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Authors
Yip, FL
Rescigno, TN
McCurdy, CW

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Fully differential single photon double photoionization of atomic magnesium

F. L. Yip,1 T. N. Rescigno,2 and C. W. McCurdy2,3

1Department of Science and Mathematics, California State University-Maritime Academy, Vallejo, CA 94590, USA
2Lawrence Berkeley National Laboratory, Chemical Sciences, Berkeley, California 94720, USA
3Department of Chemistry, University of California, Davis, CA 95616 USA

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The valence-shell double ionization of atomic magnesium is calculated using a grid-based representation of the 3s2 electron configuration in the presence of a fully-occupied frozen-core configuration of the remaining ten electrons. Atomic orbitals are constructed from an underlying finite element discrete variable representation (FEM-DVR) that facilitate accurate representation of the interaction between the inner shell electrons with those entering the continuum. Length and velocity gauge results are compared with recent theoretical calculations and experimental measurements for the total, single and triple differential cross sections, particularly at the photon energy of 55.49 eV for the latter. Comparison between the similar processes of double ionization of the ns2 atoms helium, beryllium and magnesium further illuminates the role of valence-shell electron correlation and in atomic targets with helium-like electronic configurations and symmetry.

I. INTRODUCTION

The process of double photoionization (DPI) from an atom or molecule remains one of the most sensitive probes of electron correlation and has been studied for many years to investigate the consequences of electron correlation in simple targets. The mechanism of removing two electrons via a single photoabsorption necessarily requires a correlated target, and the resulting angular distributions and possible energy sharings of the outgoing electrons from the Coulomb breakup problem requires an accurate non-perturbative treatment for theory and coincidence measurements of the fragments for experiment.

The prototypical system to study atomic double photoionization is helium and good agreement between experiment and theoretical formulations has elucidated this simplest system [1, 2]. Recent work has sought to progress double ionization investigations by examining helium-like systems, with several theoretical calculations approximating atomic targets that have ns2 helium-like configurations for removal by one photon. The alkaline earth metals represent such systems for comparison and numerous theories have been dedicated to studying these quasi-two electron targets due to the large energetic and spatial separation between the core electrons and those in the valence shell that are removed by the photoabsorption [3–12]. By extending these investigations to other targets that parallel the single photon transition of DPI from helium in the initial and final states [13], the nature of the electron correlation between the outgoing electrons has been further elucidated as to consequences of the initially bound target orbitals [14, 15] and the relative strength of the correlation energy compared to the nuclear attraction of the residual fragment left behind when two electrons are photoejected [16].

Most recently, theory and experiment have examined DPI from the valence shell of atomic magnesium, which has a neon-like core interacting with the 3s2 outer electrons. Experimental measurements [17, 18] of the triple differential cross section (TDCS), which measures the angular distributions of both electrons and their energy sharing above the double ionization threshold have been recently compared to theoretical calculations using the convergent close coupling (CCC) [18] and time-dependent close coupling (TDCC) [19] methods at a photon energy of 55.49 eV. This energy is resonant with promotion of a 2p → 3d electron embedded in the double continuum. A formulation that can fully take into account the dynamics of this resonant process currently lies beyond any theoretical treatment for examination of quasi-two electron targets. Nevertheless, there have been comparisons of the experimental results undertaken with theoretical descriptions at this photon energy approximated by non-resonant ab initio calculations that feature only two-active electrons [18, 19] or by a semi-empirical accounting of the Fano profile of the resonant transition [18]. Stimulated by these studies, we present here an examination of the TDCS for double photoionization of magnesium using exterior complex scaling (ECS) examine the three-body Coulomb breakup dynamics at this energy and compare with other theoretical calculations at 55.49 eV and for the total double ionization process in this atom, for which other experimental measurements exist [20].

The formalism we employ to study this process requires denoting the neon-like core occupancy of all but the valence electrons, which provide a closed-shell Coulomb and exchange interaction with the outer 3s2 electrons that feel the action of the photon towards the double continuum. The construction of atomic orbitals out of an underlying radial grid to represent these core electrons facilitates the approximation of holding them fixed in a configuration-interaction expansion and has been applied to atomic beryllium in both a time-independent formalism for one-photon double ionization [8, 9, 13], and in a time-dependent framework [11, 21] for consideration of two-photon processes that remove the outer electrons. This method has also been applied to examine the double photoionization process for neon [22], which unlike the systems considered here, leaves behind an open-shell
target with distinct final state couplings for the residual dication. For the second-column ns$^2$ targets, however, only a single final-state channel remains as the resulting ionic fragment has a neon-like core configuration.

Section II overviews the description of the magnesium target in a combined orbital-grid basis for resolution of the double photoionization amplitudes that describe the fully differential cross sections in photoionization. Results are presented in Section III and compared with experimental measurements and theoretical calculations, with particular focus on the double ionization process at 55.49 eV. We also examine the similarities between helium, beryllium and magnesium in the angular distributions of double ionization at 20 eV excess photon energy to highlight the nature of the photoionization process. Conclusions are presented in Section IV.

II. THEORETICAL FRAMEWORK

The methods utilized in this work for treating photoionization of two electrons from a target with additional electrons has been previously described in greater detail in the both a time-independent framework, [8, 9] and with a time-dependent treatment involving ultrashort laser pulses [11, 21]. Thus, here we provide a brief overview of the most important points in applying the method to magnesium. Atomic units are used throughout the following, unless otherwise stated.

The two-active electron approximation to magnesium relies on a frozen-core approximation for the remaining 10 electrons. The wave function in can be expanded in configurations with $N - 2$ core electrons held fixed in atomic orbitals throughout. Within this frozen-core approximation, the problem then can be regarded as an effective two-electron problem involving a full CI of the 3$s^2$ valence electrons of magnesium in the presence of the 1$s^2$2$s^2$2$p^6$ core, the full Hamiltonian being

$$H = h(1) + h(2) + \frac{1}{r_{12}}$$

(1)

where the correlation between the valence electrons to be ejected by the photoabsorption is represented by $1/r_{12}$ and the interaction of the remaining electrons on the valence shell is represented by the one-body operator $h$,

$$h = -\frac{Z}{r} + \sum_{\text{occ}} [2J_{\text{occ}} - K_{\text{occ}}],$$

(2)

where $T$ is the one-electron kinetic energy, $-Z/r$ with $Z = 12$ represents the attraction of the magnesium nucleus, and the terms in the sum over occupied orbitals, $2J_{\text{occ}}$ and $K_{\text{occ}}$, account for the direct and exchange interaction felt by the 3$s^3$ valence electrons of the fully-occupied core orbitals. Specifically, the direct operator for the $nl$ closed-shell orbital is given by

$$J_{nl}(r) = \int \frac{\varphi_{nl}(r') \varphi_{nl}(r')}{|r - r'|} dr',$$

(3)

while the non-local exchange component is defined according to its operation on the orbital $\chi(r)$ is

$$K_{nl}(r) = \varphi_{nl}(r) \int \frac{\varphi_{nl}(r') \chi(r')}{|r - r'|} dr'.$$  

(4)

The $nl$ closed-shell orbitals that provide the Coulombic screening and non-local exchange are taken here to be the 1$s$, 2$s$ and 2$p$ Hartree-Fock orbitals of neutral magnesium. They are plotted as solid lines on the left half of the schematic shown in Figure 1. The double ionization potential of the valence shell for the target Mg atom is given by the ground state energy of the valence electron Hamiltonian in Eq. 1, with the energy of the neon-like fixed-core common to both the bound and continuum states.

In order to construct these operators and describe the bound, single and double continuum representation on a radial grid, we have utilized an adapted atomic orbital discrete variable representation with finite elements (orbital FEM-DVR) that we briefly summarize here; significantly more detail is provided in Refs. [8, 11]. The transformation of an ordinary finite element DVR [23] suitable for describing the radial coordinates of a simple atomic target (e.g., helium) to account for core electrons is referenced to the occupied orbitals of the multi-electron atom. Over a suitably defined orbital region where the core electrons are significantly non-zero, the first few finite element regions are transformed to form an orthogonal complement to the radial orbitals of the core electrons. The transformation from a primitive FEM-DVR basis $\chi_j(r)$ with $M$ functions spanning the orbital region to an orthogonal set of atomic orbitals is represented by

$$\varphi_{\alpha}(r) = \sum_{j=1}^{M} U_{\alpha j} \chi_j(r),$$

(5)

where within the orbital region the atomic orbitals $\varphi_{\alpha}(r)$ built from the underlying FEM-DVR functions describe the coordinates of all electrons and facilitate the construction of appropriate atomic CI configurations with both active and frozen electrons upon which to expand the description of bound, single and double continuum states of the target.

Beyond the region of the atomic orbitals, the underlying FEM-DVR description is untransformed, providing a flexible grid-based representation of the electronic coordinates that has been used in a variety of applications involving continuum electron dynamics [24]. The regions of radial space partitioned in this manner are shown schematically in Figure 1. The Hartree-Fock orbitals of the core electrons that provide a reference for the transformation operator $U_{\alpha j}$ in Eq. 5 over the entire orbital region are shown as solid lines, along with the 3$s$ orbital of Mg, shown as a dashed line on the left of Fig. 1, that provides the first of the orthonormal atomic orbitals that can be populated in the expansion configurations. Fig. 1 also highlights another key feature of this...
With the orbital-DVR radial basis, we describe wave function for the 3s\(^2\) valence electrons of magnesium as

\[ \Psi(r_1, r_2) = \sum_{l_1l_2} \frac{1}{r_{12}} \psi_{l_1l_2}(r_1, r_2) Y_{l_1l_2}^M \hat{r}_{12}, \]  

where \( Y_{l_1l_2}^M \) is a coupled-spherical harmonic describing the angular degrees of freedom and the atomic symmetry permits the decomposition into partial-wave radial wave function \( \psi_{l_1l_2}(r_1, r_2) \).

The double ionization amplitudes that describe the ionization processes ejecting two electrons into the continuum are given by solving a driven Schrödinger equation

\[ (E - H) \Psi_+^E(r_1, r_2) = (\epsilon \cdot \mu) \Psi_0(r_1, r_2), \]

where \( E = E_0 + \omega \) is the total energy available to share by the electrons above the double ionization potential \( E_0 \), \( \omega \) is the photon energy, and the driving term on the right of Eq. 9 represents the action of the photon with linear polarization \( \epsilon \) onto the 3s\(^2\) valence state of magnesium \( \Psi_0 \) in the dipole approximation. The scattered wave solution \( \Psi_+^E(r_1, r_2) \) determined by solving this equation on a grid with exterior complex scaling imposing the outgoing wave boundary conditions contains all continuum processes at energy \( E \).

To isolate the double continuum amplitudes for electrons with momenta \( k_1 \) and \( k_2 \),

\[ f(k_1, k_2) = \sum_{l_1l_2} \left( \frac{2}{\pi} \right)^2 e^{-i(E_{l_1l_2} - E)} \left( \frac{2}{\pi} \right)^2 e^{-i(E_{l_1l_2} - E)} \]  

from the single ionization channels, we employ a surface integral formulation for the partial wave amplitudes by integrating along a surface using testing functions that are continuum states of the individual one-body Hamiltonian \( h \) of the residual dication as defined in Eq. 2. Further details of this implementation can be found in Refs. [8, 9, 24]. Here we provide the final result for these partial amplitudes,

\[ \mathcal{F}_{l_1l_2}(k_1, k_2) = \frac{\rho_0}{2} \int_0^{\pi/2} \left[ \varphi_{l_1l_2}^{k_1}(r_1) \varphi_{l_1l_2}^{k_2}(r_2) \frac{\partial}{\partial \rho} \psi_{l_1l_2}(r_1, r_2) \right. \]  

\[ - \left. \psi_{l_1l_2}(r_1, r_2) \frac{\partial}{\partial \rho} \varphi_{l_1l_2}^{k_1}(r_1) \varphi_{l_1l_2}^{k_2}(r_2) \right|_{\rho = \rho_0} d\alpha. \]

The triply-differential cross sections (TDCS), representing the most detailed information that can be known in the double ionization process can be computed from the double ionization amplitudes as

\[ \frac{d^3\sigma}{dE_1 d\Omega_1 d\Omega_2} = \frac{4\pi^2 \omega}{c} k_1 k_2 |f(k_1, k_2)|^2, \]
in the length gauge. Integrating over all angles $\Omega_1$ and $\Omega_2$ of both electrons yields the single differential cross section (SDCS), which exhibits the energy sharing of the outgoing electrons for the total continuum energy. Integration of the SDCS over the possible energy sharings yields the total double ionization cross section at the photon energy $\omega$,

$$\sigma = \int_0^E \frac{d\sigma}{dE_1} dE_1. \quad (13)$$

Having defined these cross sections and provided a brief overview of the time-independent treatment, we present in the next section the results of valence shell double ionization of magnesium.

III. RESULTS AND DISCUSSION

The ground state $^1S$ state of magnesium is determined by diagonalizing the Hamiltonian in Eq. 1, constructed on a purely real radial grid with extent $r = 32.0$ bohr. The orbital region range is determined by the extent of the core $1s$, $2s$ and $2p$ orbitals and was taken to consist of three finite elements with 16-th order DVR in each, with endpoints of 0.5, 8.0 and 16.0 bohr. The maximum electron angular momentum used to converge the TDCS results below was found to be $l_{\text{max}} = 9$. With these grid parameters, the double ionization potential of the $3s^2$ shell of magnesium is 22.8 eV, in excellent agreement with the experimentally determined value of 22.7 eV [25] and with other theoretical treatments [19].

The solution of the driven equation (Eq. 9) proceeds on a larger grid with 8.0 bohr finite elements up to an ECS radius at $R_0 = 72.0$ bohr with two additional complex-scaled finite elements (ECS angle $\theta = 30^\circ$) to impose the outgoing wave boundary conditions. TDCS results for the photon energies considered were observed to be converged with respect to the grid parameters, maximum individual electron angular momenta $l_{\text{max}}$ and amplitude extraction radius.

A. Double to single ionization cross section ratios

In Figure 2, we present the calculated ratio of the total double ionization cross section to the single ionization cross section. These results are calculated in the length gauge and compared with theoretical results from compared convergent close coupling (CCC) [14], time dependent close coupling (TDCC) [10], the R-matrix-with-pseudostates (RMPS) [10], and with experimental measurements of Wehlitz, et.al. [20]. The agreement of the double to single ionization ratios is good with the other theoretical treatments and tracks very well with the experimental measurements. All but the lowest energy considered in the present treatment lie within the experimental error bars.

B. Single differential cross sections

The energy sharing cross section (SDCS) is shown in Figure 3 for length gauge results at different photon energies and normalized to unity to plot them in a common panel. The absolute cross section scaling factors for each photon energy are noted in the caption. The profile of these single differential cross sections, peaking at the unequal energy sharing extremes with a minimum at equal energy sharing $E_1 = E_2$ is typical of double ionization from $ns^2$ helium-like atoms. The theoretical results of TDCC [19] are also shown for comparison as dashed lines, exhibiting a similar evolution of the SDCS towards more extreme energy sharings as the photon energy is increased. The magnitude of the these cross sections is lower than the TDCC results, as seen in the total double-to-single ionization ratios plotted in Fig. 2.

C. Triple differential cross sections at $\omega = 55.49$ eV

The most sensitive probe of electron correlation that can be revealed in a double ionization investigation is contained in the triple differential cross section (TDCS), which exhibits the angular distributions of both exiting electrons and their energy sharing. For these helium-like cases absorbing a single photon, the overall transition features a symmetry of $^1S \rightarrow ^1P$. The main features due to electron correlation are most constrained by symmetry at equal energy sharing. Figure. 4 shows the TDCS plotted in the coplanar geometry (with both electrons exiting in the plane of the polarization, which defines the $z$-axis for the angular measurements, i.e. $\phi_1 = \phi_2 = 0^\circ$) at a photon energy of $\omega = 55.49$ eV. Each panel plots
the cross section as a function of the angle of the second electron when the other electron (labeled electron 1) is held fixed at the angle indicated.

The results shown for both length and velocity gauges exhibit a known parity symmetry that requires the TDCS to be zero for back-to-back electron ejection when $E_1 = E_2$ for helium-like targets [26]. The TDCS plotted for the present results are absolute, with good agreement seen between the different gauges indicating a suitably converged description of the initial target state valence electrons within the frozen-core approximation. Experimental measurement points measured at Elettra [18] are also shown in each panel, as are the results from previous theoretical calculations: the time dependent close coupling (TDCC) results [19], convergent close coupling (CCC) calculations [18]. Following the comparison and discussion in Refs. [18] and [19], the non-resonant CCC calculation results are absolute while the resonant CCC calculation (employing a semi-empirical treatment to account for the resonance process populating the 3$d$ state of the target) has been scaled by a factor of $3q^2$, where $q$ is the Fano profile index, taken here with a value of 50 [18]. The TDCC results are also absolute. We note that, additionally, all of the results for comparison have be divided by a factor of 2 from the figures in Ref. [19] due to an alternate definition of the TDCS (see Ref. [8]).

The experimental results, gathered in the same measurement, are internormalized between the panels and can be compared after consistent scaling scaling (here taken to be a factor of 0.011 for consistency with

FIG. 3: (Color online) Energy differential cross section for DPI of magnesium for different photon energies. Results have been normalized to unity. Solid lines: present results calculated in the length gauge. The scaling factors to recover the absolute SDCS are (in kb/eV) 0.218 for 30 eV, 0.147 for 35 eV, 0.108 for 40 eV, 0.086 for 45 eV, and 0.057 for 55.49 eV. Dashed lines: TDCC calculation results [19], with scaling factors (in kb/eV) of 0.238, 0.173, 0.133, 0.106 and 0.073 for this list of photon energies, respectively. 1 kb=10$^{-21}$ cm$^2$.

FIG. 4: (Color online) Triple differential cross section (TDCS) for DPI from magnesium at 55.49 eV photon energy for equal energy sharing. Each panel shows plots the second electron in the coplanar geometry when the first electron is fixed at the angle shown relative to the polarization. Solid (black) curve: present results in the length gauge, solid (red) curve: present results in the velocity gauge, dashed (green) curve: TDCC results [19], dotted (brown) curve: CCC non-resonant results, dashed-dot (blue) curve: CCC resonant results [18], magenta circles: experimental measurements [18]. 1 b=10$^{-24}$ cm$^2$. 
The present results show fairly good agreement with the absolute theoretical treatments in the number of lobes and the relative sizes of the secondary peaks compared to the primary peaks. Slight differences in the predicted angles of these peaks are apparent, but the equal energy sharing results are consistent with the symmetry conditions for the helium-like double photoionization at equal energy sharing. All theories seem to produce a smaller result than the experiment at \( \theta_1 = 0^\circ \), producing better agreement of the size of the dominant peak with the experimental measurements at \( \theta_1 = 30^\circ \) and \( \theta_1 = 60^\circ \). At these angles in particular, the present results are slightly larger than the absolute TDCC and CCC (non-resonant) calculations. The secondary features along the primary lobes predicted by the semi-empirical CCC calculation that incorporates the resonant nature of the transition at this photon energy are not found in the ab initio two-active electron theories and are not particularly well-resolved in the experimental points. We note that all the theoretical results show narrower dominant peaks than observed for helium double photoionization at 20 eV above threshold, consistent with the spatially more diffuse initial state orbital of the magnesium target [15].

In Figure 5, the length and velocity gauge TDCS results for unequal energy sharing at \( \omega = 55.49 \) eV are shown in the coplanar geometry for fixed directions \( \theta_1 = 0^\circ \) and \( \theta_1 = 30^\circ \) and two different unequal energy sharing cases, representing the fixed electron at these angles carrying 31.7% of the excess energy in the top row, and the complementary \( E_1 \) carrying 68.3% of the available energy (32.8 eV) in the lower row. The relaxation of the parity selection rule away from equal energy sharing is evident in these figures compared to Fig. 4. We find good agreement in the overall shape of the TDCS lobes with different theories, being most in accord for the upper-right panel.

In Figure 6, we report the length and velocity gauge TDCS results for both equal energy sharing and the unequal energy sharings of Fig. 5 with the first electron direction fixed at \( \theta_1 = 90^\circ \). No experimental measurements were reported at fixed electron direction perpendicular to the polarization \( \theta_1 = 90^\circ \), but we compare with the theoretical descriptions. All theories show a pair of dominant and secondary lobes symmetric about \( \theta_2 = 270^\circ \). The resonant CCC results show significantly larger secondary peaks for both equal and unequal energy sharings, while all non-resonant calculations predict smaller values for the secondary lobes compared to the primary peaks. The location of the primary peaks differs slightly between the different theoretical results, with the largest variation between the present results and the other calculated results most prominent when the fixed electron is slow (middle panel). We also note less variation in the magnitude of these cross sections as a function of the fixed electron rotating away from the polarization direction than is seen for helium.

Finally, to compare the different \( ns^2 \) atomic targets that have been treated by the present method, we present in Figure 7 the normalized angular distribution of the second electron plotted outside the plane of the polarization and fixed electron for 5% energy sharing of 20 eV excess energy above the double ionization potential for helium, beryllium, and magnesium. The fixed electron is at \( \theta_1 = 60^\circ \) in each panel. Comparing these results shows a similar lobe structure in each, with slight variation in the width of the primary and secondary lobes as the initial state of the target becomes more diffuse. There is noticeable variation in the angle of the lobes across these different targets when compared at the same excess photon energies, but the general structure of the TDCS is similar. The indication of these comparisons and with those of the previous theoretical and experimental results is that the symmetry of the one-photon process in these overall \( ^1S \rightarrow ^1P \) transitions determines much of the features of the TDCS, with variations in the angles and relative sizes of the lobes of photoejection distribution slightly sensitive to the details of the correlated initial state.

### IV. CONCLUSION

We have presented single photon double ionization cross sections for removal of the valence electrons of atomic magnesium. As a function of photon energy, good agreement between the total double-to-single photoionization cross sections exists between the present results and other theoretical calculations and experiment. We also find good agreement with the energy sharing cross section profiles at different photon energies with the TDCC treatment.

Focusing on the results at the photon energy of 55.49 eV for which experimental TDCS results have been collected via a resonant process, we find fairly good agreement with other ab initio theoretical results that similarly cannot account for the resonant transition of the experimental conditions. Nevertheless, comparison of the experimental measurements with the various theoretical calculations reveals common structures in the general shape of the triple differential cross sections for several angles and energy sharings. The most substantial impact of the resonant transition for the experiment that is unaccounted for in the ab initio calculations affects the absolute magnitude of the cross sections at the relative sizes of the secondary lobes compared to the dominant photoejection angles for the cases considered. Overall, reasonable agreement is found between the experimental angular distributions and the various theoretical treatments, which is significantly determined by the overall symmetry of the helium-like \( ^1S \rightarrow ^1P \) single photon transition.
FIG. 5: (Color online) TDCS for DPI from magnesium at 55.49 eV photon energy for unequal energy sharing at two different fixed electron directions: $\theta_1 = 0^\circ$ (left panels) and $\theta_1 = 30^\circ$ (right panels) relative to the polarization. The fixed-direction electron has 10.4 eV and 22.4 eV in the upper and lower rows, respectively. Solid (black) curve: present results in the length gauge, solid (red) curve: present results in the velocity gauge, dashed (green) curve: TDCC results [19], dashed-dot (blue) curve: CCC resonant results [18], magenta circles: experimental measurements [18]. $b=10^{-24}$ cm$^2$.

FIG. 6: (Color online) TDCS for DPI from magnesium at 55.49 eV photon energy for various energy sharings with fixed electron direction $\theta_1 = 90^\circ$ relative to the polarization. From top-to-bottom, the fixed-direction electron has 16.4 eV, 10.4 eV and 22.4 eV. Solid (black) curve: present results in the length gauge, solid (red) curve: present results in the velocity gauge, dashed (green) curve: TDCC results [19], dashed-dot (blue) curve: CCC resonant results [18]. 1 b = $10^{-24}$ cm$^2$.


