Simultaneous position and state measurement of Rydberg atoms

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Abstract. We present a technique for state-selective position detection of cold Rydberg atoms. Ground state Rb atoms in a magneto-optical trap are excited to a Rydberg state and are subsequently ionized with a tailored electric field pulse. This pulse selectively ionizes only atoms in e.g. the 54d state and not in the 53d state. The released electrons are detected after a slow flight towards a micro channel plate. From the time of flight of the electrons the position of the atoms is deduced. The state selectivity is about 20:1 when comparing 54d with 53d and the one-dimensional position resolution ranges from 6 to 40 \textmu m over a range of 300 \textmu m. This state selectivity and position resolution are sufficient to allow for the observation of coherent quantum excitation transport.

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1 Introduction

Rydberg atoms are well-known for their large dipole moments. Strong dipole-dipole interaction leads to interesting novel phenomena in particular for cold atoms. Examples are the triggering of spontaneous plasma formation \cite{12} and the so called dipole blockade of optical excitation of a Rydberg atom in the vicinity of an already present Rydberg atom \cite{3,4,5,6}. Another example is the formation of macrodimers from two Rydberg atoms \cite{7,8}.

When the distances between the interacting Rydberg atoms are fixed, which is achieved by using laser cooled atoms, the dipole-dipole interaction can be coherent \cite{9,10}. For a pair of atoms with two successive principal quantum numbers, both in the highest (or lowest) energy Stark state, the interaction leads to an exchange of states between the atoms; in other words an excitation hops from one atom to the other \cite{9}. The time required for hopping of an excitation back and forth is given by:

\begin{equation}
\tau_{\text{hop}} = \frac{9\pi}{n^3} d^3 \quad (1)
\end{equation}

with \( \tau_{\text{hop}} \) in atomic units, \( d \) the distance between the atoms in atomic units and \( n \) the principal quantum number. For \( n=60 \) and \( d=20 \mu m \) the hopping time is 2.8 \textmu s. These mesoscopic numbers make a quantum information system with position-resolved Rydberg atoms feasible. An interesting system \cite{9} is for example the hopping of one excitation (e.g. \( n=61 \) or \( |1\rangle \) in qubit notation) on a lattice of lower Rydberg states (e.g. \( n=60 \) or \( |0\rangle \)). Another example is the diffusion of the \( |1\rangle \) state in a random gas of atoms in state \( |0\rangle \), which might show features of Anderson localization \cite{11}.

In this paper we present experimental results that are a first step towards exploration of the coherent evolution of systems of Rydberg atoms. We show that the position-dependent readout of \( |0\rangle \) and \( |1\rangle \) states is indeed possible. The possibility to create and measure Rydberg atoms in a position and state resolved way is demonstrated here by exciting a narrow region in a cold cloud of rubidium atoms to the 54d state. The one-dimensional position sensitive detection is done with a time-of-flight technique. Time of flight is our choice in preference to a position-sensitive multi-channel plate detector because of simplicity and compatibility with existing equipment.

The time-of-flight approach makes it difficult to combine these measurements with other time-resolved techniques. State resolution is therefore based on the threshold character of field ionization, rather than timing as in conventional ramped state selective field ionization (SFI) \cite{12}. The combination of techniques leads to conflicting optimizations. In the remainder of this paper these problems will be quantified and solved.

2 State and position determination

We first focus on the state determination by SFI \cite{12}. The electric field \( E \) in which a Rydberg atom ionizes is roughly given by the classical ionization threshold, which is at an energy of \( U = -2\sqrt{E} \) in atomic units, and corresponds to a field of \( E = 1/16n^4 \). Due to Stark shifts, however, the energy of the state changes with the electric field. Therefore the ionization field is different. In non-hydrogenic atoms many Stark states couple with each other, forming a level scheme with many avoided crossings. The best state selectivity is achieved when the crossings are traversed either all adiabatically or all diabatically. In the latter case the field has to be ramped up faster than technically convenient, so we choose to ramp the field very slowly. In the adiabatic case the energy stays approximately the same and the ionization field will just be around \( 1/16n^4 \).

For the time-of-flight method we use the same electric field that ionizes the atom. This electric field pushes the electron through one of the ionizing electrodes into a field-free flight tube. The time it takes for the electron to
fly through the tube is a measure for its kinetic energy at the beginning of the tube, and therefore a measure for the potential energy at its starting position. At the position of the atom cloud the electric potential decreases linearly with position (see Fig. 1). Consequently the electric potential, and thus the arrival time of the electron on the detector is a measure for the original position of the Rydberg atom. To magnify the arrival time differences beyond what is possible with the electric potential on axis. Also depicted are the position of the MOT (black dot) and the dye laser beam.

\[ t(x) = L \sqrt{\frac{m_e}{2e(Ex + V_0)}} \]  

(2)

with \( L \) the length of the flight tube \( e \) the charge and \( m_e \) the mass of the electron, \( E \) the electric field and \( V_0 \) the potential at \( x=0 \) minus the potential on the flight tube.

For the time-of-flight technique the time of ionization has to be well defined. This is in contradiction with the slow field ramp needed for the state selective field ionization. Therefore we use a more advanced scheme for the combination of the two techniques: a slow field ramp brings the field just below the ionization limit of the \( |1\rangle \) state, followed by a fast pulse to go above the limit of the \( |1\rangle \) state. In addition, this pulse has to stay below the ionization limit of the \( |0\rangle \) state, in order to detect the atoms in state \( |1\rangle \) exclusively.

### 3 Experimental Setup

To make sure the atoms don’t move on the relevant timescale of the experiment we use a magneto-optical trap (MOT) of \(^{85}\text{Rb} \) atoms as our cold atom source. A typical rubidium MOT has a temperature below 300 \( \mu \text{K} \), which corresponds to an average velocity of 0.3 \( \mu \text{m/\mu s} \). The atoms are loaded from a dispenser into the MOT, created at the intersection of three orthogonal pairs of counterpropagating \( \sigma^+ - \sigma^- \) laser beams at the center of a magnetic quadrupole field. The background pressure is \( 3 \times 10^{-8} \) mbar. Charged particles from the dispenser are removed by deflection in an electric field of 50 V/cm, which is shielded from its environment. The laser frequency is tuned about 13 MHz below the 5S\(_{1/2} \) (F=3) \( \rightarrow \) 5P\(_{3/2} \) (F=4) resonance. A repumping laser beam tuned to the 5S\(_{1/2} \) (F=2) \( \rightarrow \) 5P\(_{1/2} \) (F=3) resonance is added at the MOT center. The cooling laser and the repumping laser are both Toptica DL100 diode lasers at resp. 780 nm and 795 nm and are locked to the rubidium resonance with Doppler-free FM spectroscopy. During all measurements the cooling laser is blocked, so that all atoms are pumped to the 5S\(_{1/2} \) (F=3) ground state by the repumping laser.

To create atoms in a Rydberg state we excite cold atoms with an 8 ns, \( \sim 2 \mu \text{J} \) laser pulse at 594 nm by a two-photon process from the 5s state to 53d or 54d (resp. \( |0\rangle \) and \( |1\rangle \)). This light is provided by a Lambda Physik dye laser pumped with a Spectra Physics frequency-doubled, Q-switched Nd:YAG laser with a repetition rate of 10 Hz. The linewidth of the dye laser pulse is 0.15 cm\(^{-1} \), which is well below the energy spacing between 53d and 54d (1.55 cm\(^{-1} \)). A small contamination of the nearest s-state, which lies 0.32 cm\(^{-1} \) higher, can not be excluded but is of no further relevance. The dye laser beam is focused in the MOT cloud by a lens placed on a micrometer translation stage. The beam waist is determined to be approximately 23(1) \( \mu \text{m} \) (1/e diameter), measured with the knife-edge technique. The polarization of the light is parallel to the electric field that will be applied after the laser pulse.

The Rydberg atoms are field ionized by applying voltage pulses on two field plates P1 and P2 spaced by 2.5 cm (see Fig. 1). The circular stainless steel plates have a diameter of 5.5 cm and have a 1.4 cm hole in the middle for the transmission of one of the pairs of counterpropagating MOT laser beams. The released electrons go through the positively charged field plate P2 into a 40 cm long stainless steel tube, which is at negative potential and serves to slow down the electrons. After a 45 cm flight the electrons are detected on a Hamamatsu Micro Channel Plate (MCP). A copper mesh is placed in front of the MCP. The mesh is 95% open and set at +90 Volt.

Additional coils outside the vacuum chamber compensate for background magnetic field. This is important for the electrons to fly straight towards the detector. The axis of the magnetic quadrupole field for the MOT is in the same direction as the electric field between the field plates.

For the simultaneous position and state detection we use the pulse scheme depicted in the lower half of Fig. 2. Shortly after the dye laser pulse we apply a slow voltage...
ramp on the field plate P1 which goes from 0 V to -115 V in 1.2 µs. It then remains -115 V for 1.3 µs, while a 10 V pulse is applied on the other plate P2 with a duration of 30 ns and a rise time of 10 ns. After that the voltage on the plate P1 is increased to -230 V in 1.2 µs, to ionize all atoms that remained unionized. With the fast pulse we aim to ionize the upper state atoms exclusively at a well defined time. The slow ramp is provided by an Agilent 33240A arbitrary waveform generator amplified by a home-built amplifier system. The 10 V pulse is provided by an HP 8114A high power pulse generator.

For the calibration of the conversion from time-of-flight to position we shift the Rydberg-production volume along the flight path by moving the lens that focuses the dye laser beam in the MOT cloud and record the signal of the MCP for every lens position. The voltage on the flight tube is -48 V and the position of the dye laser beam is 280 µm on the scale of Fig. 3.

For the time-of-flight measurements we will focus on peak b, because this peak is most sensitive to the position of the Rydberg-production volume. Peak b1 consists of 30% of the total signal of state |1⟩. When the atoms are prepared in state |0⟩, the same measurement should ideally

4 Results

In the upper panel of Fig. 2 the MCP signal during the voltage pulses is depicted for two different dye laser wavelengths. The upper trace is the signal for λ=594.166 nm, which excites to the 54d state or the |1⟩ state and the trace below this is the signal for λ=594.193 nm, leading to excitation of the 53d state or the |0⟩ state.

In the upper trace of Fig. 3 three different peaks can be distinguished, labelled a, b and c. Apparently atoms, originally in the 54d state, can be in different states at the time of ionization. This is because the atoms have traversed the avoided crossings in the Stark map differently during the slow field ramp, ending up in a distribution over blue and red states in the Stark manifold. The fast pulse sequentially ionizes some of these states within a very short time window (during the 10 ns rise time of the pulse). As the potential at the time of electron release is different, electrons acquire different velocities depending on whether they originate from red or blue states, which results in peak a (the fast electrons released first) and peak b (the slowest electrons). Peak c results from atoms in a red state that remain unionized by the fast pulse and are ionized by the subsequent ramp. This slow ramp is used for estimating the ionization efficiency during the earlier pulses.
give no signal (b0). From our measurements we find an upper bound that is 1% or 2% of the total signal. The ratio of the genuine and spurious |1⟩ signal is therefore about 20. In future experiments we will attempt to improve this ratio further. The spurious signal could result from atoms that are actually excited to the |1⟩ state, due to the finite laser linewidth, or from atoms in the |0⟩ state that ionize in a smaller field.

Fig. 3 shows time-of-flight spectra for the state |1⟩. The peaks correspond to the peak b1 as defined in Fig. 2 for several positions of the dye laser focus. It can be seen that for faster electrons the peaks are narrower, but also start to overlap more. The peaks in the time-of-flight spectrum appeared to be narrowest when the duration of the fast pulse applied on plate P2 was 30 ns.

We transformed the time-of-flight spectra to the position domain using Eq. 2 with $L=40$ cm. As the potential is significantly more complicated (see Fig. 1) than assumed for the simple model of Eq. 2 we use an effective field $E$ and $V_0$ as fit parameters. Also we add a short offset time of 13 ns, based on numerical calculations, for the parts of the flight that are outside the flight tube. Our fit is based on more traces than depicted in Fig. 3 since the measurements were taken for lens positions 20 µm apart. The peaks in the position domain can be excellently fitted with gaussian profiles, as shown in Fig. 4. Optimal agreement of the center positions of the fitted gaussians with the known 20 micron between lens positions is obtained for $E=50.2$ V/cm and $V_0=-2.2$ V.

It can be seen that in the position domain the peaks get narrower for larger $x$ (i.e. “downhill” on the potential in Fig. 1), while the corresponding peaks in the time domain (i.e. later times) get wider. For large $x$ the broadening due to the finite width of the Rydberg-production volume $w$ (due to a finite laser beam waist) is dominant. For small $x$ the time broadening effect $\tau$ is dominant. This time width $\tau$ and not the Rydberg-production volume $w$ is the limit for the spatial resolution of our time-of-flight system. Therefore we decouple $\tau$ from the 2σ-widths $\Delta x_{TOP}$ of the gaussians by fitting the two broadening effects to the following expression

$$\Delta x_{TOP} = \sqrt{w^2 + \left(\frac{dx(t)}{dt}\right)^2 \tau^2}$$

with $x(t)$ the inverse function of Eq. 2. In the inset of Fig. 4 $\Delta x_{TOP}$ is plotted against the center position $x_{TOP}$ together with the fitted dependence. The fitted parameters are $\tau=20(1)$ ns and $w=21(1)$ µm, which is comparable to $1/\sqrt{2}$ times the measured laser waist (the $1/\sqrt{2}$ factor originates from the two-photon character excitation).

In Fig. 5 we plot the center positions of the gaussians $x_{TOP}$ against the read-off position $x_L$ of the ungated dye laser beam into the MOT cloud ($x_L$), together with the line $x_{TOP} = x_L$. The data points show an excellent agreement with the straight line, which shows that the simple model Eq. 2 is well suited to convert flight times into position. The small horizontal error bar is an estimate of the read-off error (2 µm). The vertical error bars provide the spatial resolution of our system and are obtained by transforming the time width $\tau$ to position. The spatial resolution ranges from 6 to 40 µm over a distance of 300 µm (1/e diameter), and it is better than 20 µm over a range of 150 µm.

5 Conclusions

We have demonstrated a technique for state-selective position detection of cold Rydberg atoms. The state selectivity is about 20:1 when comparing 54d with 53d and might be
improved by using a narrower spectral linewidth of the Rydberg-exciting laser or by optimizing the electric field pulse sequence. The position resolution ranges from 6 to 40 µm over a distance of 300 µm. These n-state and position resolution are sufficient to allow for the observation of long range hopping transport by coherent dipole-dipole interactions in cold Rydberg systems \[9\].

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