Non-perturbative scaling behavior for net ionization of biologically relevant molecules by multiply-charged heavy-ion impact

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Abstract

A recently developed model to describe proton collisions from molecules involving basic atoms such as hydrogen, carbon, nitrogen, oxygen and phosphorus (H, C, N, O, P) is extended to treat collisions with multiply charged ions. The ion-atom collisions are computed using the two-center basis generator method (TC-BGM), which has a proven track record of yielding accurate total cross sections for electron capture and ionization. The atomic net ionization cross sections are then used to assemble two models for ion-molecule collisions: an independent atom model (IAM) that follows the Bragg additivity rule (labeled IAM-AR), and also the so-called pixel-counting method (IAM-PCM). The latter yields reduced cross sections relative to IAM-AR near the maximum, since it takes into account the overlapping nature of effective cross sectional areas. The IAM-PCM for higher-charge projectiles leads to strong reductions of net ionization cross sections relative to the IAM-AR method, and is computed directly for projectile charges $Q = 1, 2, 3$. The scaling behavior of the IAM-PCM is investigated over a wide range of energies $E$, and at high $E$ it converges towards the IAM-AR. An empirical scaling rule is established which allows to reproduce these results based on proton impact calculations. Detailed comparisons are provided for the uracil target ($C_4H_4N_2O_2$), for which other theoretical as well as experimental results are available. Data are also shown for targets such as water ($H_2O$), methane ($CH_4$), adenine ($C_5H_5N_5$), L-valine ($C_5H_{11}NO_2$), and the nucleotide dAMP ($C_{10}H_{14}N_5O_6P$). Based on the scaling model derived from the IAM-PCM data it is shown how the experimental data for uracil and water bombarded by multiply charged ions can be reduced to effective $Q = 1$ cross sections respectively, and these are compared to proton impact data.
I. INTRODUCTION

The understanding of ionization in charged-particle impact on biologically relevant molecules is an important prerequisite for ion beam cancer therapy. In order to perform simulations of radiation damage caused by the projectile ions, the secondary electrons and molecular charged fragments produced during collisions [1] one first has to study the fundamental processes of ion collisions with molecules in the gas/vapor phase, and this has been the motivation for experimental and theoretical studies. Important target molecules in this context are water (for which theoretical studies can be performed as extensions of ion-atom collisions), and biomolecules that form the DNA and RNA. The RNA base uracil \( \text{(C}_4\text{H}_4\text{N}_2\text{O}_2) \) was chosen as a candidate for extensive experimentation and was reported on together with theoretical analyses [2, 3]. Theoretical treatment of ionization of these biomolecules escapes at this point the capabilities of sophisticated quantum-mechanical modelling, but first attempts have been made [4–6]. The net ionization cross sections for biomolecules are very large due to their size and number of available valence electrons.

In Ref. [7] an experimental (and theoretical) investigation of p-uracil collisions summarizes how the ionization cross section grows with the number of valence electrons. This idea also forms the basis of a theoretical approach that combines ion-atom scattering calculations within the continuum distorted wave with eikonal initial state approach (CDW-EIS), which also uses information from the molecular orbital energy-level structure to imprint some molecular character on the model. Another approach that was used with some success is an independent-molecule model, where parametrizations of experimental total ionization cross section data for proton collisions with small constituent molecules are used to assemble results for the uracil target [8].

Independent-atom models (IAM) were studied extensively for proton impact in our group [9–12]. On the basis of proton-atom collision calculations performed using the two-center basis generator method (TC-BGM) [13] one can form simply an estimate for the ion-molecule cross section by the Bragg additivity rule, and this was called the IAM-AR model. This somewhat naive addition rule should provide the correct high-impact-energy limit when the ionization cross section becomes small. A much more sophisticated IAM was introduced and tested in these works, which models the ion-molecule collision process for a given orientation of the molecule by considering the projectile ion as observing an
effective cross sectional area that is formed as the overlap of all atomic cross sections. This geometric overlap is easily calculated by pixelization, and therefore the method was called the pixel counting method (PCM). This method gives substantially reduced cross sections for net ionization and net capture as compared to the IAM-AR method, but goes over into it by construction when the cross sections become small and the overlap disappears, i.e., the measured area corresponds to the sum of all atomic cross sections. Quite relevant for the present work is the fact that the IAM-PCM results for net ionization in p-uracil collisions at the energies $E = 0.5, 1.0, 2.0$ MeV match the experimental data quite well, as does the CDW-EIS theory for proton impact [7]. In Ref.[11] a detailed comparison between IAM-AR, IAM-PCM and molecular-orbital energy based models is provided for proton-pyrimidine ($C_4H_4N_2$) collisions in Fig. 2a, and it is evident that the shape of the IAM-PCM cross sections begins to deviate at energies below $E = 1$ MeV/amu from AR and the other models. A comparison for the case of adenine molecules is provided in Fig. 3 of Ref. [12].

An important measure that helps to understand how theory and experiment match up is the scaling behavior of the ionization cross sections with projectile charge $Q$ and collision energy $E$ or impact velocity $v$. An experimental (and theoretical) investigation of charged-ion collisions with uracil [3] found that even in the MeV/amu collision energy regime perturbative, and even distorted-wave theory are not in good agreement. It is argued that the experimental cross section scales approximately as $\eta^{1.5}$, where $\eta = Q/v$ is the Sommerfeld parameter. The CDW-EIS theory overestimates the experimental data by more than a factor of two. Thus, it is important to investigate the situation using a theory that is not rooted in the Born series or its distorted-wave siblings. A recent IAM based on a stoichiometric argument and CDW-EIS theory for ion-atom collisions [14] makes an attempt at scaling by defining a reduced CDW-EIS cross section, which however fails to reconcile the proton impact measurements [7] with the $Q = 4−8$ data of Ref. [3] with a difference of about a factor of three (cf. Fig. 5 in Ref. [14]). This problem shows that the scaling problem has remained unresolved, and it therefore represents a major thrust for the present work.

A significant body of ionization data for atoms and molecules, including some biomolecules has been treated theoretically by a semiclassical method that represents an $\hbar = 0$ limit of quantum mechanics, namely the Classical Trajectory Monte Carlo (CTMC) method which simulates quantum mechanics by classical statistical mechanics. It allows to obtain differential electron emission cross sections and is non-perturbative. Molecules can be described by multi-
center effective model potentials adopted from quantum structure calculations. For the water molecule target bombarded by highly charged ions a detailed comparison against CDW-EIS results and experiments was carried out recently in a model that includes time-dependent screening parameters that change with electron removal from the molecule [15].

Experimental data for charged-particle impact with $Q = 1, 2, 3$ on water molecules have been obtained for a wide range of collision energies [16–20]. For higher charges $Q$ measurements are available at higher collision energies [21–23] in the form of differential electron emission cross sections together with CDW-EIS theory, and integration of the differential data determines total ionization cross sections.

Calculations based on the TC-BGM were carried out for proton-H$_2$O collisions by representing the self-consistent field molecular orbitals in terms of an expanded atomic oxygen basis generated by the optimized potential method of density functional theory [25]. Not all molecular orientations could be treated, but arguments were given how the total cross sections could be estimated on the basis of a subset of orientations. The method was applied successfully to the cases of proton projectiles [26] and a fragmentation model was developed and compared to experimental data [27]. The work was extended to He$^+$ collisions including the treatment of projectile electron loss [28]. The experimental work on Li$^{3+}$ collisions was supported using this methodology [20]. However, the extension to higher multiply charged projectiles turned out to be difficult (despite some additional encouraging results for He$^{2+}$ projectiles [29]), and the program was not pursued further in favor of the IAM-PCM approach. The main motivation for this was that TC-BGM ion-atom calculations are pushed to convergence including a proper representation of the discrete excited states of the constituent atoms. This turns out to be difficult for the ion-molecule TC-BGM calculations, in part due to the representation in terms of atomic oxygen basis states. A proper representation of the molecular discrete excitation spectrum requires a correlated quantum chemistry approach [30]. Thus, it is not clear whether a limited representation within the framework of the TC-BGM ion-molecule calculations is capable of separating accurately the continuum contributions from discrete excitations.

Other groups tested methods based on the semiclassical approach where the electronic wave function developed in a molecular orbital basis is propagated in time [31] and obtained different (higher) net ionization cross sections at low energies. A full three-center single-electron CTMC model potential was designed [32] that gained popularity within the community with
extensive sampling of molecular orientations. The potential was also used in numerical grid calculations \cite{33} at higher collision energies.

An attempt at scaling was recently made in the context of CTMC calculations \cite{33} by graphing the net ionization cross section divided by the projectile charge $Q$ against $E' = E/Q$. The CTMC data run approximately parallel to the CDW-EIS data and experimental data at high energies, and begin to deviate from experimental data as they approach $E' = 100 \text{ keV/amu.}$ from above. The question whether the cross section data can be scaled within the maximum region therefore also remains open up to now.

A recently introduced CTMC mean field model with dynamic screening on projectile and target has been used to make extensive comparisons with the data and explored the question of saturation behavior in the net cross sections by looking at them as a function of the Sommerfeld parameter $\eta = Q/v$ \cite{35}. No obvious scaling was observed in these calculations when looking at the entire energy range in the data.

The paper is organized as follows. In Sect. II we introduce the theoretical basis for the current work. Sect. IIA presents new results for TC-BGM ion-atom calculations for He$^{2+}$ and Li$^{3+}$ projectiles; for atomic hydrogen targets the results are compared with theory and experiment. In Sect. IIB we demonstrate the scaling behavior of ion-molecule collisions with respect to projectile charge $Q$ and collision energy $E$ in the IAM-AR and IAM-PCM approaches for the targets uracil and water. A parametrization of the scaling behavior is then used to generate cross section for arbitrary projectile charge on the basis of IAM-AR and IAM-PCM proton impact data. Sect. III serves to provide a detailed comparison of the scaling behavior: in Sect. IIIA for uracil and in Sect. IIIB for water. The paper ends with a few concluding remarks in Sect. IV. Atomic units, characterized by $\hbar = m_e = e = 4\pi\varepsilon_0 = 1$, are used unless otherwise stated.

II. MODEL

In this section we present arguments that lead to the empirical scaling models for the net ionization cross sections according to the IAM-AR and IAM-PCM models using the uracil target molecule C$_4$H$_4$N$_2$O$_2$ as an example. It is a good example from several points of view: on the one hand it contains four different atoms, and ionization shouldn’t be dominated by any one of them, and on the other hand it is a target for which systematic experimental
studies in the gas phase were carried out. The model will then later be confirmed by looking at the water molecule (vapor phase), for which again, a number of experiments with differently charged projectile ions are available.

A. Ion-atom collisions

Results from new TC-BGM solutions of the time-dependent Schrödinger equation (reduced to density functional theory at the level of an independent electron model) are presented here for collisions of multi-charged ions (bare charges $Q = 1, 2, 3$) from atoms that form constituents of the biologically relevant molecules, i.e., (H, C, N, O, P). The case of proton projectiles was treated explicitly in Ref. [11], so those results are not repeated here. The TC-BGM is implemented in projectile potential $W_p$ hierarchy which converges well and adds 123 pseudostates to span the complementary space (for all projectile and target combinations) to the number of shells treated explicitly for target and projectile. For the target the $n_T = 1, ..., 5$ shells are included in all cases. For the projectile we include $n_P = 1, ..., 6$ in the case of He$^{2+}$ projectiles and $n_P = 1, ..., 7$ for Li$^{3+}$. For ionization we observe a simple trend for high collision energies, namely $\sigma_C = \sigma_N = \sigma_O = 4\sigma_H = 2\sigma_P/3$, as was found previously for proton impact [11].

Fig. 1 displays the net ionization cross sections that form the ingredients for direct application of the IAM-AR and IAM-PCM calculations described in the next subsection, from which then a scaling model is derived and verified for ion-molecule collisions. The results demonstrate the non-perturbative nature of these calculations which were obtained using an exchange-only density functional (optimized potential) method for the target atoms other than hydrogen.

While a detailed comparison of charge-state correlated cross sections would benefit from dynamical screening, the net ionization cross sections are deemed reliable from such a frozen-potential approach for proton impact. For C$^{4+} −$ Ne collisions at $E = 20$ keV/amu we found in Ref. [36] that the total ionization cross sections are reduced by 25% when response is included for both the target and projectile potentials. For higher energies the effect is less pronounced. It can be debated whether IAM-PCM can be based on atomic calculations with response, and our current approach is that we use frozen-potential calculations even for higher projectile charges.
The basic shapes of these cross sections roughly do not change with increasing projectile charge $Q$, and there are differences in shape for the various target atoms. In the limit of high energies we observe the known scaling with $Q^2$, but in the vicinity of the maxima in the cross sections this scaling factor is reduced. There is a small, but noticeable shift in the position of the maxima as $Q$ increases, and then at low collision energies ionization becomes less efficient with increasing projectile charge so that the net ionization cross sections become comparable even though $Q$ is increased. At even lower energies one can understand the inefficiency of ionization by multiply charged projectiles not only as the competition with capture channels, but also on the basis of (avoided) adiabatic energy level crossings. The TC-BGM calculations are deemed capable of describing such behavior reasonably well. A detailed comparison of shapes for $Q = 2, 3$ then shows a steepening of the rise in the cross section starting from low $E$ values.

The ion-atom collision cross section calculations are challenging and require significant computing resources. Comparison with the experimental data for atomic hydrogen is possible (for He$^{2+}$ and Li$^{3+}$ projectiles the data are from Ref. [38, 39] and [40] respectively) The comparison shows that converged theory is slightly higher than experiment. For p-H collisions (cf. Fig. 6 in Ref. [11]) a similar situation arises. The agreement of our calculation with the convergent close coupling approach (CCC) for He$^{2+}$ impact [37] is as good as can be expected from two complementary large-scale approaches.

B. Scaling model for ion-molecule collisions

Fig. 2 forms the basis for the discussion. The empirical scaling models aim to capture the essence of the data in the non-perturbative regime where the net ionization cross sections have their maxima. The left panel of Fig. 2 displays the net ionization cross sections for the two models, which have the following characteristics: (i) with increasing projectile charge $Q = 1, 2, 3$ the difference between IAM-PCM (solid lines) and IAM-AR (dashed lines) results grows dramatically. This is caused by the increase in the atomic cross sections with projectile charge $Q$ (except at the lowest collision energies) and thus an increase in the overlap effect which the IAM-PCM takes into account; (ii) the position of the maxima in these cross sections changes very similarly with $Q$ for the two models, which implies that a common energy scaling can be applied, particularly if one is interested in scaling the cross sections in
FIG. 1. In the left panel the total (net) ionization cross sections for He$^{2+}$ ions colliding with atoms H, C, N, O, P is shown (solid lines) as calculated with the TC-BGM using basis sets described in the text, while the right panel is for the case of Li$^{3+}$ projectiles. The dashed lines indicate how in the limit of high $E$ the merging results for the heavier target atoms can be related to the values for atomic hydrogen by using a scale factor. The He$^{2+} - H$ results are compared with the convergent close coupling calculation of Ref. 37, and the experimental data are from Ref. 38, 39 in this case, and from Ref. 40 for Li$^{3+} - H$ collisions.

in the vicinity of the maximum.

For the IAM-AR (middle panel of Fig. 2) the energy scaling for the maximum implies that it is useful to introduce

$$E' = E \frac{Q}{Q_{Q=1}}$$

(1)

in order to find a common curve, and that the net cross sections should be scaled in accord with

$$\sigma_{Q=1}^{\text{AR}}(E') = \sigma_{Q}^{\text{AR}}(E) \frac{Q}{Q_{Q=1}}$$

(2)

in order to obtain a single universal curve for $\sigma_{Q=1}^{\text{AR}}(E')$ where the only $Q$-dependence is through the scaled energy $E'$. Here $\sigma_{Q}^{\text{AR}}(E)$ is the IAM-AR net ionization cross section for a projectile with charge $Q$ and energy $E$ colliding with the molecule.

In the right panel of Fig. 2 the IAM-PCM data are presented while applying the same
FIG. 2. The left panel shows the total (net) ionization cross sections produced from uracil by charged-ion impact for $Q = 1, 2, 3$ as calculated in the IAM-PCM (solid lines), and in the IAM-AR (dashed lines). The middle panel shows a scaled net ionization cross section for the IAM-AR model (dashed lines falling onto a single curve to within 5% for $E > 20$ keV/amu), and the the IAM-PCM calculations which do not obey this scaling behavior in the energy range shown. The right panel shows a scaled net ionization cross section that works for the IAM-PCM (solid lines agreeing to within 5% in a range enclosing the maximum, i.e., for $30 \leq E \leq 500$ keV/amu).

energy scaling. Since in the vicinity of the maximum the IAM-PCM cross section grows differently with $Q$ than the IAM-AR result quite a different scaling factor is needed: $Q^{-0.75}$ vs $Q^{-4/3}$. We then arrive at

$$\sigma_{Q=1}^{\text{PCM}}(E') = \sigma_{Q}^{\text{PCM}}(E) \cdot Q^{-0.75}.$$  \hspace{1cm} (3)

If the scaling works the reduced cross sections $\sigma_{Q=1}^{\text{AR}}(E')$ and $\sigma_{Q=1}^{\text{PCM}}(E')$ must correspond to the calculated cross sections for proton impact at energy $E'$, and should merge in the limit of high $E'$. We note that others have made attempts at scaling ion-molecule cross sections using classical-trajectory calculations [34] or distorted-wave models [14], and these attempts have so far not led to an understanding of scaling in the regime of the cross section maxima, but have focused on the high-energy behavior.

The present calculations obey the following behavior in the limit of high energies: the IAM-PCM goes over into the IAM-AR in this limit because the cross sections become small. Where exactly this occurs depends on the value of the projectile charge $Q$. The left panel of Fig. 2 indicates that for protons this merge happens at about $E = 2000$ keV/amu, and at substantially higher energies for $Q > 1$ to the point where for these higher projectile charges
the overlapping effect contained in the IAM-PCM cannot be ignored in the regime of interest (below 10 MeV/amu).

The scaling behavior of the IAM-AR is compatible with the behavior known for the ion-atom cross sections, i.e., the Bethe-Born limit which is obeyed by the TC-BGM calculations\cite{10,11}. Since this limit applies to proton-atom collisions we can argue that for any molecular target the IAM-AR net ionization cross section for a projectile of charge $Q$ obeys

\[ \sigma_{Q}^{\text{AR}}(E) \rightarrow Q^{4/3} \frac{A \ln E' + B}{E'} = Q^{4/3} \frac{A \ln \left( EQ^{-2/3} \right) + B}{EQ^{-2/3}} = \frac{Q^2}{E} \left( A \ln E + B' \right), \tag{4} \]

and therefore is proportional to the square of the projectile charge.

The right panel of Fig. 2 displays an interesting phenomenon: while the IAM-PCM results do go over into the corresponding IAM-AR ones at the highest collision energies there is a cross-over phenomenon connected with the shape change. While the IAM-PCM result for protons merges with the IAM-AR result smoothly, for the cases of $Q = 2$ (green) and $Q = 3$ (red) we find extended regions around $E = 1$ MeV/amu with a characteristic slope that is not as steep as the perturbative high-$E$ result. This will turn out to be of even bigger importance discussed below in the context of $Q = 4−8$ projectiles at $E = 1−6$ MeV/amu. At high energies the pure IAM-PCM scaling has to fail because it takes into account some saturation in order to describe the fact that the height of the cross section at maximum does not increase with $Q$ as rapidly as at high energies. Therefore, a parametrization is introduced to switch from the IAM-PCM form of scaling which works at intermediate energies $E$ to the IAM-AR scaling.

Now that we understand the scaling behavior of the IAM-PCM cross sections we proceed with developing a parametrization that will allow to predict cross sections for collisions with higher projectile charges $Q$ for which the computation of ion-atom collisions using the TC-BGM becomes more challenging. The idea is to find a parametrization that allows to derive $Q$-fold charged projectile collisions from molecules on the basis of proton-impact collisions (at scaled energies), confirm its validity for $Q = 2, 3$ and then compare with experiment which is available for $Q = 4−8$. We can expect such empirical scaling to work at energies $E > 40$ keV/amu on the basis of the data shown in the right panel of Fig. 2. The scaling of the IAM-PCM cross section maxima with $Q^{0.75}$ is an empirical result for uracil. In general, we assume scaling with $Q^\beta$, with the implication that $\beta = 0.75$ for uracil and that it can have different values for other molecules where the cross sectional area overlap effects
are different.

In order to accommodate the change in behavior which is apparent in the right panel of Fig. 2 we parametrize the representation of the IAM-PCM cross section by a switching function which keeps the IAM-PCM scaling in the vicinity of the maximum, but then with increasing collision energy shifts over to the known IAM-AR scaling of the cross sections. This is not an attempt to modify the IAM-PCM result, however, it is just a convenient way to represent it making use of both scaling regimes obeyed by the IAM-PCM calculations.

The switching function is defined as

\[ a(E) = \left[ 1 + \left( \frac{E/E_0}{Q-1} \right)^{\frac{1}{\sqrt{Q}}} \right]^{-1}, \]

(5)

with the switching point chosen as \( E_0 = 2000 \text{ keV/amu} \) and \( Q \geq 2 \). The parametrization of the IAM-PCM net ionization cross section scaling now takes the form

\[ \sigma_{PCM}^Q(E) = a(E)Q^\beta \sigma_{PCM}^{Q=1}(E') + (1 - a(E))Q^{4/3} \sigma_{AR}^{Q=1}(E') . \]

(6)

This parametrization of the IAM-PCM cross sections for \( Q \)-fold charged ion impact in terms of the proton-impact cross sections evaluated at \( E' \) depends on a single parameter \( \beta \) whose value is connected with the ‘density’ of independent atoms, and which needs to be determined by tracking the movement of the maxima with charge in the IAM-PCM cross sections. A discussion of detailed results for the uracil target (for which \( \beta = 0.75 \) was shown to work) follows, and a table of values of \( \beta \) for other molecules is given further below.

In Fig. 3 the effectiveness of the parametrization for energies \( E > 40 \text{ keV/amu} \) is demonstrated. The IAM-AR results are reproduced by the simple scaling law, as suggested by data shown in the middle panel of Fig. 2. The IAM-PCM scaling results (crosses) track the data well around the maxima in the cross sections, but show some weakness at low energies (\( E < 40 \text{ keV/amu} \)) for reasons discussed before. Ionization by highly charged ions becomes ineffective at low energies as one moves towards the quasi-molecular regime and due to increasing competition from capture channels.

Given that the IAM-PCM cross section for proton impact converges to the IAM-AR cross section at energies of scale \( E_0 = 2000 \text{ keV/amu} \) it is possible to simplify the parametrization and eliminate the IAM-AR cross section \( \sigma_{AR}^{Q=1} \) while admitting a small error. This is due to the fact that the switching function (5) deviates significantly from unity only at very high
FIG. 3. Net ionization cross sections for projectile charges $Q$ impinging on uracil. Solid and dashed lines are the IAM-PCM and IAM-AR calculations respectively for $Q = 1, 2, 3$. Solid circles: IAM-AR scaling prediction based on $Q = 1$ IAM-AR calculations. Crosses: IAM-PCM scaling predictions based on $Q = 1$ IAM-PCM calculations which track the respective maximum in the cross section but fail at high energies $E$. Open circles: Parametrization results for the IAM-PCM cross sections which switch from IAM-PCM scaling for the maximum to IAM-AR scaling for the high-$E$ regime. The radius of the open circles corresponds to a deviation of 5% in the cross section.

collision energies $E \gg E_0$. This simplified parametrization then becomes

$$\sigma^\text{PCM}_Q(E) = \left[ a(E)Q^\beta + (1 - a(E))Q^{4/3} \right] \sigma^\text{PCM}_{Q=1}(E') .$$

This form allows one to take experimental data for $\sigma^\text{net}_Q(E)$ and relate it directly to the proton impact result $\sigma^\text{net}_{Q=1}(E')$ in order to test whether the scaling prediction from the IAM-PCM theoretical model translates into measured physical evidence. The only required additional piece of information for a given molecule is the $\beta$ parameter which has to be found on the basis of IAM-PCM calculations for $Q = 1, 2, 3$. A list of parameter values is given in Table I for a few characteristic examples. The value of $\beta$ is larger for simple molecules such as water
FIG. 4. Same as in Fig. 3 but for the water molecule target.

or methane, and decreases when the number of constituent atoms increases.

| formula    | name       | group         | β     |
|------------|------------|---------------|-------|
| H₂O        | water      |               | 1.1   |
| CH₄        | methane    | alkane        | 1.05  |
| C₄H₄N₂O₂  | uracil     | pyrimidine    | 0.75  |
| C₅H₅N₅   | adenine    | purine        | 0.7   |
| C₅H₁₁NO₂  | L-valine   | amino acid    | 0.65  |
| C₁₀H₁₄N₅O₆P | dAMP | nucleotide   | 0.6   |

We conclude this section by a demonstration of the effectiveness of the parametrization for the water molecule target in Fig. 4 for which the β parameter is larger than for uracil. We begin by noting that the p – H₂O IAM-PCM cross sections have been corrected compared to our previous work [9, 10] in that the correct bond length has been implemented here.
We have also tested the IAM-PCM calculated cross sections against an exact geometric calculation of the overlapping circular areas which is possible for a triatomic molecule, and found that the IAM-PCM implementation matches the geometric calculation to excellent precision.

As a consequence of the corrected bond length error the IAM-PCM net ionization cross section now is further reduced around the maximum as compared to the IAM-AR calculation and the IAM-PCM and IAM-AR results now merge only at an energy of \( E = 1000 \text{ keV/amu} \).

We also verified that \( \text{H}_2\text{O} \) was the only target molecule for which a bond length error happened (it was an angstrom vs atomic unit problem, and the previously made error is apparent in Figs. 2a and 3a of Ref. [10]).

Concerning the comparison of scaling behavior for uracil (\( \beta = 0.75 \)) and water (\( \beta = 1.1 \)), i.e., comparing Fig. 3 to Fig. 4 we can provide the following comments. In the case of water the larger \( \beta \)-parameter value implies that the feature associated with the maximum in the net proton-molecule collision (for which IAM-PCM, i.e., atomic cross sectional area overlap is giving the strongest effect) acquires a higher scaling with \( Q \). The progression of cross section values at maximum is (in units of Å\(^2\)) \{4.6, 10, 15\} for \( Q = \{1, 2, 3\} \). For the uracil target the corresponding progression is \{19, 32, 44\}. The decrease of the \( \beta \) value with the complexity of the molecule is connected with the fact that overlap effects become more important with the growing number of atoms and the IAM-PCM takes this into account.

### III. COMPARISON WITH EXPERIMENTS

#### A. Collisions with uracil (\( \text{C}_4\text{H}_4\text{N}_2\text{O}_2 \))

Total ionization cross section measurements for uracil targets have a bit of a history. Net ionization can be obtained by collecting all electrons produced during the collision, a measurement process known from ion-atom collisions [41] (called \( \sigma_- \)), but that is not how all the data were obtained. Time-of-flight mass spectroscopy was used to determine the fragment yields, and some analysis went into finding the fragment distributions following electron capture versus direct ionization [42] for proton impact at velocities that match those of the valence electrons (20-150 keV/amu). Doubly charged uracil is not observed, so it is likely to fragment before the mass analysis as observed by coincidence spectroscopy of
fragments [1]. This must also be the case for higher-projectile charge impact, such as C^{4+} and O^{6+} [2, 3], where doubly-charged uracil is absent from the mass spectra. Tabet et al. [43] found very large values for the direct ionization and capture cross sections in this energy range, for $E = 80$ keV proton impact direct ionization is reported to reach $177 \pm 35 \text{ Å}^2$. Itoh et al. [7] determined absolute differential cross sections at higher energies by detecting the electron yield and by comparing with calculations, they demonstrated that the high cross section value of Ref. [43] was in conflict with their findings for $\sigma_-$.

The heavy ion measurements in the same velocity regime [2, 3] were normalized by using features in the differential electron emission: KLL Auger lines from the constituent atoms C, N, O were observed. Absolute normalization at a high energy (3.5 MeV/amu) was obtained with an uncertainty estimated at $\pm 20\%$, and relative uncertainties for the data were assessed at $\pm 12\%$. The total cross sections were obtained by measuring the fragment yields with some uncertainty coming from $N_2^+$ contributions from collisions with the background gas. Unlike the proton impact measurements of Ref. [43], however the projectile charge state after the collision was not determined. To ensure a proper separation from ionization and fragment production by pure capture processes coincidence with an ejected electron was imposed. This procedure includes electron production from transfer ionization processes, but if one then adds up all charged fragments one overestimates the contribution from transfer ionization to net ionization (by a factor of two for the process where one electron is captured, and one transferred to the continuum, and by more for higher-order capture plus higher-order ionization). At the higher velocities ($v = 7 - 15$ Bohr units) this problem does not exist, since capture plays a small role (except for capture from innermost shells, but these are small contributions).

We now confront the predictions from the IAM-PCM scaling model which was established on the basis of $Q = 2, 3$ calculations with experimental and theoretical data for higher projectile charges $Q = 4, 6$ for which experimental data exist both below the predicted maximum and above [2]. The comparison provided in Fig. 5 reveals the following: at the high-energy end, i.e., for $E > 1000$keV/amu the scaled IAM-PCM cross section does agree with the experimental data for both projectile charges almost within error bars, but shows a somewhat different energy dependence. The energy dependence of the experimental data follows the trend given by CTMC, CB1 (first Born model with correct boundary condition), and CDW-EIS in this regime, although these theories yield higher cross section values. We
note that both projectiles do have fully populated K-shells, i.e., we do not expect major anomalies in the cross sections from Auger processes.

At the lower end of collision energies, below $E = 100 \text{ keV/amu}$ we find the comparison to be more complicated. For the case of $Q = 4$ the present results join the two regimes of experimental data reasonably well, but fall short at the lower end where the observed scaling behavior becomes less accurate as discussed in Section 11B. All other theories than scaled IAM-PCM provide estimates higher by at least a factor of four, with CDW-EIS doing better, but showing an energy dependence that differs from the experimental and from the scaled IAM-PCM data trends. Clearly, the situation calls for experiments to map out the location of the maximum in the net ionization cross section.

The comparison with the case of $Q = 6$ seems to warrant a different conclusion: the experimental data are consistently higher than the scaled IAM-PCM results by a factor of two in the regime to the left of the maximum in the IAM-PCM data. The experimental data trend is supported by the CTMC calculation, which however failed in the case of $Q = 4$. Again, we call for experiments in the regime that connects the low-$E$ and high-$E$ data shown in Fig. 5. As explained at the beginning of the subsection the low-energy data in Ref. [2] may be affected by over-counting of transfer ionization contributions, and this may be a bigger problem for the system with the higher projectile charge, but we have no further support from calculations to make this case. Therefore, the scaling of the net ionization cross section with $Q$ for impact velocities that match the valence shell orbital speeds remains a mystery.

Another non-perturbative calculation that could be trusted in this regime, the CTMC calculation of Ref. [44] matches the $Q = 6$ experimental data (right panel) at low energies, then joins the CDW-EIS result which is a factor of two above the high-energy data. For $Q = 4$ (left panel) this calculation overestimates the data over the entire energy range, and still matches the CDW-EIS calculation at high $E$. The comparison with the CTMC calculation is interesting, since (as explained in Ref. [45]) the calculation makes use of an effective multi-center potential that matches quantum chemistry calculations, uses molecular orbital energies, and performs random molecular orientation sampling.

In Fig. 6 we present a comparison with many available experimental ion-uracil ionization cross sections at high impact energies in order to explore how the data scale. The most striking feature is the difference in the energy dependence between the case of $Q = 1$ and the higher projectile charges in the IAM-PCM. While the proton impact data are described well
FIG. 5. Net ionization cross sections for projectiles with charges $Q = 4$ (left panel) and $Q = 6$ (right panel) impinging on uracil. Solid and dashed blue lines are the IAM-PCM and IAM-AR calculations respectively obtained by scaling the proton impact cross sections. The experimental data (diamonds with error bars) are for $C^{4+}$ projectiles (left panel) and for $O^{6+}$ projectiles (right panel) [2]. Also shown are theoretical results from the classical trajectory method (CTMC) [44], the first-order Born method with corrected boundary conditions (CB1), and from distorted wave theory CDW-EIS (both from Ref. [2]).

by both our results and the CDW-EIS model [14], the latter retains the energy dependence for the higher charges $Q$. The difference in the energy dependence between $Q = 1$ and the higher projectile charges for IAM-PCM is a consequence of the overlap effect when the effective cross sectional area is computed: with increasing cross sections the areas representing the individual atoms increase, thereby causing more overlap. Thus, we observe that while in the proton impact case the dependence still follows somewhat the proton-atom cross section energy dependence a turnover happens between $Q = 1$ and $Q = 4$ in the IAM-PCM results. Note that all $Q > 1$ cross sections can be obtained simply from Eq. (7) based on the input for
FIG. 6. Experimental net (total) ionization cross sections for uracil bombarded by projectiles with charges $Q = 1$ (Ref. [7]) and $Q = 4 - 8$ (Ref. [2, 3]) are compared to the present IAM-PCM results. The proton impact data are multiplied by a factor of four for comparison purposes and are shown in grey. The highly charged projectiles are distinguished by color (red, blue, green, magenta, black) for $Q = 4 - 8$ respectively, and the projectile ions C, O, F are represented by diamonds, triangles and squares respectively. For clarity the experimental data are shown without uncertainties which are estimated to be $\pm 25\%$ for the $Q \geq 4$ data.

IAM-PCM which are given in Table A1 of Ref. [11]. The slightly more accurate data based on Eq. (6) require the IAM-AR cross sections which can be assembled from ion-atom cross sections (there is no orientation dependence in this IAM, very much like in the CDW-EIS based calculations used, e.g., in Ref. [14]). Proton-atom cross sections that are input to our IAM-AR result can be read off Fig. 6 in Ref. [11].

How well do the IAM-PCM results compare to the experimental data? Given that absolute normalization of ionization cross sections is a challenging task the error estimates of the data ($\pm 25\%$) are dominated by the normalization procedure. For some of the data the slope of the IAM-PCM results matches experiment, particularly for $O^{5+}$ and $C^{4+}$ projectiles over a wide
FIG. 7. The experimental net (total) ionization cross sections for uracil bombarded by projectiles with charges $Q = 4 - 8$ (Ref. [2, 3]) are turned into a reduced cross section $\sigma_{Q=1}$ using the simplified IAM-PCM scaling formula (7) and compared to the IAM-PCM (solid line) and IAM-AR (dashed line) proton-uracil calculations. The three experimental proton impact data points from Ref. [7] are also shown. The symbols follow the same patterns as used in Fig. 6.

range of energies. At the highest energies measured the experimental data are sharply lower which does not help with the assessment as to whether the data follow the present IAM-PCM trend or not. Concerning the $Q$ dependence of the magnitude of the cross sections we observe that IAM-PCM matches the experimental data reasonably well, while the CDW-EIS and CTMC models show serious discrepancies, which is highlighted in greater detail in Fig. 8 below.

We now use the scaling behavior found on the basis of explicit IAM-PCM calculations for $Q = 1, 2, 3$ (with some confirmation from $Q = 6$, which is not shown) to extract a reduced $Q = 1$ cross section from the experimental data for $Q = 4 - 8$, and compare the result with the IAM-PCM $Q = 1$ cross section and show the IAM-AR $Q = 1$ result for reference, as well. Fig 7 reveals a few rather interesting facts: the proton impact data at
E = \{0.5, 1.0, 2.0\} \text{ MeV} \text{ span the entire range of high-energy data from Ref. [2], namely 1 to 6 MeV/amu. A good part of the high-energy } Q = 4 - 8 \text{ experimental data is mapped into a band more or less between the IAM-PCM and IAM-AR } Q = 1 \text{ cross sections, but the highest points near } E' = 2 \text{ MeV/amu (from } \text{C}^{4+} \text{ and } \text{C}^{5+} \text{) fall below. Data which would cover the } E' = 50 - 400 \text{ keV/amu would be most welcome in order to settle the question whether the IAM-PCM prediction (solid line, which appears to work for } \text{C}^{4+} \text{ projectiles, but not for the case of } \text{O}^{6+} \text{) is indeed correct. Proton impact data below } E = 500 \text{ keV/amu would also be most welcome.}

In order to provide more clarity concerning the comparison between theories and experiment at high energies we show in Fig. 8 the data separated by projectile charge using a semi-
logarithmic representation. This figure establishes how well the experimental data are reproduced on an absolute scale by the IAM-PCM, and how far they are from the IAM-AR scaling, which matches the behavior of the CDW-EIS and CTMC theories. The figure also demonstrates how the data really do depend on the charge state \( Q \) with very little dependence on the nuclear charge of the projectile. Some experimental data appear to deviate from the IAM-PCM results at high energies (as commented upon before), notably for \( C^{5+} \), but no such deviation is visible in the same energy range for \( O^{7+} \). It remains an open question whether there are physics reasons for such a deviation from the proposed scaling behavior for particular projectiles or not.

B. Collisions with water vapor (\( \text{H}_2\text{O} \))

In Fig. 9 the IAM-PCM data for proton impact, and the scaled IAM-PCM data for higher \( Q \) are compared to available measurements. The proton impact data (shown as a black curve) which form the input for the scaled cross sections are consistent in shape with the three available sets of measurements [16–19] for impact energies \( E > 30 \text{ keV/amu} \). They agree very well in magnitude with the data of Luna et al [19], and are a bit lower than the other data.

For smaller energies \( E < 30 \text{ keV} \) they do not follow the experimental data but drop faster as \( E \) decreases - a property they share with another non-perturbative, albeit classical calculation [32]. At these lower energies the IAM cross sections drop not only more rapidly than the experimental data, but also than TC-BGM ion-molecule calculations [26], and even more so than the molecular-orbital based semiclassical calculation of Ref. [31], which exceeds the experimental data in this range. One may draw the conclusion that molecular effects due to the orbital energy level structure are unlikely to be captured by an IAM.

The scaled IAM-PCM net ionization cross sections for \( Q = 2 \) (shown in blue) display a similar behavior, and fall below the experimental data for \( \text{He}^{2+} \) more markedly for \( E < 80 \text{ keV/amu} \). Nevertheless, it should be noted that the position of the maximum in the cross section agrees well. Again, the TC-BGM ion-molecule calculation [29] does not suffer from this drop at low \( E \). At the highest energies the IAM-PCM results match very well with the two measured points of Ref. [47].

For \( Q = 3 \) (shown in green) there is agreement with the \( \text{Li}^{3+} \) experimental data con-
FIG. 9. Net ionization cross sections for projectiles with charges $Q = 1, 2, 3, 6, 8, 13$ impinging on water vapor. Solid lines are the scaled IAM-PCM results (except for $Q = 1$). The experimental data are for: $Q = 1$ shown as black solid squares [16], circles [46], triangles [19], diamonds [18]; $Q = 2$ shown as blue solid squares [16], circles [17], open diamonds [47]; $Q = 3$ shown as green solid triangles [20]; $Q = 6$ shown as open red circles (6 MeV/amu): [48], (4 MeV/amu): [23]; $Q = 8$ shown as open purple triangles [22, 24]; and $Q = 13$ shown as an open magenta square [23].

cerning the magnitude of the cross section for for $E > 80$ keV/amu, while the shape of the experimental data pattern differs somewhat from the scaled IAM-PCM results. The fact that the experimental data for He$^{2+}$ and Li$^{3+}$ projectiles show such similar magnitudes cannot be explained with scaling behavior, and should be resolved by further experimentation. At the lowest energies the experiments of Luna et al [20] may suffer from a shortfall due to transfer ionization processes yielding two protons as fragments (only one of them would be detected).

For higher-$Q$ projectiles comparison with the experimental data for C$^{6+}$ [23, 24, 48], O$^{8+}$ [22, 24], and Si$^{13+}$ [23] is very satisfactory with some tension with the O$^{8+}$ data. Overall we find that the proposed IAM-PCM scaling with projectile charge and energy works well for the water molecule target and is strongly supported by experiment for medium to high
FIG. 10. Reduced ($Q = 1$) net ionization cross section obtained from experimental data for multiply charged ions colliding with H$_2$O which are obtained on the basis of the simplified IAM-PCM scaling formula (7). The solid and dashed lines are the IAM-PCM and IAM-AR results for $Q = 1$ respectively. The experimental data are from the sources cited with Fig. 9 and share the symbol patterns, but uncertainties are omitted here.

energies. Note that for $Q \geq 3$ the IAM-PCM (and experimental) data have not yet reached the $Q^2$-proportional scaling of the Bethe-Born limit at the right end of Fig. 9.

In Fig. 10 the IAM-PCM scaling behavior of Eq. (7) is tested and compared against the IAM-PCM calculation for $Q = 1$, while the IAM-AR result is provided for reference. As was the case for the uracil target (Fig. 7) the experimental data at high energies show scatter, but follow the trend well - especially since there are discrepancies for different data taken for identical projectiles. At energies below 200 keV/amu the two IAM results separate as one approaches the maximum. The Li$^{3+}$ impact data 20 which were compared against TC-BGM ion-molecule calculations (and which are also compared with in the recent CTMC mean-field model work of Ref. 35) provide some support for the proposed scaling. At energies below $E' < 50$ keV/amu one may want to be critical of the IAM approach and of CTMC, since
these models cannot describe the molecular effects associated with the electronic structure of the particular molecule in question. At higher energies one can think of IAM-AR results as representing an upper bound to the reduced cross section, while IAM-PCM may be on the low side, since the effect of the overlapping cross sectional areas does reduce the contributions from certain multiple processes, namely those associated with separate constituent atoms. The reduction of multiple electron contributions may be viewed as somewhat similar to what is contained in the dynamical mean-field CTMC model [35], which in contrast to the frozen-potential CTMC calculations shows comparable behavior to what we find in the IAM-PCM results.

IV. CONCLUSIONS

We have presented new basis generator method calculations for scattering of projectile ions with charges $Q = 2, 3$ from fundamental atoms that are the building blocks for biologically relevant molecules. The calculated net ionization cross sections were then used to compute independent atom model cross sections for molecular targets using both the simple additivity rule (IAM-AR, which should represent the correct high-energy limit) and an independent atom model that is based on the idea that the projectiles only experience the overlap of atomic cross sections taken as geometric areas (carried out by a pixel counting method or PCM). The scaling behavior of the IAM-AR and IAM-PCM results is established, and a parametrization is presented that is capable of reproducing the computed IAM-PCM net ionization cross sections in terms of the proton-molecule IAM-PCM and IAM-AR results. To a good approximation this can also be obtained in terms of the IAM-PCM proton-molecule cross section alone. These scaled results are then compared to experimental data for two targets, namely uracil and water vapor. While the agreement is not perfect a number of trends are reproduced better than by any other theory applied previously to the problem. The theoretical scaling model was used to compute reduced $Q = 1$ cross sections from $Q \geq 2$ experimental data for uracil and water and compare them to the proton impact case. We also pointed out a few inconsistencies in the experimental data - this fact implies that further experiments are needed. For the uracil target, in particular, systematic measurements for at least $Q = 1, 2$ in the tens of keV/amu energy range would help to establish the entire domain where the scaling model applies in nature. Such investigations together with the presented
results will pave the way towards reliable simulations for cancer therapy with highly charged ions.

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[1] M. E. Alcocer-Ávila, M. A. Quinto, J. M. Monti, R. D. Rivarola, and C. Champion, *Scientific Reports* **9**, 14030 (2019).

[2] A. N. Agnihotri, S. Kasthurirangan, S. Nandi, A. Kumar, M. E. Galassi, R. D. Rivarola, O. Fojón, C. Champion, J. Hanssen, H. Lekadir, P. F. Weck, and L. C. Tribedi, *Phys. Rev. A* **85**, 032711 (2012).

[3] A. N. Agnihotri, S. Kasthurirangan, S. Nandi, A. Kumar, C. Champion, H. Lekadir, J. Hanssen, P. F. Weck, M. E. Galassi, R. D. Rivarola, O. Fojón, and L. C. Tribedi, *Journal of Physics B: Atomic, Molecular and Optical Physics* **46**, 185201 (2013).

[4] P. López-Tarifa, M.-A. Hervé du Penhoat, R. Vuilleumier, M.-P. Gaigeot, I. Tavernelli, A. Le Padellec, J.-P. Champeaux, M. Alcamí, P. Moretto-Capelle, F. Martín, and M.-F. Politis, *Phys. Rev. Lett.* **107**, 023202 (2011).

[5] C. Covington, K. Hartig, A. Russakoff, R. Kulpins, and K. Varga, *Phys. Rev. A* **95**, 052701 (2017).

[6] A. B. Salo, A. Alberg-Flojborg, and I. A. Solov'yov, *Phys. Rev. A* **98**, 012702 (2018).

[7] A. Itoh, Y. Iriki, M. Imai, C. Champion, and R. D. Rivarola, *Phys. Rev. A* **88**, 052711 (2013).

[8] S. Paredes, C. Illescas, and L. Méndez, *Eur. Phys. J. D* **69**, 178 (2015).

[9] H. J. Lüdde, A. Achenbach, T. Kalkbrenner, H.-C. Jankowiak, and T. Kirchner, *Eur. Phys. J. D* **70**, 82 (2016).

[10] H. J. Lüdde, M. Horbatsch, and T. Kirchner, *Eur. Phys. J. B* **91**, 99 (2018).

[11] H. J. Lüdde, M. Horbatsch, and T. Kirchner, *J. Phys. B* **52**, 195203 (2019).
[12] H. J. Lüdde, M. Horbatsch, and T. Kirchner, The European Physical Journal D 73, 249 (2019).

[13] M. Zapukhlyak, T. Kirchner, H. J. Lüdde, S. Knoop, R. Morgenstern, and R. Hoekstra, J. Phys. B 38, 2353 (2005).

[14] A. M. P. Mendez, C. C. Montanari, and J. E. Miraglia, “Ionization of biological molecules by multicharged ions using the stoichiometric model,” (2019), arXiv:1909.13847 [physics.atm-clus]

[15] A. Jorge, M. Horbatsch, C. Illescas, and T. Kirchner, Phys. Rev. A 99, 062701 (2019).

[16] L. H. Tobureu, W. E. Wilson, and R. J. Popowich, Radiation Research 82, 27 (1980).

[17] M. E. Rudd, T. V. Goffe, and A. Itoh, Phys. Rev. A 32, 2128 (1985).

[18] M. A. Bolorizadeh and M. E. Rudd, Phys. Rev. A 33, 888 (1986).

[19] H. Luna, A. L. F. de Barros, J. A. Wyer, S. W. J. Scully, J. Lecointre, P. M. Y. Garcia, G. M. Sigaud, A. C. F. Santos, V. Senthil, M. B. Shah, C. J. Latimer, and E. C. Montenegro, Phys. Rev. A 75, 042711 (2007).

[20] H. Luna, W. Wolff, E. C. Montenegro, A. C. Tavares, H. J. Lüdde, G. Schenk, M. Horbatsch, and T. Kirchner, Phys. Rev. A 93, 052705 (2016).

[21] D. Ohsawa, H. Tawara, F. Soga, M. E. Galassi, and R. D. Rivarola, Physica Scripta T156, 014039 (2013).

[22] S. Bhattacharjee, S. Biswas, C. Bagdia, M. Roychowdhury, S. Nandi, D. Misra, J. M. Monti, C. A. Tachino, R. D. Rivarola, C. Champion, and L. C. Tribedi, Journal of Physics B: Atomic, Molecular and Optical Physics 49, 065202 (2016).

[23] S. Bhattacharjee, S. Biswas, J. M. Monti, R. D. Rivarola, and L. C. Tribedi, Phys. Rev. A 96, 052707 (2017).

[24] Bhattacharjee, Shamik, Bagdia, Chandan, Chowdhury, Madhusree Roy, Monti, Juan M., Rivarola, Roberto D., and Tribedi, Lokesh C., Eur. Phys. J. D 72, 15 (2018).

[25] H. J. Lüdde, T. Spranger, M. Horbatsch, and T. Kirchner, Phys. Rev. A 80, 060702(R) (2009).

[26] M. Murakami, T. Kirchner, M. Horbatsch, and H. J. Lüdde, Phys. Rev. A 85, 052704 (2012).

[27] M. Murakami, T. Kirchner, M. Horbatsch, and H. J. Lüdde, Phys. Rev. A 85, 052713 (2012).

[28] M. Murakami, T. Kirchner, M. Horbatsch, and H. J. Lüdde, Phys. Rev. A 86, 022719 (2012).

[29] T. Pausz, H. J. Lüdde, M. Murakami, M. Horbatsch, and T. Kirchner, Journal of Physics: Conference Series 488, 102013 (2014).

[30] M. Rubio, L. Serrano-Andrés, and M. Merchán, The Journal of Chemical Physics 128, 104305 (2008) https://doi.org/10.1063/1.2837827.
[31] L. F. Errea, C. Illescas, L. Méndez, and I. Rabadán, Phys. Rev. A 87, 032709 (2013).
[32] C. Illescas, L. F. Errea, L. Méndez, B. Pons, I. Rabadán, and A. Riera, Phys. Rev. A 83, 052704 (2011).
[33] L. Errea, C. Illescas, L. Méndez, I. Rabadán, and J. Suárez, Chemical Physics 462, 17 (2015).
[34] Otranto, Sebastian, Bachi, Nicolás, and Olson, Ronald E., Eur. Phys. J. D 73, 41 (2019).
[35] A. Jorge, M. Horbatsch, and T. Kirchner, “Multi-charged ion-water molecule collisions in a classical-trajectory time-dependent mean-field theory,” (2020), arXiv:2001.06539 [physics.atom-ph].
[36] T. Kirchner, M. Horbatsch, and H. J. Lüdde, Phys. Rev. A 64, 012711 (2001).
[37] J. Faulkner, I. B. Abdurakhmanov, S. U. Alladustov, A. S. Kadyrov, and I. Bray, Plasma Physics and Controlled Fusion 61, 095005 (2019).
[38] M. B. Shah and H. B. Gilbody, Journal of Physics B: Atomic and Molecular Physics 14, 2361 (1981).
[39] M. B. Shah, D. S. Elliott, P. McCallion, and H. B. Gilbody, Journal of Physics B: Atomic, Molecular and Optical Physics 21, 2455 (1988).
[40] M. B. Shah and H. B. Gilbody, Journal of Physics B: Atomic and Molecular Physics 15, 413 (1982).
[41] M. E. Rudd, Y. K. Kim, D. H. Madison, and J. W. Gallagher, Rev. Mod. Phys. 57, 965 (1985).
[42] J. Tabet, S. Eden, S. Feil, H. Abdoul-Carime, B. Farizon, M. Farizon, S. Ouaskit, and T. D. Märk, Phys. Rev. A 81, 012711 (2010).
[43] J. Tabet, S. Eden, S. Feil, H. Abdoul-Carime, B. Farizon, M. Farizon, S. Ouaskit, and T. D. Märk, Phys. Rev. A 82, 022703 (2010).
[44] L. Sarkadi, Journal of Physics B: Atomic, Molecular and Optical Physics 49, 185203 (2016).
[45] L. Sarkadi, Phys. Rev. A 92, 062704 (2015).
[46] M. E. Rudd, T. V. Goffe, R. D. DuBois, and L. H. Toburen, Phys. Rev. A 31, 492 (1985).
[47] D. Ohsawa, Y. Sato, Y. Okada, V. P. Shevelko, and F. Soga, Phys. Rev. A 72, 062710 (2005).
[48] C. D. Cappello, C. Champion, O. Boudrioua, H. Lekadir, Y. Sato, and D. Ohsawa, Nuclear Instruments and Methods in Physics Research Section B: Beam Interactions with Materials and Atoms 267, 781 (2009).