Double ionization mechanisms of magnesium driven by electron impact

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We study double ionization of Mg by electron impact through the vantage point of classical mechanics. We consider all electron-electron correlations in a Coulomb 4-body problem, where three electrons belong to the atom and the fourth electron causes the impact ionization. From our model we compute the double ionization probability of Mg for impact energies in [15, 125] eV. Double ionization occurs through eight double ionization mechanisms, which we classify into four categories: Inner Shell Capture, Direct, Delay and Ionized Inner Shell mechanisms. We show that Delay and Ionized Inner Shell mechanisms require electron-electron correlations among the four electrons, and are responsible for the second increase in the double ionization probability. Furthermore, we show that our theoretical prediction about the relative prominence of certain double ionization mechanisms is in agreement with experimental results on the relative prominence of non-first over first order mechanisms.

I. INTRODUCTION

Multiple ionization by an electron impact is a phenomenon wherein a free electron collides with bound electrons in a target (an atom, a molecule, an ion and so forth) and ionizes them. This phenomenon is used to study energy exchanges and electron-electron correlations in many-electron systems, electron dynamics and the structure of the target. Multiple ionization of atoms by electron is also relevant in X-ray astronomy because it is a cosmic source of X-rays 11. Knowledge about multiple ionization of atoms by electron impact is useful for modelling in fusion plasma 2. Moreover, double ionization of atoms by electron impact may help interpreting photon-atom interactions 3,4.

In this study, we consider the double ionization of Mg by electron impact. The probability of double ionization of Mg by electron impact was experimentally measured in Ref. 5–8. In all the experiments, two drastic increases are observed in the probability of double ionization as a function of the initial energy of the impact electron (impact energy). The first increase occurs for impact energies greater than 25 eV, which corresponds to the sum of the first two ionization potentials of Mg. The second increase occurs for impact energies between 40–70 eV. The second increase lends a ‘knee’ structure to the probability of double ionization as a function of the impact energy. A similar knee structure is observed in the probability of double ionization versus impact energy for Boron ion 9, Barium 10,11, Calcium and Strontium 5,12, Argon and Xenon ions 7,13, and other target species 14. Ref. 15 attributes the knee structure to the emergence of different double ionization mechanisms. In particular, Ref. 15 conjectures that the knee structure marks the emergence of the Auger effect. The conjecture, in turn, suggests the prominent role of the inner shell electrons (electrons other than the valence shell electrons) in the double ionization of targets by electron impact for impact energies greater than a target specific energy value.

In principle, the double ionization of a target by electron impact is a Coulomb n-body problem, with n ≥ 3, but with n ≥ 4 to accommodate the Auger effect. Therefore, the simplest model, which accounts for the role of the inner shell electron in the double ionization of a target by electron impact, yields a Coulomb 4–body problem. That model contains one inner shell and two valence shell electrons in the target and one impact electron. A quantum mechanical description of the simplest model may reveal the electron dynamics that cause the knee structure in the probability of double ionization as a function of the impact energy. However, a complete quantum mechanical description of a Coulomb 4–body problem is computationally too complex to be solved with present technology. Therefore, several approximate quantum mechanical approaches have been developed, of which the reduced non-perturbative methods are Time-Dependent Close Coupling (TDCC), R-Matrix with Pseudo-States (RMPS), and Convergent Close Coupling (CCC) methods 17–20, and the perturbative method is the Distorted Wave (DW) method 21,22. The non-perturbative methods have successfully tackled the Coulomb 3–body problem 20. So, the present quantum approaches reduce the Coulomb 4–body problem, of the simplest model of double ionization of a target by electron impact, to a series of Coulomb 3–body problems. For instance, Ref. 23 selectively applies the TDCC and DW methods on different sets of electrons and in different domains of impact energies to reproduce the knee in the probability of double ionization. Similarly, Ref. 24 and Ref. 25 perform two separate TDCC calculations on two distinct sets of electrons and add the probability contributions from the two calculations to recover the knee structure in the probability of double ionization. The piece-wise treatments in Ref. 23,24 serve as evidence that the double ionization of a target by electron impact is indeed not a Coulomb 3–body problem, but a more complicated, fully coupled, Coulomb n–body problem for n ≥ 4. A comparison be-

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between theory and experiment further supports the idea that correlations among more than three electrons are required to explain the probability of double ionization of a target by electron impact [29]. Therefore, the present quantum mechanical approaches are limited in that they approximate electron-electron correlations in their model of double ionization of a target by electron impact [29–25] [27].

Classical mechanics provides a complementary approach to the present quantum mechanical approaches. Ref. [28] [30] treat electron-target collisions in the framework of classical mechanics and derive the formulas for the probability of double ionization of targets by electron impact. The results agree qualitatively and quantitatively well with experimental results. The classical mechanical approach is not only computationally more efficient than quantum mechanical approaches, but also presents two advantages over quantum mechanical approaches. First, classical mechanical approaches describe electron dynamics locally in phase space, while quantum mechanical approaches describe electron dynamics globally in phase space. A corollary benefit is that increasing the system size (number of electrons in the model or the spatial dimensions) is much more advantageous in classical mechanical treatments as opposed to quantum mechanical treatments. The second advantage is that classical mechanical trajectories provide intuition for making approximations, such as the binary encounter approximation [12] [25] [30] [31], in the complementary quantum mechanical approaches. Thus, classical mechanical approaches are relevant in interpreting a truly quantum phenomenon, we employ a classical mechanical approach to describe the double ionization of Mg by electron impact.

In this article, we consider three bound electrons and a static ionic core in the target Mg atom, and one impact electron. One of the three bound electrons in the atom resides closest to and most tightly bound to the ionic core in the ground state configuration of the atom. That electron is referred to as the inner shell electron. The other two electrons are further away from and more loosely bound to the ionic core as compared to the inner shell electron. They are referred to as the valence shell electrons. We consider Coulomb 4–body problem that accounts for all electron-electron correlations among the four (one inner shell, two valence shell and one impact) electrons. We show that our purely classical mechanical treatment of the problem reproduces the knee structure in the probability of double ionization of Mg by electron impact. We use classical trajectories to illustrate and understand the mechanisms behind the knee structure.

In Sec. [II] we introduce the Hamiltonian model for the double ionization of Mg by electron impact and the choice of interaction potentials. In Sec. [III] we present the resulting probability of double ionization of Mg by electron impact. Next, we describe and illustrate the double ionization mechanisms, and the change in the contribution of the different mechanisms to the probability of double ionization. In Sec. [IV] we conclude with two points of discussion; first, about the similarities and differences between our three-active electron target atom and a two-active electron target atom [22] for Mg, and, second, about the experimental link of the mechanisms predicted by this model.

II. HAMILTONIAN MODEL

In our model, the Mg atom is composed of four components: one static ionic core, one inner shell electron and two valence shell electrons. The static ionic core consists of a fixed nucleus and five dynamically frozen electrons, screening the charge of the nucleus. The three-active electrons interact among themselves and with the impact electron.

A. Hamiltonian

In what follows, all quantities are in atomic units (a.u.), unless explicitly stated otherwise. The electrons in the target atom are labeled by \(i \in \{1, 2, 3\}\). The position of the \(i\)-th electron is denoted \(\mathbf{r}_i\) and its canonically conjugate momentum is denoted \(\mathbf{p}_i\). The Hamiltonian of the isolated target atom comprised of a static ionic core and three-active electrons reads

\[
H_{\text{Mg}} = \sum_{i=1}^{3} \left[ \frac{\mathbf{p}_i^2}{2} + V_{\text{ee}}(\mathbf{r}_i) + \sum_{j=i+1}^{3} V_{\text{ee}}(\mathbf{r}_i - \mathbf{r}_j) \right].
\]  (1)

In the outer summation, the first term is the kinetic energy and the second term is the electron-core potential, \(V_{\text{ee}}(\mathbf{r}_i)\), for each electron. The second summation encapsulates the electron-electron potential, \(V_{\text{ee}}(\mathbf{r}_i - \mathbf{r}_j)\), between the \(i\)-th and the \(j\)-th electrons. Initially, the Mg atom is in the ground state of energy \(E_0\), defined as the sum of the first three ionization potentials.

The impact electron is labeled by \(i = 0\). The impact electron induces a perturbation in the target atom. The Hamiltonian for this perturbation reads

\[
H_1 = \frac{\mathbf{p}_0^2}{2} + V_{\text{ee}}(\mathbf{r}_0) + \sum_{j=1}^{3} V_{\text{ee}}(\mathbf{r}_0 - \mathbf{r}_j),
\]

where the first term is the kinetic energy and the second term is the electron-core potential, \(V_{\text{ee}}(\mathbf{r}_0)\), of the impact electron. The summation encapsulates the electron-electron interactions between the impact electron and the three-active electrons in the target atom, \(V_{\text{ee}}(\mathbf{r}_0 - \mathbf{r}_j)\). The initial energy of the impact electron is \(H_1 = E_0\) and is referred to as the impact energy. The total Hamiltonian describing the electron dynamics is given by

\[
H = H_{\text{Mg}} + H_1.
\]

Here, we consider two-spatial dimensions. We note that it is still possible to carry out the calculations in
three dimensions. However, for the same number of initial conditions, calculations in three spatial dimensions yield smaller double ionization probabilities as compared to calculation in two spatial dimensions. Therefore, to achieve a good resolution in the trend for probability of double ionization as a function of $\epsilon_0$, a greater number of initial conditions is required, which makes the calculations computationally more expensive. In addition to that cost, the mechanisms are not as clear in three spatial dimensions as they are in two spatial dimensions because of a higher dimensional phase space corresponding to three spatial dimensions.

### B. Choice of Potentials

The electron-core potentials, $V_{ec}(r)$, and the electron-electron potentials, $V_{ee}(r_i - r_j)$, in the Hamiltonian in Eq. (1), must satisfy the following three criteria to ensure a stable ground state Mg atom in our model:

1. **No self ionization.** Electrons should not ionize from the atom in the absence of the impact electron.

2. **Existence of an inner shell electron.** This condition requires that one of the three electrons in the atom remains the most tightly bound and the closest electron to the ionic core at all times.

3. **Non-empty ground state.** The ground state is classically defined as the set of $\{|r_i|, \{|p_i|\} \}$ such that

$$H_{Mg}(|r_i|, |p_i|) = E_g. \tag{2}$$

A non-empty ground state demands that there exist at least one point in phase space $(r_1, p_1, r_2, p_2, r_3, p_3)$ which satisfies Eq. (2).

Conditions 1 and 3 are required to have a stable atom, and Condition 2 emerges from the experimentally observed ionization potentials. The first and second ionization potentials, $E_1 = -0.28$ a.u. and $E_2 = -0.55$ a.u., are rather close to each other, whereas the third ionization potential $E_3 = -2.95$ a.u., is much more negative than the first two ionization potentials. Therefore, as compared to the other two electrons, one of the electrons in the atom is more tightly bound to the ionic core.

#### 1. Soft-Coulomb potentials

A widely used form of the electron-electron and electron-core potentials is the soft-Coulomb potentials. Soft-Coulomb potentials are given by

$$V_{ec}^{(SC)}(r) = \frac{-3}{\sqrt{||r||^2 + a^2}}, \tag{3}$$

and

$$V_{ee}(r) = \frac{1}{\sqrt{||r||^2 + b^2}}. \tag{4}$$

![FIG. 1: Effective potentials for the ion-electron interaction as a function of the distance from the ionic core.](image)

The role of the effective charge in the electron-core soft-Coulomb potential is to create an energy barrier, the height of which is greater than the sum of the first two ionization potentials of the atom. This energy barrier ensures that bound electrons that are on one side of the barrier cannot move to the other side of the barrier by any redistribution of energy among the bound electrons in the ground state. Since the proximity of an electron to the ionic core determines the strength of the bond between the two, it follows that Condition 2 can be satisfied.

In order to remedy this problem, we keep the electron-electron soft-Coulomb potential given in Eq. (3), and introduce an effective charge, $Z_{eff}(r)$, in place of the constant charge in the electron-core soft-Coulomb potential in Eq. (3). The resulting electron-core soft-Coulomb potential with effective charge is

$$V_{ec}(r) = \frac{-Z_{eff}(r)}{\sqrt{||r||^2 + a^2}}. \tag{5}$$
We model the effective charge as

\[ Z_{\text{eff}}(r) = 3 + 3(\chi - 1) \exp \left( -\alpha \left( \|r\|^2 - R^2 \right)^2 \right), \tag{6} \]

where the parameters \( \chi, \alpha \) and \( R \) govern, respectively, the height, width and position of the energy barrier. Although we choose \( Z_{\text{eff}}(r) \) given by Eq. (6), we note that any function with the general features highlighted above may have been used and would lead to the same results, at least qualitatively. Figure 1 depicts the effective charge as a function of the distance from the ionic core by a dashed blue line. The corresponding electron-core soft-Coulomb potential, given by Eq. (5), is depicted in solid black line. The local maximum at \( \|r\| = 2 \) a.u. is the peak of the energy barrier. To highlight the role of the effective charge and the energy barrier, we have depicted the soft-Coulomb potential without effective charge given by Eq. (4) in dashed gray line in Fig. 1.

In sum, the total Hamiltonian is given by

\[ H = \sum_{i=0}^{3} \left( \frac{p_i^2}{2m_i} - \frac{Z_{\text{eff}}(r_i)}{\sqrt{\|r_i\|^2 + a^2}} \right) + \sum_{j=i+1}^{3} \frac{1}{\sqrt{\|r_i - r_j\|^2 + b^2}}, \tag{7} \]

with \( Z_{\text{eff}} \) given by Eq. (6). We choose soft-Coulomb parameters \( a = 0.95 \) and \( b = 0.1 \). Our choice allows us to find non-empty ground states and allows significant energy exchanges among the electrons. The parameters, \( \chi, \alpha \) and \( R \) are taken as 0.3, 0.4, and 2 respectively.

### C. Initial Conditions

We integrate Hamilton’s equations obtained from Hamiltonian (7), to compute trajectories. In this section, we discuss our procedure to assign initial conditions to the electrons for the integration.

#### 1. Impact Electron

We start the integration at an initial time \( t_{\text{in}} < 0 \). The initial position of the impact electron is denoted by \( r_{0,\text{in}} = x_{0,\text{in}} \hat{x} + y_{0,\text{in}} \hat{y} \). The initial \( x \)-coordinate of the impact electron \( x_{0,\text{in}} < 0 \) is fixed for all impact energies. The initial \( y \)-coordinate \( y_{0,\text{in}} \), which is referred to as the impact parameter, is randomly chosen from a uniform distribution in \([0, 5]\) a.u. Initially, the impact electron is sent along the \( \hat{x} \)-direction with impact energy \( \epsilon_0 \). The initial momentum of the impact electron is given by

\[ p_{0,\text{in}} = \hat{x} \sqrt{2\epsilon_0}. \]

The initial time \( t_{\text{in}} \) is chosen such that the impact electron would reach \( x = 0 \) at \( t = 0 \) if the target atom were absent. Therefore,

\[ t_{\text{in}} = \frac{x_{0,\text{in}}}{\|p_{0,\text{in}}\|}, \]

and the first collision between the impact electron and any bound electron occurs approximately at time \( t = 0 \). Both \( x_{0,\text{in}} \) and the distance beyond which an electron is considered ionized is fixed at 200 a.u.

#### 2. Isolated atom

In this section, we describe the procedure to assign the initial positions and momenta to the bound electrons in the isolated atom.

First, the configuration of the inner shell electron is determined in absence of valence electrons. The inner shell electron is randomly positioned such that \( \|r_{3,\text{in}}\| < 2 \) a.u. and

\[ V_{\text{ec}}(r_{3,\text{in}}) < \mathcal{E}_3, \]

where \( \mathcal{E}_3 \) is the third ionization potential of Mg. Then, the kinetic energy of the inner shell electron, \( T_3 \), is given by

\[ T_3 = \mathcal{E}_3 - V_{\text{ec}}(r_{3,\text{in}}). \]

Following which, the initial momentum of the inner shell electron becomes

\[ \|p_{3,\text{in}}\| = \sqrt{2T_3}. \]

The momenta along \( \hat{x} \)- and \( \hat{y} \)-direction are randomly decomposed to determine the initial momentum of the inner shell electron.

Next, we determine the configuration of the two valence shell electrons in the presence of the ionic core and the inner shell electron. The initial position of the two valence shell electrons are randomly generated such that \( \|r_{1,\text{in}}\| \in [2, 8] \) a.u., \( \|r_{2,\text{in}}\| \in [2, 8] \) a.u. and

\[ V_{\text{ec}}(r_{1,\text{in}}) + V_{\text{ec}}(r_{2,\text{in}}) + V_{\text{ee}}(r_{1,\text{in}} - r_{2,\text{in}}) + V_{\text{ec}}(r_{1,\text{in}} - r_{3,\text{in}}) + V_{\text{ec}}(r_{2,\text{in}} - r_{3,\text{in}}) < \mathcal{E}_1 + \mathcal{E}_2, \]

where \( \mathcal{E}_1 \) and \( \mathcal{E}_2 \) are the first two ionization potentials, respectively. Then, the sum of the kinetic energies of the two valence shell electrons, \( T_1 + T_2 \), becomes

\[ T_1 + T_2 = \mathcal{E}_1 + \mathcal{E}_2 - \left[ V_{\text{ec}}(r_{1,\text{in}}) + V_{\text{ec}}(r_{2,\text{in}}) + V_{\text{ee}}(r_{1,\text{in}} - r_{2,\text{in}}) + V_{\text{ec}}(r_{1,\text{in}} - r_{3,\text{in}}) + V_{\text{ec}}(r_{2,\text{in}} - r_{3,\text{in}}) \right]. \]

The initial momenta of the two valence electrons are determined using a parametrization of the 4-sphere by three randomly chosen angles such that

\[ \|p_{1,\text{in}}\|^2 + \|p_{2,\text{in}}\|^2 = 2(T_1 + T_2). \]
D. Integration scheme

We employ the 4th order Runge-Kutta numerical integration scheme. We fix the ending time of integration at \( t_f = 1500 \) a.u. for all impact energies. We integrate \( 3 \times 10^7 \) initial conditions for each \( \epsilon_0 \in \{10, 11, 12, ..., 125\} \). We fix the step size in the integration scheme at 0.1 a.u. Further reducing the time step size presents no perceptible changes in the double ionization probability and increases the computational time. The integration scheme was developed in C/C++ and CUDA C, and deployed on NVIDIA Tesla V100 GPU and Intel Core i9 8950HK CPU. The double ionization probability was calculated from the results of the computation done on the GPU.

III. DOUBLE IONIZATION PROBABILITY

The probability of double ionization of Mg by electron impact as a function of the impact energy \( \epsilon_0 \) computed from our model is represented by the thick black line in Fig. 2a. We observe that our model reproduces the knee structure in the double ionization probability for low impact energies as observed in experiments [5–8]. Classical mechanics allows visualizing trajectories, which unveil the different double ionization mechanisms. The probability of different mechanisms sums up to the total double ionization probability. Our model predicts that the double ionization of Mg by electron impact proceeds through eight main mechanisms in the studied range of impact energies. We classify those eight mechanisms into four categories: Inner Shell Capture, Direct, Delay and Ionized Inner Shell. We show the probabilities of the four categories of mechanisms by thick color lines in Fig. 2a. Inner Shell Capture in green, Direct in blue, Delay in red and Ionized Inner Shell in yellow. In the Inner Shell Capture mechanisms, the impact electron gets momentarily trapped in the inner shell region and ionizes later. As expected, these mechanisms occur for low impact energies. The Direct mechanisms are characterized by the absence of a time delay between impact and ionization. Roughly speaking, the Direct mechanisms proceed via collisions like the collisions of balls in the game of billiards. On the contrary, the Delay mechanisms are characterized by a time delay between impact (around \( t = 0 \)) and ionization. Consequently, an intermediate metastable state of the target forms during the period of delay. The last but not the least, the Ionized Inner Shell mechanisms group together all the mechanisms in which the inner shell electron is ionized.

At about \( \epsilon_0 = 50 \) eV, we observe a drastic increase of the probability of Delay mechanisms, depicted with the thick red line. This increase is responsible for the increase in the probability of double ionization. Then, at about \( \epsilon_0 = 100 \) eV, the increase in the probability of the Ionized Inner Shell mechanisms, represented by the thick yellow line, counters the decrease in the probability of the Delay mechanisms, creating a plateau in the probability of double ionization till \( \epsilon_0 = 125 \) eV.

In summary, the knee shape in the double ionization probability is mostly due to the cross-over between the Inner Shell Capture and the Delay and Direct mechanisms. Even if there are no visible qualitative changes in the double ionization probability for impact energies...
around 100 eV and higher, the probability curve is hiding the appearance of additional mechanisms such as the Ionized Inner Shell mechanisms.

Figure 2 illustrates the different contributions of the different mechanisms within each category: TS2 and TS1 are the mechanisms within the Direct category, and are represented by thin dark blue and light blue lines respectively in Fig. 2a. Similarly, D1 and D0 are the sub-mechanisms within the Delay category, and are represented by thin dark red and light red lines, respectively in Fig. 2a. The mechanisms TS2*, TS1* and D1* are associated with the Ionizer Inner Shell category, and are represented in Fig. 2b by thin dark blue, light blue and dark red lines respectively. Each mechanism is explained in more detail below.

A. Format for the figures of the mechanisms

Before explaining each double ionization mechanism identified in our model, we introduce the format of Figs. 3 which accompany the text to explain the mechanisms. In each of these figures, e.g., Fig. 3 there is a schematic diagram in the top left corner, two trajectory plots (e.g., in Figs. 3a and 3b), and a plot of the electron-electron potential versus time (e.g., Fig. 3c).

a. Schematic diagram In the schematic diagrams, the labels ‘X’, ‘V’, ‘O’ and the gray circular region correspond to the impact electron, valence shell electron, inner shell electron and the inner shell region, respectively. The arrows correspond to ionization processes which occur on short timescales (time shorter than a certain threshold $t_{\text{Th}}$), and a wiggly line corresponds to an ionization process which occurs on long timescales (time larger than $t_{\text{Th}}$). We choose a threshold $t_{\text{Th}} = 30$ a.u. for which there is a clear separation between the Delay and Direct processes.

b. Panels (a) and (b) In panels (a), we plot the electron trajectories covering 0 to 80 a.u. from the ionic core, and in panels (b), the same electron trajectories are plotted in a range from 0 to 8 a.u. around the ionic core. In both panels, the black and yellow lines represent the impact electron and inner shell electron trajectories, respectively. The red line represents the trajectory for that valence shell electron that ionizes and leaves the boundary before the other valence shell electron. The blue line represents the trajectory for the other valence shell electron. We indicate the direction of approach of the impact electron with a black arrowhead. We plot the arrowheads to indicate the direction of the valence shell electrons in panels (b). The arrowheads in panels (b) correspond to the same instant of time for all electrons.

c. Panel (c) In panels (c), we depict the six electron-electron potentials, $V_{ee}(r_i - r_j)$, for $i, j \in \{0,1,2,3\}$, with an inset focusing on a time interval around the collision, i.e., around $t = 0$ a.u. The electron-electron potentials are colored as follows: $V_{ee}(r_0 - r_1)$ is red, $V_{ee}(r_0 - r_2)$ is blue, $V_{ee}(r_0 - r_3)$ is yellow, $V_{ee}(r_1 - r_2)$ is purple, $V_{ee}(r_1 - r_3)$ is brown, and $V_{ee}(r_2 - r_3)$ is green.

Lines in panels (a), (b) and (c) are thin and translucent until the time the impact electron first enters the 8 a.u. boundary, thick and solid until the last electron in double ionization leaves the 8 a.u. boundary for the last time, and thin and translucent for the time after.

Using this typology, we examine all the mechanisms leading to double ionization we found by a close inspection of the trajectories.

B. Inner Shell Capture Mechanism

An example of the Inner Shell Capture mechanism is plotted in Fig. 3. In Fig. 3a, the trajectory of the impact electron (black line) shows that the impact electron falls into the inner shell region, $|r| < 2$ a.u. In Fig. 3c, the yellow electron-electron potential shows that the impact electron and the inner shell electron interact for a time duration greater than $t_{\text{Th}}$. Meanwhile, the steep decrease in all but the yellow electron-electron potentials, at about time $t = 0$ a.u., implies that the two valence shell electrons leave the atom right after the impact. Finally, Fig. 3b illustrates that the impact electron eventually leaves the atom, completing the double ionization mechanism.

The schematic diagram in the top left in Fig. 3 summarizes this process: The impact electron ‘X’ falls into the inner shell region (gray circle) and stays there for a long time before it leaves the atom while the two valence shell electrons ‘V’s leave the atom before $t = t_{\text{Th}}$ (arrows only).

The impact electron exchanges energy with the valence electrons, which leave the atom. During this process, the energy of the impact electron falls below the energy barrier (see Sec. 1B2), and the impact electron gets captured in the inner shell region. The capture persists and strengthens the electron-electron correlation between the impact electron and the inner shell electron. This mechanism is characterized by a series of energy exchanges between the impact electron and the inner shell electron, after which the impact electron finally gets enough energy to overcome the potential barrier and leaves the atom. The Inner Shell Capture mechanism requires an inner shell electron dynamics and the inner shell region to occur, and the mechanism proceeds through strong electron-electron correlation between the inner shell and the impact electrons.

This mechanism dominates the probability of double ionization for impact energies between 10 eV and 40 eV (see Fig. 2).

C. Direct Mechanism

Direct mechanisms involve the ionization of the two valence shell electrons before time $t = t_{\text{Th}}$. There are
two types of Direct mechanisms: TS1 and TS2, which are depicted in Fig. 4 and Fig. 5 respectively, and are described in this section. In the nomenclature of the mechanisms, ‘TS’ indicates that the mechanisms lead to double ionization in two steps (TS), and ‘1’ or ‘2’ refers to the number of collisions between the impact electron and valence shell electrons, which trigger ionization. In TS1 mechanisms, the impact electron collides and ionizes one valence shell electron, which on its way out of the atom collides with and ionizes the other valence shell electron. While, in TS2 mechanism the impact electron collides sequentially with and ionizes both the valence shell electrons.

1. TS1

A typical TS1 mechanism is depicted in Fig. 4. In Fig. 4b, the impact electron (black line) collides with and ionizes only one valence shell electron (blue line), and leaves the atom. This is the first step in the mechanism. Then, that valence electron (blue line) on its way out of the atom, collides with the other valence shell electron (red line), and both leave the atom. This is the second step in the mechanism. Figure 4c shows the timescale of the electron-electron interactions during TS1 processes. In the inset in Fig. 4c, we observe that the first peak in the electron-electron potential (blue) corresponds to the first collision between the impact electron and a valence shell electron (blue line in Figs. 4a and 4b). The second peak in the electron-electron potential (green line) corresponds to the rapid transit of this valence shell electron through the inner shell region. The third peak in the electron-electron potential (purple line) corresponds to the last collision between the two valence shell electrons.

A schematic diagram of TS1 mechanisms is depicted in the top left corner of Fig. 4a. In TS1 mechanisms, the impact electron provides energy, which is greater than the sum of the first two ionization potentials of Mg, to one valence shell electron. This sets that valence shell electron on a trajectory towards ionization. In its transit through the inner shell region, the blue valence shell electron provides energy to the inner shell electron, but keeps enough energy to remain on its path towards ionization. Then, the collision between the two valence shell electrons leads to a redistribution of energy carried by the two electrons, such that a valence shell electron is set on a trajectory towards ionization (red line), while the other valence shell electron (blue line) is reset onto a new trajectory towards ionization.

2. TS2

A typical TS2 mechanism is depicted in Fig. 5. In Fig. 5b, the impact electron (black line) collides sequentially with and ionizes both valence shell electrons (blue and red lines) and leave the atom. These are the two steps in the mechanism. Just as TS1 mechanisms, TS2 mechanisms occur on short time scales (typically, a few atomic units). In Fig. 5c, the first peak in the electron-electron potentials (red line) corresponds to the first collision between the impact electron and a valence shell electron (blue line) through the inner shell region. The second peak in the electron-electron potential (green line) corresponds to the rapid transit of this valence shell electron through the inner shell region. The third peak in the electron-electron potential (purple line) corresponds to the last collision between the two valence shell electrons.
electron. The second peak in the electron-electron potential (yellow line) corresponds to the transit of the impact electron through the inner shell region. The third peak in the electron-electron potential (blue line) corresponds to the second collision between the impact electron and the other valence shell electron.

In the top left corner of Fig. 5a, a schematic diagram of TS2 mechanisms is depicted. In TS2 mechanisms, the impact electron provides energy sequentially to the two valence shell electrons via collisions, which cause the valence shell electrons to ionize.

D. Delay Mechanism

Delay mechanisms are characterized by three features; first, the mechanisms occur on long time scales (typically, few tens of atomic units); second, the mechanisms do not involve any capture in the inner shell; and third, all valence shell electrons ionize. There are two types of Delay mechanisms, D1 and D0, that are represented in Fig. 6 and in Fig. 7, respectively. In the nomenclature of the mechanisms, ‘D’ refers to Delay mechanisms. ‘1’ or ‘0’ refers to the number of electrons that are ionized before the threshold time $t = t_{Th}$, or, in other words, the number of electrons which are ionized on a short timescale.

1. D1

A typical D1 mechanism is depicted in Fig. 6. In Fig. 6a, the impact electron (black line) first collides with one valence shell electron (red line). Then, the impact electron transits through the inner shell region and leaves the atom. In the inset in Fig. 6c, the electron-electron potential between the impact electron and the red valence shell electron (red line) peaks first. It indicates that the first collision occurs between the impact electron and the red valence shell electron. This peak is followed by the peak in the electron-electron potential between the impact electron and the inner shell electron (yellow line). It indicates the rapid transit of the impact electron through the inner shell region. The steep decrease in the red, blue, yellow, brown and purple electron-electron potentials implies that the impact electron and the red valence shell electron leave the atom. Thus, the atom is reduced to a two-electron ion. Several peaks in the green electron-electron potential between the blue valence shell electron and the yellow inner shell electron imply that the blue valence shell electron collides several times with the inner shell electron, with rather large excursions from the ionic core. The blue valence shell electron eventually ionizes.

The electron-electron potential (in green) between the blue valence shell electron and the inner shell electron (in yellow) after $t = t_{Th}$ highlights the three features of the D1 mechanism; first, the initial collision between the impact electron and the red valence shell electron ionizes the later before $t = t_{Th}$; second, the impact electron excites the inner shell electron, when it transits rapidly through the inner shell region. Therefore, the two-electron ion that forms during the mechanism has net positive energy, which is redistributed between the inner shell electron and the blue valence shell electron through collisions; And third, the D1 mechanism relies
on a strong electron-electron correlation between the inner shell electron and valence shell electron, and, thus, without the existence of an inner shell electron, the \( D1 \) mechanism cannot occur. The schematic diagram in the top left of Fig. 6a summarizes the important features of the mechanism: one valence shell electron ionizes before \( t = t_{\text{Th}} \), while the other ionizes with a delay because of the electron-electron correlation with the excited inner shell electron.

2. \( D0 \)

A typical \( D0 \) mechanism is depicted in Fig. 7. In Fig. 7b, the impact electron (black line) first enters the inner shell region and collides with the inner shell electron. As a result, the impact electron gets deflected and leaves the atom. In Fig. 7c, this collision appears as a peak in the yellow electron-electron potential between the impact electron and inner shell electron. The ionized impact electron leaves behind an excited three-electron atom. The two valence shell electrons in the atom undergo multiple collisions with the inner shell electron, potentially with large excursions from the ionic core. In Fig. 7c, these collisions appear as successive peaks in the brown electron-electron potential between red valence shell electron and inner shell electron, and green electron-electron potential between the blue valence shell electron and the inner shell electron. Through collisions inside the excited atom, the red valence shell electron ionizes first and leaves behind an excited two electron ion. Eventually, the blue electron also ionizes.

The electron-electron potential (in brown) between the red valence shell electron and the inner shell electron, and the electron-electron potential (in green) between the blue valence shell electron and the inner shell electron beyond \( t = t_{\text{Th}} \), brings the three significant features of the \( D0 \) mechanism to light; first, none of the electrons ionizes from the atom before \( t = t_{\text{Th}} \), but the atom is left in an excited state, that is, with net positive energy.; second, the ionization occurs because of a redistribution of the net positive energy, through multiple collisions among the valence shell and inner shell electrons; and third, the magnitude of the electron-electron potentials of the valence shell electrons with the inner shell electron (brown and green electron-electron potentials) is greater than electron-electron potential between the two valence shell electrons (purple), which implies that the mechanism mainly relies on the electron-electron correlations of the valence shell electrons with the inner shell electron. Therefore, just like \( D1, D0 \) requires the existence an inner shell electron to take place. Moreover, \( D0 \) involves a complex (chaotic) choreography between the three electrons of the target.

E. Ionized Inner Shell Mechanisms

Ionized Inner Shell mechanisms involve the ionization of the inner shell electron. There are two types of Ionized Inner Shell mechanisms: Direct* and Delay* mechanisms, which are in turn composed of \(\text{TS1}^* \) and \(\text{TS2}^* \) mechanisms, and \(\text{D1}^* \) mechanisms, respectively. The electron dynamics in \(\text{TS1}^*, \text{TS2}^* \) and \(\text{D1}^* \) mechanisms are similar to the electron dynamics in \(\text{TS1}, \text{TS2} \) and \(\text{D1} \) mechanisms, respectively, with the exception that the inner shell electron is ionized in \(\text{TS1}^*, \text{TS2}^* \) and \(\text{D1}^* \) mechanisms. We use an asterisk in the nomenclature to mark the difference between the processes which do not (or very weakly) involve the inner shell electron.

The contribution of \(\text{TS1}^*, \text{TS2}^* \) and \(\text{D1}^* \) mechanisms to the probability of Ionized Inner Shell mechanisms is depicted in Fig. 2b. We notice that we have not mentioned any \( D0^* \) mechanisms because they occur with negligible probability in our calculations. This also explains why the probability of \(\text{D1}^* \) mechanisms overlaps the probability of Delay* mechanisms in Fig. 2b. Figure 2a illustrates that the Inner Shell Ionized mechanisms begin to occur at about \( \epsilon_0 = 80 \text{ eV} \), and are dominant over \(\text{TS1} \) and \(\text{D0} \) mechanisms at \( \epsilon_0 = 125 \text{ eV} \).

1. \(\text{TS1}^*, \text{Direct}^* \)

A typical \(\text{TS1}^* \) mechanism is depicted in Fig. 8. From the zoomed-out view of the electron trajectories in Fig. 8a, we see that this is an Ionized Inner Shell trajectory, because it leads to the ionization of the inner shell electron and a valence shell electron, trajectories of which are traced in yellow and red respectively. Fig-
FIG. 8: TS1\* mechanism at $\epsilon_0 = 125$ eV (see Sec. III A for details).

ure 8b shows that the impact electron travels into the inner shell region, undergoes a strong collision with the inner shell electron and the blue valence shell electron, and deflects out of the atom. As a result of its collision with the impact electron, the inner shell electron escapes the inner shell region, and, on its trajectory towards ionization, collides with the red valence shell electron. That collision sets the red valence shell electron also on a path towards ionization, and the collision resets the inner shell electron’s trajectory towards ionization. The schematic diagram in the top left corner of Fig. 8a summarizes the main features of the mechanism.

2. TS2\*, Direct\*

A typical TS2\* mechanism is depicted in Fig. 9. Again, the zoomed-out view of the electron trajectories in Fig. 9a illustrates that the trajectory corresponds to an Ionized Inner Shell mechanism, because it leads to the ionization of the inner shell electron and a valence shell electron, trajectories of which are traced in yellow and red, respectively. Figure 9b shows that the impact electron first collides with the red valence shell electron before traveling into the inner shell region. In the inner shell region, the impact electron collides with the inner shell electron. Next it collides with the blue valence shell electron, before deflecting out of the atom. As a result of their collisions with the impact electron, the inner shell electron and red valence shell electron ionize. In Fig. 9c, the collision between the impact electron and the inner shell electron appears as a peak in the electron-electron potential (in yellow) between the impact electron and the inner shell electron. The sequence and duration of the events in this mechanism are schematically depicted in the diagram in the top left corner of Fig. 9a.

3. D1\*, Delay\*

A typical D1\* mechanism is depicted in Fig. 10. From the zoomed-out view of the electron trajectories in Fig. 10a, we conclude that trajectory depicts an Ionized Inner Shell mechanism, because it leads to the ionization of the inner shell electron and a valence shell electron, which are traced in yellow and red respectively. As inferred from Fig. 10a, the impact electron travels into the inner shell region, collides with the inner shell electron and leaves the atom. As a result of that collision, the inner shell electron (yellow line) is set on trajectory towards ionization. On its path, the inner shell electron collides with the red valence shell electron and then leaves the atom. A two-electron ion results, in which both the electrons are valence shell electrons. When the red valence shell electron passes near the 2 a.u. boundary, the energy is redistributed between the two valence shell electrons, with a greater share belonging to the red valence shell electron. Therefore, the red valence shell electron ionizes. Information about the main actors in the collisions and the time scales involved in these collisions is contained in the schematic diagram in the top left corner of Fig. 10a.

As a side note, since this mechanism involves a direct ionization of the inner shell electron, followed by a relaxation towards a stable state by a subsequent ionization,
this is a classical equivalent of the Auger effect. From our analysis, we see that this is a very probable mechanism at impact energies above 100 eV, but it is not the dominant mechanism, since it appears that direct processes (as the combination of the two direct processes \(TS1^*\) and \(TS2^*\)) are at least as present as \(D1^*\).

F. Selecting mechanisms by varying the impact parameter \(y_{0,\text{in}}\)

*Inner Shell Capture, D1, D0, TS1*, TS2* and D1* mechanisms rely on electron-electron correlations with the inner shell electron. These correlations are pronounced because the impact electron excites or ionizes the inner shell electron when it transits through the inner shell region in these mechanisms. Therefore, by increasing the impact parameter, \(y_{0,\text{in}}\), the probabilities of occurrences of the six mechanisms that rely on electron-electron correlations with the inner shell electron might decrease. We decompose the probability curves in Fig. 11 for three ranges of impact parameters. We show that decomposition of probabilities for each of the three sets of impact parameters in the three panels of Fig. 11 where the panels Figs. 11a, 11b and 11c represent the probabilities for impact parameter \(y_{0,\text{in}}\) in \([0, 2)\) a.u., \([2, 3)\) a.u. and \((3, 5]\) a.u., respectively. Each panel in Fig. 11 follows the legend of Fig. 2.

The upper panels in Fig. 11 show that the probabilities of *Inner Shell Capture, D1* and *D0* mechanisms are progressively suppressed by one order of magnitude from the leftmost to rightmost panel. In other words, the probabilities of *Inner Shell Capture, D1* and *D0* mechanisms are significantly reduced when the impact parameter is increased from \([0, 2)\) a.u. to \((3, 5]\) a.u. Moreover, the lower panels in Fig. 11 show that the probabilities of *TS1*\(^*\), *TS2*\(^*\) and *D1*\(^*\) mechanisms are suppressed by four orders of magnitude from the leftmost panel to the rightmost one. This indicates that the probabilities of *TS1*\(^*\), *TS2*\(^*\) and *D1*\(^*\) mechanisms are drastically suppressed when the impact parameter increases from \([0, 2)\) a.u. to \((3, 5]\) a.u.

According to Fig. 11c, our model predicts that *Direct* mechanisms dominate the probability of double ionization of Mg by electron impact for large impact parameters. Furthermore, our model predicts that *TS1* mechanisms dominate all other mechanisms in large impact parameter scenarios. The impact electron misses the inner shell when it arrives at large impact parameters. All electron-electron correlations with the inner shell electron remain unchanged, and electron-electron correlations with the valence electrons drive the ionization mechanisms. Furthermore, the two valence shell electrons are likely to occupy opposite sites in the atom because of electron-electron repulsion. And, when the impact electron hits the atom at large impact parameters, the impact electron is likely to collide with only one of the valence shell electron, closest to it. Therefore, *TS1 Direct* mechanisms become dominant. By the same logic, if the impact electron arrives at small impact parameters, then electron-electron correlations with inner shell electron become significant and mechanisms other than the *Direct* mechanisms become dominant. Also, the impact electron is likely to collide with both valence shell electrons in small impact parameter scenarios. Consequently, *TS2* mechanism becomes the dominant *Direct* mechanism for small impact parameters, which is observed in Figs. 11a and 11b.

In general, different mechanisms are dominant at different impact energies as depicted in Fig. 2. Moreover, we find that different mechanisms can be made dominant by appropriately selecting the impact parameters. Therefore, the impact energy, \(\epsilon_0\), and the impact parameter, \(y_{0,\text{in}}\), may be tuned to selectively trigger or suppress certain double ionization mechanisms.

IV. DISCUSSIONS

A. Comparison with a two electron model

Up to now we have described our model and its two salient features: first, a successful qualitative reproduction of the probability of double ionization of Mg by electron impact, and second, the prediction about the underlying double ionization mechanisms. In this section, we compare our model and its implications with the two bound electron model of Mg in Ref. [32]. This comparison will highlight the importance of considering an inner shell electron and the importance of fully accounting for
Ref. [32] considers a two-active electron model to study the single and double ionization of Mg by electron impact. Ref. [32] also uses a purely classical mechanical approach to describe the double ionization processes and all electron-electron correlations taken into account. However, the model in Ref. [32] differs from the model in this paper in two ways; first, the atom consists of an ionic core and two valence shell electrons only, and second, a soft-Coulomb potential without effective charge governs the electron-core potential. Appropriately adjusting the soft-Coulomb parameters \( a \) and \( b \) alone guarantees no self ionization and non-empty ground states in the two-electron model in Ref. [32]. Our model considers an inner shell electron in addition to an ionic core and two valence shell electrons, and the existence of the inner shell electron necessitates an effective charge in the soft-Coulomb electron-core potential in our model.

The two-electron model in Ref. [32] explains the first increase in the double ionization probability and predicts that only TS1 and TS2 Direct mechanisms contribute to the double ionization probability. It also predicts that TS2 mechanisms are dominant at small impact parameters. The model reproduces the double ionization probability of Mg by electron impact for \( 20 < \epsilon_0 < 50 \) eV, but it fails to explain the knee in the double ionization probability. This indicates that the knee in the probability of double ionization of Mg by electron impact must be a consequence of electron-electron correlations with the inner shell electron(s). Figure 2, which shows the contribution of the Delay and the Ionized Inner Shell mechanisms to the increase in the double ionization probability for \( \epsilon_0 > 50 \) eV, substantiates this claim.

Also as a result of the differences between a two-electron and three-electron model, the former can only predict TS1 and TS2 Direct mechanisms, which require electron-electron correlations among an impact electron and two valence shell electrons only. Our model predicts six more mechanisms, other than TS1 and TS2, because our model accounts for all electron-electron correlations among an impact electron, two valence shell electrons and an inner shell electron.

Another consequence of the differences between a two-electron and three-electron models is the nature of mechanisms which underlie the probability of double ionization for \( 20 < \epsilon_0 < 50 \) eV. Ref. [32] finds that only the TS1 and TS2 mechanisms contribute to the double ionization probability in that domain of impact energies. However, we find that Inner Shell Capture mechanisms also contribute substantially to the total double ionization probability in the same domain of impact energies. For large impact parameters and in the intermediate range of impact energies, we find Direct mechanisms dominate over all electron-electron correlations with the inner shell electron.

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Delay mechanisms, which agrees with Ref. [32].

Altogether, the models in Ref. [32] and in this paper are different with respect to the number of components in the models and the structure of the soft-Coulomb electron-core potentials used in the models. The differences result in different sets of possible electron-electron correlations. The different sets of electron-electron correlations, in turn, affect the possible double ionization mechanisms, their probabilities and the probability of double ionization of Mg by electron impact. Some agreement can be found between these models in the intermediate range of impact energies and relatively high impact parameters (in order to suppress the role of the inner shell electron).

B. Relevance of the model with respect to experimental results

So far, the discussion focused on the double ionization of Mg by electron impact for $\epsilon_0 < 125$ eV. However, experiments on double ionization by electron impact are also conducted for $200 < \epsilon_0 < 5000$ eV. In order to check how our model compares with these experiments, we compute the probabilities of the eight double ionization mechanisms predicted by our model at $\epsilon_0 = 700$ eV. In what follows, we first explain the relevant experiments, and then link results from our model with experimental observations.

In the experimental measurements reported in Ref. [33], we observe an asymmetric differential cross-section in electron impact double ionization of Helium. In Ref. [34], it is suggested that this asymmetry implies the prominence of second or higher order mechanisms (all mechanisms but $TS1$ and $TS1^*$) over first order mechanisms ($TS1$ and $TS1^*$). We refer to Refs. [35][36] for a more detailed discussion on this issue. Moreover, in Ref. [37], it is suggested that non-first order mechanisms, other than $TS2$, may be responsible for the disagreement between their theoretical and experimental results. Hence, experiments on double ionization by electron impact suggest prominence of non-first order over first order double ionization mechanisms. So far such experiments probing the prominence of first and non-first order mechanisms have been conducted for Argon, Neon, molecular Nitrogen and Helium for impact energies approximately between 600 and 700 eV [36][38][41].

The probability of first and non-first order mechanisms at high impact energies of Hamiltonian (7), for $\epsilon_0 = 700$ eV, is shown in the form of a bar graph in Fig. 12. The vertical axis measures the contribution of each mechanism as a fraction of the total count of double ionization of Mg by electron impact. The left panel in Fig. 12 differentiates first order mechanisms, $TS1$ and $TS1^*$ (blue) from non-first order mechanisms, the remaining six mechanisms (black). The non-first order mechanisms are broken up into constituent mechanisms in the right panel in Fig. 12. The hatched sections in the bars represent the contribution from those mechanisms in which the inner shell electron is ionized, that is, from those mechanisms with '*' in their nomenclature. The left panel in Fig. 12 exhibits a clear agreement between results from our model and experimental observation, because our model predicts, as is observed in the experiments, that non-first order mechanisms are more probable than, and, thus, dominant over, first order mechanisms at 700 eV.

![FIG. 12: Relative prominence of double ionization mechanisms at $\epsilon_0 = 700$ eV. The first order mechanisms are the $TS1$ and $TS1^*$ mechanisms. The hatched section in the bars represents the mechanisms whose denominations contain '$*$'.](image)

V. CONCLUSION

We have introduced a model to study the double ionization of Mg atom by electron impact. In this model, we consider two valence shell electrons, one inner shell electron and an ionic core. The electrons are bounded to the core using a soft-Coulomb electron-core potential with an effective charge. We find that the probability of double ionization of the Mg atom as a function of the initial energy of the impact electron calculated from is qualitatively similar to that obtained from experiments. Our model enables us to analyze electron trajectories of double ionization mechanisms, and based on visual inspection of the electron dynamics in trajectories, we find eight dominant mechanisms - Inner Shell Capture, $TS1$, $TS2$, $D1$, $D0$, $TS1^*$, $TS2^*$, and $D1^*$. We group the eight mechanisms into four broad categories - Inner Shell Capture, Direct, Delay and Ionized Inner Shell mechanisms. We find that Delay and the Ionized Inner Shell mechanisms underlie the knee in the double ionization probability. We provide a detailed description of the electron dynamics in all the listed mechanisms, and end with two points of
and correlations with the inner shell electron give rise to the double ionization. Therefore, electron-electron amplitudes are consequences of electron-electron correlations with the inner shell electron. Both points of discussion may help motivate future experiments in the field of electron impact ionization. The Delay and the Ionized Inner Shell mechanisms are consequences of electron-electron correlations with the inner shell electron. Therefore, electron-electron correlations with the inner shell electron give rise to the Delay and Ionized Inner Shell mechanisms, which in turn contribute to building up the knee in the probability of double ionization of Mg by electron impact.

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