Young-type interference in projectile-electron loss in energetic ion-molecule collisions

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Abstract

Under certain conditions an electron bound in a fast projectile-ion, colliding with a molecule, interacts mainly with the nuclei and inner shell electrons of atoms forming the molecule. Due to their compact localization in space and distinct separation from each other these molecular centers play in such collisions a role similar to that of optical slits in light scattering leading to pronounced interference in the spectra of the electron emitted from the projectile.

PACS numbers: PACS:34.10.+x, 34.50.Fa

The wave–particle duality, which states that all atomic objects exhibit particle as well as wave properties, is one of the basic concepts of quantum mechanics. Proposed initially by Louis de Broglie [1] in 1923, this concept has been confirmed few years later in the electron diffraction experiments [2]–[3]. Since then, a large number of investigations have been performed in order to observe the wave nature of not only electrons but also heavier particles such as, for example, neutrons, atoms, dimers and even fullerenes C_{60} [4]. Most of these measurements were aimed at a demonstration of Young’s double–slit phenomena, in which the coherent addition of the amplitudes of two (or many) paths, leading to interference, is related to the wave–like particle behavior.

In the atomic world the natural analog of the Young’s slits is represented by diatomic molecules. Starting with the works [5]–[9], especially significant interest has been focused on studying interference phenomena involving homo-nuclear molecules [7]–[22].

These studies were dealing with two principally different interference scenarios. In one of them the attention was focused on interference in the spectra of electrons emitted from the molecule in the course of photoionization [6]–[13] and consequent Auger decay [14], as well as in ionization by electrons [12] and heavy ions [16]–[18]. Note that in such a scenario, unlike the Young’s experiment, the wave is not diffracted by the ”slits” but rather emerges from them. In the second scenario, which was realized in [19]–[22] for electron capture and proton scattering and is a more direct analog of the Young’s optical experiment, interference is caused by coherent scattering of the incident projectile on the atomic centers of the molecule.

In this letter we propose yet another way to collision-induced interference. It falls into the second scenario but, similarly to [16]–[18], deals with interference in electron emission spectra. It is realized in collisions of molecules with partially stripped multiply-charged projectile-ions, in which the electron(s) of the projectile is emitted. Compared to the electron emission, studied in [16]–[18], the present case possesses important differences. In particular, in the situation, considered in [16]–[18], the electron wave is launched from the ”slits”, which are not really separated and well localized since the electrons of molecules like H_2 occupy the whole volume of the molecule and are mainly located not on the atomic nuclei but rather between them. As a result, the corresponding interference pattern is rather smooth. In contrast, as will be shown below, the emission from the projectile occurs due to a coherent scattering of the electron of the projectile on the nuclei of the molecule (partially screened by the inner shell electrons) and, therefore, the ”slits” are very well separated and localized in space that can lead to very pronounced interference effects in the emission pattern.

Below, based on the relativistic time–dependent perturbation approach, we shall derive the cross section for electron loss in collisions with homo-nuclear dimers. The possibility of interference effects will be demonstrated by calculating the cross section for fast hydrogen–like magnesium Mg^{11+}(1s) and S^{15+}(1s) ions colliding with N_2 dimers.

Atomic units are used throughout except where otherwise stated.

Since the collision between an ion carrying an electron and a molecule in general represents a very complex many-body problem, our consideration will be based on a simplified model which, however, takes into account all essential physics of the collision process in question. Within this model, in order to describe electron transitions in the projectile we shall use the first order perturbation theory in the interaction between this electron and the molecule. Such an approximation is a good one, provided Z_p \gtrsim Z_A, where Z_p and Z_A are, respec-
tively, the nuclear charges of the ion and the atoms in the molecule, and one merely wishes to describe projectile-electron transitions, without paying attention to what happens with the molecule in such collisions.

Further, we shall only consider molecules whose atoms have relatively large atomic numbers, \( Z_A \gg 1 \). Under the simultaneously fulfilled conditions \( Z_A \gg 1 \) and \( Z_p \sim Z_A \) the main contribution to the projectile-electron transitions in collisions with the molecule is given by the screening target mode, in which the projectile electron interacts with the molecule \(^8\) “frozen” during the short collision time in its initial state \(^9\).

Moreover, provided the condition \( Z_p \approx v \) is fulfilled (\( v \) is the collision velocity) the momentum transferred in the collision becomes so large (on the molecular scale) that the outer electrons of the molecule are not able to screen the nuclei of the molecule. Therefore, the main contribution to the electron loss arises from the interaction with the nuclei of the molecule partially screened by the inner shell electrons. Thus, the projectile electron undergoes transitions due to the interaction with well localized centers of force which, in addition, are well separated in space. Besides, since the inner electrons are basically atomic electrons, one can treat the molecule as a sum of free atoms and use the atomic parameters for the description of the field produced by the molecule in the collision.

Taking all this into account, the scalar potential describing the field of the molecule in its rest frame \( K' \) can be written as

\[
\Phi_M(r') = \sum_{j=1}^{2} Z_j \phi_j(|r' - R'_j|),
\]

where \( r' \) is the observation point of the field and \( R'_j \) is the coordinate of the nucleus of the \( j \)-th atom, \( Z_j \) the charge of the nucleus and

\[
\phi_j(x) = \sum_l A^{j,l}_l \exp(-\kappa_j^l x)
\]

with the screening parameters \( A^{j,l}_l \) \((\sum_l A^{j,l}_l = 1)\) and \( \kappa_j^l \) tabulated in \(^{24} \) and \(^{25} \).

It is convenient to treat the projectile-electron transitions using the reference frame \( K \) in which the nucleus of the projectile is at rest. We take the position of the nucleus as the origin of \( K \) and assume that in this frame the center of mass of the molecule moves along a straight-line classical trajectory \( \mathbf{R}(t) = \mathbf{b} + \mathbf{v}t \), where \( \mathbf{b} = (b_x, b_y, 0) \) is the impact parameter, \( \mathbf{v} = (0, 0, v) \) is the collision velocity and \( t \) is the time. Using Eqs. \(^11\)–\(^12\) and the Lorentz transformation for the potentials we obtain that the electromagnetic field of the molecule in the frame \( K \) is described by the potentials

\[
\Phi_M(r, t) = \gamma \Phi_M(s_t),
\]

\[
\mathbf{A}_M(r, t) = \left( 0, 0, \frac{v}{c} \Phi_M \right),
\]

where \( r = (\mathbf{r}_\perp, z) \) with \( \mathbf{r}_\perp \cdot \mathbf{v} = 0 \) is the coordinate of the point of observation of the field in the frame \( K \), \( c \) is the speed of light and \( \gamma = 1/\sqrt{1 - v^2/c^2} \) is the collisional Lorentz factor. Further,

\[
s_j = (\gamma(z - vt_j), \mathbf{r}_\perp - \mathbf{b}_j),
\]

where \( \mathbf{b}_j = \mathbf{b} + \delta \mathbf{b}_j \) is the impact parameter for the nucleus of the \( j \)-th atom of the molecule, \( t_j \) is the time of its closest approach to the origin and \( s_j \) is the vector connecting the position of the \( j \)-th atomic nucleus of the molecule and the electron of the ion (as is viewed in the rest frame of the molecule).

Using the first order perturbation theory in the interaction of the electron of the ion with the molecular field, described by the potentials \(^{13} \), one can show that the cross section \( \sigma_{fi} \) for the projectile-electron transitions occurring in collisions with the molecule is given by

\[
\sigma_{fi} = 4\sigma_{fi}^{(A)} \cos^2 \left( \frac{\mathbf{q}' \cdot \mathbf{l}_0}{2} \right).
\]

Here, \( \sigma_{fi}^{(A)} \) is the cross section for the projectile-electron transitions occurring in collisions with the corresponding single atom, \( \mathbf{q}' = (q_{\perp}, \frac{q\cdot s_t}{\gamma v}) \) is the momentum transferred to the projectile-ion (as viewed in the rest frame of the molecule) with \( s_t \) being the transition frequency for the electron of the ion, and \( \mathbf{l}_0 = (l_0, \vartheta_M, \varphi_M) \) is the vector connecting the positions of the atomic nuclei of the molecule in its rest frame. In what follows we shall count the polar orientation angle \( \vartheta_M \) of the molecule from the direction of the projectile velocity \( \mathbf{v} \). Besides, we set \( \varphi_M = 0^\circ \).

In figure 1 we present the electron loss cross section, \( d^3\sigma/dp_{\perp}dp_{tr}d\varphi_p \), differential in the longitudinal \( (p_{\|} = \mathbf{p} \cdot \mathbf{v}/v) \) and transverse \( (p_{tr} = \sqrt{p_{\perp}^2 - p_{\|}^2}) \) momenta and the azimuthal emission angle \( \varphi_p \) of the electrons emitted from 7.8 MeV/u Mg\(^{11+}\)(1s) projectiles in collisions with N\(_2\) molecules. The cross section is obtained by integrating over the vector of the transverse momentum transfer \( \mathbf{q}_{\perp} \). In the figure this cross section is given in the target frame as a function of \( p_{\|} \) and \( p_{tr} \) for the emission into the plane spanned by the molecular axis and projectile velocity (i.e. for \( \varphi_p = 0^\circ \)). The molecular polar orientation angle is \( \vartheta_M = 90^\circ, 20^\circ \) (the upper row, from left to right), \( 15^\circ, 10^\circ \) (the second row, from left to right) and \( 5^\circ, 0^\circ \) (the lower row, from left to right). At small \( \vartheta_M \) the spectra exhibit very clear structures, which arise due to interference caused by the coherent interactions between the electron of the projectile and the two atomic centers of the molecule \(^{20} \).

At an impact energy of 7.8 MeV/u (\( v = 17.6 \) a.u.) the typical momentum transfer to the electron of the ion, which is necessary for its removal out of the ion, is \( \sim 6-8 \) a.u.. This magnitude is substantially larger than the typical momenta of the outer electrons of nitrogen. This means that within the screening target mode the projectile-electron transitions are governed mainly by
the interaction between this electron and the target nuclei (partly screened by the K-shell electrons). Moreover, since the momentum transfers are large, the relative contribution of the collision mode, in which the target is excited to the projectile-electron loss process is by about $Z_A = 7$ times smaller than that due to the screening mode. Thus, the outer target electrons have a minor effect on the projectile-electron transitions and, therefore, the latter ones can indeed be regarded as occurring due to the interaction with two "slits", which are well localized and well separated from each other within the space occupied by the molecule.

At small polar orientation angle of the molecule the spectrum displays clear ring-like structures. The center of the rings is located at the point $\mathbf{p}_C = (p_{tr} = 0; p_{tg} = m_e v)$, where $m_e$ is the electron mass, implying that each ring is formed by electrons which in the rest frame of the projectile have close energies. Indeed, the origin of these structures can be traced back by considering the energy spectrum of the emitted electrons in the rest frame of the projectile. Such a spectrum is shown in figure 2. It is seen that in this frame the energy spectrum exhibits oscillations (especially pronounced at very small $\vartheta_M$) due to the alternation in the energy spectrum of the parts with constructive and destructive interferences. It is not difficult to convince oneself that the ring-like structures in the momentum spectrum originates namely from these oscillations.

For more information in figure 2 we present the same cross section as in figure 1 but for the electron loss from $20\text{ MeV/u } \text{S}^{15+}(1s)$ projectiles colliding with $\text{N}_2$. Like in the previous case, the interference pattern in figure 2 is caused by the coherent scattering of the electron of the projectile on the two "slits", which are
very well localized in space and are distinctly separated from each other.

Comparing the emission patterns in figures 1 and 3 we see that the range of the molecular orientation angle $\vartheta_M$, at which the interference effects are clearly visible in the emission pattern, decreases when the charge $Z_p$ of the projectile nucleus increases. This can be easily understood if we recall that the size of the electron orbit in the initial state scales as $1/Z_p$. Therefore, a more tightly bound electron can interact simultaneously with both molecular centers only if the transverse size of the molecule $l_T = l_0 \sin \vartheta_M$ becomes smaller.

As seen in figures 1–3, the most pronounced interference pattern in the emission spectrum arises at small orientation angles of the molecule. Therefore, in order to verify predicted effects in an experiment, it is very desirable to single out those loss events, which occur at small orientation angles, from the rest. This can be achieved by the determination of the molecular orientation ex post, which has been successfully applied in many experimental situations where molecular targets dissociated or Coulomb exploded after photo- and strong-field ionization or due to electron or ion impact induced ionization.

In the collisions, considered above, by far a dominant contribution to the total electron emission is given by electrons ejected from the target. Therefore, an important question to address is whether in the momentum transfer and the nuclei of the atomic centers (partially screened by the inner shell atomic electrons). This means that the interference arises from the scattering of the projectile electron on atomic "slits", which are well localized in space and distinctly separated from each other, playing a role rather similar to that of the optical slits in the Young-type experiments with photons. Owing to recent advances in the experimental techniques it has become feasible to test the above theoretical predictions. In particular, this is planned to be done in forthcoming experiments at MPI-K (Heidelberg, Germany).

In conclusion, we have considered interference effects in the electron emission accompanying energetic collisions of ionic projectiles with molecular targets. In contrast to all the previous studies of this subject, which were focused on interference in the electron emission from the target, we were searching for signatures of the interference effects in the electron emission from the projectile. We have shown that this emission may possess very clear interference structures which are caused by the coherent interactions between the electron of the projectile and the atomic centers of the molecule. Under certain conditions (which were discussed in detail above) this interaction is basically the one between the electron of the projectile and the nuclei of the atomic centers (partially screened by the inner shell atomic electrons). This means that the interference arises from the scattering of the projectile electron on atomic "slits", which are well localized in space and distinctly separated from each other, playing a role rather similar to that of the optical slits in the Young-type experiments with photons.

Note that in the 90°-case the emission pattern is practically identical to that in collisions with two independent nitrogen atoms.

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