Absolute frequency measurements of $^{85}\text{Rb} \, n \, F_{7/2}$ Rydberg states using purely optical detection

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Abstract. A three-step laser excitation scheme is used to make absolute frequency measurements of highly excited $nF_{7/2}$ Rydberg states in $^{85}\text{Rb}$ for principal quantum numbers $n = 33$–100. This work demonstrates the first absolute frequency measurements of rubidium Rydberg levels using a purely optical detection scheme. The Rydberg states are excited in a heated Rb vapour cell and Doppler-free signals are detected via purely optical means. All of the frequency measurements are made using a wavemeter that is calibrated against a Global Positioning System (GPS)-disciplined self-referenced optical frequency comb. We find that the measured levels have a very high frequency stability, and are especially robust against electric fields. The apparatus has allowed measurements of the states to an accuracy of 8.0 MHz. The new measurements are analysed by extracting the modified Rydberg–Ritz series parameters.

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1. Introduction

The accurate measurement of highly excited Rydberg level energies in alkali atoms plays an important role in improving the accuracy of atomic models [1]. In most Rydberg spectroscopy experiments, atoms are detected via field ionization. However, in this study we use a method of purely optical detection in an ordinary vapour cell, which has been demonstrated in [2]–[5]. A vapour cell is a convenient and straightforward solution for finding Rydberg levels, which could potentially permit rapid advances in Rydberg spectroscopy. This technique presents a method of finding Rydberg states quickly, with a large signal-to-noise ratio and an apparent insensitivity to electric fields [3, 4], which makes it particularly well-suited for studying high \( \ell \) Rydberg states with large polarizabilities. It is therefore important to verify the ability to perform precision spectroscopy in such a setup.

Although there is a large body of work on precision interval and fine structure measurements of the different rubidium Rydberg series [6]–[10], measurements of the absolute energies of these levels are more difficult to carry out, and are therefore mainly limited to the lower \( \ell \) states [11]–[14]. It appears that absolute measurements of the \( ^{85}\text{Rb} n F \) series have only been made once by Johansson in 1961 [15] for \( n = 4–8 \). However, as new tools are now available in laser spectroscopy, such as the optical frequency-comb technique, it is interesting to return to such measurements. In this work we wish to demonstrate that precision laser spectroscopy measurements of Rydberg states can be made effectively using purely optical detection with a vapour cell sample.

During the experiment, \( nF_{7/2} \) Rydberg states between \( n = 33–100 \) were excited in \( ^{85}\text{Rb} \) using a three-step laser excitation scheme identical to that outlined in [4, 14]. The three-step level system, shown in figure 1, consists of a 780.24 nm transition \( 5S_{1/2} F = 3 \) to \( 5P_{3/2} F = 4 \), a 775.98 nm transition \( 5P_{3/2} F = 4 \) to \( 5D_{5/2} F = 5 \) and finally a 1260 nm transition \( 5D_{5/2} \) to \( nF_{7/2} \).

To observe excitations to Rydberg states, the first two-step lasers are fixed at their respective transition frequencies and the absorption of the 780 nm laser is monitored while the 1260 nm laser is swept across the transition of interest. This technique involves the quantum amplification effect; due to the large differences in the decay lifetimes of the three excited states of the system, the excitation of a single atom by the third step laser will hinder many absorption–emission cycles on the second step transition. This in turn will hinder a large amount of cycles on the strong first step cycling transition, which can cause a measurable decrease in...
Figure 1. The three-step level scheme used to excite $^{85}$Rb $nF_{7/2}$ Rydberg states in this experiment.

Laser excitation is carried out using $\sigma^+$ polarized light on all three steps. Optical pumping on the strong first step transition, along with dipole selection rules, ensures the second step laser only excites to the $m_F = 5$ sublevel of the 5D$_{5/2}$ $F = 5$ hyperfine state and the third step laser only excites a single transition, the 5D$_{5/2}$ $F = 5$ to $nF_{7/2}$ $F = 6$. Having a well-defined pathway to the Rydberg states is important because of the relatively small $\sim 10$ MHz hyperfine splitting of the 5D$_{5/2}$ level [16]. Successful optical selection in this scheme was previously demonstrated by one of our co-authors in [4].

2. Apparatus

In the experimental setup, shown in figure 2, all three steps are excited using commercial tunable external cavity diode lasers and associated electronics. The third step laser is broadly tunable across a range of 110 nm using a precision stepper motor; this allows a large range of $nF$ states to be accessed. The first and third step lasers are superimposed and co-propagate through an uncoated rubidium vapour cell of length 80 mm. The second step laser travels through the same cell; it counter-propagates and overlaps with the first and third step lasers. Absorption of the first step laser is monitored using a conventional photodiode as the third step laser is swept across the 5D$_{5/2}$ to $nF_{7/2}$ transition of interest. Removal of the first step laser from the other two laser paths is carried out using a polarizing beam splitter. The first two steps are circularly polarized using quarter wave plates, and the third step laser is circularly polarized using a broadband Fresnel rhomb. All three lasers are focused to a beam waist of $\sim 100 \mu$m inside the cell, which increases the available third step laser power density, allowing an interaction time of about 1 $\mu$s. The first
Figure 2. Experimental setup used for measuring Rydberg state frequencies. The first step is phase locked to a self-referenced optical frequency comb and the second step is frequency locked using a separate rubidium reference cell. The first and third step laser light is transported to the comb and wavemeter using single-mode optical fibres.

and second step transitions are driven just below saturation intensity. We were unable to saturate the third step with the 3 mW of available laser power. The laser powers are chosen to give the largest Rydberg excitation rate, with minimal power broadening. The vapour cell is heated to a temperature of 60 °C to increase the atomic density in the cell and to therefore enhance the first step absorption.

In this experiment the first step laser Doppler selects those atoms that take part in subsequent excitations; therefore, it is important that the first step frequency is well known and well stabilized. Hence we stabilize this laser to a self-referenced frequency comb, by phase locking the beat note between the laser and a comb line to a stable direct digital synthesizer. The frequency comb repetition rate is adjusted such that the laser frequency is stabilized to 384 229 242.8 MHz, corresponding to the first step transition frequency from [17]. All locking circuits are referenced to a GPS-disciplined rubidium frequency standard. The comb system allows laser frequencies to be measured with an absolute accuracy of $10^{-11}$. Fast feedback for the offset lock is supplied using a field-effect transistor connected to the laser diode. The stability of the first step lock was measured as less than 100 Hz over all time scales relevant to this experiment. However, the absolute accuracy is limited to the measurement uncertainty of 750 kHz from Barwood et al [17], who observed anomalous offsets on this particular transition, which limits its accuracy as a standard.

Before adding the third step laser to the system, we verified that efficient optical pumping was occurring on the first step transition by scanning the second step laser across the $5D_{5/2}$ manifold, with the first step laser locked. The first step laser selects only zero-velocity atoms, and therefore the second step laser scan showed a single and symmetric Doppler-free peak in the first step absorption. This single peak, with a full width at half maximum (FWHM) of 11.5 MHz, corresponds to the reduced absorption of the first step laser as the second step laser excites...
the 5P_{3/2} F = 4 to 5D_{5/2} F = 5 transition, singled out by the dipole-selection rules. To confirm this we measured the absolute frequency of this transition using our frequency comb and added it to the first step locked frequency to get 770 570 284(1) MHz. This agrees with 770 570 284 734(8) kHz from [16], obtained from two-photon spectroscopy. This therefore demonstrates that the excitation pathway of the first two steps is well understood. This scheme is also used to stabilize the second step laser with a separate room temperature vapour cell. By adding a small frequency modulation to the second step laser, and monitoring the first step absorption via a lock-in amplifier, an error signal is extracted. Using our frequency comb we verified that this second step frequency lock was repeatable to an absolute accuracy of 1 MHz on a daily basis.

We found that it is possible to detect lower n states with a very good signal-to-noise ratio. Therefore, to verify the line shape of the detected signals, the photodiode was monitored directly on an oscilloscope during a fast scan across the 5D_{5/2} to 33F_{7/2} transition. The trace is displayed in figure 3. The scan was carried out in 10 ms and the frequency axis was calibrated using a Fabry–Pérö resonator. The data fit a Lorentzian function with a linewidth of 20 MHz. This linewidth prevents the nF_{7/2} and nF_{5/2} fine structure splitting from being resolved, which for n = 33 to 100 is 4.35 MHz to 0.16 MHz, respectively [9]. However, the use of σ+ light on the third step ensures that only the nF_{7/2} level is excited in this case. This is confirmed by the symmetry of the trace in figure 3, and the absence of any visible effect of the nF_{5/2} level. Even if there is a complete redistribution of the m_F levels in the 5D_{5/2} F = 5 state due to the Earth’s magnetic field, we expect the transition to the nF_{5/2} level to contribute at most 5% to the signal, which can be calculated from the relevant 6j symbols.

To improve the detection sensitivity of third step transitions, a frequency modulation is added to the third step laser via the injection current, with a modulation amplitude of 15 MHz.

**Figure 3.** A scan of the third step laser across the 33F_{7/2} Rydberg state from an oscilloscope. The vertical axis is the first step transmitted intensity from the photodiode; an offset of 10.1 V has been subtracted for convenience. The frequency axis was calibrated with a Fabry–Pérö resonator at 1268 nm. The fitted curve is a Lorentzian with an FWHM of 20 MHz.
and a frequency of 6 kHz. Detection of the first step absorption is carried out at the first harmonic 
using a lock-in amplifier with a time constant of 1 s. The free-running third step laser is scanned 
by applying a linear voltage ramp to the laser piezo using computer software and a digital to 
analogue converter interface. The free-running laser stability was measured as less than 1 MHz 
over 1 s, which is sufficient to carry out slow scans across the Rydberg transitions. As the third 
step laser is scanned, its absolute frequency is monitored using a WS7 High Finesse wavemeter. 
The wavemeter readings are recorded simultaneously using the same computer software.

We used our frequency comb to check the wavemeter’s accuracy and stability across the 
range of third step laser wavelengths used in this experiment. This was carried out by stabilizing 
the third step laser to the comb at arbitrary frequencies across the 1254–1268 nm range, where 
there was a coincident comb line, and measuring its frequency with the wavemeter. We found 
that the wavemeter’s stability stayed below 2 MHz for times of \( \sim 1000 \) s. We also found that the 
wavemeter was able to maintain a day-to-day absolute accuracy of 6.2 MHz across this range, 
when regularly calibrated at 780 nm. Therefore, throughout this experiment the wavemeter is 
calibrated every 30 min to the comb-locked first step laser, to supply a direct frequency link 
with the comb.

3. Results

The third step transition absolute frequencies were collected for \( n = 33–50 \) in intervals of one, 
and from \( n = 50–100 \) in larger intervals of five. Fitting to the transition data was done using a 
Wahlquist first derivative function \[ (1) \]. The function is given by

\[
f(H_δ) = \frac{H_δ}{|H_δ|} \left( 2 \frac{H_ω}{H_δ} \right)^2 \frac{\sqrt{2γ - u}}{2\sqrt{u - 2(u - γ)}},
\]

where \( γ = 1 + β^2 + α^2, \ u = γ + \sqrt{γ^2 - 4α^2}, \ α = H_δ / H_ω \) and \( β = (\frac{1}{2} H_{1/2} / H_ω) \). \( H_{1/2}, H_ω \) and \( H_δ \) 
are the FWHM, modulation amplitude and frequency detuning, respectively. Figure 4 shows a 
typical scan across a Rydberg transition with the fitted profile from (1).

Ten traces were taken for each state in order to understand the repeatability of the 
measurements. It was found that on average the standard deviation of each set of ten scans was 
2 MHz with an accuracy limited by the short-term drift of the wavemeter during the time taken to 
collect each set. The mean transition frequencies of the third step are summarized in the second 
column of table 1. The third column of this table displays the total 5S \( 1/2 \) to \( nF_{7/2} \) frequency, 
measured from the centre of mass of the 5S \( 1/2 \) ground states. These values were calculated by 
adding a constant value of 770 571 549.6 MHz to the third step transition frequencies in column 
two; this frequency was computed from \[ (16) \] and \[ (19) \].

Rydberg atom interactions, for example the dipole–dipole interaction, scale strongly with 
inter-atomic spacing \[ (20) \]. It was therefore important to eliminate any pressure-dependent shifts. 
We estimate an average inter-atomic Rydberg atom spacing of \( \sim 10 \) \( μm \), which is where we 
expect to see observable shifts of a few MHz for the 100F \( 7/2 \) state \[ (20) \]. As well as causing 
a broadening, increased power of any of the three lasers could also cause systematic shifts of 
Rydberg signals. We expect both Rydberg atom density and \( m_F \) state selection efficiency to have 
some dependence on laser power, both of which could result in a peak pulling. There could also 
be a contribution from the light shift \[ (16) \]. To target systematics from pressure and laser power, 
we took measurements of the 33F \( 7/2 \) and 100F \( 7/2 \) transitions with a range of cell temperatures 
from 20 to 65 °C and a range of first, second and third step laser powers, respectively. In all cases
we were unable to detect any repeatable trends in the transition frequencies. Therefore, potential offsets from both temperature- and power-dependent shifts were not added as corrections, but instead the spread of measurements was used to estimate the maximum possible error in each individual case.

Using a fluxgate magnetometer, the magnitude of stray magnetic fields was measured along the axis of the vapour cell as $<20 \mu T$. The Zeeman shift of the $nF_{7/2} F = 6 m_F = 6$ level is 0.06 MHz $\mu T^{-1}$, which gives a Zeeman shift of $<1.1$ MHz. However, because of the four-level nature of the excitation we actually expect Zeeman shifts of the transitions to be equal to the difference in Zeeman shifts between subsequent levels of the excitation scheme, in this case 14 kHz $\mu T^{-1}$ for each of the three steps. We therefore estimate a maximum Zeeman shift of 840 kHz. To check for this shift, measurements of the $33F_{7/2}$ state were made with both $\sigma_+\sigma_+\sigma_+$ and $\sigma_-\sigma_-\sigma_-$ polarized light on the three excitation steps, respectively. Equal and opposite Zeeman shifts are expected for these two cases. We were unable to detect any difference within the short-term drifts of the wavemeter, and therefore we assume that the contribution to the total uncertainty from this effect is negligible. As an additional test of the optical arrangement, and the magnetic field sensitivity, we applied magnetic fields of 100 $\mu T$ along the axis of the cell. In this case we were able to measure shifts of the Rydberg signals of up to 5 MHz.

We also checked for errors from hysteresis and time delays in the data acquisition process by scanning the third step laser across the same transition in opposing directions. We found no difference in these two measurements. The summarized error estimates are displayed in table 2. The wavemeter calibration is a source of random error and all other uncertainties are systematic in nature. The errors add in quadrature to give a total error of 8.0 MHz.

Rydberg $nF$ states are highly polarizable in external electric fields, with polarizabilities scaling as $n^2$ [21]. However, in [3] and [4] no dc Stark shifts were observed when detecting Rydberg atoms in a vapour cell, for electric fields of up to 100 V cm$^{-1}$. This makes a cell...
Table 1. Third step transition frequencies $\nu_3$ and total $5S_{1/2}$ to $nF_{7/2}$ transition frequencies $E_n$ for $n = 33–100$. The total frequencies are measured from the centre of mass of the two $5S_{1/2}$ hyperfine ground states. The total accumulated error on all frequencies is 8.0 MHz.

| $n$ | $\nu_3$ (MHz) | $E_n$ (MHz) |
|-----|---------------|-------------|
| 33  | 236 429 214   | 1007 000 764|
| 34  | 236 604 549   | 1007 176 099|
| 35  | 236 765 078   | 1007 336 627|
| 36  | 236 912 402   | 1007 483 952|
| 37  | 237 047 954   | 1007 619 503|
| 38  | 237 172 932   | 1007 744 481|
| 39  | 237 288 417   | 1007 859 967|
| 40  | 237 395 343   | 1007 966 892|
| 41  | 237 494 542   | 1008 066 092|
| 42  | 237 586 734   | 1008 158 283|
| 43  | 237 672 570   | 1008 244 119|
| 44  | 237 752 610   | 1008 324 159|
| 45  | 237 827 379   | 1008 398 929|
| 46  | 237 897 325   | 1008 468 875|
| 47  | 237 962 850   | 1008 534 399|
| 48  | 238 024 325   | 1008 595 874|
| 49  | 238 082 056   | 1008 653 605|
| 50  | 238 136 367   | 1008 707 917|
| 55  | 238 364 972   | 1008 936 522|
| 60  | 238 538 826   | 1009 110 376|
| 65  | 238 674 124   | 1009 245 673|
| 70  | 238 781 461   | 1009 353 011|
| 75  | 238 868 053   | 1009 439 602|
| 80  | 238 938 927   | 1009 510 477|
| 85  | 238 997 658   | 1009 569 208|
| 90  | 239 046 866   | 1009 618 416|
| 95  | 239 088 516   | 1009 660 066|
| 100 | 239 124 074   | 1009 695 624|

an excellent spectroscopic sample, by eliminating a potential systematic. To check that this experiment was also insensitive to electric fields, we applied electric fields of up to 30 V cm$^{-1}$ across the vapour cell and checked for frequency shifts of both the $33F_{7/2}$ and $100F_{7/2}$ transitions. In each case there was no measurable deviation.

4. Analysis

Rydberg level energies are very well described by the Rydberg formula

$$E_n = E_i - \frac{R_X}{[n - \delta(n)]^2} = E_i - \frac{R_X}{n^2},$$

(2)
Table 2. Estimated errors.

| Type                | Source                      | Error |
|---------------------|-----------------------------|-------|
| Random              | Wavemeter calibration       | 6.2 MHz |
| Systematic          | First step frequency        | 750 kHz |
|                     | Second step frequency       | 1.0 MHz |
|                     | Pressure shifts             | 2.7 MHz |
|                     | Power-dependent shifts      | 4.0 MHz |
|                     | Total                       | 8.0 MHz |

where $E_i$ is the ionization energy, $E_n$ is the excitation energy from the ground state to a state with principal quantum number $n$, $R_X$ is the Rydberg constant for the atom of interest, $\delta(n)$ is the quantum defect and $n^*$ is the effective quantum number. The quantum defect can also be written as a Ritz expansion

$$\delta(n) = \delta_0 + \delta_2 t_n + \delta_4 t_n^2 + \cdots,$$

where

$$t_n = \frac{1}{[n - \delta(n)]^2} = \frac{E_i - E_n}{R_X}.$$

The data from this experiment were analysed using three different fitting methods. The first two methods follow the same theme as [22], while the third method is a consistency check of the data with previous work. These methods are outlined in sections 4.1, 4.2 and 4.3. To aid in the analysis, five values of $E_n$ for $n = 4–8$ were added to the data set from [15]. Weighted fitting was important to take account of the larger uncertainties on these older measurements. Throughout the analysis the Rydberg constant for rubidium 85 was taken as $R_{Rb} = 10,973,660.672,249 \times c$ from [14].

4.1. Method 1

In method 1 the energy levels $E_n$ were fitted using a least squares fitting procedure to the formula

$$E_n = E_i - \frac{R_{Rb}}{[n - \delta_0 - \delta_2 t_n - \delta_4 t_n^2 - \cdots]^2}.$$

The fit algorithm balanced both sides of (5) to find the optimum parameters for $E_i, \delta_0, \delta_2, \delta_4, \ldots$. The results from this fit are displayed in table 3 and the residuals are shown in figure 5. Drake and Swainson [1] describe in great detail how the series parameters extracted from this type of fit can explain physical properties of the Rydberg atom, such as core polarization.

4.2. Method 2

To remove the recursive nature of (3), it is common to make the approximation

$$I_n \approx \frac{1}{(n - \delta_0)^2},$$

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Table 3. Fit parameters from the method 1, 2 and 3 fitting routines. The method 1 fit gives the series parameters $\delta_0$, $\delta_2$ and $\delta_4$; see [1]. The method 2 and 3 fits, and [9], give the equivalent series parameters $\delta_0$, $a$ and $b$; see [22]. Uncertainties are statistically derived from the fitting. The errors on the $E_i$ values include possible systematic contributions outlined in table 2. The parameters from [9] are shown for comparison with the method 3 results.

| Method | $E_i$ (MHz) | $\delta_0$ | $\delta_2$, $a$ | $\delta_4$, $b$ |
|--------|-------------|------------|-----------------|-----------------|
| 1      | 1010024719(8) | 0.016473(14) | $-0.0783(7)$ | 0.028(7) |
| 2      | 1010024719(8) | 0.016473(14) | $-0.0784(7)$ | 0.032(7) |
| 3      | 1010024717(8) | 0.01640(8)  | 0.00(9)        | $-$ |
| Jianing et al [9] | $-$ | 0.0165437(7) | $-0.086(7)$ | $-$ |

Figure 5. Residuals for the $n = 33–100$ states from the method 1 fitting routine. The error bars show the total accumulated error on each data point of 8.0 MHz.

which when substituted into (5) gives a Rydberg–Ritz expression that can be evaluated with greater simplicity [22]:

$$E_n = E_i - \frac{R_{Rb}(n-\delta_0^2-a(n-\delta_0)^3-b(n-\delta_0)^4-\cdots)}{[n-\delta_0-a(n-\delta_0)^2-b(n-\delta_0)^3]^2}.$$  (7)

The method 2 fit involved a direct least squares fit of (7) to the energy levels $E_n$. The results from this fit are displayed in table 3 where the $a$ and $b$ parameters are placed underneath the equivalent $\delta_2$ and $\delta_4$ parameters from the method 1 fit. It can be seen that the values of $E_i$ and the series parameters extracted from the first two fitting methods agree to well within the uncertainties. The value of $E_i$ from this work also lies within $2\sigma$ of the previous value from [14]. An analysis of the residuals shown in figure 5, from the method 1 fit, shows that the points are scattered around a mean of zero with a standard deviation of 4.4 MHz. The states were measured
across several days and therefore this spread comes mainly from the long-term accuracy of the wavemeter.

The Rydberg–Ritz formula in (7) has the significant advantage that it allows any energy level $E_n$ to be calculated with knowledge only of the principal quantum number $n$. In this manner, (7) can be used with the relevant parameters in table 3 to predict the absolute energies of other rubidium $nF_{7/2}$ states outside the range of this experiment.

4.3. Method 3

As a consistency check of these data, we compared the Ritz series parameters extracted from our absolute measurements with those from the most recent relative interval measurements [9]. For this fit we used an abridged version of (7),

$$E_n = E_i - \frac{R_{\text{Rb}}}{[n - \delta_0 - \frac{a}{(n-\delta_0)^2}]}.$$  

(8)

This is the equivalent function that was used for fitting in [9] and is an accurate approximation for $n \geq 20$. For this reason we restricted this fit to the $n \geq 33$ levels. The parameters from this fitting method are shown in table 3 with the values from [9]. The $a$ parameter is placed underneath the equivalent $\delta_2$ parameter from the method 1 fit.

It can be seen that the $\delta_0$ and $a$ parameters from this fit agree at the $2\sigma$ level with those from the previous work [9]. Our parameters are extracted from absolute measurements. Therefore, one does not expect as high a precision as from interval measurements; however, absolute measurements do have the advantage that the ionization energy $E_i$ can also be extracted. The larger errors on the series parameters from this fit, as compared to the method 1 and 2 fits, arise because of the absence of lower $n$ states. This makes extracting higher order series parameters more difficult. For example, in (7), for lower $n$ states the parameters $\delta_0$ and $a$ make a bigger contribution than for higher $n$ states, where $E_n$ becomes dominated by $E_i$. As displayed in table 3, the addition of the lower $n$ states from [15] greatly aided in the reliable extraction of the higher order parameters in the method 1 and 2 fitting routines.

5. Conclusion

We have presented absolute frequency measurements of $nF_{7/2}$ Rydberg states in rubidium 85 to an accuracy of 8.0 MHz. This is a factor 40 improvement over previous measurements of the $n = 4–8$ $nF_{7/2}$ states [15], and gives measurements for a range of $nF_{7/2}$ states between $n = 33–100$ for the first time. The Rydberg–Ritz series parameters that have been extracted from this work allow absolute energies of $nF_{7/2}$ states with higher or lower principal quantum number $n$ to be predicted with comparable accuracy. Our new measurements also show consistency with the results from recent microwave spectroscopy experiments [9]. This work demonstrates that methods of Rydberg spectroscopy involving purely optical detection can be used very effectively to carry out precision measurements of Rydberg states in a simple way, and with extraordinary robustness to dc stark shifts. Not only is the setup simple to construct and maintain but it is easier to use than beam experiments, and Rydberg signals can be monitored in real time on an oscilloscope. We believe that this experiment could be readily adapted to study other alkali metal atoms and could even be used to study unusual features such as Rydberg–Rydberg interactions and molecular states.
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