{q^2}{4\pi\varepsilon_0 ak_B T} \gg 1$$

Here, the charge and the temperature of the plasma constituents are given by $q$ and $T$, respectively, while $a = \sqrt{3/(4\pi\rho)}$ is the Wigner–Seitz radius, which characterizes the separation between particles in a plasma of density $\rho$. To date ultracold plasmas with $G \approx 0.1–2$ have been realized [83, 84]. However entering deep into the strongly coupled regime has remained out of reach despite the ultralow temperatures possible with laser cooling. Rapid temperature rises were observed during the plasma formation, which is partly attributed to the initially random distribution of the atoms [85–87]. This disorder-induced heating arises when the Coulomb interaction energy of the initial spatial distribution is converted into kinetic energy of the ions, posing one of the primary limits to the attainable coupling strengths in current experiments.

One solution to overcome disorder-induced-heating is to produce the plasma from a Rydberg blockaded sample, in which the constituent particles are spatially ordered, with the probability to find close-by pairs strongly suppressed [17, 18]. This approach is illustrated in Fig. 10 and works as follows: (i) An initially randomly dis-
tributed gas of ultracold neutral atoms is excited to repulsively interacting Rydberg states via a continuous and doubly resonant two-photon laser coupling. (ii) Due to the Rydberg blockade, each Rydberg-excited atom blocks further excitations within the blockade radius, leading to spatial correlations, resembling those of a strongly coupled plasma [18]. After a short time, the Rydberg density reaches steady state, while at later times Rydberg atoms start to decay through a combination of blackbody photoionization [88] and ionizing collisions [89, 90], both leading to a successive increase of charged particles. (iii) Once a critical number of ions has accumulated the resulting space charge starts to trap subsequently produced electrons. At this critical time $t_{\text{crit}}$, rapid electron–Rydberg collisions trigger an ionization avalanche, which ultimately produces an ultracold plasma. Interestingly, the plasma forms rapidly compared to the motional timescales of the ions, since the avalanche is triggered by the fast moving electrons. Therefore the original density–density correlations of the Rydberg gas should be preserved, which is expected to significantly reduce the effects of disorder-induced heating [18, 91].

We have recently performed experiments along these lines, in which we observed the spontaneous ionization dynamics for Rydberg gases of different densities at temperatures around 230 $\mu$K [17]. This was possible by independently measuring the number of ground state atoms, Rydberg atoms, and ions within the sample via the combination of absorption imaging and ionization detection. Evidence for the ionization avalanche deep in the blockade regime is given in Fig. 10(c), which shows the number of plasma ions as a function of continuous excitation time. For an atomic density of $\rho = 5 \times 10^{10} \text{cm}^{-3}$ we witness a sharp increase of plasma ions around $t_{\text{crit}} \approx 10 \mu$s, which is strongly density dependent. We observe that repulsive Rydberg–Rydberg interactions delay the ionization avalanche by several micro seconds. We found that the typical plasma formation time scales are around 2 $\mu$s, which is short compared to the motional dynamics of the ions. During this time the ions only move by approximately 600 nm, which is an order of magnitude smaller than the length scale imposed by the blockade radius $R_{\text{bl}}$. This suggests that the initial correlations should be preserved in the plasma phase, putting the production of strongly coupled ultracold plasmas within reach.

Recent molecular dynamics simulations for a gas of Sr atoms by Bannasch et al. predict that sudden ionization of a pre-structured Rydberg gas would suppress the effects of disorder-induced heating by more than one order of magnitude, giving rise to coupling parameters of up to $G \approx 30$ [18]. Future studies of these systems will benefit from independent tuning of plasma parameters through appropriate choice of atomic densities, Rydberg quantum numbers and laser detunings. This would provide the means to reveal a proposed universal scaling between the coupling parameter $G$ and the number of Rydberg blocked atoms [18] or to explore the phase diagram of ultracold plasmas including strongly coupled liquid phases and liquid-vapor phases [92].

### 3.2 Sub-Poissonian counting statistics of strongly-interacting Rydberg-Polaritons

For high densities the number of Rydberg excitations in the gas should be determined by the maximum number

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*Fig. 10* Generation of an ultracold plasma from a Rydberg-blockaded gas. (a) An initially uncorrelated gas of ultracold atoms is prepared in an optical dipole trap (i). By a doubly resonant two-photon laser excitation some atoms are excited to a Rydberg state. The strong Rydberg–Rydberg interactions prevent the excitation of close-by pairs, leading to spatial correlations (ii). After a time $t_{\text{crit}}$ the Rydberg gas is observed to spontaneously ionize into an ultracold plasma (iii). (b) Qualitative dynamics of the involved populations: ground state atoms $N_g$ (solid line), Rydberg atoms $N_r$ (dashed line), and ions $N_i$ (dash-dotted line). (c) Plasma ion signal taken at a ground state density of $\rho = 5 \times 10^{10} \text{cm}^{-3}$ in a volume of $\approx 1 \times 10^5 \mu$m$^3$. The rapid ionization avalanche starts after $t_{\text{crit}} \approx 10 \mu$s of continuous excitation.
of blockade spheres that fit into the excitation volume. Therefore, the total number of Rydberg excitations saturates and is only subjected to small fluctuations between consecutive experimental runs. This can be seen in the statistical number distributions of Rydberg excitations which changes from Poissonian to sub-Poissonian character in the transition from the weakly-interacting to the strongly-interacting regime [see Figs. 11 (a) and (b)] [22, 71, 72]. This crossover can be quantified by the Mandel Q parameter [93]

\[
Q = \frac{\langle N^2 \rangle - \langle N \rangle^2}{\langle N \rangle} - 1
\]  

(6)

where \( N \) denotes the number of Rydberg atoms. Values of \( Q > 0 \) correspond to super-Poissonian, \( Q = 0 \) to Poissonian and \( Q < 0 \) to sub-Poissonian fluctuations. The latter corresponds to a narrowing of the Rydberg excitation number distribution. The extreme case, a number- or Fock-state is characterized by the total absence of fluctuations with \( Q = -1 \). Therefore the Q parameter is an excellent measure for the effectiveness of the Rydberg blockade on the many-body system.

The Mandel Q parameter can also be related to the spatial correlation function \( g^{(2)}(R) \) of the Rydberg atom distribution by

\[
Q = \int \rho_r \left[ g^{(2)}(R) - 1 \right] d^3R
\]  

(7)

with \( \rho_r \) being the Rydberg density in the excitation volume [94]. Even though (7) is only valid for homogeneous densities and isotropic interactions, it nonetheless underlines the fact that sub-Poissonian counting statistics are intimately connected with the emergence of spatial correlations, reflected by \( g^{(2)}(R) \). Statistical methods for observing the transition to strongly-correlated or even crystalline states have recently been discussed in a number of theoretical papers [8, 95, 96].

We recently reported experimental measurements of sub-Poissonian statistics of dark-state polaritons (DSPs) propagating in a gas of Rydberg-dressed atoms [22]. DSPs arise under the conditions of electromagnetically induced transparency [59], in which a strong optical control field \( \Omega \) renders an otherwise absorbing medium transparent for a weak probe field. At the few photon level, EIT can be described by the propagation of individual dark-state polaritons, which simultaneously possess the properties of both the photonic and the atomic degrees of freedom, which can be interchanged in a fully coherent and reversible way [97].

The DSP wave function for a single probe photon can be written as

\[
|D, 1\rangle = \cos \theta |G, 1\rangle - \sin \theta |R, 0\rangle
\]  

(8)

where the collective quantum states involving \( N \) atoms and the light field are

\[
|G, 1\rangle = |g_1, \cdots, g_i, \cdots, g_N\rangle |1\rangle
\]  

(9a)

\[
|R, 0\rangle = \frac{1}{\sqrt{N}} \sum_i |g_1, \cdots, r_i, \cdots, g_N\rangle |0\rangle
\]  

(9b)

The mixing ratio \( \tan(\theta) = \frac{\sqrt{\rho_c c}}{\sqrt{\Omega}} \) determines the properties of the polariton depending on the atom density \( \rho \), the resonant scattering cross-section \( \sigma \), the speed of light in vacuum \( c \), the decay rate \( \Gamma \) of the intermediate \( |5P_{3/2}\rangle \) state, and the coupling Rabi frequency \( \Omega \).

In our experiments, \( |R\rangle = |55S_{1/2}\rangle \) and the DSPs are almost entirely matter-like with \( \cos^2(\theta) \approx 10^{-4} \). Therefore, measuring the Rydberg population serves as a projective measurement of the number of polaritons inside the cloud. This is achieved through single-atom sensitive detection of the Rydberg population which gives direct access to the matter part of the polariton wave function inside the atomic cloud. Rydberg–Rydberg interactions give rise to an effective interaction blockade for DSPs, which we observe as a transition from Poissonian to sub-Poissonian statistics for the polariton number, indicating the emergence of spatial and temporal correlations between dark-state polaritons.

Figures 11(c) and (d) show a comparison of the statistical distributions for low and high polariton densities corresponding to low and high atomic densities. For low
atomic densities of \( \rho \approx 2 \times 10^{10} \, \text{cm}^{-3} \), we obtain a distribution compatible with a Poissonian distribution and Mandel \( Q \) parameter of \( Q = 0.05^{+0.11}_{-0.10} \) [see Fig. 11 (c)]. In contrast, for \( \rho \approx 8 \times 10^{11} \, \text{cm}^{-3} \), we observe a substantial narrowing of the polariton number distribution, with \( Q = -0.31^{+0.04}_{-0.05} \) [see Fig. 11 (d)], where the experimental uncertainties have been estimated using the bootstrap method. The true distribution \( (Q_{\text{DSP}} = Q/\eta) \) is even narrower as one also has to take into account the ion detection efficiency \( \eta_{\text{MCP}} \approx 0.4 \) [71, 72, 98, 99]. This indicates that the Rydberg–Rydberg interactions give rise to significant spatial correlations between DSPs.

Effective polariton–polariton interactions mediated by the atomic admixture can lead to highly nonlinear [20, 22, 60, 100] and nonlocal optical effects [23], which allow the engineering of novel photonic many-body states. By detuning from the intermediate state, the dissipative character of the effective polariton–polariton interactions can be suppressed while maintaining the giant nonlinearities [101]. This would give rise to elastic scattering of DSPs, providing means to implement deterministic photonic quantum logic gates [102, 103]. As the nonlinearity can be controlled via the choice of the Rydberg state, repulsive, attractive and long-range interactions would become possible [23]. This could pave the way to create and study new types of quantum gases such as dipolar Bose-Einstein condensates of dark-state polaritons [104, 105]. The ability to produce and coherently control the propagation of quantum fields using interacting dark-state polaritons will open up new applications including few-photon nonlinear optics [3], nonclassical light sources [19–21, 106, 107] and photonic quantum logic gates [108, 109].

4 Conclusion and outlook

In this paper we have presented a new experimental system for the investigation of many-body effects in dense ultracold gases of Rydberg atoms. The setup combines high atomic densities with excellent control of external fields and precise Rydberg atom detection. Fast experimental duty cycles are made possible using a high flux 2D-MOT followed by efficient evaporative cooling in an optimized three-beam optical dipole trap, which allows for the production of all-optical \(^{87}\text{Rb}\) BECs. We have described a specially designed electrode structure for electric field control and field ionization and detection of Rydberg atoms. This was then used to study the effects of correlations induced by Rydberg–Rydberg interactions in two vastly different physical regimes.

The first study exploited Rydberg–Rydberg interactions as a means to imprint spatial correlations into an otherwise randomly arranged gas. Subsequent formation of an ultracold plasma was evidenced by a rapid avalanche ionization, suggesting a new route to the production of strongly coupled plasmas [17]. The second study used measurements of the counting statistics to reveal temporal and spatial correlations between Rydberg-interacting dark-state polaritons. Here field ionization measurements served as a projective measurement of the polariton wave function inside the Rydberg medium [22].

In future experiments field ionization of Rydberg states, as described here, could be combined with optical probing of Rydberg states using absorption imaging under EIT conditions [57, 58] (compare Fig. 1). The combination of both techniques will provide information about Rydberg state populations and optical coherences with spatial resolution [45], thereby giving dynamical access to macroscopic and microscopic properties of the system.

Another future goal will be to explore strongly-correlated phases at the single particle level and with high spatial resolution. With this in mind we recently proposed a new imaging technique exploiting the large level shifts induced by a Rydberg impurity atom on a background gas of atoms, that is coupled to an auxiliary Rydberg state via an EIT resonance [110]. Associated with this is a new characteristic radius within which the impurity–probe interactions destroy the transparency, thereby changing the optical properties for many atoms surrounding each impurity. This method should make it possible to image individual Rydberg atoms with high spatial and temporal resolution. This technique is fundamentally nondestructive, thus allowing for time-resolved images of many-body dynamics and can be implemented in existing experiments. Numerous applications of this imaging technique can be foreseen, for example the possibility to directly image the formation (and melting) of crystalline states [10–12]. Other studies could aim at spatially resolving individual dark-state polaritons propagating through the EIT medium and to directly reveal the nonlocal character of the optical nonlinearity [23]. Ultimately, the ability to exploit Rydberg–Rydberg interactions to control both light and atomic fields, and to image the resulting new quantum phases will open a new chapter of Rydberg physics with many new and intriguing possibilities.

Acknowledgements We acknowledge the tremendous support of the mechanical and electronic workshops and the construction office of the Physics Institute of the University of Heidelberg. In particular we want to thank J. Gerhäuser for the mechanical design and S. Rabenecker for the assembly of the electrode structure, as well as K. Stumpf for the mechanical design of the 2D-MOT cage system. This work is supported in part by the Heidelberg Center for Quantum Dynamics and the Deutsche Forschungsges.
meinschaft under WE2661/10.2. M.R.D.S.V. (grant number FP7-PEOPLE-2011-IEF-300870) and S.W. (grant number PERG08-GA-2010-277017) acknowledge support from the EU Marie-Curie program.

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