ENERGY APPROACH TO ELECTRON CAPTURE AND IONIZATION PROCESSES IN ION-ATOMIC COLLISION SYSTEM

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Abstract. Energy approach is generalized to calculate the electron capture cross sections in the H++ H(1s) collision system. The numerical results are presented for collision energies 10 and 100 keV

Keywords: ion-atomic collision system, energy approach

1. Introduction

Studying characteristics of the electron, ion-atomic collision processes attracts traditionally an intense interest because of the important applications in the astrophysics, plasma and laser physics etc. [1-24]). One of the recent actual problems is investigation of the collisions dynamics between the atoms and surfaces, walls and nanostructures. Modern technological advances have made it possible to perform experiments with full control of the ion-atomic collision systems [1,2,7]. Naturally, the collision processes are to be studied for understanding emission spectra of the plasma [2,10,17,22]. An important application of the theory of atomic collision theory of plasma is search of the optimum plasma excitation condition for lasing and discovery of new pumping approaches. In addition, these investigations are important to understand the plasma processes themselves. Given systematic data about intensities of spectral lines and respective gins, one can establish basic rules of plasma motion in phase space. The history of plasma spectrum modeling ex-
tends for decades, starting with the simplest models (see, for instance [2,10]). Different atomic levels are populated in laboratory plasma by different physical processes. This results in a different dependence of each line intensity on the plasma parameters. It had been understood for a long time that this phenomenon could be used for plasma diagnostics. The general principles of such a diagnostic for the simple H-like and He-like ions have been formulated [2]. These diagnostic principles have proven to be useful for understanding the physics of the system and for planning new experiments. In the last years a special interest attracts a studying the electron, ion-atomic collisions in the Debye plasma [3,6,21]. Nevertheless, a consistent theory of accounting for effects of the screened Coulomb interaction on the collision dynamics in multielectron atomic systems is absent hitherto. Traditionally a majority of papers devoted to collision processes in relatively simple systems. In particular, one could mention the proton—H and hydrogen-like ion, H-He collisions, which are accompanying by different impact-excitation, collision ionization and capture phenomena. The usually used theoretical approaches to these problems are the standard quantum-mechanical perturbation theory (PT), the R-matrix approach, the classical trajectory Monte Carlo method etc. [1-6]. For example, in ref. [2] the Monte Carlo method approach has been used to calculate the electron capture and ionization cross sections in hydrogen atom, fully striped ion collisions determined in the Debye-Hückel potential. In ref. [3] the two-centre atomic orbital close-coupling method has been used to calculate the cross-sections of the excitation and electron capture processes in the H^++ H(1s) collision system in a Debye plasma. In the cited paper it has been shown that the dynamics of electron capture and excitation processes is significantly affected by the effect of interaction screening on the direct and exchange electronic couplings and reducing the number of reaction channels. The more information regarding the considered topics can be found in the references [3,6,21]. It is obvious that studying collision dynamics between heavy atoms, multicharged ions and fully striped ions requires a relativistic generalization of the cited methods. Besides, one could mention the cooperative electron, ion-atomic collision processes when the different decay channels (including the production of new particles, the electron-positron pair production (EPPP) in collision of heavy and super heavy ions and nuclei) are opened and must be taken into account simultaneously [2,11,14,15]. From this point of view above cited and other methods are dealing with known fundamental theoretical and computational problems [2,15]. In our opinion, generally speaking, the problem of adequate treating the ion-atomic collision with electron capture and other accompanying processes requires a development consistent quantum electrodynamics (QED) approach or some approximation to QED theory (energy approach). The energy approach (the fundamental aspects) has been developed in refs. [9-14] and then modified and applied to studying a number of problems, namely, studying the electron-ion collision dynamics in the Debye plasma, the resonant states of compound super-heavy nucleus (ions) and EPPP in heavy ions collisions, the recoil induced excitation and ionization in atoms during capture of neutron, the discharge of metastable nuclei during the negative muon capture and collisions of the ultracold atoms with walls and nanostructures etc. [15-23]. In fact it had been applied previously in the study of the purely electronic, electron-nuclear processes in atoms and meso-atomic systems. In this paper the energy approach is generalized to calculate the electron capture cross sections in the H^++ H(1s) collision system. The numerical results are presented for collision energies 10 and 100 keV. It is important to note that the presented theory is initially relativistic and, in our opinion, can be used in studying the collision dynamics of heavy atoms, multicharged ions and fully stripped ions. Let us recall very interesting task, connected with investigation of the collision dynamics for H-like and Li-like ions U^{+91}, U^{+89}[14].

2. The energy approach to collision problem and model potentials

In the theory of the non-relativistic atom a convenient field procedure is known for calculating the energy shifts ΔE of degenerate states because of the interparticle interaction or interaction with an external field [9,12]. This procedure is connected with the secular matrix M diagonalization. In constructing M, the Gell-Mann and Low adiabatic formula for ΔE is used. A similar energy approach, using the Gell-Mann and Low formula with the QED scattering matrix, is applicable in the relativistic atom theory [9-14]. The method is a consistently electrodynamics one, allowing for the uniform consideration of a variety of induced and spontaneous
processes different by their physical nature and with any number of photons. In contrast to the non-relativistic case, the secular matrix elements are already complex in the PT second order (first order of the inter-electron interaction). Their imaginary parts are connected with the radiation decay (collision decay, decay in an external electromagnetic field etc.) possibility. The total energy shift of the state is usually presented in the form [12]:

\[ \Delta E = \text{Re}\Delta E + i \text{Im}\Delta E \quad \text{Im}\Delta E = -\Gamma/2, \tag{1} \]

where \( \Gamma \) is interpreted as the level width, and the decay possibility \( P = \Gamma \). The whole calculation of the energies and decay probabilities of a non-degenerate excited state is reduced to calculation and diagonalization of the complex matrix \( M \). The calculation procedure for \( \text{Re}\Delta E \) may be generalised for the case of nearly degenerate states, whose levels form a more or less compact group. One of these variants has been previously used by us [17,18] for a system with a dense energy spectrum, a group of nearly degenerate states is extracted and their matrix \( M \) is calculated and diagonalized. If the states are well separated in energy, the matrix \( M \) reduces to one term, equal to \( \Delta E \). To start with the QED Gell-Mann and Low formula one must choose the zero-order approximation. Usually one uses for this purpose a one-electron Hamiltonian with a central potential that can be treated as a bare potential in the formally exact QED PT. The bare potential includes the electric potential of the atomic nucleus and some model potential of the inter particle interaction. Let us underline that the QED approach is useful in our task as a tool for explaining approximations and a regular method for generalizations in order to take into account additional physical effects. It is obvious that a non-relativistic approximation is quite acceptable in treating the \( \text{H}^+ + \text{H} \) collision dynamics.

In general case let us consider a collision of two similar ions (atoms) with nuclear charge \( Z \) and mass \( M \). Following the general formalism we may introduce the bare interaction which can be treated as the zeroth approximation in some formally exact QED PT. As bare potential it is natural to choose non-relativistic electron-nuclear interaction \( W(r,R) \) and the inter-nuclear interaction \( V(R) \). Here \( R \) is distance between the nuclei (ions) and \( r \) is the electron coordinate in a centre of the nuclear masses. In a case of the heavy ions (nuclei) collision these potentials include the corresponding terms, connected with account for the finite nuclear size effects and possible strong inter nuclear interaction for small \( R \) [14-16]. The smallness parameter of the QED PT is the fine structure constant \( \alpha = 1/137,034 \). Under given choice of the bare interaction in the PT zeroth approximation we are dealing with quantum mechanical multi-body problem with known interaction potentials. Such task may be solved within the operator PT [11,13,17], which is called by quantum mechanical PT (QMPT) in a difference from the QED PT [12]. In the QMPT zeroth order a movement of nuclei (nuclear subsystem) is treated independently upon the electron subsystem and described by some equation with potential \( V(R) \). One may choose the Dirac equation (DE) for nuclear subsystem in our case only from the point of view of the theoretical consistency and analogy with the electron subsystem, which is described by the DE too. Such an approach is surely more exact than it is necessary for solving the \( \text{H}^+ + \text{H} \) collision problem.

As the potential \( W(r,R) \) it is considered the potential of two point Coulomb centres. The full electron-ion interaction can be represented as follows:

\[ W(r,R) = W_s(r,R) + \sum_x U_{2x}(r,R)P_{2x}(r,R), \tag{2} \]

where a spherically symmetric part is formally as follows: \( W_s(r,R) = U(R) \), if \( r < R \) and \( U(r) \), if \( r > R \) and \( P_{2x}(r,R) \) are the Legandre polynomials and \( U_{2x}(r,R) \) are the radial parts of the non-spherical interaction. We use here a terminology from the one-centre theories of molecules and quasimolecules (see [2,14]). The \( U(r) \) can be defined as follows: \( U(r) = W(r, R = 0) \). Naturally in a case of the heavy ions, this potential is defined by a charge distribution in the corresponding nucleus. Surely in a case of the heavy ions, it will contain two continuums (up and down) with boundaries \( \pm a^2 \), where \( a = \alpha \cdot Z \) (see [11,14]).

In the QMPT zeroth approximation the Hamiltonian of the electron-nuclear system is in a representation of the second quantization as follows:

\[ H_0 = \sum_{i,j} (H_{0i})_i a_i^\dagger a_j + \sum_{i,j} (H_{0ij})_i a_i^\dagger A_j, \tag{3} \]

where \( a^\dagger, a \) are the electron creation and annihilation operators and \( A^\dagger, A \) are the same for the nuclear particles (generally speaking quasi-particles corresponding the collective coordinate \( R \); see below); \( (H_{0i})_i \) and \( (H_{0ij})_i \) the one-particle matrix elements of the corresponding zeroth-order Hamiltonians for the electron and nuclear subsystems. Formally, the representation for nuclear subsystem
is given by a spectrum of the DE with some inter
uclear potential \( V(R) \) and for the electron subsystem
it is given by a spectrum of solutions of the DE
with potential of the zeroth approximation \( U_0(r) \)
\((U(r))\). The corresponding system for radial parts
of the Dirac function is as follows (in the Coulomb
units):

\[
F = -\frac{F(\omega + |\omega|)}{T} - G(E + 2M|\alpha|^{-2} - U)|\alpha|,  \\
G = \frac{G(\omega - |\omega|)}{T} + F(E - U)|\alpha|,  \\
\]

where \( \alpha \) is the Dirac angular quantum number, \( E \)
is the state energy, \( F, G \) being the large and small
radial components correspondingly \((F=dF/dT; \ G=dG/dT)\); \( T \) is the general argument for all sys-
tem of differential equations (generally speaking for the
electron and for the nuclear subsystems). The QMPT perturbation is as follows:

\[
\sum_{i,j} (V_{int e}) a_i^* a_j + \sum_{i,j,k,l} (V_{int EN}) a_i^* A_j a_k a_l,  \\
\]

where \((V_{int e})\) is the matrix element of the one-elec-
tron operator \( U(r) - U_0(r) \), \((V_{int EN})\) is the matrix
element of the potential \( W(r,R) - U(r) \).

The differential cross-section for electron cap-
ture \( nl \) (or ionization \( \varepsilon s \)) in the collision in the low-
est QMPT approximation is proportional to the square of the matrix element:

\[
M_{1s,F\varepsilon s} = \int dR d\varepsilon s \psi_F^* (1s|\varepsilon s) \psi_F^* (1|\varepsilon s) \times \\
\sum_{i,j} (V_{int e}) a_i^* a_j + \sum_{i,j,k,l} (V_{int EN}) a_i^* A_j a_k a_l,  \\
\]

where \( \psi_F \) are the initial and final state functions for nuclear subsystem (see below). After integ-
ration on the electron coordinate the matrix ele-
ment (5) becomes:

\[
M_{1s,F\varepsilon s} = \int dR J_{IF}(R) \Phi_{1s}(R),  \\
\Phi_{1s} = \int_0^\gamma dr \psi_{1s}^* (W_F(r,R) - U(r)) \psi_{1s} + \\
\int_0^\gamma d\varepsilon s H_{\varepsilon s} \psi_{1s}^\varepsilon s.  \\
\]

Here \( H_{\varepsilon s} \) is the total Hamiltonian with potential
\( U(R) \). The notation for the production of the nu-
clear state functions is introduced:

\[
J_{IF}(R) = \psi^*_F(R) \psi_F(R).  \\
\]

This function includes the total information about
the model describing the nuclear subsystem.
The function \( \Phi_{1s} \) includes the total information
about the electron subsystem and the model describ-
ing the electron-nucleus interaction. Naturally, the
last term in (6) is not needed for the present task,
as it is formally corresponding to the cooperative
processes in a system (EPPP etc.) \([11,14]\). Follow-
ing the general receipt of the operator PT \([11,13]\,
we suppose that the eigen functions and eigen ener-
gies for the potentials \( U(r) \) and \( U_0(r) \) coincide
and they are defined by its energy spectrum and the set
of the eigen functions without specifying the ana-
lytic form of the zero order potential \([10]\). Such a sche-
me treats the widely known distorted waves ap-
proximation as the zeroth order approximation in
the formally exact QMPT allowing for application
of the well developed stationary-state methods to
the collision problem with variable number of par-
ticles and further successive refinement of calcula-
tion \([11,13]\). The final electron scattering function
\( \varphi_\varepsilon s \) is constructed as the quadratically integrable ei-
gen function and being the orthogonal complement
to set of the discrete state functions in a full analogy
with \([13]\). As it is indicated above, such an approach
treats the widely known distorted wave approxima-
tion in the zeroth order approximation.

Now we should define the imaginary part of the
energy shift provided by a collision process as fol-
lows:

\[
\text{Im } \Delta E = -\Gamma/2 = \text{Im } \sum_{n,s} \frac{(M_{1s,Fs})^2}{E_F + \varepsilon(ns) - E_F - \varepsilon s}.  \\
\]

Here \( \Sigma \) means summation over the discrete and
integration over the continuum parts of spectra. In-
deed, the imaginary contribution is associated with
the poles on the energy surface. The individual poles
present the concrete channels (capture to bound state,
ionization etc.) of reaction with definite final states of the electron (and indeed, nuclear) system.
The non-stationary feature of the collision problem manifests itself in the way of the normalization
of the nuclear initial and final state functions. Initially
the nuclei are free. The final state function \( \psi_F \) must
be normalized to momentum the same as the func-
tions of the virtual states in the formulae (7). The
normalization of the nuclear initial state function
\( \psi_i \) is determined by the flow of the initially free
nuclei. The value \( \Gamma \) given by formula (7) equals to the
ionization (capture) cross-section if this function
is normalized to unit flow at \( R \rightarrow \infty \). As the ze-
roth approximation the initial state of the nuclear
subsystem is described by the plane wave, which is
expanded on the spherical harmonics in order to
use the symmetry of task of the zeroth QMPT ap-
proximation. In the general case (arbitrary charges
of ions, ions with high \( Z \) etc.) surely one should
introduce the special inter nuclear interaction and exchange the trivial Coulomb potential (as in our task), for example, as it has been done in ref. [22] by introducing the differential equation for the potential \( V(R) \) in the EPPP problem in the heavy nuclei collisions and using the special quantization procedure or in ref. [21,22] by introducing the Yukawa type potential to the electron-ion collision task. This block is used by us in the present paper. In particular, we used the Debye–Hückel potential:

\[
U(r) = -\frac{Z_a \cdot Z_b}{r} \exp(-r/\lambda_D),
\]

where the Debye screening parameter \( \lambda_D \) is connected with the plasma parameters such as the temperature \( T \) and the charge density \( n \) as follows:

\[
\lambda_D = \sqrt{k_B T / 4\pi n},
\]

and \( k_B \) is the Boltzmann constant (the electron charge \( e=1 \); besides, in our case \( Z=1 \)).

Let us make several important notations. Firstly, naturally in the case of the potential \( U(r) \) there are no continuums [11]. Besides, the EPPP channel [14] is not present and naturally the nuclear structure is not detailed (indeed, in our case the nuclei are naturally considered as the point-like charges; \( Z_a=Z_b=1 \)). The relativistic effects are not important in our case of the \( H^+ + H \) collision problem. Nevertheless, we use the relativistic DE as these equations are the basic equations in the corresponding atomic numerical PC code “Superatom” (see description in ref.[8]), which is used by us in this work. Let us note that this code has been earlier used to solve a majority of the atomic and nuclear tasks, including the collision problem of heavy ions [8-24]. At last, the relativistic collision theory is corresponding to our interest to studying the collision dynamics for heavy H-like and Li-like ions such as \( U^{+89} \) and \( U^{+89}[14] \), where the role of the relativistic effects is obviously very high.

3. Results and discussion.

Below we present the results of our studying the electron capture dynamics in the \( H^+ + H \) collision system. Calculation has been carried out on the basis of the PC code “Superatom”. The calculated cross sections (7) for electron capture to the lowest states at collision energies of 10 keV and 100 keV are presented in tables 1 and 2. The two cases are considered: firstly, the unscreened (pure Coulomb interaction) case and, secondly, for the screened case with the interaction screening parameter \( \lambda_D = 4a_0 \) (\( a_0 \) is the Bohr radius). From physical point of view in the second case one may deal with quite dense plasma (the thermonuclear reactor plasma etc.). In table 1 we compare the our data on the electron capture (state \( 1s \)) cross sections for unscreened and screened (\( \lambda_D = 4a_0 \)) cases with the analogous data from ref. [3]. In table 2 we present our data on cross-sections of the electron capture (state \( nl: n=2,3; l=0,1,2 \)) cross sections for the unscreened and screened (\( \lambda_D = 4a_0 \)) cases for the same collision energies \( E=10 \) and \( E=100 \) keV.

### Table 1

| \( E (\text{keV}) \) | \( 1s \) [3] | \( 1s \) |
|---------------------|--------------|--------------|
| No screening        |              |              |
| 10                  | 7.870 57     | 7.871 42     |
| 100                 | 0.083 54     | 0.084 31     |
| With screening      |              |              |
| 10                  | 7.314 61     | 7.316 27     |
| 100                 | 0.063 94     | 0.065 04     |

### Table 2

| \( E (\text{keV}) \) | \( 2s \) | \( 2p \) | \( 3s \) | \( 3p \) | \( 3d \) |
|---------------------|--------|--------|--------|--------|--------|
| No screening        |        |        |        |        |        |
| 10                  | 0.209 98 | 0.353 12 | 0.019 08 | 0.046 21 | 0.026 15 |
| 100                 | 0.018 82 | 0.004 22 | 0.006 84 | 0.002 33 | 0.000 64 |
| With screening      |        |        |        |        |        |
| 10                  | 0.108 99 |          |          |          |          |
| 100                 | 0.003 16 |          |          |          |          |

In a whole, our results on the cross-sections of capture to the \( 1s \) state for the unscreened case and the case with screening are in a good agreement with the similar data from ref. [3]. A little difference between the presented results is probably explained by using the different approaches in the present
paper and ref. [3]. In fact, it is connected with using the different atomic orbital basis’s and different formula for the corresponding cross-section. In any case, we can conclude that the presented energy approach is successfully tested for the H++ H(1s) collision system in the Debye plasma. We believe that our approach can be effectively used for studying the electron capture processes in the more complicated ion-atomic collision systems, where an application of other standard theoretical approaches [2-7] can deal with the serious fundamental and numerical problems (see discussion in ref. [2]).

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