Efficient, high-resolution resonance laser ionization spectroscopy using weak transitions to long-lived excited states

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Laser spectroscopic studies on minute samples of exotic radioactive nuclei require very efficient experimental techniques. In addition, high resolving powers are required to allow extraction of nuclear structure information. Here we demonstrate that by using weak atomic transitions, resonance laser ionization spectroscopy is achieved with the required high efficiency (1-10%) and precision (linewidths of tens of MHz). We illustrate experimentally and through the use of simulations how the narrow experimental linewidths are achieved and how distorted resonance ionization spectroscopy lineshapes can be avoided. The role of the delay of the ionization laser pulse with respect to the excitation laser pulse is crucial: the use of a delayed ionization step permits the best resolving powers and lineshapes. A high efficiency is maintained if the intermediate level has a lifetime that is at least of the order of the excitation laser pulse width. A model that describes this process reproduces well the observed features and will help to optimize the conditions for future experiments.

I. INTRODUCTION

The study of the fundamental ground-state properties of exotic nuclei is one of the challenges in contemporary nuclear physics research [1]. Laser spectroscopic methods contribute to this ongoing research endeavor by providing information on the nuclear electromagnetic moments, spins, and changes in mean-squared charge radii. These observables provide key input towards a theoretical description of the nucleus, as illustrated e.g. in [2]. Experimental measurements on very exotic nuclei are challenging, reflecting a combination of short half-lives and production in only minute quantities, accompanied by a large amount of unwanted contamination. Furthermore, facilities that produce these exotic nuclei only allot a limited time to a given experiment, which means that the study of the most exotic cases requires techniques that are both very selective and efficient. In addition, such measurements are often performed only once, so systematic uncertainties must be understood, removed, or at least minimized.

Many laser spectroscopic techniques have been applied in nuclear physics research, each with their strengths and weaknesses [3–5]. Resonance Ionization Spectroscopy (RIS) methods, which rely on multi-step laser ionization (RIS) methods, are typically very sensitive, motivating the development of numerous RIS experiments at online isotope separators [6]. To achieve efficient laser ionization, pulsed laser systems are typically used, often operating at high powers. This leads to a drawback for spectroscopy, including unwanted lineshape distortion and/or line broadening. These effects can become apparent when performing high-resolution RIS, as illustrated recently in [7]. Given the current developments towards high-resolution RIS of exotic nuclei in e.g. collinear RIS [7] and in-gas-jet laser spectroscopy [8, 9], a detailed understanding of the interaction of atoms with pulsed lasers is vital.

This article will present a model that describes the RIS process using CW or pulsed lasers for the resonant excitation and non-resonant ionization step (section IV). Spontaneous decay of the intermediate level is taken into account and time delays between the two excitation steps are investigated. Through both model simulations and experimental verification, this article will address how some of the aforementioned detrimental line distortion effects can be understood and avoided by delaying the ionization laser pulse. This will be presented in section V. Furthermore, through the same delayed-ionization approach, it is possible to remove virtually all power broadening due to both lasers in a two-step RIS scheme, which could be important for future high-precision studies on radioactive isotopes. This will be illustrated experimentally and through simulations in section VI.

The delayed ionization method is greatly enhanced by using a weak transition to a long-lived excited state. Firstly, with short-lived excited states, a significant fraction of the excited state population would decay before the ionization can take place, reducing the efficiency of the method. Secondly, long-lived states result in intrinsically narrower linewidth since their natural width is smaller. The feasibility of using weak transitions for efficient resonance laser ionization spectroscopy using both continuous wave and pulsed lasers will be addressed in section V.

II. A MODEL FOR RIS

The evolution of the population of a two level system irradiated by a laser tuned close to resonance can be cal-
culated by solving the Schrödinger equations with the following Hamiltonian:

\[ H = \frac{\hbar}{2} \begin{pmatrix} 0 & \Omega(t) \\ \Omega(t) & 2\Delta \end{pmatrix}, \]  

(1)

where \( \Omega(t) \) is the coupling parameter of the two states, also called the Rabi frequency, and \( \Delta \) is the laser-atom detuning. Defining the laser frequency as \( \omega_e \), the ground state level energy \( \hbar\omega_0 = 0 \) and the excited state energy \( \hbar\omega_1 = \Delta = \omega_1 - \omega_e \). When using linearly polarized light, the coupling parameter \( \Omega(t) \) can be calculated using

\[ \Omega = \sqrt{AP_e} \left( \frac{c}{\omega_1 - \omega_0} \right)^{3/2} (2F_1 + 1) \times \sum_{m_{F_0},m_{F_1}} \left( \frac{F_1}{m_{F_1}} 1 \frac{F_0}{m_{F_0}} 0 m_{F_0} \right) \left\{ J_i F_1 I \right\}, \]  

(2)

with \( F_i \) the total angular momentum of state \( i \), \( () \) and \( \{ \} \) respectively Wigner 3J and 6J symbols, \( A \) the Einstein \( A \) coefficient of the transition and \( P_e(t) \) the power of the laser.

A second laser with laser power \( P_1(t) \) can ionize excited atoms at a rate of \( \Gamma = P_1(t) \sigma \), with \( \sigma \) the non-resonant photo-ionization cross section of the excited state at the wavelength of the ionization laser. Photo-ionization requires modeling population loss, since population has to flow out of the two-level system, into the continuum. This requires a non-Hermitian Hamiltonian, given by

\[ H = \frac{\hbar}{2} \begin{pmatrix} 0 & \Omega(t) \\ \Omega(t) & 2\Delta + 2S(t) - i\Gamma(t) \end{pmatrix}, \]  

(3)

In this Hamiltonian \( S \) is the net dynamic Stark shift induced by both the ionization laser and excitation laser, with each laser contributing a shift proportional to the laser power \[10, 11\]. If there are no relaxation processes, the evolution of the system can be calculated using the time-dependent Schrödinger equation

\[ \dot{\rho} = \frac{i}{\hbar} (H\rho - \rho H^\dagger). \]  

(4)

The right-hand side of this equation reduces to the more familiar commutator \([H, \dot{\rho}]\) for Hermitian \( H \). The populations of the hyperfine levels are the diagonal elements of the density matrix, \( \rho_{ii}(t) \). Usually the spontaneous decay of the excited state to the ground state cannot be neglected. Including incoherent relaxation processes into equation \(4\) can be done as follows:

\[ \dot{\rho} = \frac{i}{\hbar} (H\rho - \rho H^\dagger) + L(\rho). \]  

(5)

For the simplified two-level system, \( L \) is given by

\[ L = \begin{pmatrix} A_{\rho 11} & -\frac{i}{2} A_{\rho 10} \\ -\frac{i}{2} A_{\rho 10} & -A_{\rho 11} \end{pmatrix}, \]  

(6)

This additional term in the equations of motion causes decay of the excited state to the lower-lying state, and exponentially damps the coherence terms.

These equations of motion can be generalized to systems with multiple hyperfine levels in a ground- or excited state multiplet (see Fig. 1) in a relatively straightforward manner. Using Greek indices for levels in the excited state multiplet and roman indices for the ground state multiplet, the Hamiltonian for such a system can be written down as

\[ H_{ii}(t) = \hbar\omega_i \]  

(7)

\[ H_{jj}(t) = 0 \]  

(8)

\[ H_{\alpha\alpha}(t) = \hbar (S_\alpha(t) + \Delta_\alpha - \frac{i}{2}\Gamma(t)) \]  

(9)

\[ H_{\alpha i}(t) = H_{i\alpha}(t) = \frac{\hbar}{2} \Omega_{\alpha i}(t) \]  

(10)

\[ H_{\alpha\beta}(t) = -\frac{\hbar}{2} \Gamma(t)(q + i), \]  

(11)

with \( \hbar\omega_i \) and \( \hbar\omega_\alpha \) the energy of the atomic states of the ground- and excited-state hyperfine multiplet, and \( \Delta_\alpha = \omega_\alpha - \omega_e \). The off-diagonal terms in \(11\) are due to the embedding of structure into the continuum by the high-power ionization laser, and are characterized by a Fano parameter \( q \) \[12\]. This \( q \) parameter plays a role in laser-induced continuum phenomena and can be calculated from first principles (see e.g. \[13,16\]). It induces a coupling between the excited state multiplet levels by the ionization laser via the interaction with the continuum. This coupling can influence of RIS line profiles, but only for either very large values of \( q \) or for unrealistically high laser powers. For the purpose of this article the Fano parameter will be taken to be zero.

The generalized form of the matrix \( L \) can be written
down using the partial decay rates defined as
\[
\gamma_{\alpha i} = \frac{4\alpha |\omega_\alpha - \omega_i|^3}{3e^2}(2F_i + 1)(2J_\alpha + 1)(2J_i + 1)
\]
\[
\times |\langle \alpha_L, L_\alpha | r | i, L_i \rangle|^2 \left\{ \begin{array}{cc} J_\alpha & 1 \\ F_\alpha & I_\alpha \end{array} \right\}^2 \left\{ \begin{array}{cc} L_\alpha & 1 \\ J_\alpha & S_J \end{array} \right\}^2 .
\]

These rates can be calculated using the observation that the partial decay rates of an excited hyperfine level should sum up to the Einstein \(A\) coefficient. Using this definition of the partial rates, \(L\) can be written down as
\[
L(\rho)_{ii} = \sum_\alpha \rho_{\alpha\alpha} \gamma_{\alpha i}
\]
(13)
\[
L(\rho)_{\alpha\alpha} = -\sum_i \rho_{\alpha\alpha} \gamma_{\alpha i}
\]
(14)
\[
L(\rho)_{\alpha i} = -\frac{\rho_{\alpha i}}{2} \sum_j \gamma_{\alpha j}
\]
(15)
\[
L(\rho)_{i\alpha} = -\frac{\rho_{i\alpha}}{2} \sum_j \gamma_{\alpha j}
\]
(16)
\[
L(\rho)_{\alpha\beta} = -\frac{\rho_{\alpha\beta}}{2} \sum_j \gamma_{\alpha j} + \gamma_{\beta j}.
\]
(17)

These are the generalized equations that will be used for the simulations presented throughout this article. The computer code developed in Python used to run the simulations, is available upon request to the authors.

III. POWER BROADENING AND DELAYED IONIZATION

A. Model Predictions

In a two-step RIS scheme, both lasers can broaden the resonance line profiles. Power broadening due to the excitation laser in a closed two-level system using continuous-wave (CW) laser light is well understood in the steady-state limit. In this case, the linewidth of the optical resonance increases with the laser power [17]:
\[
\text{FWHM} = A\sqrt{1 + 2(\Omega/A)^2}. \tag{18}
\]

In other words, population is only excited efficiently to the excited state when
\[
|\Delta| \lesssim A\sqrt{1 + 2(\Omega/A)^2}. \tag{19}
\]

However, for pulsed laser excitation, this relationship is not always valid [18]. It can be derived that for a Gaussian shaped excitation laser pulse (in absence of ionization and spontaneous decay), the detuning range that results in excited state population after the action of the excitation pulse is given by [19]
\[
|\Delta| \lesssim \frac{\sqrt{\log(\Omega/\Delta)}}{T}, \tag{20}
\]
with \(T\) the length of the laser pulse in time. The different power dependence of the line widths for CW and pulsed laser excitation is illustrated in Fig. [2] using \(A = 1\ MHz\) in [19] and \(T = 50\ ns\) in [20]. The linewidth predicted by equation [20] for pulsed laser excitation scales much more favorably with the laser power. Since this reduced linewidth is only obtained after the excitation pulse has passed (as illustrated further in Fig. [3]), the significantly reduced power broadening presented in Fig. [2] can only be obtained by using a subsequently delayed ionization pulse. The considerable reduction of the resonance linewidth provides a strong argument in favor of using an ionization step that is delayed with respect to the pulsed excitation laser step, such that the narrower line shape is probed.

Figure 2: Comparison of equations (19) and (20), representing the line broadening \(\Delta\) due to the excitation coupling \(\Omega\), for an Einstein \(A\)-coefficient of 1 MHz for continuous-wave lasers and \(T=50\ ns\) for pulsed lasers.

Besides power broadening due to the excitation laser, the interaction of the system with the ionization laser, if applied during the excitation laser pulse, can also further broaden the level. This can be understood in an intuitive way. Since the ionization laser couples the excited state to the continuum, the lifetime of this excited state is reduced. By virtue of the Heisenberg energy-time uncertainty principle, this implies an increase in the energy uncertainty of the excited state. Indeed, the resonant excitation is probed by looking at the ions that are created by subsequent excitation of the atoms from the excited level towards the ionization continuum. The energy uncertainty induced in the intermediate level translates into a broadening of the resonance observed in the spectrum. If the ionization laser is delayed with respect to the excitation laser step, this broadening does not occur, since the perturbing ionizing laser field is not present when the resonant excitation happens.

Fig. [3] illustrates these observations by presenting nu-
Figure 3: Two simulations for a two-level atom, using the parameters in table I. For each of the figures, the central surface plot shows the excited state population as a function of laser detuning and time. The diagram to the left of these central plot schematically displays the laser pulse sequence. On the top of each figure the resonance ionization spectrum is shown.

(a) Excited state population when using simultaneous laser pulses (see diagram on the left). The resonance ionization spectrum (red curve in the top panel) is power broadened by both laser pulses.

(b) As Fig. 3a but with a delayed ionization pulse. For comparison, the red ionization spectrum in the uppermost plot is taken from the simulation in Fig. 3a. Delaying the ionization step removes the power broadening.

In Fig. 3a, it is shown that population is transferred to the excited state as the excitation laser builds up. When the ionization laser fires, the accumulated population is removed from the excited state. Upon comparing the ionization spectra shown at the top of figures 3a and 3b, it becomes apparent that delaying the ionization laser until the excitation laser has ended considerably reduces the linewidth of the final optical resonance obtained through the resonance ionization process. This is due to the transient nature of the population transfer outside of the narrow region governed by equation (20); only in that region will population remain in the excited state after the action of the excitation laser. The width of the resonance ionization signal will therefore be narrower when using a delayed ionization laser. Note also in Fig. 3a how the presence of the ionization laser leads to additional broadening of the excitation spectrum, an additional source of line broadening that is avoided by using a delayed ionization stage.

The goal of the experiments that are described below is to illustrate how power broadening can be mitigated by using a delayed ionization step. In this demonstration, the use of a weak transition to a long-lived excited state is crucial, since spontaneous decay from the excited state is minimal even with delayed ionization. This means that the resonance ionization process still occurs...
efficiently, which is critical for applications on exotic radioactive beams.

B. Experimental verification

1. Description of the experiment

$^{63,65}$Cu atoms have been laser-ionized using a two-step resonance ionization process depicted in Fig. [4] The experiment was performed at the JYFL laboratory in Jyväskylä, Finland. The resonant 244.237 nm line from the $3d^{10}4s^2 S_{1/2}$ ground-state to the $3d^94s4p^4 P_{1/2}^{o}$ state at 40943.78 cm$^{-1}$ was followed by a 441.679 nm transition to the auto-ionizing $3d^94s5s^4P_{1/2}^{o}$ state at 63584.65 cm$^{-1}$. Given the long lifetime of the excited bound state (479(28) ns [20]), this system is well suited to study the behavior of power broadening for pulsed lasers and the role of the delay of the ionization laser on the line shape and ionization efficiency. Furthermore, the laser system used to excite the transition had a narrow bandwidth ($\approx$ 20 MHz) and could deliver an order of magnitude more laser power than the required saturation power density, resulting in clear power broadening effects.

A description of the atomic beam unit used for this work is presented in [21], but the essential features of the device are repeated here. The copper atoms were produced by resistively heating a tantalum tube containing a sample of copper. The resulting atomic beam passed through a collimation slit and then orthogonally crossed the laser beams. Electrostatic ion optics extracted the laser ionized copper atoms from the interaction region and guided them to an electron multiplier tube which served as the particle detector.

The laser system used for this work is described in detail in [22]. For the 244.237 nm line, an injection-locked Ti:sapphire laser system produced narrowband laser light (bandwidth $\approx$ 20 MHz), which was then frequency tripled. The master laser for this seeding cavity was a CW Matisse Ti:sapphire laser, which can be scanned continuously. The fundamental output of the seeded laser was 2.8 W at 10 kHz repetition rate, which after beam transport losses resulted in 300 mW/cm$^2$ of tripled UV light entering into the atomic beam unit. A maximum of 1.6 W/cm$^2$ of 411.679 nm light for the ionization step was produced using an intra-cavity frequency doubled pulsed Ti:sapphire laser. The two lasers were pumped using different Nd:YAG lasers, which introduces a jitter in the timing synchronization of both Ti:sapphire lasers of about 10 ns. This time jitter was of no consequence for the experiment. The pulse length of both lasers is typically 50 ns. The wavelength of the injection seeded Ti:sapphire was recorded using a High Finesse WS6 wavemeter and further monitored with a Topica scanning Fabry Perot Interferometer FPI-100-0750-y with a free spectral range of 1 GHz. This interferometer was used to more precisely determine the wavelength of the laser as it was scanned. An example of a resonance ionization spectrum of $^{63,65}$Cu is shown in Fig. [5]

![Figure 5: Ionization spectrum of $^{63,65}$Cu. The black box indicates the peaks presented in the zoom-in in Fig. 6.](image)

2. Discussion of results

Ionization spectra of $^{63,65}$Cu were obtained at different UV laser powers and for several time delays of the ionization step with respect to the excitation step. The linewidth of the Gaussian component of the fitted Voigt profiles was found to vary between 40 and 60 MHz for all experimental conditions. The contributions to this Gaussian component from the remaining Doppler broadening and the laser linewidth could not be separated, but...
are likely of a similar magnitude. The measurement performed at the lowest UV laser power (3 mW/cm$^2$) and using a temporally overlapped excitation and ionization laser resulted in a Lorentzian component with a width of 53.8(4) MHz. Increasing the power of the laser to 150 mW/cm$^2$ increased the linewidth of the Lorentzian component to 124.3(3) MHz. The two right-most hyperfine transitions in the spectrum of $^{63,65}$Cu measured with this larger laser power are shown in Fig. 6. The spectrum in red is measured with the excitation and ionization lasers firing simultaneously, while the green spectrum was obtained by delaying the ionization laser 40(10) ns. Symbols are the experimental data points, the full line is the fit with a Voigt line shape. As can be clearly seen by comparing the two spectra, keeping the laser power fixed and delaying the ionization laser drastically reduces the width of the resonance lines.

Note that this reduced Lorentzian linewidth is much less than the 53.8(4) MHz linewidth obtained at the lowest laser power of 3 mW/cm$^2$ (with temporally overlapping laser pulses). This provides direct evidence for the absence of power broadening not only from the excitation laser but also from the ionization laser.

This sharp reduction in the experimental linewidth as the laser is delayed is further illustrated in Fig. 7. This figure shows the linewidth of the Lorentzian component as a function of the delay time of the second laser pulse. The red star corresponds to the red spectrum in Fig. 6, the green triangle corresponds to the green spectrum in Fig. 6. Errors are smaller than the symbols. Also shown is a theoretical calculation of the linewidth as function of the laser delay, using the experimental parameters described in the text.

Note finally that the efficiency loss due to spontaneous decay is negligible for delays below 50 ns due to the long lifetime of the excited state of 479(28) ns. This is the key advantage offered by using the weak transition to a long-lived state rather than a stronger transition.

IV. EXPERIMENTS ON LINESHAPE DISTORTIONS AND DELAYED IONIZATION

A. Model Predictions

In addition to the power broadening effects discussed in the previous section, there is another effect to consider: the possibility of lineshape distortions induced by a high-power ionization laser. This effect is illustrated in Fig. 8. This figure repeats the simulations presented earlier in Fig. 3 but this time includes a Stark shift induced by the ionization laser ($S$ in equation (3)). In Fig. 8a, a clear asymmetry can be seen in both the population of the excited state and the final ionization spectrum. By contrast, when using a delayed ionization laser, this asymmetry is naturally absent.
The next section will discuss some experimental data which demonstrate this kind of lineshape distortions. We will also show that the model for laser ionization introduced in section I can be used to qualitatively explain these distortions. Delaying the ionization laser in time with respect to the excitation laser removes the unwanted effects, since the distortion is induced after the atomic structure is already probed by the excitation laser.

**B. Experimental verification**

1. **Description of the experiment**

The possibility of ionization-related lineshape distortions, and how they can be removed by using a delayed ionization step, has been illustrated using the Collinear Resonance Ionization Spectroscopy (CRIS) experiment at ISOLDE-CERN [7, 23], using a radioactive beam of $^{221}$Fr. Details on the experimental set-up and measurement procedure can be found in [7]. The ionization scheme that was used is presented in Fig. 9 and consists of an excitation step from the $7s \, ^2S_{1/2}$ ground state to the $8p \, ^2P_{3/2}$ state at 23658.306 cm$^{-1}$ (422.685 nm), and an ionization step that non-resonantly ionizes from the $8p \, ^2P_{3/2}$ state using pulsed 1064 nm light. The lifetime of the excited atomic state is 83.5(1.5) ns, sufficiently long to justify the use of a delayed ionization pulse.

![Ionization scheme used to excite and ionize $^{221}$Fr.](image)

**Figure 9:** Ionization scheme used to excite and ionize $^{221}$Fr. The size of the hyperfine splittings was calculated using the values in [24].

A $^{221}$Fr ion beam was produced by the ISOLDE facility at CERN by impinging 1.4 GeV protons onto a uranium carbide target. Francium atoms diffuse out of this target and are then surface ionized in a hot capillary tube. After mass separation from the other francium isotopes, the $^{221}$Fr ion beam is then guided to a gas-filled linear Paul trap, where it is cooled and bunched. This beam is then
accelerated to 30 kV and transported towards the CRIS experiment. The first stage of the CRIS experiment consists of neutralizing the ion beam through collisionless charge exchange with a hot potassium vapor [25]. This is often required, since suitable transitions are usually easier to find for neutral atoms rather than ions. The non-neutralized fraction of the beam is electrostatically deflected into a beam dump while the neutralized fraction is temporarily and spatially overlapped with the laser beams in an ultra-high vacuum (UHV) interaction region. The laser frequency of the UV excitation laser is scanned across the hyperfine resonance transitions, and the resonantly ionized $^{221}\text{Fr}$ ions are then deflected onto a copper dynode. The secondary electrons emitted from the dynode are detected using a microchannel plate (MCP) electron detector. The UHV is required to minimize collisional ionization that would otherwise result in a constant background in the hyperfine spectra.

Because of the combination of an accelerated beam and the collinear overlap of the atom and laser beams, Doppler broadening is reduced to the point where it only contributes a few MHz to the total linewidth of the hyperfine structure spectra. The laser light for the first step was produced by frequency doubling the output of a Matisse TS cw Ti-sapphire laser with a Wavetrain external cavity frequency doubler. This continuous light was chopped into pulses of variable length through the use of a pockels cell and subsequent polarization sensitive beam optics, described in detail in [7]. This experimental configuration was used to create light pulses with a pulse length of 100 ns, at a repetition rate of 100 Hz. The 1064 nm light for non-resonant ionization was produced using a dual-cavity Litron LPY 601 50-100 PIV laser system, operated at 100 Hz and with a pulse length of 13 ns. After beam transport losses, 250 mW/cm$^2$ of continuous wave laser light and 32 mJ/pulse of 1064 nm laser light reached the entry of the CRIS beamline.

2. Discussion of results

Fig. 10 shows two measurements of the low-frequency component of the hyperfine structure of $^{221}\text{Fr}$. The red (broad) spectrum is obtained with the ionization laser temporally overlapped with the 100 ns wide excitation laser pulse, as illustrated in the inset of Fig. 10. The green spectrum was obtained with an ionization pulse delayed by 100 ns from the start of the excitation pulse. Using simultaneous laser pulses distorts the high-frequency side of the peaks, which displays a clear asymmetry. This asymmetry disappears when the ionization laser is delayed, which indicates that the tailing is induced by the ionization laser.

The figure also shows simulations using the model introduced in section [11]. The ionization cross section $\sigma$ was taken to be 1 Mb, which should at least be of the correct order of magnitude (see e.g. [20, 27] for cross sections in Rb and Cs). The Fano $q$ parameter was taken to be zero. The effective Stark shift $S$ was tuned to give the best match with the experimental data; a final value of $S(t) = 4\Gamma(t)$ provided good agreement. The simulations were also rescaled to match the intensity of the highest peak in the experimental data.

Using these parameters, the asymmetric tail of the peaks is well reproduced, supporting the idea that the observed asymmetry is due to a Stark shift caused by the strong electric field of the high-power ionization laser. The intensity of the smallest resonance in the spectrum is not well reproduced by the model. The reason for this discrepancy is unclear.

![Figure 10: Resonance ionization spectra of the leftmost components of the hyperfine structure of $^{221}\text{Fr}$, obtained with simultaneous laser pulses (red) and with a delayed ionization step (green). The solid lines are fits using the model for RIS presented in section [11]. The asymmetry of the line disappears when the ionization laser is delayed, while the total ionization efficiency does not decrease significantly.](image)

As with the data on the copper isotopes, delaying the ionization step does not result in significant loss in efficiency, since the excited state is long-lived. The linewidth of the resonance is 20(1) MHz. This linewidth could only be reached due to the removal of power broadening and the lineshape-distorting AC Stark shift by delaying the ionization laser. In a two-step resonance ionization scheme, this can only be done efficiently with a weak transition to a sufficiently long-lived excited state.

V. EFFICIENT LASER EXCITATION AND IONIZATION WITH WEAK TRANSITIONS

Weak transitions have shown desirable features for laser spectroscopy purposes. In addition to the inherently small linewidth, weak transitions show no sign of efficiency loss when delaying the ionization pulse. This section will further argue that weak transitions to long-lived
states can be excited with very high absolute efficiencies, comparable to efficiencies obtained with stronger lines. Applying the model of section II, one obtains the steady state population of an excited level in a two-level approximation as:

\[ P_{\text{exc}}(\Delta = 0) = \frac{\Omega^2}{A^2 + 2\Omega^2} \]

(21)

\[ \propto \frac{I/A}{1 + 2I/A}, \]

(22)

since \( \Omega \propto \sqrt{IA} \). Since, for a fixed laser intensity, the equilibrium population is a monotonically decreasing function of \( A \), weak transitions can achieve higher steady state population in the excited state. However, the irradiation time required to reach this steady state is longer than for strong transitions, though it also decreases with laser power. Therefore, there are two strategies to consider when maximizing the efficiency of excitations using weak transitions.

First of all, one can use high power pulsed laser systems which increase the rate at which the equilibrium population is reached, resulting in higher efficiency for short pulse lengths. This is the approach used for the first dataset in this article (see section III): a high-power pulsed laser can saturate the excitation step and therefore efficiently excite the system.

On the other hand, employing low power chopped CW laser light with long interaction times, as in the experiment of IV also allows for high efficiency (>1-10%). Indeed, the efficiency in the experiment described above using chopped CW laser light \( I_{\text{p}} \) is similar to that obtained in an earlier experiment using a pulsed high-power laser for the excitation step \( I_{\text{p}} \). In both experiments, the total efficiency was 1%, where the detection efficiency was 80%, beam transport efficiency <30%, neutralization efficiency <50%, and the laser ionization efficiency therefore >8%.

The ability to reach high laser ionization efficiencies with chopped CW laser pulses is also illustrated in Fig. 11. This figure shows simulated ionization efficiencies for weak and strong transitions for a system with a ground state doublet and a single excited state, as a function of the pulse length of the excitation step, and using overlapping excitation and ionization lasers. The laser power was set to 10 mW, which is usually easily achieved with modern CW lasers. With sufficiently long interaction times, weak transitions can be excited very efficiently, with simulated ionization efficiencies better than or comparable to the efficiency obtained for short pulses and strong transitions. For time-separated laser beams, the ionization efficiency for a strong transition never reaches that of the weak transition, due to decay losses.

VI. CONCLUSIONS

This article has shown that efficient resonance ionization spectroscopy with high resolving powers can be achieved with pulsed (or chopped) laser beams using weak transitions to long-lived states. This was demonstrated using simulations, supported by experimental observations and illustrates the twofold advantages that weak transition to long-lived states offer.

Firstly, it is possible to remove virtually all power broadening due to both lasers in a two-step RIS scheme by choosing a suitable delay between the excitation and the ionization laser pulses. Secondly, lineshape distortions due to the presence of a strong ionizing laser field when using non-resonant ionization can cause significant lineshape distortions, which can also be removed by delaying the ionization step. Similar arguments can be made in the case of RIS schemes that use more than two lasers, but this lies outside of the scope of the work presented here.

The long lifetime of the excited state ensures that no significant efficiency losses occur due to the delay of the ionization laser. Experimental evidence for both advantages was presented and compared to simulations with a theoretical model for resonance ionization spectroscopy. The experimental data also illustrate that high efficiencies can be obtained using weak transitions.

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