3smd\(^1\)D to 3png\(^1\)F autoionization resonances of Mg

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Intense ultranarrow resonances have been observed in the photoionization of Mg from the singly excited 3smd\(^1\)D \((m=3,4,5,6)\) states to the doubly excited 3png\(^1\)F \((n=5,6,7)\) autoionization states using stepwise laser excitation and time-of-flight mass spectrometry detection. The reported experimental energy resolution at an order of \(10^{-7}\) of the photon energy represents a considerable improvement over the best photon energy resolution of about one part in \(10^5\) for the synchrotron radiation light sources. The ultra-narrow 3png\(^1\)F resonances acquire a significant strength for the normally weak 3smd\(^1\)D\(\rightarrow\)3png\(^1\)F two-electron transition due to a strong configuration interaction with an overlapping broad 3pnd\(^1\)F autoionization series and the 3s\(\sigma\)F\(^1\)F continuum. The measured line positions and transition widths are in close agreement with the theoretical results derived from a B-spline-based configuration-interaction calculation.

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I. INTRODUCTION

The spectra of the alkaline-earth atoms immediately above the first ionization threshold are dominated by resonances resulting from the strongly energy-dependent doubly excited ionization states. The photoabsorption/photoionization spectra of these divalent systems are characterized typically by two asymmetric autoionizing series, one broad and one narrow in width, caused by the simultaneous change of electronic orbitals of the two outer electrons in a double-excitation process. Theoretically, the spectrum between the first and second ionization thresholds of Mg is one of the most often studied doubly excited resonant structures for a lighter alkaline-earth atom [1–5]. In addition to photoionization from the Mg ground state, calculations have also been extended to photoionization from the bound excited states of Mg [1,2] and Mg-like ions [5]. These calculations quantify the photoionization of these species from their bound excited states. They are a part of a concerted effort to provide precise physical interpretation of multielectron interaction in many-electron systems. One of the most interesting physical features suggested by these studies is the interference between the overlapping 3pnd\(^1\)F and 3png\(^1\)F autoionization series. Chang and his co-workers, using a B-spline-based configuration interaction (BSCI) approach [2,5,6], showed that the strong 3smd\(^1\)D to 3pnd\(^1\)F transitions are dominated by an isolated-core excitation (or shake-up) of the outer md electron following a bound-bound 3s to 3p excitation [3]. Their calculations also showed the presence of the extremely narrow 3png\(^1\)F \((n=5,6,7)\) resonances located on top of broad 3pnd\(^1\)F resonances. These features result primarily from the energy variation of the bound component of the final-state wave function in the vicinity of the 3png\(^1\)F resonance. The effect due to transition from the initial bound state to the continuum component of the final-state wave function is relatively small. The width of the 3png\(^1\)F resonances is expected to be similar to the He 2pnd\(^1\)P resonances as both can be estimated qualitatively from a \(\Delta/\gamma = +1\) Coulomb interaction, i.e., interaction between 1s\(\sigma\)p and 2pnd\(^1\)P for He and 3s\(\sigma\)f and 3png\(^1\)F for Mg. The calculated width of Mg 3png\(^1\)F at an order of \(5\times 10^{-8}\) Ry [2–4] is consistent with the estimated width of the He 2pnd\(^1\)P resonances at an order of \(10^{-8}\) Ry (\(\sim 33\) MHz) [7].

Gallagher and his co-workers observed experimentally the 3pns and 3pnd doubly excited autoionizing Rydberg series of Mg using an isolated-core excitation method with stepwise laser excitation [8,9]. Their studies provided much insight into the n dependence of Mg 3pnd \((J=1,3)\) high Rydberg autoionization series converging to the \(2P_{3/2}\) or \(2P_{1/2}\) limit of Mg\(^+\). The lasers used in their experiments had an energy resolution of \(\sim 1.0\) cm\(^{-1}\). Consequently, they have limited their study to the broad 3pnd\(^1\)F series, for \(n=9\) to 40. Complementing their early works, in this paper, we report in detail a multistep multicolor laser study in Mg for transitions from the singly excited 3smd\(^1\)D \((m=3–6)\) states to the ultranarrow doubly excited 3png\(^1\)F \((n=5,6,7)\) autoionization states [10]. These results provide a precise quantitative comparison of experiments and theoretical calculations for the narrow 3png\(^1\)F series.

II. EXPERIMENTAL APPROACH

The experimental approach follows that initially employed by Cooke et al. in what they describe as an isolated-core excitation scheme [11]. One of the two valence electrons in the \(3s^2\) I\(^1\)S ground electronic state of Mg is excited to a 3smd\(^1\)D state by a two-step sequential absorption process. A single-longitudinal-mode pulsed laser \((I_1)\) is tuned to the \(3s^2\) I\(^1\)S to 3s3p\(^1\)P resonance and excites the atom to the 3s3p\(^1\)P state. After a fixed time delay, a pulse from a second laser \((I_2)\) transfers a portion of this population from the 3s3p\(^1\)P state to a 3smd\(^1\)D state. A third pulsed laser \((I_3)\) then photoionizes the atom from the \(1D\) state. The laser \(I_3\) is scanned to probe for autoionization resonances resulting from the 3smd\(^1\)D to 3sng\(^1\)F transitions. Figure 1 illustrates the relevant energy states of Mg and the three-excitation steps used.
The experimental apparatus is comprised of three components: a laser system, a vacuum chamber consisting of a time-of-flight (TOF) mass spectrometer connected to a Mg atomizing oven, and a data-acquisition system. The schematic diagram of the system is shown in Fig. 2.

The laser system consists of three lasers. The first laser \( I_1 \) is a CW single-frequency ring dye laser (Coherent model 699-29) that is pulse-amplified through a multistage dye amplifier pumped by a Nd:YAG laser [12]. The output energy is about 10 mJ and is 5 ns in duration. This output is frequency doubled in a KDP crystal to generate the 285.3 nm radiation needed to excite the \( 3s^2 \, ^1S \) to \( 3s3p \, ^1P \) resonance of neutral Mg. The second laser \( I_2 \) is a grating-tuned dye laser (Lumonics model HD-300) pumped by the same Nd:YAG laser. \( I_2 \) is used to generate pulses at the wavelengths needed to excite Mg from \( 3s3p \, ^1P \) to the \( 3smd \, ^1D \) states. The value of these wavelengths is calculated using published data of the Mg energy levels that are accurate to 0.01 cm\(^{-1} \) [13]. The third laser \( I_3 \) is a CW Ti:sapphire single-frequency ring laser (Coherent model 899-29) that is amplified through a dye cell-Ti:sapphire crystal amplifier arrangement [14]. In order to maintain the highest resolution for the output, we removed the phase-conjugate mirror in the laser system described in Ref. [14] and used only four passes in the Ti:sapphire amplifier. The resulting pulse is about 10 ns long and has an energy of 5 mJ. This output is frequency-tripled or quadrupled by harmonic generation crystals to generate a tunable UV output of \( \approx 50 \) \( \mu \)J per pulse for the experiment. The CW laser wavelength is calibrated using the laser’s built-in wavemeter and an external wavemeter to an accuracy of 0.01 cm\(^{-1} \). All three lasers are linearly polarized.

An effusive beam of Mg atoms is produced from a resistive oven that is isolated from the main chamber and maintained at a temperature of 450°C. A diffusion pump in the main chamber and a turbo molecular pump in the TOF mass spectrometer maintain a background vacuum of less than \( 10^{-6} \) Torr in the vacuum system. The three lasers are each collimated to \( \approx 5 \) mm in diameter. The center portion of each beam then is passed through an iris aperture with a diameter of 1 mm. This ensures a uniform (±5%) intensity profile so that the excitation of the atoms in the overlapped region has a uniform spatial and concentration distribution in this multiple photon experiment. The lasers \( I_1 \) and \( I_2 \) are copropagating while \( I_3 \) is counterpropagating into the experimental chamber. The three beams overlap and cross the atomic beam at right angle at the entrance to the mass spectrometer. Ion extraction is done using two electric field plates separated by 11.5 mm. A +150 V extraction voltage pulse is applied to the front plate after the Mg atoms are photoionized to drive the ions through a 4-mm-diameter opening in the rear plate towards the TOF mass spectrometer. The flight path is 1 meter long. This distance is sufficient to distinguish easily the three stable isotopes of Mg. A microchannel plate detects the resulting photoions. Only the signal from the ions of the dominant isotope (\(^{24}\)Mg) is recorded.

The pulse from \( I_2 \) is delayed by 12 ns from \( I_1 \). \( I_3 \) is further delayed by 50 ns from \( I_2 \). These delays are necessary to ensure that there is no unwanted optical interference between the three laser pulses and no quantum interference between these pulses with the energy states of the atom [15]. The value of these delays is determined by experiments designed to achieve the best frequency resolution in the ionization spectrum and the least background ion interference on signal detection. Prior to the recording of each high-resolution spectrum, the intensities of \( I_1 \) and \( I_2 \) are adjusted using neutral density filters as described in the next section until \( I_1 \) and \( I_2 \) do not produce any one-color or two-color ions and there is no ion pulse broadening due to Coulomb expulsion of the ions produced by \( I_3 \). Under these conditions, the pulse energies of \( I_1 \) and \( I_2 \) are well below the detectivity of our pyroelectric energy meter of 1 \( \mu \)J per pulse. \( I_3 \) is operated at the full pulse energy of 50 \( \mu \)J. A delay generator is used to set the timing and to trigger the two Nd:YAG lasers at 30 Hz. The signal from the microchannel plate is amplified and fed into a boxcar integrator. Finally the data are stored in a personal computer, which is used to manage all phases of the experiment.

**III. RESULTS AND DISCUSSION**

The first step in the present experiment is to determine the spectral resolution. The laser \( I_1 \) excites ground state Mg to
the 3s3p 1P state. Population in the 3s3p 1P state is then ionized by resonance-enhanced 1+1 ionization via the 3s8d 1D state by the laser \( I_3 \). The intensities of \( I_1 \) and \( I_3 \) are attenuated and the time delay between \( I_1 \) and \( I_3 \) is adjusted until the width of the intermediate resonant state in the scanned spectrum will not reduce upon further reduction of the laser intensities. The theoretical width of the 3s3p 1P to 3s8d 1D transition is given by the sum of the inverse of the lifetimes of the initial and the final energy states. The lifetime of 3s3p 1P is well-documented and is equal to 2 ns [16]. The lifetime of the 3s8d 1D state is about 94 ns [17]. For excitation using linearly polarized light, the only decay from the 3s3p 1P level in a collision-free environment is via fluorescence to the 3s2 1S ground state. The theoretical \( ^1P \) to \( ^1D \) transition width \( \Gamma \) is therefore primarily determined by the \( ^1P \) to \( ^1S \) decay and can be estimated from these lifetimes to be 81 MHz. Figure 3 shows the ion signal recorded as the probe laser is scanned in wavelength. This spectral line can be fitted very well to a Lorentzian profile with a spectral full width at half maximum (FWHM) of 107 ± 2 MHz. Since the spectral profile of the Fourier-transform-limited laser is well represented by a Gaussian distribution, a deconvolution procedure detailed in the Appendix gives a spectral resolution \( \Omega \) of the present experiment at 84 MHz (FWHM).

Our experimental \( ^1D \) to \( ^1F \) work was focused on the transitions from initial states of \( m=3 \) to 6 to final states of \( n=5 \) to 7. There are a total of 12 possible transitions. The transition frequency and the peak photoionization cross section of these transitions obtained from a BSCI calculation are listed in Table I. A search was performed for eight of these transitions in consideration of the laser wavelengths that were readily obtainable. In order to assist each initial search, the intensity of \( I_1 \) was increased by a factor of 10 from that established in the spectral resolution determination experiment, and \( I_3 \) was scanned for a few hundred wavenumbers to the right and to the left of the theoretically predicted spectral position of the transition. From theory, it is seen that the absorption cross section of the 3smd 1D to 3png \( ^1F \) transition is 10–100 times stronger than the corresponding 3smd 1D to 3pnd \( ^1F \) transitions. It is thus expected that the scans would yield a weak but broad baseline feature resulting from ion signal produced due to transition to the broad 3pnd state and background ions, and a sharp narrow peak that corresponds to transition to the 3png \( ^1F \) state. This is the criterion used to identify the existence of the 3png state. Indeed, in every successful spectrum, only one clearly identifiable intense but narrow peak appeared for the entire width of the scan. When a peak was observed, the intensity of \( I_1 \) was reduced to the predetermined levels, and a power dependence on \( I_3 \) of the peak strength and the peak width was measured to ensure that the signal was linear to \( I_3 \) and that the width was constant of \( I_3 \). Typically, the highest number of ions detected at the peak wavelength was about 50 per pulse. Altogether, a total of five resonances were observed. Figure 4 is a representative of the observed spectrum near the region of the sharp resonance. The resonance shown is that of the 3s4d 1D to 3p5g \( ^1F \) transition. From the spectrum, the position of the peak of the 3p5g \( ^1F \) state can be determined precisely using the wavelength of the laser and the known position of the lower state of the transition. The width of the transition is determined by fitting the measured resonance line to a Lorentzian profile. The assumption of a

\[
\Delta \pm 24638 \text{ cm}^{-1}
\]

FIG. 3. High-resolution resonance-enhanced 1+1 photoionization spectrum of the 3s3p 1P state in the vicinity of the 3s8d 1D intermediate state. The solid curve is the observed data. The dotted curve is a Lorentzian fit. The laser (\( I_3 \)) pulse frequency \( \nu = \Delta + 24638 \text{ cm}^{-1} \).

\[
\nu = \Delta + 38776 \text{ cm}^{-1}
\]

FIG. 4. Spectra of the 3s4d 1D to 3p5g \( ^1F \) autoionization transition. The solid curve is recorded ion signal from the frequency scan of laser \( I_3 \). The dotted curve is a theoretical BSCI calculation of the autoionization cross section that has been convoluted with the experiment's spectral resolution. The two curves are displayed by matching the peak height and the position of the resonance. The laser (\( I_3 \)) pulse frequency \( \nu = \Delta + 39776 \text{ cm}^{-1} \). The experimental spectral resolution \( \Omega \) of 84 MHz (i.e., 0.0028 cm\(^{-1}\)) is indicated by the two vertical bars shown.

<table>
<thead>
<tr>
<th>Initial state</th>
<th>( n=5 )</th>
<th>( n=6 )</th>
<th>( n=7 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( m=3 )</td>
<td>( E )</td>
<td>( \sigma )</td>
<td>( E )</td>
</tr>
<tr>
<td>3</td>
<td>46660.1</td>
<td>1.55</td>
<td>47983.7</td>
</tr>
<tr>
<td>4</td>
<td>39928.5</td>
<td>8.85</td>
<td>41252.1</td>
</tr>
<tr>
<td>5</td>
<td>36754.8</td>
<td>0.50</td>
<td>38078.4</td>
</tr>
<tr>
<td>6</td>
<td>35039.9</td>
<td>5.78</td>
<td>36363.5</td>
</tr>
</tbody>
</table>

TABLE I. Calculated transition energy \( E \) (cm\(^{-1}\)) and photoionization cross sections \( \sigma \) (10\(^3\) Mb) for the 3smd 1D (\( m=3,4,5,6 \)) states to the 3png \( ^1F \) (\( n=5,6,7 \)) autoionization states of Mg.
Lorentzian line shape in place of a Fano line shape is justifiable since the observed spectral line shows a high degree of symmetry. Also displayed in Fig. 4 is the result of a BSCI calculation of the ionization cross section from $3s4d^1D$ to the same region of the spectrum. For comparison purposes, the calculated peak of the ionization cross section is matched to the position and the height of the ion signal peak. The figure shows that the measured line profile and the linewidth are in very good agreement with the calculated results.

The spectra of the four other observed transitions are shown in Fig. 5. The measured transition frequencies and the experimental widths and the corresponding quantities obtained from the various calculations are shown in Table II. As is shown in the table, the measured line positions are generally in close agreement with the calculated values. However, the experimental values are uniformly to the red of the calculated values. The absolute difference is small, of the order of 100 cm$^{-1}$ to 150 cm$^{-1}$, and is only about 0.2% of the total energy of the observed state. This difference in value is within the expected accuracy of the BSCI calculation that results from the use of a model potential. More work will need to be performed to determine if the persistent difference to the red is incidental or systematic. Since the low-$n$ states included in our study are well described by LS coupling as pointed out by Gallagher [8], the quantum defect is fairly small and the effective principal quantum number $n^*$ of these low-$n$ states is close to integer as expected (see, e.g., Table I of Ref. [2]).

The experimental linewidths $\Gamma$, derived from Eq. (A5) of the Appendix with a measured width $\Gamma_c$ at an energy resolution $\Omega$ determined earlier, are also listed in Table II. For the $3p5g^1F$ and $3p6g^1F$ states, they are $(281 + 169)/2 = 225 \pm 50$ MHz and $344\pm35$ MHz, respectively, in substantial agreement with the BSCI results, i.e., 249 MHz and 318 MHz, respectively. For the $3p7g^1F$ level, the experimental value is larger by about 30% than the calculated value, still in acceptable agreement with theory. Although great care was exercised during the experiments to eliminate the effect of various broadening mechanisms such as collisions, saturation, and Coulomb expulsion, it is not surprising that small contributions from these mechanisms remain, giving a larger experimental value than the actual value. From these linewidths, a total lifetime of about 0.5 ns is obtained for the $3p5g^1F$ states. This is quite long for energy states situated 4 eV above the ionization threshold. The radiative lifetime of the $3p^1g$ to the $3sng^1G$ transitions is expected to be similar to the $3s3p^1P$ to $3s^21S$ decay (i.e., about 2 ns). Hence, radiative damping [18], if it is included in the present theoretical estimate, could reduce the calculated ionization cross section significantly. The fluorescence decay from the narrow $3p^1g$ transitions could possibly be observed. However, experimentally, it is a lot easier to observe the equivalent radiative fluorescence from the $3s3p^1P$ resonances, since the $2pnd^1P$ radiative decay (i.e., the strong 21.2 eV He resonance line) is expected to occur with a lifetime between 0.5 and 0.6 ns [19].

<table>
<thead>
<tr>
<th>Transition</th>
<th>Final-state position $(\text{cm}^{-1})$</th>
<th>Linewidth $\Gamma$ (MHz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$3s3d^1D$ $\rightarrow$ $3p5g^1F$</td>
<td>93063.22 $^a$</td>
<td>249</td>
</tr>
<tr>
<td>$3s3d^1D$ $\rightarrow$ $3p6g^1F$</td>
<td>93053.26 $^b$</td>
<td>1296</td>
</tr>
<tr>
<td>$3s4d^1D$ $\rightarrow$ $3p5g^1F$</td>
<td>93074.66 $^c$</td>
<td>211</td>
</tr>
<tr>
<td>$3s4d^1D$ $\rightarrow$ $3p6g^1F$</td>
<td>93211.57(3) $^d$</td>
<td>281(30)</td>
</tr>
<tr>
<td>$3s5d^1D$ $\rightarrow$ $3p5g^1F$</td>
<td>93063.22 $^a$</td>
<td>249</td>
</tr>
<tr>
<td>$3s5d^1D$ $\rightarrow$ $3p6g^1F$</td>
<td>93053.26 $^b$</td>
<td>1296</td>
</tr>
<tr>
<td>$3s6d^1D$ $\rightarrow$ $3p5g^1F$</td>
<td>93074.66 $^c$</td>
<td>211</td>
</tr>
<tr>
<td>$3s6d^1D$ $\rightarrow$ $3p6g^1F$</td>
<td>93291.60(3) $^d$</td>
<td>169(18)</td>
</tr>
</tbody>
</table>

$^a$BSCI calculation, Ref. [2].
$^b$L2-based CI calculation, Ref. [3].
$^c$B-spline-based stabilization method, Ref. [4].
$^d$Present experiment, numbers inside the brackets are uncertainties of the most significant digit(s). The uncertainty in line position is cumulative uncertainty of the position of the intermediate state and of the wavelength of laser $I_1$. Uncertainty of the linewidth, resulting from the fitting of the noisy ion signal amplitude and the background, is estimated at 10% of the fitted width.
It is clear in comparing the theoretical curves with the measured curves in Figs. 4 and 5 that there is a large off-resonance contribution in the measured signal not predicted by theory, which includes only the far wings of a single $3 s m d \rightarrow 3 p n d$ transition. In the experiment, the observed ion signal is the total number of ions produced by the probe laser. In addition to producing ions from the $3 s m d$ state, the photon energy of the probe laser $I_3$ is sufficient to ionize the Mg atom from every other excited state. Because of the time delay imposed by the experimental conditions, several states are populated due to radiative decay from the initial $3 s m d^1D$ state. For example, the states $3s3p^1P$, $3s4p^1P$, and $3s4s^1S$ are energetically below the initial $3s4d^1D$ state and are likely to be populated before $I_3$ arrives. Ionization from these states and from any residual population in the $3s3p^1P$ state not excited by $I_2$ is due to the featureless continuum and is likely the major contributor to the larger than predicted observed baseline ion signal. We note that in the most adverse case the strength of this baseline signal is as big as 10% of the on-resonance ion signal. This residual ion signal has made it difficult to obtain a better analysis of the line shape of these transitions.

The close agreement between experimental and theoretical results in this study supports the physical interpretation of the autoionization of the doubly excited $3p n d^1F$ and $3p n g^1F$ states manifested by the BSCI calculation. Photoionization from the $3s m d^1D$ states is dominated by the shake-up of the outer $d$ electron following the $3s\cdot3p$ transition. This shake-up process contributes substantially to the oscillator strengths for transitions from $3s m d^1D$ states to a $3p n d^1F$ broad resonance and a $3p n g^1F$ narrow resonance. The experimental results reported here, however, are not sufficient to determine when the LS coupling employed in the BSCI calculation begins to take on the $jj$ character as $n$ increases. Further theoretical work in conjunction with these precise measurements may provide new insight into the transition from the LS to $jj$ character for the high angular momentum autoionization series. To summarize, the multistep multicolor laser ionization, with its ultrahigh energy resolution, offers a powerful approach to study ultranarrow resonances in the VUV region not possible even with the most advanced synchrotron radiation light sources.

**ACKNOWLEDGMENTS**

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**APPENDIX**

In general, an isolated spectral line $L(E)$ can be represented by a Lorentzian profile,

$$L(E) = \frac{C}{\pi} \frac{\left(\frac{1}{2}\Gamma\right)}{(E - E_0)^2 + \left(\frac{1}{2}\Gamma\right)^2} \quad (A1)$$

centered at an energy $E_0$ with a width $\Gamma$, where $C$ is an appropriate constant. If we assume that its observed spectrum $L'(E; \Omega)$ can also be approximated by a Lorentzian profile,

$$L'(E; \Omega) = \frac{C}{\pi} \frac{\left(\frac{1}{2}\Gamma_c\right)}{(E - E_0)^2 + \left(\frac{1}{2}\Gamma_c\right)^2} \quad (A2)$$

at the same $E_0$ with a measured width $\Gamma_c$, then $L'(E; \Omega)$ is a convoluted spectrum of $L(E)$ using a laser spectral function in terms of a Gaussian distribution $G$, i.e.,

$$L'(E; \Omega) = \int_{-\infty}^{+\infty} L(E') G(E' - E; \Omega) dE', \quad (A3)$$

where

$$G(E; \Omega) = \frac{e^{-E^2/\delta^2}}{\sqrt{\pi} \delta^2}. \quad (A4)$$

The energy resolution $\Omega$ is represented by the full width at half maximum (FWHM) of the distribution function $G$ and $\delta = \Omega/(2\sqrt{\ln 2})$.

Similar to the deconvolution procedure given in [20], by projecting on both sides of Eq. (A3) to the ground-state eigenfunction of a harmonic oscillator, i.e., $\Psi_0 = e^{-\rho^2/\pi}^{1/4}$, where $\rho = \sqrt{2(E - E_0)/\Omega}$, a straightforward integration will lead to a simple relation

$$F_v(u)e^{v^2} = \sqrt{2} F_c(u)e^{u^2}, \quad (A5)$$

where $v = \sqrt{\ln 2}(\Gamma/\Omega)$, $u = \sqrt{\ln 2}(\Gamma_c/\Omega)$, and $F_c(x) = 1 - (2/\sqrt{\pi}) e^{-y^2}dy$ is the complementary error function. The energy resolution $\Omega$ can be readily obtained from Eq. (A5) when the width $\Gamma$ and the measured width $\Gamma_c$ are known.


